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Part 2
Sections IX–XVIII

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ELSEVIER

Low-threshold injection lasers based on vertically coupled quantum dots

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Abstract

We have fabricated and studied injection lasers based on vertically coupled quantum dots (VECODs). VECODs are self-organized during successive deposition of several sheets of (In,Ga)As quantum dots separated by thin GaAs spacers. VECODs are introduced in the active region of a GaAs–AlGaAs GRIN SCH lasers. Increasing the number of periods (N) in the VECOD leads to a remarkable decrease in threshold current density (\(\sim 100\) A/cm\(^2\) at 300 K for \(N = 10\)). Lasing proceeds via the ground state of the quantum dots (QD) up to room temperature. Placing the QD array into an external AlGaAs–GaAs quantum well allows us to extend the range of thermal stability of threshold current density (\(T_0 = 350\) K) up to room temperature. Using (In,Ga)As–(Al,Ga)As VECODs in combination with high temperature growth of emitter and waveguide layers results in further reduction of threshold current density (60–80 A/cm\(^2\), 300 K) and increase in internal quantum efficiency (70\%). Room temperature continuous wave operation (light output 160 mW per mirror) and lasing via the states of QDs up to \(I = (6-7) I_{th}\) have been demonstrated.

PACS: 42.55.P; 42.60; 42.60.L; 68.55.B; 68.55.J; 78.55.C

Keywords: Molecular beam epitaxy; Injection lasers; Quantum dots

1. Introduction

Application of quantum dots (QDs) as an active region in a semiconductor injection laser should lead to ultra low threshold current density and extremely high thermal stability, due to delta function density of states [1]. However, complete practical realization of these theoretical predictions still remains a challenge owing to severe technological problems in synthesizing absolutely uniform defect-free dense arrays of objects exhibiting three-dimensional quantum confinement in an appropriate
higher bandgap matrix. In the present work we use the method of direct QD formation based on a self-organization process at the initial stages of strained layer (In,Ga)As-on-GaAs heteroepitaxy [2, 3]. The quantum pyramids formed under appropriate deposition conditions exhibit a delta function like density of states, good structural perfection, reasonable size uniformity, and intense room temperature (RT) luminescence [3]. Introducing these (In,Ga)As QDs in the active region of a GRIN SCH GaAs-(Al,Ga)As laser allowed us to realize lasing via the ground state of quantum dots at low temperatures. Threshold current density was found to be practically temperature insensitive in the temperature range up to \( \sim 100 \) K [4]. However, at higher temperatures we observed the steep increase in the threshold current density accompanied by a blue shift of lasing wavelength. Room temperature lasing energy was close to optical transition energy in wetting layer (WL) [5]. In the present work we report on important improvements in lasing characteristics due to modification of the structure design and molecular beam epitaxy (MBE) growth regimes. We will show that using the concept of vertically coupled quantum dots (VECODs) [6, 7], high temperature growth of emitter and waveguide layers, and the effect of external quantum well or AlGaAs barriers on energy spectrum of a quantum dot lead to remarkable reduction in threshold current density, ground state lasing and \( T_0 = 350 \) K up to room temperature, and continuous wave (CW) operation of quantum dot injection laser at 300 K.

The structures for optical and transmission electron microscopy (TEM) studies were grown on GaAs (1 0 0) semi-insulating substrates. QDs were inserted into the middle of a 0.2 \( \mu \)m thick GaAs layer confined by 10 period AlAs (20 Å) / GaAs (20 Å) superlattices.

The laser structures were grown on (1 0 0) Si-doped GaAs substrates. They were composed of 1.5 \( \mu \)m thick Al\(_{0.4}\)Ga\(_{0.6}\)As emitters, 0.4 \( \mu \)m GRIN waveguide region and 0.6 \( \mu \)m contact layer. The lasers were fabricated in a four cleaved facet or a shallow stripe (\( W = 60 \mu \)m) geometry. Laser characteristics were studied under pulsed (pulse duration 100 ns, pulse frequency 5 kHz) and CW excitation in the 80–300 K temperature range.

TEM studies were performed on a high voltage JEOL JEM 1000 (1 MV) microscope.

PL was excited with an Ar\(^+\) laser and detected with a cooled Ge photodetector. Calorimetric absorption (CAS) spectra were taken at 500 mK.

3. VECOD lasers

3.1. Structural and optical properties of vertically coupled quantum dots

Vertically coupled (In,Ga)As/GaAs quantum dots are formed as the result of successive deposition of (In,Ga)As quantum dot sheets and thin GaAs spacers. The mechanisms of VECOD formation for the cases when the GaAs spacer is thicker or thinner than the QD height are discussed in details in Refs. [6, 7], respectively. Typical cross section TEM image of a 3-period VECOD structure is given in Fig. 1. The dot density is approximately \( 10^{11} \) cm\(^{-2}\). Each dot is well separated from the others and is composed of three individual InAs islands stacked in vertical direction. Since the GaAs spacers are very thin, the wavefunctions of neighboring islands are essentially overlapped and each VECOD can be considered as a one unified quantum-mechanical object. PL data in favor of this assumption is reported in [8]. It is important that no dislocations or large relaxed clusters are formed.

Fig. 2a shows typical PL and CAS spectra of a VECOD structure [9]. At low excitation densities PL spectrum contains intense line due to ground
state exciton transition. Increasing the excitation density or observation temperature leads to the appearance of additional peaks due to transitions involving WL states [9]. The indicated PL lines shift toward long wavelengths with increasing the temperature following the GaAs bandgap. Ground- and excited-state peaks in the PL spectrum well agree to similar peaks in calorimetric absorption spectrum indicating the presence of high-density uniform QDs.

Fig. 2b shows PL spectra of structures with different number of QD sheets, \( N \), under high excitation density. One can see that the sample with \( N = 1 \) demonstrates considerable saturation of luminescence associated with the ground-state transition in a QD, and an intense line due to transitions involving WL states is observed. For the \( N = 3 \) sample the effect of saturation of ground-state luminescence is pronounced to a less extent, and no saturation is observed for the \( N = 6 \) sample which is extremely important for laser applications [9].

### 3.2. Lasing characteristics

Remarkable progress in PL properties of VECODs as compared to single-sheet QDs led to important improvements in corresponding lasing characteristics. Fig. 3 shows the near-threshold electroluminescence (EL) spectrum of a laser structure based on the \( N = 3 \) VECOD. Lasing starts at the long-wavelength side of EL peak and the lasing wavelength is in the vicinity of the PL maximum recorded under low excitation density in the entire 77−300 K temperature range. Fig. 4 represents the temperature dependences of lasing wavelength for the VECOD and single-sheet QD based lasers. One can see that these dependences are essentially
For InGaAs QDs it decreases approximately one order of magnitude from 950 A/cm² ($N = 1$) to 98 A/cm² ($N = 10$) most probably due to the increase in optical confinement factor. The range of $J_{th}$ thermal stability is extended up to 140–150 K in VECOD lasers, as shown in Fig. 4. The reason for this is presumably the increase in localization energy of carriers in VECODs [8]. It is worth mentioning that low-temperature $J_{th}$ also decreases with $N$ reaching the value as low as 18 A/cm² (120 K, $N = 10$).

Differential efficiency ($\eta_D$) for the structures studied increases with increasing $N$, Fig. 5a [10]. However, only moderate values of $\eta_D$ were measured for the structure with $N = 10$. The reason for this is the relatively low internal quantum efficiency ($\eta_I$) characteristic for these structures. Linear extrapolation of the $1/\eta_D$-vs-cavity length ($L$) dependence (Fig. 5b) gives the value of $\eta_I \approx 0.5$. It means that a significant number of the carriers recombine nonradiatively, most probably in the GaAs layer covering the VECODs which is grown at low substrate temperature (490°C).

Improving the quality of layers adjacent to the QDs through the structure heat-treatment and increasing the separation between energy levels in QDs and barriers through inserting QDs in AlGaAs/GaAs QW or directly in AlGaAs matrix are considered below.

3.3. Effect of heat treatment on structural and optical properties of (In,Ga)As–GaAs quantum dots

A possible way to reduce the concentration of point defects due to low-temperature growth is to perform a post-growth annealing. Moreover, high-temperature AlGaAs growth ($\sim 700°C$) is essential for light emitting devices based on the AlGaAs–GaAs heterostructures to ensure good device performance and operation lifetime.

TEM studies have shown that indium composition in QDs decreases with increasing the lateral dot size, while the dot shape remains essentially the same. This leads to the noticeable blue shift of the PL peak position, as shown in Fig. 6. The increase in integrated PL intensity is presumably due to annealing of the point defects in the process of heat treatment [11].

**Fig. 3.** PL (dashed curve) and electroluminescence spectra of a VECOD laser at 80 and 300 K.

**Fig. 4.** Temperature dependence of lasing wavelength and threshold current density for the three-period VECOD (solid squares and circles) and single-sheet QD lasers (open squares and circles).

different. Unlike the case of the single-sheet QD laser, the wavelength of the VECOD laser follows the temperature dependence of the GaAs bandgap. Taking into account the lack of ground-state PL saturation described above we may conclude that in VECOD structures lasing proceeds via the ground state of quantum dot up to room temperature.

The threshold current density ($J_{th}$) in a VECOD gradually decreases with the number of QD sheets.
Room temperature values of $J_{th}$ for the QD laser with AlGaAs layers grown at 700°C are significantly lower than those for the 600°C one: 250 and 700 A/cm², respectively. We attribute this to the reduction in the concentration of point defects in a low-temperature GaAs covering QDs due to annealing during the deposition of upper layers of laser structure.

3.4. Effect of external quantum well and aligaas barriers on characteristics of (In,Ga)As QD lasers

Inserting the InAs QD array into an external AlGaAs/GaAs quantum well (QW) is a possible way to increase the localization energy of the QD relative to the WL and the GaAs matrix. Calculations within the zero radius potential approximation [12] show that in this case, the WL level shift stronger than the QD level because the latter is more localized than the former. Moreover, the three-dimensional density of states in GaAs is replaced by the two-dimensional one. These features should lead to the extension of the range of high temperature stability of the threshold current density.

In Fig. 7 we show temperature dependence of $J_{th}$ for the laser whose active region consists of three 100 Å AlGaAs/GaAs QWs separated by 50 Å barriers with the InAs QDs in the middle of each well. The three-well design of the recombination region was chosen to increase optical confinement factor. It is clearly seen that the temperature dependence of $J_{th}$ is described by $T_0 = 350$ K in the entire 77–300 K range, whereas the typical value for an InGaAs QW laser is 130–150 K.

Another way to increase carrier localization energy in a QD is to increase the barrier height. Calculations show that when the GaAs matrix is substituted by AlAs, the energy separation between electron (hole) levels in QD and WL is drastically increased [13]. In our experiments we used (In,Ga)As–Al$_{0.15}$Ga$_{0.85}$As quantum dots as the active region and Al$_{0.3}$Ga$_{0.7}$As emitters. The growth
of the emitter and waveguide layers at 700°C allowed us to overcome the problem of poor crystalline quality of (Al,Ga)As grown at 490°C through point defect annealing similar to the case of GaAs matrix described above. Injection lasers based on (In,Ga)As-(Al,Ga)As VECODs show RT threshold current density as low as 63 A/cm² and internal quantum efficiency as high as 70%.

3.5. Room temperature continuous wave (CW) operation

The considerable reduction in threshold current density allowed us to realize RT CW operation of a QD injection laser. In this work we report on characteristics of the VECOD laser based on 10 sheet (In,Ga)As QDs separated by 50 Å GaAs spacers. We used 60 μm wide laser diodes. Light output vs. current characteristics, spontaneous electroluminescence, and lasing spectra at various pumping currents are shown in Fig. 8. The maximum power per facet obtained was 160 mW and the differential efficiency was 45%. The lasing spectrum is somewhat broadened and only slightly shifted toward short wavelengths with increasing pumping current. Consequently, lasing proceeds via the states of quantum dots up to \( I = (6-7) I_{th} \), the maximum value of CW operation achieved in the present experiment.

4. Conclusions

Injection lasers based on vertically coupled quantum dots have been shown to exhibit considerable advantages over the single-sheet quantum dot lasers studied previously. They demonstrate remarkable reduction in threshold current density with increasing number of dot sheets (100 A/cm², \( N = 10 \times 300 \) K) and extended range of thermal stability due to increase in optical confinement factor and carrier localization energy. An external AlGaAs/GaAs quantum well increases the separation between the energy levels in the quantum dots, wetting layer, and GaAs matrix, leading to further extension of the range of \( I_{th} \) thermal stability up to room temperature. \( T_0 \) as high as 350 K was obtained in the entire 77–300 K range. Inserting (In,Ga)As QDs into an AlGaAs matrix in combination with high-temperature growth of emitter and waveguide layers allowed us to obtain RT threshold current density as low as 60–80 A/cm² and internal quantum efficiency as high as 70%. Room temperature CW operation of QD injection laser and lasing via the states of quantum dots up to \( I = (6-7) I_{th} \) have been demonstrated.
Acknowledgements

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References

Overgrowth of InGaAs quantum dots formed by alternating molecular beam epitaxy

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Abstract

Alternating molecular beam epitaxy is used to grow InGaAs quantum dots on (1 0 0) GaAs by utilizing the Stranski–Krastanow growth transition. A growth pause during the overgrowth with GaAs that is used to bury the quantum dots improves the photoluminescence intensity and line width compared to continuous growth of GaAs. The optimum thickness of GaAs to grow at the quantum dot growth temperature of 515°C is 9.0 nm. Multiple layers of quantum dots have been grown, and transmission electron micrographs show vertical alignment between the layers. Intensity-dependent, room-temperature photoluminescence shows that excited-state luminescence can be reduced by using multiple layers of quantum dots, but growing too many layers leads to degradation of the photoluminescence intensity.

Keywords: Quantum dots; Indium gallium arsenide; MBE

1. Introduction

Molecular beam epitaxy (MBE) is generally used to grow heterostructures of III–V semiconductors because of the atomically flat interfaces that are required for quantum well structures and devices. However, highly lattice-mismatched layers grown by MBE do not grow in a layer-by-layer fashion. Instead, these lattice-mismatched layers grow in the layer-plus-island (Stranski–Krastanow) growth mode due to a reduction in the strain energy from the island formation. The islands formed are quantum-sized, and this layer-plus-island growth mode has been successfully used to grow arrays of quantum dots (QDs) in the In(Ga)As/GaAs system [1–5], as well as many other III–V semiconductor combinations.

Several groups have recently demonstrated room-temperature operation of QD lasers [6–8] with just a single layer of QDs. Just as in quantum well (QW) lasers, though, it is desirable to obtain higher gain by stacking multiple layers of QDs. Multiple layers of QDs have also recently been demonstrated [1, 9–11], and QD lasers and vertical cavity lasers have been formed with multilayer structures [12, 13]. However, little attention has

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been paid to the overgrowth and burial of these strained, nonplanar QDs [14], although it has been shown that it is possible to have rough interfaces at subsequent layers due to the underlying islands [9]. Furthermore, the overgrowth thickness has been shown to be very important in determining the structural and optical properties of thin InAs QWs on GaAs [15]. Thus, this GaAs overgrowth is important not only for smoothing the surface so that multiple layers of QDs can be grown with good layer-to-layer uniformity, but also for minimizing the number of nonradiative defects, as is required for optimal devices. Strong interaction between layers via roughness or strain fields can lead to vertical alignment of islands [9–11]. However, this interaction can also lead to layer-to-layer non-uniformity. In this paper, we will describe our experimental techniques for improving the optical properties of QDs and the QD uniformity, both within a single layer and layer-to-layer.

2. Continuous growth versus growth pause

2.1. Sample growth

The samples are grown on semi-insulating (1 0 0) GaAs substrates in a Varian Gen II MBE machine equipped with an EPI valved arsenic cracking cell. The As$_2$ flux is kept constant throughout the growth and has a beam equivalent pressure (BEP) of 9.0 x 10$^{-9}$ Torr. This BEP is converted to an arrival rate of about 2.0 ML/s of atomic arsenic by measuring As-induced reflection high-energy electron diffraction (RHEED) oscillations on a Ga-saturated, GaAs surface [16]. A 0.25 μm GaAs buffer is grown at 590°C at a rate of about 0.93 ML/s. The sample is then cooled to 515°C under As$_2$ flux for the growth of the InGaAs quantum dots.

The shutter sequence for the growth of the InGaAs is as follows: 0.265 ML In, 2.0 ML As, 0.62 ML Ga, 3 s growth pause, and 6.0 ML As. This cycle is repeated 20 times, giving an epilayer that nominally has 30% In mole fraction and a thickness of 17.7 ML. The 2D–3D RHEED transition occurs between 7.1 and 8.0 ML of deposition. The actual In mole fraction of the quantum dots is larger than 30%, though, due to In segregation to the growth front under these growth conditions.

GaAs is then deposited using conventional MBE (simultaneous beams of Ga and As$_2$). Samples CG (continuous growth) and GP (growth pause) have 8.0 nm of GaAs deposited at the InGaAs growth temperature of 515°C. Then, the substrate temperature of sample CG is raised to 590°C while an additional 50 nm of GaAs is deposited. Sample GP has a growth pause under As$_2$ flux in order to raise the substrate temperature to 570°C. When the temperature reaches 570°C, an additional 50 nm of GaAs is deposited while the temperature is increased to 590°C. Both samples then have 50 nm of Al$_{0.4}$Ga$_{0.6}$As and 50 nm of GaAs deposited at 590°C.

2.2. Optical characterization

Room-temperature photoluminescence (RTPL) spectra from samples CG and GP are shown in Fig. 1. The peak emission wavelength from sample CG is at about 1260 nm, compared to about 1220 nm from sample GP. The full-width at half-maximum (FWHM) from sample CG is 60.0 meV versus 41.5 meV for sample GP. The peak emission intensity from sample GP is about 3 times that of sample CG. The integrated PL intensity from sample GP is almost double that of sample CG.

The shift in peak emission wavelength between the two samples can be attributed to In segregation into the GaAs overgrowth, and possibly In evaporation, during the growth pause while the sample

![Fig. 1. Room-temperature photoluminescence spectra show the differences between continuous growth and a growth pause to raise the substrate temperature during the overgrowth of InGaAs islands.](image-url)
temperature is raised to 570°C. Indium segregation leads to shallower confinement, and the ground state of the electron is raised in energy as the electron becomes more characteristic of the barrier, thus blue-shifting the emission wavelength.

Since the surface is nonplanar due to the InGaAs islands, the 8.0 nm of GaAs will not uniformly bury the islands. When the substrate temperature is raised above about 540°C, In near the surface may evaporate from the islands. Note that is especially important on the non-(1 0 0) oriented sidewalls of the islands, as it has been shown that In segregation is enhanced along these directions [15]. As the In evaporates, the islands become smaller, and the bandgap of the island increases due to the larger Ga mole fraction. The consequence of the smaller islands is that the quantization energy increases, and the emission wavelength decreases. As the bandgap increases, the emission wavelength decreases. Thus, In segregation and evaporation from the islands qualitatively explains the decrease in emission wavelength from sample GP.

3. Optimization of overgrowth thickness

3.1. Sample growth

Three samples are grown to determine the optimum thickness of the low-temperature (515°C) GaAs overgrowth layer immediately adjacent to the InGaAs islands. Samples OG3, OG6, and OG12 have the same epilayer structure as sample GP, except the thickness of GaAs grown at 515°C is 3.0, 6.0, and 12.0 nm, respectively. Next comes a growth pause under As₂ flux while the substrate temperature is raised to 570°C. Then, additional GaAs is deposited to give a total of 58 nm for each sample. Finally, 50 nm of Al₀·₄Ga₀·₆As and 50 nm of GaAs are deposited at 590°C.

3.2. Optical characterization

RTPL spectra from these three samples are shown in Fig. 2. OG6 has the highest peak intensity of the three samples. The integrated intensity from OG6 and OG12 is approximately equal, and both are about twice the integrated intensity of OG3.

The FWHM of OG6 is only 42 meV, compared to 53 meV from OG3 and 60 meV from OG12. The peak emission wavelength from OG12 is at 1234 nm, compared to 1213 nm from OG6 and 1129 nm from OG3.

As with the previous example, the wavelength shift can be attributed to In segregation and evaporation. It is likely that In evaporation has occurred in OG3, with its thin (3.0 nm) GaAs overgrowth, since the wavelength shifts by about 100 nm compared to OG12. The wavelength shift between OG12 and OG6 is only about 20 nm, so it is unclear if this is due only to segregation or if evaporation is also occurring.

A second set of samples is grown in order to further study the effect of the overgrowth. In this set of samples, the identical epilayer structure is used, and the low temperature overgrowth thicknesses used are 4.5, 6.0, 7.5, and 9.0 nm. The sample with the 9.0 nm overgrowth has a peak intensity three times greater than any of the other samples, and a FWHM that is only 38 meV, whereas all the other samples have line widths of at least 45 meV. Thus, we conclude that 9.0 nm is the optimum thickness of GaAs to grow at the InGaAs quantum dot temperature of 515°C.

4. Multiple quantum dot layers

4.1. Sample growth

The same alternating MBE technique described above is also used to grow multiple layers of QDs.
Samples M17-1 and M17-2 have one and two layers of 17.7 ML In_{0.3}Ga_{0.7}As QDs, respectively. The GaAs barrier in the two layer sample is 25.0 nm. The first 9.0 nm are grown at the InGaAs growth temperature of 515°C. Then there is a growth pause while the substrate temperature is increased to 570°C, and the remaining 16.0 nm of GaAs are deposited as the substrate temperature is increased from 570°C to 590°C. The temperature is then decreased to 515°C to grow the second layer of QDs. Samples M13-1, -3, and -5 have one, three, and five layers of 13.3 ML In_{0.3}Ga_{0.7}As QDs, respectively. The same growth procedure used for M17-2 is used for the samples with multiple layers of QDs. The 2D-3D RHEED transition occurs between 7.1 and 8.0 ML for the first two layers, but occurs between 6.2 and 7.1 ML for the subsequent three layers for the five layer sample.

4.2. Optical characterization

RTPL spectra at various pump powers are shown for M17-1 (Fig. 3a) and M17-2 (Fig. 3b). The lowest pump power is approximately 0.22 mW, and the highest pump power is approximately 169 mW. Note that the vertical scales differ by one order of magnitude, with the single QD layer sample showing greater intensity. This is not the case for a similar group of InGaAs QW samples, where the double QW sample has much brighter luminescence than does the single QW sample.

At 0.22 mW pump power, M17-1 has a peak emission wavelength of 1250 nm and a FWHM of about 51 meV. However, as the pump power is increased the line width increases rapidly as excited-state luminescence becomes more pronounced. At a pump power of about 27 mW, the ground-state luminescence is saturated and the excited-state luminescence intensity surpasses that of the ground state.

At 0.22 mW pump power, M17-2 has a peak emission wavelength of 1276 nm and a FWHM of about 28 meV. This line width is improved compared to M17-1. It has been predicted theoretically that subsequent layers will have improved uniformity [17]. However, there is some ambiguity in this case because the intensity has degraded sharply compared to the single layer of QDs, which suggests that the second layer of QDs in M17-2 has nonradiative defects. Also, the carrier concentration is lower in M17-2 than in M17-1 because M17-2 has two layers of QDs. Since state-filling effects are very strong in these samples, as indicated by the excited-state luminescence, a lower carrier concentration should lead to a narrower line width.

RTPL spectra at various pump powers are shown for M13-1 (Fig. 4a), M13-3 (Fig. 4b), and M13-5 (Fig. 4c). The peak emission wavelength at low pump power from M13-1, -3, and -5 is at 1154, 1188, and 1180 nm, respectively. The peak intensity and the integrated intensity from the three layer QD sample is comparable to that of the single layer QD sample. However, both of these quantities are decreased in the five-layer QD sample. The line width from M13-1 is slightly lower than that of M13-3 at low pump power, but increases more rapidly as the excitation power is increased and excited-state luminescence becomes more pronounced. The line shape remains approximately constant for the three-layer QD sample over the range of pump powers used in this experiment. The line width of the five-layer QD sample increases.
rapidly as the pump power increases, and several shoulders are observed around the main peak.

The shift in peak emission wavelength from the multilayer samples indicates some layer-to-layer nonuniformity. This correlates with the RHEED observations that the 2D–3D transition occurs when fewer monolayers have been deposited for subsequent layers, which indicates either insufficient smoothing by the GaAs barrier or strain-field interaction with the surface. This allows the islands to grow taller than the previous layers, and thus have lower quantization energy and longer wavelength emission. The relatively poor intensity of the five-layer sample suggests that nonradiative defects have been introduced into the uppermost layers of QDs, which should be the tallest layers and thus have the longest emission wavelength. Consequently, the peak emission wavelength from the five-layer sample is about the same as that of the three-layer sample.

4.3. Transmission electron microscopy

Transmission electron microscopy (TEM) images of M13-1, -3, and -5 are shown in Fig. 5a–Fig. 5c, respectively. The images are taken under two-beam, (2 0 0) dark field conditions to maximize chemical contrast. The multiple layers samples both show vertical alignment of the islands, as has been demonstrated by others [1, 9–11]. The starting surface for each dot layer appears to be flat, so it is likely that the strain field from the underlying islands interacts with the surface to cause this alignment [10].

The subsequent layers of islands get progressively larger, both in the vertical direction and in the lateral direction. This agrees with the RHEED observations that subsequent layers undergo the 2D–3D transition at a slightly lower deposited thickness. It also agrees with the PL spectra, which show a longer emission wavelength for multiple layer samples. Finally, the TEM image of the five-layer sample shows some unusual contrast directly under some of the island stacks. This contrast is not observed in the one-layer and three-layer samples. No threading dislocations are observable under

![TEM images](image-url)
various beam conditions, but we speculate that this contrast is caused by some defects that are also the cause of the reduced emission intensity from the five-layer sample.

5. Conclusions

We have demonstrated that the overgrowth of GaAs on highly nonplanar, strained InGaAs islands plays an important role in determining the optical properties of these islands. A growth pause after partial burial of the islands improves the PL intensity and line width from the array of islands. There is an optimum thickness of this overgrowth that prevents excess In evaporation from the islands, but still allows high-quality GaAs to be deposited. Multiple layers of QDs have been studied by RHEED, PL, and TEM. The RHEED transition gives a good indication of the layer-to-layer uniformity from the QDs. PL peak emission wavelengths and TEM images correlate with the RHEED observations. Although the formation of islands provides some strain relief, there is still a limit to the number of layers of QDs that can be stacked, as indicated by the decrease in PL intensity and the defects seen from some multilayer samples. The number of layers that can be deposited depends on the composition and thickness of the islands. These results have important consequences for the growth of quantum dot lasers and multilayer quantum dot lasers.

Acknowledgements

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Self-assembling InP quantum dots for red lasers

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Abstract

Self-assembling InP quantum dots are prepared by solid source molecular beam epitaxy. The dots have a diameter of 15 to 50 nm and a height of 5 to 15 nm depending on the nominally deposited InP layer thickness between 1.5 and 7 ML. Transmission electron microscopy measurements are presented to provide information about the structural properties. The InP quantum dots are embedded in Ga\textsubscript{0.52}In\textsubscript{0.48}P lattice matched to the GaAs (1 0 0) substrate and show a strong and narrow photoluminescence at room temperature in the visible range. Laser structures are prepared with one layer of InP quantum dots within a 160 nm thick Ga\textsubscript{0.52}In\textsubscript{0.48}P waveguide region. 0.7 \textmu m thick AlInP layers are used as cladding layers below and above the waveguide. We observe lasing of the InP quantum dots at room temperature in optically pumped cleaved samples with a length of 500 \textmu m.

Quantum dots based on III/V semiconductor heterostructures are interesting for fundamental research and future device applications. The discrete energy spectrum in zero-dimensional systems promises advantages especially for laser diodes [1]. The concept of self-assembling quantum dots is based on the Stranski–Krastanov growth mode in strained layer heteroepitaxy of materials with a large lattice mismatch [2]. This method offers the possibility to prepare extremely small quantum dots in a simple process without lithography. The materials which have been investigated mainly are InAs and InGaAs on GaAs or AlGaAs [3–9], and InP on Ga\textsubscript{0.52}In\textsubscript{0.48}P [9–12].

We are interested in the preparation of InP/Ga\textsubscript{0.52}In\textsubscript{0.48}P quantum dots. Ga\textsubscript{0.52}In\textsubscript{0.48}P is lattice matched to the GaAs substrate and has a direct band gap at 1.95 eV at 4 K. The lattice mismatch of InP to Ga\textsubscript{0.52}In\textsubscript{0.48}P is 3.7%. We have already demonstrated that the energy of the photoluminescence (PL) from the InP islands is adjustable by the amount of InP. An energy shift is observed from 1.85 to 1.53 eV as the nominally deposited thickness increases from 2 monolayers (ML) to 10 ML [11]. Intense PL is obtained even at room temperature as shown in Fig. 2, which makes

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it a promising material system for quantum dot laser diodes. The open question is how much does the size distribution of the dots deteriorate the laser properties, and what is the influence of the wetting layer. In this contribution we report on the preparation of optically pumped self-assembling InP/ Ga$_{0.52}$In$_{0.48}$P quantum dot structures which emit in the visible range at room temperature.

The samples were grown on GaAs (1 0 0) substrates by solid source molecular beam epitaxy (MBE). The phosphorus compounds were prepared by using a GaP decomposition source which provides a very clean and stable P$_2$ beam [13]. For all samples a 200 nm thick GaAs buffer layer was deposited on semi-insulating GaAs substrates at a temperature of $T_s = 580^\circ$C. The Ga$_{0.52}$In$_{0.48}$P, Al$_{0.51}$In$_{0.49}$P and InP layers were deposited at a substrate temperature of $T_s = 470^\circ$C with a beam equivalent P$_2$ pressure of about $5 \times 10^{-6}$ Torr. The growth rate for InP and Ga$_{0.52}$In$_{0.48}$P was 0.5 and 1 ML, respectively.

Fig. 1 shows a plan view transmission electron microscopy (TEM) image from an InP/In$_{0.52}$Ga$_{0.48}$P quantum dot structure. One layer of dots with nominally 3 ML InP is grown on top of a 50 nm Ga$_{0.52}$In$_{0.48}$P layer. The dots are overgrown by another 50 nm thick Ga$_{0.52}$In$_{0.48}$P cap layer. The larger dark spots clearly show the InP island formation. Misfit dislocations have not been detected in this and in other cross-sectional TEM micrographs for the 2 and 3 ML InP dot samples [14]. The size of the islands is 20 to 30 nm in diameter and the density is about $5 \times 10^{10}$ cm$^{-2}$. This is comparable to the data obtained from atomic force microscopy from 3 ML islands directly on the surface of a Ga$_{0.52}$In$_{0.48}$P layer.

Fig. 2 shows PL spectra measured at (a) 10 K and (b) room temperature (RT) for samples with InP islands embedded in Ga$_{0.52}$In$_{0.48}$P. The nominal InP thickness is 2.5, 3, 5.5 and 7.8 ML. The typical PL line width is 30 meV at 10 K and 60 meV at RT.

![Fig. 1. TEM plan view micrograph from an InP/Ga$_{0.52}$In$_{0.48}$P quantum dot structure. The dot layer with nominally three ML InP is embedded in 50 nm Ga$_{0.52}$In$_{0.48}$P grown on GaAs (1 0 0) substrate. The size of the InP dots is about 20 to 30 nm in diameter and the density is about $5 \times 10^{10}$ cm$^{-2}$.](image-url)

![Fig. 2. PL spectra measured at (a) 10 K and (b) room temperature (RT) for samples with InP islands embedded in Ga$_{0.52}$In$_{0.48}$P. The nominal InP thickness is 2.5, 3, 5.5 and 7.8 ML. The typical PL line width is 30 meV at 10 K and 60 meV at RT.](image-url)
Ga$_{0.52}$In$_{0.48}$P barrier layer. At 10 K the InP dot peak is a factor of 35 more intense than the Ga$_{0.52}$In$_{0.48}$P peak. The strong blue shift of the InP dot PL against bulk InP is essentially due to the confinement effect and the compressive strain within the island.

A shift of the PL energy with increasing nominal layer thickness was also observed for MBE grown InAs dots on GaAs [15]. In contrast to this, MOCVD grown InP/Ga$_{0.52}$In$_{0.48}$P structures show an energetically constant PL at 1.64 eV for InP coverages above 2.5 ML [12]. Beside this they observe another PL line around 1.9 eV. The reason of the different results for MBE and MOCVD grown samples is probably the higher surface diffusion in the MOCVD process which leads to the formation of larger islands. Especially for the InP/Ga$_{0.52}$In$_{0.48}$P material system there seems to be a pronounced 2-fold size distribution. In comparison to MOCVD grown self-assembling InP dots on Ga$_{0.52}$In$_{0.48}$P it seems that the MBE samples prepared at relatively low temperatures provide a more homogeneous size distribution and smaller dots with higher confinement energy.

The room temperature PL spectra in Fig. 2b show an asymmetric shoulder towards higher energies. For ideal quantum dots with a large energy separation between ground state and excited states the line width should not depend on temperature. Possible explanations for the observed broadening of the line width are phonon scattering processes, electron–light hole transitions, and transitions to higher excited states within individual islands. The existence of an excited state has already been reported in photoreflectance measurements on these samples [14]. Another possible mechanism is thermal reemission of carriers from larger dots into coupled smaller dots. Phonon assisted coupling of dots with different size has been observed in resonant time resolved PL measurements [16].

To summarize, the PL properties of the InP quantum dot samples are the following: The PL energy is in the visible range and can be varied by adjusting the nominal InP layer thickness. The integral intensity is very high and dominates even at room temperature for dots formed from 3 ML InP or more. Consequently, the InP/Ga$_{0.52}$In$_{0.48}$P self-assembling quantum dot system is a promising candidate for laser diodes.

After carefully optimizing the growth parameters in order to get maximum PL intensity and minimum line width we prepared laser structures for optical gain measurements and optical pumping. The 3 ML InP dots are used for the laser structures because they provide a strong PL signal at room temperature and because they are very small and have a high density within one layer of dots as shown in Fig. 1. Fig. 3 shows a cross-sectional TEM micrograph from an InP/Ga$_{0.52}$In$_{0.48}$P quantum dot laser structure with 0.9 µm thick Al$_{0.51}$In$_{0.49}$P cladding layers. The 120 nm thick Ga$_{0.52}$In$_{0.48}$P waveguide contains one layer of nominally 3 ML InP quantum dots. During the InP deposition the self-assembling islands are formed on top of a 1 to 1.5 ML thick wetting layer. The inset in the upper left corner shows a magnification of the dot layer which is right in the middle of the waveguide. The laser structure is grown on a GaAs substrate and covered with a 50 nm thick GaAs cap layer. The quantum dot related PL from this sample is observed at 1.76 eV with a FWHM of 60 meV at room temperature. The line width at 4.2 K is 28 meV. Optical gain measurements have been reported in Ref. [17].

We have optically pumped a cleaved sample similar to the one shown in Fig. 3 with a cavity length of 500 µm. The waveguide in this sample was 160 nm and the thickness of the cladding layers was 0.7 µm. Fig. 4a shows the intensity of the optical output as a function of the pumping intensity for the peak energies of the quantum dots (QD) and the wetting layer (WL). The wetting layer was not observed in PL; however, it is clearly identified in the gain measurements [17]. Fig. 4b shows the corresponding emission spectrum for 43 kW/cm² pumping intensity at room temperature. Above a pumping intensity of about 25 kW/cm² the optical output of the quantum dots increases due to lasing activity while the emission from the wetting layer remains relatively weak. The threshold of this laser structure is comparable to values obtained in the same experimental set-up with GaInP/AlGaInP double quantum wells in a similar structure.
The behaviour of getting stimulated emission from both the wetting layer and the dots can be explained only if the carriers are not in thermal equilibrium. One explanation is spatial separation of the dots and the wetting layer. A thinner or vanishing wetting layer around the dots has been observed for InAs dots [15]. The consequence of that would be a separated transition rate from the waveguide into the dots and the wetting layer. Further details about the optical characterisation of the laser structure are given in Ref. [17].

In summary, we have demonstrated lasing activity from InP quantum dots at room temperature. This is the first quantum dot laser structure based on InP/Ga$_{0.52}$In$_{0.48}$P. The observation of an intense laser line from the InP quantum dots is surprising, considering that self-assembling dots have a relatively broad size distribution and that the active volume is extremely small compared to a quantum well laser. Future experiments will concentrate on the introduction of several layers of dots into the waveguide and on removing the wetting layer by in situ etching techniques as demonstrated in Ref. [18].

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Fig. 4. (a) Intensity of the optical output as a function of the pumping intensity for InP dots/Ga0.51In0.49P/Al0.51In0.49P laser structures with 5 0 0 µm length. (b) shows the corresponding emission spectrum for 43 kW/cm² pumping intensity (QD: quantum dots, WL: wetting layer. The measurements are performed at room temperature.

References

Increased size uniformity through vertical quantum dot columns

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Abstract

The effects of stacking strain-induced InAs quantum dot layers on the size uniformity and density of dots in the uppermost layer have been investigated using single-dot layers, and stacks of 10 and 20 quantum dot layers with a 56 Å space region of GaAs. In these multilayered structures dots from different layers are aligned in vertical columns. X-ray diffraction indicates diffraction from a superlattice of InAs and GaAs, where the InAs contribution to the diffraction is from the quantum well region which is an integral part of the dot formation process. Thus, these quantum well regions remain intact and planar throughout the stacking process. The quantum well thickness is estimated to be 1.9 InAs monolayers. Atomic-force microscopy indicates the dot density decreases with dot layers, while the average dot height and in-plane dot diameter increase with dot layers. The uniformity of dot height and in-plane diameter also increases with stacking of dot layers as has been predicted [J. Tersoff, C. Teichert and M.G. Lagally, Phys. Rev. Lett. 76 (1996) 1675].

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Keywords: Self-assembly; InAs; Quantum dots

The precise monolayer (ML) control of current epitaxial crystal growth techniques and the highly developed understanding of the III–V and group IV semiconductors has led to a well-controlled heterointerface and a quantum well (QW) structure that has had a dramatic impact in semiconductor research and mainstream semiconductor technology. The heterointerface control is responsible for the 1-D carrier confinement in the growth direction, and while a significant effort has been concentrated towards extending this control to 2-D and 3-D confinements with quantum wires and quantum dots, the results have been encouraging but not resounding. This is generally so because lithography techniques are needed for increased confinement and the resolution in lithography is not of the monolayer-like resolution available through epitaxial growth techniques. Recently, a purely epitaxial technique has been used to produce quantum dots in the InAs/GaAs [1] and Ge/Si [2] semiconductor systems. This technique

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utilizes the strain-induced islanding of the Stranski-Krastonow (SK) growth mode in which the growth surface islands, compensating an increase in energy due to extra surface for a decrease in accumulated strain energy. Unfortunately, the large surface migration processes common in SK growth lead to large inhomogeneous island size distributions so that the spectral features are broadened with respect to QWs, and the narrow, atomic-like transitions are lost in the ensemble broadening. This research is still active and with the fine tuning of growth processes which were needed in the development of successful QW technology, the inhomogeneous broadening can be reduced.

We have extended the strain-induced growth of quantum dots by layering the quantum dots, separating them by spacer layers of GaAs. Transmission-electron microscopy results show that the dots from different layers are arranged in vertical columns [3, 4]. Furthermore, photoluminescence results indicate that the dots within these columns are electronically coupled [4], and this coupling can be adjusted by varying the GaAs spacer layer [5]. A surprising consequence of this layering process was recently discovered by Tersoff et al. [6]. The layering process should decrease the size distribution of the top layer dot ensemble as the layering process continues. We believe that the combination of electronic coupling within the columns and a decreased size variation in the upper dot layers will lead to an improved strain-induced dot system with interesting new features.

In this paper we investigate the structural ordering of the quantum dot columns. Single, 10, and 20 dot layers are investigated each with 50 Å of GaAs spacer layers. Using X-ray diffraction (XRD), we find that the wetting layer continues to be present as the dot layers are stacked, and these wetting layers and GaAs spacer layers are uniform enough to combine to produce superlattice diffraction. Atomic-force microscopy (AFM) is used to investigate the size distribution as various numbers of quantum dot layers are deposited. We observe an increase in the average height and in-plane diameter of the dots, resulting in slightly more hemispherical dots after 20 dot layers are deposited. A decrease in the size variation of the dots with increasing number of quantum dot layers is also seen, as well as an ordering of dots in the (1 0 0) direction.

The crystal growth was conducted using a Gen II MBE machine with GaAs and InAs growth rates of 0.1 µm/h. The As flux consists of cracked As₄ (As₂) and the V/III flux ratio was 9. The substrate growth temperature was 460°C measured by band edge absorption. A 2500 Å GaAs buffer layer was used and deposited at 600°C. The InAs island layers consist of the equivalent of 3 layer-by-layer MLs of InAs. In the vertically stacked quantum dot layers the GaAs thickness between adjacent dot layers is approximately 56 Å, measured by TEM. X-ray diffraction was conducted using a Philips 4 crystal X-ray diffractometer. A Digital Nanoscope II atomic-force microscopy was used to image the surface structure and these measurements were conducted in ambient conditions.

In Fig. 1 XRD results are shown for a sample consisting of 10 InAs dot layers. Superlattice diffraction resulting from a supercell consisting of the GaAs spacer and the InAs are observed. Peaks for the n = 0, ±1, and ±2 reflections are present where the (0 0 4)GaAs substrate peak resides.

![intensity_plot](image)

**Fig. 1.** X-ray diffraction of 10 periods of 3 MLs of InAs and 56 Å of GaAs. Superlattice diffraction indicates an InAs thickness of only 1.9 MLs and is due to the InAs wetting layer region and not the InAs dots.
between the \( n = 0 \) and \( n = 1 \) reflections. When fit
using a dynamical diffraction model the diffraction in
Fig. 1 results in a strained InAs thickness of
1.9 MLs (± 0.2 MLs) and a GaAs thickness of
52 Å. Our interpretation of these results is that only
the wetting layers of the dots–wetting-layer system
are contributing to the superlattice diffraction.
Since Fig. 1 represents symmetric diffraction about
the (0 0 4)GaAs reflection, the dot regions, with
their varying InAs and GaAs vertical sections,
should not contribute to superlattice diffraction.
The only extensive regions with periodic structure
in the growth direction are the regions of the
sample without InAs dots where only the wetting
layers are present. The best fit to the XRD data
occurs when the majority of the InAs is distributed
over approximately 9 Å. This diffusion of InAs in
GaAs has been observed by other researchers in
thin InAs quantum wells in GaAs [7]. This measured
value of 1.9 MLs is close to the 1.8 MLs
wetting layer thickness observed by other re-
searchers [8] using reflection high energy electron
diffraction (RHEED). The RHEED measurements
determined the equivalent amount of material de-
posited leading to the dot transition, while this
XRD measurement determines the wetting layer
thickness after the dot transition. Because the
transition to island growth results in a overall
change in the surface topology, it is interesting to
note that at least in the regions observed in this
type of XRD, the wetting thickness is unchanged
by the transition to islanding.

It is of interest to know how the stacking of these
InAs dot layers changes the size distribution of the
dot ensemble. AFM images of the InAs dot surface
are shown in Fig. 2. In these images the growth was
terminated after the deposition of the InAs dot
layer. Although this is necessary to directly observe
the dots, the GaAs layer that normally covers these
InAs dots modifies the structure of the dots. In
addition, the AFM measurement is done ex situ in
an ambient environment and adsorbed materials,
as well as the effects of the AFM tip lead to some
additional measurement error. In Fig. 2a, the sur-
face topology after a single layer of InAs dots is
observed with the AFM. The dots are approximately
40 Å high (in the growth direction) and 200 Å
in the in-plane direction. In Fig. 2b, the surface
topology after 20 layers of InAs dot layers have
been deposited is shown. The dots height and in-
plane diameter have increased. The dot density is
decreased between the single layer and 20 layer
samples from 8.4 to \( 5.7 \times 10^{10} \) dots/cm\(^2\). In addi-
tion to the decreased island density two other fea-
tures are apparent in Fig. 2. First, in the 20 layers
sample the dots are aligned in the (100) directions.
This alignment is different from the alignment of
dots along the (110) directions observed in single
dot layer samples, which is due to steps on the
growth surface. Second, there is new surface rough-
ness present in the 20 layers sample.

The decrease in dot density with stacking of individ-
ual quantum dot layers, has been predicted if the initial dot density is higher than some equi-
librium value which minimizes the overlapping strain
fields of the adjacent dots [6]. If the dot density is
large then the strain fields of two adjacent dots
overlap and only one dot will form above the dots
and spacer layer. In the opposite case in which the
initial dot density is lower than some equilibrium
value, dots will form above the lower dots as well as
in the regions where no dots are found below the
spacer region. If an equilibrium state can result, the
strain field should be periodic over the growth
surface, assisting surface diffusion, and the nearest-
neighbor distance between nucleating dots will be
constant, resulting in uniform nutrient fields for
each dot, and thus a more uniform size distribution.
Since we have adjusted our growth parameters to
increase the dot density in single InAs dot layer
samples [9] for optoelectronics applications it is
not surprising, in the context of the above dis-
cussion, that the dot density should decrease with
layering. We believe that the density of dots in an
isolated dot layer is determined by the uniformity
of the wetting layer [9]. If the wetting layer was
ideally uniform then the transition from layer-by-
layer growth to island growth would occur uni-
formly over the surface, leading to a dense array of
dots with competing nutrient fields. We believe that
this is a local minimum which reduces surface dif-
sion and even though the dots density is large in the
single layer, late stage ripening, the growth of larger
dots at the expense of smaller ones, is inhibited. The
stacking of dot layers overcomes this problem. The
more uniformly periodic strain fields which lead to
the reduced InAs dot density should also lead to a more uniform dot size distribution. This is indeed observed and discussed below.

A statistical evaluation of the AFM images in Fig. 2 is shown in Figs. 3 and 4. In Fig. 3, a comparison of the variation in height in the single and 20 layers samples is shown. The average height increases from 43 Å in the single sample to 65 Å in the 20 layers sample. The FWHM of the dot height is reduced from 17 to 10 Å between the single and
20 layers samples even though the average dot height has increased in the 20 layers sample. In Fig. 4, a comparison of the variation in the in-plane dot diameter between the two samples is shown. The in-plane area is not circular but elliptical and the two differently shaded regions in Fig. 4 represent the orthogonal directions with maximum and minimum values. Similar to the average dot height, the dot in-plane diameter has increased after 20 dot layers. The average in-plane diameter of the single dot layer is 218 Å, while the average in-plane diameter after 20 dot layers is 273 Å. The distribution of the in-plane diameter has also narrowed in the 20 dot layer structure. The FWHM of the diameter distribution is reduced from 50 Å (single layer) to 41 Å (20 dot layers). There is a small but observable difference between the height increase and diameter increase in the 20 layer dot distribution. The average height increases more than the average diameter so that the ratio of the height to in-plane radius changes from 0.4 to 0.48 after 20 layers, indicating that these dots are more hemispherical. The increase in the dot height and diameter is in agreement with the reduced island density observed in Fig. 2. Since the dot density is reduced and the wetting layer thickness is unchanged (Fig. 1), the dot size must increase. The reduction in the size variation in height and diameter is due to the more uniform local dot environment resulting from stacking of the dot layers. Since the strain field in the dot layers becomes periodically more uniform with each dot layer, dots are nucleated with more uniform nearest-neighbor distances and hence more uniform nutrient fields. These more uniform surroundings lead to more uniform dots size.

In conclusion, we have shown that the stacking of quantum dot layers with a GaAs spacer layer can be used to reduce the large size distribution in
strain-induced InAs quantum dot layers. Using XRD, we show that through the layering process the wetting layer structure is maintained and remains flat. The 20 layer dot structure produces less dense dot arrays than the single dot structure, and the dots at the top of the 20 layers structure are larger and more uniform than the isolated single dot layer. This increased uniformity has been predicted and is due to the elimination of dot columns with overlapping strain field. Thus, these vertically aligned dot columns are interesting experimentally because of new effects such as electronic coupling within a column, as well as a mechanism to increase the size uniformity of the InAs dots.

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References

Self-assembled structures of closely stacked InAs islands grown on GaAs by molecular beam epitaxy

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Abstract

Closely stacked Stranski–Krastanow (S–K) growth islands were investigated. InAs islands of nominal 1.8 monolayer thickness were grown periodically with GaAs intermediate layers of less than 3 nm by molecular beam epitaxy. Reflection high-energy electron diffraction and atomic force microscopy revealed that S–K growth islands were formed even with stacked intervals of 2 nm, but the upper island size expanded slightly as the number of stacked layers increased. Transmission electron microscopy revealed that upper islands grew closely just on the lower islands aligning vertically on the first layer islands. Drastic decrease in photoluminescence full-width at half-maximum less than 30 meV was obtained from this structure.

1. Introduction

Low-dimensional carrier-confined nanostructures such as quantum wires and dots are of great interest in high-performance electronic and photonic device applications [1–3]. For precise fabrication of these structures, much attention has been paid to self-organizing and self-assembling phenomena in epitaxial growth. In particular, Stranski-Krastanow (S–K) growth islands formed at the initial stage in the highly mismatched heteroepitaxy is one of the most promising structures [4, 5]. Both theoretical and experimental studies have shown these islands to be coherent and dislocation-free [6–8]. Attempts to fabricate quantum dot lasers [9–12] and wavelength-domain-multiplication (WDM) optical memories [13] using S–K growth InAs islands on GaAs have been reported. However, the luminescence from these S–K growth islands were broad—the reported full-width at half-maximum (FWHM) of photoluminescence (PL) from InAs islands on GaAs were typically about 80 meV [14–16], showing large fluctuation in island size. Islands of such size fluctuation are applicable to WDM memories [3], but often lead to a decrease of optical gain in the lower-order subbands and an increase of threshold current in lasers [11]. According to our estimation, if we can achieve a PL FWHM of around 20 meV, lasing at the ground state can be realized even in as-cleaved devices [11]. Broadening of luminescence was mainly due to fluctuation in island size. Since the

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S–K growth islands previously produced were disk-like, 20–25 nm in diameter and 3–5 nm in height [17], the broadening of luminescence were possibly dominated by fluctuation in island height. Therefore, if island height can be accurately controlled, luminescence broadening should be greatly reduced.

Recently, in stacked structures, vertical alignment of S–K growth islands due to strain induced by lower-layer islands was reported [15, 16, 18]. In these structures, if the upper islands could be stacked closely just on the lower-layer islands with fairly thin intermediate layers, due to vertical extension of wave function over islands [12], the effective island height can be controlled by the number of stacked layer, while keeping island diameter and density almost the same as those of the first-layer islands.

In this paper, we describe closely stacked S–K growth InAs islands grown with thin GaAs intermediate layers on (001) GaAs substrates. The structure was evaluated by reflection high-energy electron diffraction (RHEED), atomic force microscopy (AFM), transmission electron microscopy (TEM). We demonstrate highly reduced PL FWHMs of smaller than 30 meV.

2. Growth of closely stacked island structure

We stacked InAs islands with GaAs intermediate layers of less than 3 nm thickness by conventional molecular beam epitaxy (MBE) using metallic In, Ga, Al and As as the source materials. The substrates used were (001) oriented GaAs. Before growth, the substrates were thermally cleaned at about 680°C for 1 min under an As pressure of 1.2 × 10^{-3} Torr. Stacked InAs islands were grown on a GaAs (100 nm)/AlGaAs (200 nm)/GaAs (400 nm) buffer layer and covered with a GaAs (50 nm)/AlGaAs (50 nm)/GaAs (100 nm) cap layer. The substrate temperature for growth of InAs islands was 510°C and was 65°C for growth of the buffer layer. They were monitored by a pyrometer calibrated by melting of In–Al alloys. The nominal InAs thickness for island formation was fixed at about 1.8 monolayer (ML). The nominal thickness of GaAs intermediate layers was set at 2 and 3 nm. Prior to and following InAs island growth, the sample was annealed for 2 and 1 min, respectively. The growth rates of InAs and GaAs were approximately 0.1 and 0.75 μm/h. For the 1.8 ML InAs growth, it took about 18 s. The As pressure used for InAs island growth was 6 × 10^{-6} Torr, and for buffer later growth was 1.2 × 10^{-5} Torr. These were measured by an ion gauge at the substrate position. We used two As cells both set at 6 × 10^{-6} Torr and changed the As pressure abruptly at the interface by switching one of these cells on and off.

During the growth of the buffer layer, surface was As-stabilized 2×4 reconstruction, and then transited c(4×4) reconstruction while the substrate temperature dropped to 510°C. Fig. 1a is the [1 1 0] azimuthal RHEED pattern obtained before InAs growth. The 1/2 ordered fractional reflections were clearly observed. Just after InAs growth started, the fractional order reflections disappeared immediately, and the transition from streaks to spots started as the growth of InAs reached around 1.6 ML, indicating a two-dimensional (2D) growth transited to a three-dimensional (3D) island growth then. Fig. 1b is obtained after 1.8 ML InAs growth. The RHEED pattern recovered gradually to streaky as the growth of the subsequent GaAs intermediate layer started, although the fractional ordered reflections previously observed were barely visible during growth of the 2 to 3 nm thick GaAs intermediate layer. Fig. 1c is the RHEED pattern obtained after a 3 nm thick GaAs intermediate layer grown over the 1st island layer. By supplying InAs to such surface, the streaky pattern transformed to a spot pattern, but the transition time, (or in terms of the critical InAs thickness for the 2D growth to 3D island growth transition) was shorter (smaller) than for the growth of 1st layer island. Fig. 1d shows RHEED intensity transition observed at the area indicated by a square in the inset patterns during the growth of InAs islands. The 3rd- and 5th-layer islanding started when the growth of InAs reached about 1.0 ML, which is about 63% of the 1st-layer islanding. Presumably, the strain induced by the lower-layer islands or segregation of In atoms in the lower-layer islands affected subsequent islanding. However, the transition of growth mode from a 2D growth to a 3D island growth, and the existence of wetting layers as
will be shown later in TEM evaluation both indicate that S-K growth islands were formed even on such thin GaAs intermediate layers. Therefore, the formation of coherent and dislocation-free islands can be expected also in these stacked structures.

3. Evaluation of closely stacked island structure

3.1. Structural features

We evaluated island size and density by ex situ AFM operated in the ambient atmosphere (air). Fig. 2a--Fig. 2d show AFM images of the islanding surfaces at (a) the 1st, (b) the 3rd, (c) the 5th and (d) the 10th layers stacked at 3 nm intervals. We note that these images refer to different epilayers grown under the same conditions. The scanned area is 250 × 250 nm². It is evident from these images that islands form even at 3 nm stacking intervals. The upper-layer islands, however, expanded slightly as the number of stacked layers increased. Fig. 3 shows the island size dependence on the number of stacked layers. The average diameter of the 10th-layer islands was about 33 nm which was 140% larger than the 1st-layer islands (about 24 nm in diameter), while the island density decreased from 1 × 10¹¹ cm⁻² for the 1st-layer islands to 8 × 10¹⁰ cm⁻² for the 10th-layer islands.

In order to clarify the structural features, we examined them by TEM. Fig. 4 is a (1 1 0) cross-sectional TEM image of a 5-stacked island structure grown at 2 nm intervals. Each island layer was accompanied by a wetting layer, indicating that the upper-layer islands formed by S-K growth as did the 1st-layer islands. The upper-layer islands grew close just on the lower-layer islands, aligning vertically with the 1st-layer islands. These islands were stacked almost in columnar of about 22 nm in diameter and 13 nm in height shown in the image as dark megaphone-like strained regions. Islands in individual layer were seen to be spatially isolated in the vertical direction, but the distance between the bottom of the upper-layer islands and the top of the lower-layer islands was seen to be 3 to 4 ML or less, which is so thin that the wave functions can be distributed along the vertical column. This
Fig. 2. AFM images of InAs islanding surfaces of (a) the 1st layer, (b) the 3rd layer, (c) the 5th layer, and (d) the 10th layer stacked at 3 nm thick GaAs intervals.

Fig. 3. Dependence of InAs island diameter on number of stacked layers measured from AFM images.

implies that the islands were electronically coupled, behaving as a single quantum dot. We note here that similar structure was first reported in an InAs/GaAs short period superlattice grown using atomic layer epitaxy (ALE) technique [19].

Fig. 5 is a plan view of TEM image obtained from a 5-stack island structure grown at 2 nm intervals. The density of the islands was $8 \times 10^{10} \, \text{cm}^{-2}$. This value agreed well with the AFM results. Fluctuation of the lateral size seemed to be smaller than for a single island layer. Although we attempted precise control in the vertical dimension of the islands, somehow uniform lateral dimension was achieved in these closely stacked structures as reported previously [12, 20].

3.2. PL properties

We evaluated the PL properties of these structures at 77 K. Fig. 6 shows the PL spectra of a single island layer and closely stacked 3- and 5-layer island structures grown at 3 nm intervals. The broad PL spectrum of a single island layer transformed to the sharper spectra with the number of stacked layers increased. The peak energy shifted to the lower energy by about 90 meV. This energy shift can be ascribed to reduction of quantum size effects due to the increase in effective island heights.
The FWHM of the 5-layer stacked structure was 27 meV. This is less than one-third of the single island layer structure. The peak intensity was about three times higher compared to the single island layer structure. We also obtained an FWHM of 24 meV from a 5-stack island structure grown at 2 nm intervals [21]. These narrow FWHMs are attributable mainly to the suppression of relative island size fluctuation in the vertical direction accompanied possibly by the high uniformity of lateral dimension as shown in Fig. 5. We also confirmed that PL FWHM was independent on temperature, and that diamagnetic energy shift of the PL was considerably small (less than 3 meV even at 12 T). Fig. 7 shows diamagnetic shift of PL peak at 4.2 K. Magnetic field was applied
perpendicular and parallel to the sample surfaces. Diamagnetic shifts in an In$_{0.15}$Ga$_{0.85}$As (8 nm thick)/GaAs quantum well emitting at 0.94 μm are superimposed. We found that the diamagnetic energy shifts in closely stacked islands was considerably smaller than that in the quantum well. These results indicate that three-dimensional quantum confined structures were formed. Although we have demonstrated laser oscillations at room temperature from the quantum dots grown by usual S–K growth InAs islands, lasing was not at ground state in as-cleaved devices, and threshold current was much higher than expected [11]. Those optical features obtained here are expected to improve these laser characteristics.

4. Summary

Closely stacked InAs islands were grown with fairly thin intermediate layers on (0 0 1) GaAs substrates. RHEED and TEM results revealed that S–K growth islands formed even at stacked intervals of 2–3 nm. The upper-layer islands formed closely just on the lower-layer islands aligning vertically on the first-layer islands. Although island of adjacent layers appeared to be spatially isolated, the separation was so small (3–4 ML or less) that the islands can be electronically coupled, behaving effectively as a single quantum dot. We confirmed that for these stacked structures, three-dimensional quantum confined structures were formed, and PL FWHM decreased drastically to 27 meV and the peak intensity increased to about three times of a single island layer structures. These improved properties are expected to be useful for laser applications.

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References

Room temperature luminescence from self-organized
In\(_x\)Ga\(_{1-x}\)As/GaAs (0.35 < x < 0.45) quantum dots
with high size uniformity

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Abstract

Self-organized growth of In\(_x\)Ga\(_{1-x}\)As/GaAs quantum dots (QD) for 0.5 < x are studied by time-integrated, time-resolved and temperature-dependent photoluminescence (PL). Low-temperature photoluminescence demonstrates that the critical composition 'x' needed for the formation of 3D islands is between 0.3 and 0.35. This is also supported by reflection high-energy electron diffraction (RHEED) measurements and agrees well with our theoretical predictions that a misfit of > 2% is needed, from thermodynamic considerations, for the onset of 3D growth. The presence of fine excitonic features superimposed on the emission peaks confirms that the second peak observed in the PL spectra of quantum dots is from an excited state transition. We believe that the saturation of the ground-state emission observed under moderately high excitation photon densities is due to the long decay time constant (~ 2.2 ns) as measured by time-resolved photoluminescence. Temperature variation of luminescence intensity shows a sharper decay for the ground-state emission than for the excited-state emission. Room-temperature luminescence is thus dominated by the excited state emission. Broad area lasers made from separate confinement heterostructure (SCH) laser structures with In\(_{0.3}\)Ga\(_{0.7}\)As quantum dots as the active region showed lasing from excited state, with \(J_{th} = 650\) A/cm\(^2\), and the single-mode ridge lasers gave a modulation bandwidth of 5 GHz at 2.5\(J_{th}\).

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Keywords: Molecular beam epitaxy; Self-organized growth; Quantum dot; Luminescence; Laser

1. Introduction

Formation of three-dimensional (3D) islands during the molecular beam epitaxial (MBE) growth
of highly strained layers, either in the Vollmer–Weber growth mode (direct formation of 3D islands) or in the Stranski–Krastanow growth mode (where growth changes from layer-by-layer to 3D islands) have been studied extensively. Although the formation of islands was observed in highly strained (7%) InAs/GaAs superlattices when

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the thickness of InAs is more than 2 monolayers (ML) [1], it was predicted that a lattice mismatch of more than 2% is enough, from thermodynamic considerations, to form coherent 3D islands [2]. This corresponds to an In composition of ~30% in the InGaAs/GaAs system. Most of the work in this system has been on binary InAs [3–5] or ternary In_{0.5}Ga_{0.5}As [6–8] on GaAs. We have studied the formation of 3D islands for indium composition <50%. In the InGaAs/GaAs system the formation of 3D island occurs by the Stranski–Krastanow growth mode, wherein the initial layer-by-layer (2D) growth degenerates into 3D islands. The thickness of the initial 2D wetting layer before the onset of 3D growth depends on the strain in the growing layer. It has been reported that in the case of InAs on GaAs the wetting layer is 1.57 ML thick [9], whereas for In_{0.5}Ga_{0.5}As it is 4 ML [5]. A thicker wetting layer, obtained by reducing the In composition, is expected to provide efficient carrier injection into the 3D quantum dots eliminating the problems associated with large relaxation time constants in pure quantum boxes.

Here we report results of a photoluminescence (PL) study of In_{x}Ga_{1-x}As/GaAs quantum dots for x < 0.5. Change in the reflection high-energy electron diffraction (RHEED) spectra from 2D to 3D pattern is used in conjunction with the PL results to estimate the minimum composition x required for the formation of 3D islands. Dependence of PL spectra on excitation intensity and temperature was also studied. Time-resolved PL measurements were made to get an insight into the carrier dynamics in these quantum dots. Broad area and single-mode lasers made from an SCH laser structure with InGaAs QDs in the active region were fabricated and analyzed.

2. Experimental procedure

Heterostructures grown for PL measurements consist of a single layer of In_{x}Ga_{1-x}As (x = 0.3, 0.35, 0.4 and 0.45) with 0.1 μm GaAs on each sides. A superlattice buffer was grown in all the samples, to smoothen the growth front, before growing the actual structures. Growth rate of GaAs was 0.72 μm/h and that of InGaAs was 1 ML/s. Substrate temperatures used were 620°C and 520°C for GaAs and InGaAs, respectively, and the V/III flux ratio was maintained at ~5. The number of monolayers for the transition of the RHEED spectra from streaky (2D) to spotty (3D) pattern were found to be 6, 8 and 11 for x = 0.45, 0.4 and 0.35, respectively. No transition in RHEED spectrum was observed for x = 0.3 even after a growth of 20 ML, indicating that no quantum dots are formed. Thicknesses of the grown layers were 8, 10, and 15 ML for x = 0.45, 0.4, and 0.35, respectively. A sample with 15 ML quantum well with x = 0.3 will was also grown for comparing the PL emission from QW and QDs.

Photoluminescence spectra were measured at different temperatures from 16 K to room temperature. A 632.8 nm HeNe laser with a variable attenuator was used as an excitation source. A 0.75 m scanning spectrometer, liquid-nitrogen-cooled photomultiplier, and lock-in amplification were employed to scan the emission spectra with a resolution of better than 0.1 nm. For time-resolved measurements a Ti-sapphire laser was used as the excitation source and the spectrally dispersed time-resolved image was obtained with a 0.5 m monochromator, Hamamatsu streak camera, and a high-speed CCD camera. The temporal resolution of the system is 27 ps.

Broad-area and single-mode lasers were fabricated from an SCH laser structure with In_{0.4}Ga_{0.6}As self-organized quantum dots as the active layer. The laser emission spectra were scanned with 1 m spectrometer, thermoelectrically cooled Ge-detector and a Boxcar averager. Modulation bandwidth measurements were made with HP8350B sweep oscillator and HP8593A spectrum analyzer.

3. Results and discussion

Photoluminescence spectra from quantum dots are expected to be broader than those from quantum wells due to the statistical nature of size fluctuation of a large number of quantum dots sampled. The spectral shape is found to be approximately Gaussian for low excitation. At higher excitation levels, contributions from excited-state transitions
gives rise to a second peak due to the saturation behavior of the ground-state transition [5]. Fig. 1 shows the PL spectra obtained from two different samples at various excitation levels. At higher excitation levels, the excited state transition peak at around 1.24 eV is observed in the sample with In$_{0.35}$Ga$_{0.65}$As layer, indicating the formation of self-organized quantum dots, whereas there is only a slight broadening of the excitonic emission in the sample with a In$_{0.3}$Ga$_{0.7}$As layer as is expected for a 2D quantum well. We can thus infer that the critical In composition for the onset of 3D growth is between 0.3 and 0.35 for our growth conditions, which is also supported by our RHEED studies. This agrees very well with our earlier theoretical predictions [2] that a strain of >2% is necessary for the formation of 3D islands in the Stranski-Krastanow growth mode. The energy separation between the two peaks of the In$_{0.35}$Ga$_{0.65}$As sample is 62 meV. Moreover, there are fine excitonic features superimposed on the emission peaks from QDs, which represent groups of dots with identical dimensions. This is clearly resolved by a derivative spectrum in Fig. 2. The overall linewidth of the ground-state emission of the QDs is 33 meV, which is lower than the 40–50 meV normally reported. This indicates the high degree of uniformity in the QD size. The fine structure is observed in the PL emission from both the ground-state and the excited-state transitions. This provides the argument to preclude the possibility of contribution from wetting layer emission. Furthermore, the transition energy for the wetting layer is estimated to be around 1.358 eV for this structure which is very different from the observed 1.24 eV for the second peak. We have observed that decreasing the growth temperature (<500°C)

![Diagram](image1)

**Fig. 1.** PL spectra (a) from GaAs/(15 ML)In$_{0.3}$Ga$_{0.7}$As/GaAs and (b) from GaAs/(15 ML)In$_{0.35}$Ga$_{0.65}$As/GaAs at various excitation levels. (c) potential well and different transitions in self-organized quantum dots.

![Diagram](image2)

**Fig. 2.** Sharp peaks superimposed on the excitonic emission from quantum dots observed in PL spectrum (a) and resolved by a derivative spectrum (b). The derivative spectrum for the quantum-well emission peak of Fig. 1a is also shown for comparison.
decreases the size of the quantum dots (<20 nm base width), at the same time decreasing the PL intensity. Growth at a higher temperature improves the PL intensity and increases the size of the quantum dots, ultimately resulting in a large size fluctuation for >550°C. We have used a substrate temperature of 520°C for the best PL intensity, and we expect the base width of the quantum dots to be 20–30 nm. Theoretical calculations [10] assuming a biaxial strain tensor predict a second energy level in the conduction band for these dimensions. The excited-state emission can be either from e1–hh2 transition, which is not forbidden in the pyramidal shape of these type of quantum dots [5], or from e2–hh2 transition, which is possible only in relatively large quantum dots. These transitions are shown in Fig. 1c. The measured energy difference of 62 meV, between ground-state and excited-state transitions, is found to agree with the energy difference between e1–hh1 and e2–hh2 in QDs with a pyramidal base width of 21 nm and a height of 10 nm. The calculations also indicate that the e1–hh2 transition is very weak, compared to the e2–hh2 transition.

The temperature dependence of the PL intensity from In0.35Ga0.65As/GaAs QDs is shown in Fig. 3 for both low and high excitation conditions. The wavelength-integrated intensity remains constant up to about 75 K and then decays due to the ionization of the localized exciton [11]. The intensity of the ground-state emission drops off dramatically, whereas the excited-state transition first decays at a fast rate, followed by a slow decay till room temperature. The fast decay indicates non-radiative transitions associated with a defect level while the slow decay of the excited state reflects the normal thermal quenching. The room-temperature PL spectrum (Fig. 4) is dominated by the excited-state transition since the ground-state transition saturates at much lower excitation levels than that at low temperatures. This implies that if we have such QDs in the active region of a laser, the gain provided by the ground-state transition, before it saturates, may not be sufficient for the onset of lasing. We have made broad-area lasers from an SCH laser structure with In0.4Ga0.6As QDs in the active region, which had a threshold current density of 650 A/cm² for 1 mm cavity length. The emission spectra from these lasers showed [12] that the ground-state transition saturates with increasing injection current and lasing indeed takes place from the excited state transition.

In order to study the carrier dynamics in these quantum dots, we have made low-temperature time-resolved PL measurements. Fig. 5 shows the time-resolved PL intensity at different wavelengths obtained from spectrally dispersed image at 8 K. It is found that the decay time constants for the ground state and excited state are ~2.2 ns and ~200 ps, respectively. The rise time of the signal is limited by the resolution of the system to 27 ps. We believe that the slow decay of the ground state may be responsible for the saturation behavior observed in the time-integrated PL measurements. It is interesting to note that the excited state has an order of magnitude shorter decay time constant than the ground state. This predicts a better high-speed performance of the lasers emitting from excited states.
compared to that from ground states. We made single-mode ridge lasers in the ground–signal–ground configuration for bandwidth measurement. The threshold current was relatively high, around 30 mA for a cavity length of 415 μm, which we hope to bring down by optimizing the growth conditions and by having multiple layers of coupled quantum dots. The measured bandwidth was around 5 GHz for a current of 2.5I_{th}. Increasing the current further reduced the optical power and the bandwidth due to heating. We believe that higher bandwidths can be obtained by having lower threshold current levels.

4. Conclusion

We have found the critical composition for the formation of self-organized quantum dots in In_{x}Ga_{1-x}As/GaAs system to be 0.3 < x < 0.35. This agrees well with our earlier theoretical prediction that a lattice mismatch of >2% is necessary, from thermodynamic consideration, for the growth of 3D islands. Photoluminescence measurements show the saturation of the ground-state transition in quantum dots giving emission due to excited-state transition at high excitation levels. The temperature variation of the integrated intensity shows a slower variation for the excited state transition than for the ground-state transition, indicating that the ground-state transition saturates at much lower excitation levels at higher temperatures. Time-resolved measurements show that this may be due to the large decay time constant (≈ 2.2 ns) for the ground state. Broad area lasers fabricated from an SCH laser structure with In_{0.4}Ga_{0.6}As QDs as the active layer showed that the lasing occurs from excited states with J_{th} = 650 A/cm², which has a photoluminescence decay time constant of 200 ps. A modulation bandwidth of 5 GHz measured with a single-mode ridge laser at a current level of 2.5I_{th} indicates that phonon bottleneck problem is not significant for the laser operating from excited-state transition.

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References

Growth and characterization of self-organized InSb quantum dots and quantum dashes

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Abstract

We have grown self-organized InSb quantum dots on semi-insulating InP (0 0 1) substrates by molecular beam epitaxy. The size dependency of the uncapped InSb quantum dots on the nominal thickness of the deposited InSb was studied by atomic force microscopy. The dot size has a pronounced minimum at about 2.2 monolayers of InSb. After a nominal thickness of 3.2 monolayers we observe a drastic change of the dot shape, from quantum dots to quantum dashes. From thereon the dots grow in a quasi-cylindrical shape aligned in the (1 1 0) direction. The photoluminescence emission of a series of quantum dots was studied, the emission energy being independent of the dot size. When the dots partially relax, the photoluminescence is blue-shifted, which can be explained by a type-II band alignment.

There have been extensive studies on the formation of self-organized quantum-dots (QD) by several growth techniques, such as molecular beam epitaxy (MBE) [1–3], metal-organic chemical vapor deposition [4] and metal-organic vapor-phase deposition [5]. One of the remaining problems of the self-organization of the QDs is the inhomogeneity of the island sizes that broaden the observed PL peaks, and their random distribution over the surface. Also the dot geometry has been studied on (0 0 1) and other surfaces [6].

In this paper we present the growth of InSb QDs on semi-insulating InP (0 0 1) substrates grown by MBE in a pulsed mode, where the group V element is pulsed to enhance the group III surface migration. The self-organized dots were studied by atomic force microscopy (AFM) and photoluminescence measurements. Samples were grown in a conventional solid-source MBE system. After desorption of the InP (0 0 1) oxide at 490°C, we grew a 500 ML thick InP buffer layer, giving a streaky (2 × 4) reconstruction in the reflection high-energy electron diffraction (RHEED) pattern. Afterwards, the InSb layer was deposited. The surface stoichiometry during growth was controlled by measuring the surface reflection difference signal of the (1 1 0) and (1 – 1 0) directions using a HeNe laser (641.3 nm) at normal incidence to detect the absorption of In-dimers at the growth front reported elsewhere [6]. The InP buffer as well as the InSb islands were grown at a rate of 0.5 ML/s at a growth temperature of 400°C. The growth rate was calibrated by means of RHEED oscillations. After the deposition of 1.2 ML of InSb we observe
the typical growth mode transition from two-di-
dimensional layer-by-layer growth to three-di-
dimensional island growth, indicated by the onset of a spotty RHEED pattern. This “critical thickness”
is very small and it is not quite sure if the islands
grow in the Stransky–Krstanow or in the Vol-
mer–Weber mode, i.e. whether they have a wet-
ting-layer or not. During and after the annealing,
the samples were kept under Sb flux until the tem-
perature has fallen below 300°C to ensure that no
Sb was lost from the surface.

We took atomic force microscope (AFM) images
from various spots on each sample, in order to
detect a possible inhomogeneity of the dot sizes
over the sample. Typical variations of the mean size
in different regions of the same sample were less
than 15%. The island size and shape of the uncapped
QDs were studied by means of AFM. Fig. 1
shows InSb islands of a sample with nominal thick-
ness of 2 ML. The InSb QDs seem to be randomly
distributed and have a quite homogeneous size.
Their density amounts to $1 \times 10^{19}$ QDs per cm$^2$.
The size distribution for the dot diameter has a
mean value of $24 \pm 4$ nm. The height distribution
has a mean value of $6 \pm 3$ nm. (The error values
given here and in the following refer to standard
deviations of the distributions.) We grew further
samples from nominal InSb thicknesses of
1.4–2.8 ML. The statistics of these samples can be

seen in Fig. 2 (left part). At small nominal InSb
thicknesses the dot diameter is about $80 \pm 3$ nm
and decreases strongly with increasing number of
InSb monolayers. The dot height is similarly affec-
ted by the nominal InSb layer thickness. The dot
density shows opposite behavior. As the diameter
reaches its minimum at 2.2 ML, the density has
a maximum of $4 \times 10^{10}$ cm$^{-2}$. After further InSb
deposition up to 2.8 ML of InSb, the dot volume
rises again, while the density declines. The surpris-
ing feature is that the curve in Fig. 2 shows a min-
imum. The behavior of the dot sizes and density
versus the number of InSb monolayers can be ex-
plained qualitatively by material exchange pro-
cesses. At the onset of the 3D growth, when the
QD-density is still low, the islands grow more or
less independently from each other. Because the
mean inter-dot distance diminishes very quickly
(about a factor 3 within 0.4 ML) after the first 3D
formation, the QDs interact with each other, ex-
changing material until they reach their optimal

![AFM image](image)

**Fig. 1.** AFM image of a sample with 2 ML of InSb on InP
substrate. The QDs seem to be randomly distributed and have
a quite homogeneous size. The density amounts to $1 \times 10^{19}$
cm$^{-2}$. The mean diameter and height can be seen in Fig. 2.

![Graph](image)

**Fig. 2.** Statistics on diameters, heights and dot density for vari-
ous InSb thicknesses. The statistics are made on uncapped
samples by AFM. A minimum in size can be observed at around
2.2 ML, where the density is highest. On the right-hand side the
quantum dash regime can be seen. Here the islands grow in
a quasi-cylindrical shape (see also Fig. 3). The curve for the
diameter deviates into a curve for length and one for width,
respectively. The lines are only a guide for the eye.
size, which is smaller because in this way the surface area is enhanced, which is energetically favorable because this partially relaxes the strain without the formation of dislocations. Similar behavior has been reported for InAs QDs on GaAs [7, 8]. One has to remember that the samples are grown in an enhanced migration mode. This optimal size seems to be reached at 2.2 ML. Once achieving the optimal size, further InSb deposition results in new island growth, because from then on, the QDs not only interact by material exchange but also through the elastic strain field around each dot (remember the InP substrate next to a QD is expanded) results in a repulsive force between them [9]. Therefore, once having reached the minimum dot distance at 2.2 ML, further deposition can only result in growth of bigger dots, but not more. Furthermore, the density declines with bigger dots because the repulsive force rises with the dot size, which enhances the inter-dot spacing.

If we deposit more than 3.2 ML of InSb we observe a drastic change of the dot geometry (Fig. 3). The QDs no longer have a round shape, but an elongated one. After that point, the QDs maintain their quasi-cylindric shape in all samples with more than 3.2 ML of InSb forming quantum dashes (Q-dash). All Q-dashes are aligned along the (110) direction and their length to width ratio is approximately 2.5. The main Q-dash size is 100 nm long and 40 nm wide, but there are also some bigger ones with length 125 nm and width 66 nm (Fig. 3a). In regions where there is an enhanced surface roughness of the InP substrate, the dots grow all along it. This roughness cannot be explained by the growth conditions because they were the same for all samples (Fig. 1 does not show this roughness), but might be an evidence for an anisotropic strain field due to the Q-dashes, due to the different dimension in length and width. For that reason, we do not believe that the origin of this preferred growth in the (110) direction lies in the surface morphology, but vice versa. Our assumption is supported by the fact that the Q-dashes can grow closer to each other in the lateral than in the longitudinal direction, probably because the repulsive train field in the substrate, originated by the QDs, depends in each direction on the square of the corresponding QD-dimension in that direction. The relative frequencies of the lateral inter-dash spacing \( w \) and the longitudinal inter-dash spacing \( l \) of the Q-dashes on one sample can be seen in Fig. 4. There are minimum values for \( w \) and \( l \) of

![Fig. 3. AFM images of a sample with 3.5 ML of InSb. Image (a) was taken in the height mode, while image (b) was taken in the error signal mode to emphasize the dash edges. The circular QDs have transformed to rectangular quantum dashes. The longer edge lies along the (1 1 0) direction and is about 2.5 times longer than the other one (b). On the left-hand side (a) a higher number of quantum dashes can be seen with dimensions of 100 × 40 nm, but there are also a few bigger ones, measuring 125 × 66 nm.](image-url)
7 and 23 nm, respectively. The quotient, \( l/w = 3.3 \), that should depend on the square of the longitudinal and the lateral size of the islands, is somewhat smaller than the theoretical value of 6.3. Assuming an exponential distribution, we obtain a quotient of the mean ranges of the repulsive forces of 2, which is closer to \( l/w \), but the difference to the theoretical value could mean a slight amount of relaxation through dislocations at the interface in the (1 1 0) direction. Only a very small percentage of the Q-dashes are really orientated collinear with others, most of them are displaced with respect to their next neighbors due to the higher strain in that direction, which supports the model of an anisotropic strain field. Thus, the minimum surface energy in the Q-dash regime is obtained with a non-collinear geometry as shown in Fig. 3a. According to Tersoff [10], when the quantum dots pass a critical size, they grow in order to minimize their energy in the dash geometry, as observed in our samples. But his model does not explain why we cannot observe "infinite" large islands instead of many short ones and why the (1 1 0) direction is preferred. Furthermore, we can observe a distribution of the island widths, while the model predicts a constant width. On the other hand, the width distribution is much narrower than the length distribution, suggesting a preferred width value. The constant width in that model is also calculated at constant height, which is probably not the case as seen in Fig. 2, although the error bars are relatively large. Another explanation for the non-constant width could be that wider Q-dashes are formed by two individual ones growing together, which was actually observed on a few occasions on AFM images. In order to understand this unexplained behavior, one has to include the different energies of the different crystal surfaces of the Q-dashes and the interaction of different islands through their strain field in the substrate material.

We studied the photoluminescence of these samples at 12 K, exciting with a Ar⁺-laser. The spectra of three samples with 2, 2.8, and 10 ML of InSb dots, respectively, are shown in Fig. 5. The peak at 1.4 eV is attributed to InP, the other two peaks that can be seen at about 1 and 1.2 eV are attributed to two QDs. As more InSb is grown the peak at 1 eV loses intensity, while the one at 1.2 eV gains. Furthermore, the observed blue-shift with growing dot sizes is quite contrary to the expected behavior. The confinement effect of bigger dots should be smaller than that of the smaller ones and therefore a red-shift is expected. But this behavior can be explained supposing a type-II band alignment between the compressed InSb QDs and the InP matrix. Because of the strain applied to the QDs, the InSb
Fig. 5. PL spectra of samples with 2, 2.8 and 10 ML of InSb, respectively. Two peaks at 1 and 1.2 eV can be observed. The one at 1 eV is attributed to strained QDs, while the other one at 1.2 eV is attributed to partially relaxed ones. The inset shows the proposed band alignment for these two cases.

Conduction band rises above the InP. On the other side also the InP next to a QDs is expanded, which reduces its band gap in a way that a deep well at the InP/InSb interface is formed in the InP, as seen in the inset of Fig. 5. A type-II alignment has been observed, for example, in GaSb/GaAs QDs [11]. So, when the QD is strained, its emission should be indirect in real space. When the dot reaches a size sufficient to relax, all the strain disappears and a type-I band alignment is obtained again. Therefore, the peak at 1 eV should come from compressed QDs and the other one at 1.2 from partially relaxed ones. In this manner, one can explain why the emission energy of the strained QDs does not shift until about 2.5 ML InSb, although the dot-sizes differ significantly.

The initial stages of growth of InSb QDs on InP substrate have been studied by AFM. Typical QD formation can be observed until a InSb thickness of 3.2 ML. From thereon, the quantum-dots drastically change their shape forming Q-dashes along the (110) direction. This behavior can qualitatively be understood, taking into account the elastic energy and surface energy of the QDs and the kinetics of the growth predominantly at the island edges and the repelling force between the islands due to their anisotropic strain field in the substrate. PL measurements of samples with 2–10 ML of InSb reveal that in the strained state, the band alignment between the QDs and the InP matrix material is of type II.

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References

MBE growth of novel GaAs/n-AlGaAs field-effect transistor structures with embedded InAs quantum traps and their transport characteristics

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Abstract

Novel GaAs/n-AlGaAs FETs structures have been grown by molecular beam epitaxy (MBE) by placing InAs quantum dots near the channel. We show that the threshold voltage can be shifted by the trapping or detrapping of electrons in these dots. The number of electrons trapped by each InAs dot is estimated to be one or two, representing the finiteness of quasi-stable trap states. The concentration of trapped electrons at 4.2 K is identical to that at 77 K, suggesting that trapped electrons stay in the ground level up to relatively high temperatures.

1. Introduction

Owing to the development of semiconductor technology, it has recently become possible to fabricate laterally-defined nano-structures, such as quantum wires and dots [1–6]. Especially properties of zero-dimensional electrons confined in quantum dot structures have attracted a wide attention because of their importance both in physics and device applications, such as nonlinear transport devices [2, 3], improved laser performance [4] and ultra-high-density of semiconductor memory devices.

One of the promising methods to form 10 nanometer (nm)-scale dots is to use the Stranski–Krastanov mode of epitaxial growth. In this approach, a material such as InAs or InGaAs is deposited on a lattice mismatched GaAs substrate beyond its critical thickness to form very small dot structures [5, 6]. In addition to these systems, InP dots on InGaP [7] and InSb/GaSb/AlSb dots on GaAs [8] so on have been formed by molecular beam epitaxy (MBE)[5, 6] and metalorganic chemical vapor deposition (MOCVD)[11]. Structures of these self-assembled dots have been studied to find out the dot formation process [5, 6] as well as the strain distribution [9, 10]. Optical properties have been also studied to clarify quantum states in these dots [12, 13] and also explore the possibility of quantum dot lasers which are expected to have
superior properties over quantum well lasers [4]. Other new insights have been also obtained on details of quantum states by using such methods as capacitance spectroscopy [14], far-infrared spectroscopy [15], and the resonant tunneling spectroscopy of electrons via InAs quantum dots [16].

Although much work has been done on the carrier confinement in the core part of quantum dots, a small number of papers have dealt with trapping and detrapping process of electrons and the subsequent influence of the dot-induced potential on the electron transport. In this paper, we report on the MBE growth on GaAs/n-AlGaAs inverted heterojunctions (inverted HEMT), in which InAs dots are embedded near the electronic channels and investigate how InAs quantum dots serve as electron trap centers for two-dimensional electron gas.

2. Growth and sample fabrication

First, to prepare selectively-doped GaAs/n-AlGaAs inverted heterostructures of Fig. 1, we grew by MBE a 200 nm GaAs buffer layer on a seminsulating GaAs (1 0 0) substrate, and then 11 periods of a 20 nm \( \text{Al}_{0.25}\text{Ga}_{0.75}\text{As} \) and 2 nm GaAs layer at the substrate temperature \( T_s \) of 600°C. At this point, \( 1 \times 10^{12} \text{ cm}^{-2} \) Si was \( \delta \)-doped so as to minimize deleterious effects of DX centers [17]. The growth rates for GaAs and \( \text{Al}_{0.25}\text{Ga}_{0.75}\text{As} \) are 0.954 and 1.28 ML/s, respectively. We grew a rather thick (70 nm) undoped \( \text{Al}_{0.25}\text{Ga}_{0.75}\text{As} \) spacer layer to avoid the degradation of electron mobility [18] that may be induced by the segregation of Si atoms toward the sample surface. Then a GaAs channel layer was deposited to the total thickness of 600 nm. During this phase of growth, InAs quantum dots were formed by the following procedure; after growing a 200 nm-thick GaAs spacer layer, the substrate temperature was decreased immediately to 450°C. Then 1.75 ML InAs was deposited with the rate of 0.1 ML/s to form dot structures. We monitored the growth of InAs by reflection high-energy electron diffraction (RHEED) to confirm the formation of dots [19] when the streaky (2×4) reconstruction pattern transformed to a spotty pattern, as soon as the dots are formed. Then 2.5 nm GaAs was grown at 450°C without interruption to cover the dots. Then the substrate temperature was heated up again to 600°C for further overgrowth of a 400 nm-thick GaAs. Note that InAs dots of this sample S1 are 200 nm from the heterointerface, suggesting that strain effects of dots are negligible. In addition, the ground level of InAs is well above the Fermi level as shown in Fig. 1 unless the surface potential is lowered by the gate electrode.

In advance, another sample of nearly identical structure was prepared except that GaAs overgrowth was dispensed. Atomic force microscopy (AFM) was performed on this sample in air to show that 1.75 ML InAs deposited under our growth condition results in dot structures of typically 5 nm in average height and 20 nm in average diameter at the bottom. Their average concentration is about \( 5 \sim 10^{10} \text{ cm}^{-2} \). We also prepared, for comparison, two more samples, S2 and S3, in which we inserted a 1 ML InAs and no InAs layer, respectively. These three samples were grown in one day.
We processed these wafers to produce field-effect transistors by forming an Al gate, Hall voltage probes, as well as the source and drain electrodes. The carrier concentration \( N_s \) and mobility \( \mu \) of two-dimensional electron gas (2DEG) were measured as a function of the gate voltage \( V_g \) by using the Hall effect [20].

3. Transport and trapping properties

First, we measured \( N_s \) and \( \mu \) of all three samples at 4.2 K in the dark and the results are plotted in Fig. 2. Note in Fig. 2a that the threshold voltages of all three samples S1 (filled circle), S2 (solid line) and S3 (dashed line) are about \(-0.6 \text{ V}\), from which \( N_s \) increases linearly up to \( V_g = 0.8 \text{ V} \). Note that the slopes \( dN_s/dV_g \) of these three samples are almost the same. The slope is determined by the gate capacitance between the gate and the channel and is \( 20 \text{ nF/cm}^2 \) for all samples, which agrees well with the designed value.

Electron mobilities \( \mu \) were also measured for \( V_g \) between \(-0.5 \text{ V} \) and \( +0.8 \text{ V} \). \( \mu-N_s \) characteristics are shown in Fig. 2b. Note that S1, S2 and S3 show nearly identical characteristics. This proves that InAs dots play small roles in this region, since InAs dots are well displaced from the 2DEG channel. Hence, the electrons scarcely overlap with the cores of InAs dots and a surrounding strain induced by lattice mismatched InAs. Note that the observed dependence of \( \mu \) on \( N_s \) suggests that the dominant scatterers for all samples are a small number of the coulombic charges coming from background impurities or other impurities originating from nominally \( \delta \)-doped Si in the AlGaAs [21]. Note here also that \( \mu \) of electrons are more than \( 70 \times 10^4 \text{ cm}^2/\text{Vs} \) at \( N_s \sim 2 \times 10^{11}/\text{cm}^2 \) in GaAs/AlGaAs inverted heterojunctions, which indicates very good quality of our MBE growth [18, 22].

Once \( V_g \) exceeds \( 0.8 \text{ V} \), leakage current begins to flow between the gate and the channel. This anomalous region will be discussed later. As mentioned earlier, the gate leakage current remains small for \( V_g < 0.8 \text{ V} \). This is reasonable as the Schottky barrier of \( n \)-GaAs is about \( 0.8 \text{ eV} \). For \( V_g \geq 0.8 \text{ V} \), the leakage substantially increases in sample S2 and S3, while it remains rather low in sample S1. The electron concentration \( N_e \) of sample S1 stays constant for \( V_g \geq +0.8 \text{ V} \), indicating that the gate electric field is screened by some charges. This saturation of \( N_e \) and reduction of the leakage current in S1 is due to the appearance of an extra potential barrier, which is induced by the trapping of electrons by InAs dots. This situation is shown in Fig. 3a and Fig. 3b. When the potential of the surface layer is pulled down by the gate bias \( (V_g = 0.8 \text{ V}) \) as in Fig. 3b, some electrons will be trapped by the dots, which raises the potential. This charging process of dots prevents the leakage current for the gate voltage range between \( +0.8 \text{ V} \) and \( +1.9 \text{ V} \). When \( V_g \) exceeds \( +1.9 \text{ V} \) the leakage current begins to flow again, since the InAs dots are then fully occupied.

In reducing \( V_g \) from \( +1.9 \text{ V} \), \( N_e \) of S1 is found to decrease with the same slope, as shown by open circles in Fig. 2. The \( N_e-V_g \) characteristics are shifted by \( 4.5 \times 10^{10} \text{ cm}^{-2} \). This shift is due to the discontinuous change of electric field \( \delta F \), which is
induced by the trapped charges in the InAs dot layer. As shown in Fig. 3c, \( \delta F \) can be expressed as

\[
\delta F \equiv F_2 - F_1 = \frac{\delta V}{W_{GaAs}},
\]

where \( W_{GaAs} \) is the thickness (\( = 0.4 \mu m \)) of the upper surface GaAs layer and \( \delta V \) is the voltage shift of \( N_s-V_g \) characteristics. From such data, we can estimate the number, \( N_{\text{Dot}} \), of captured electrons in InAs quantum dots, using Gauss's law.

\[
N_{\text{Dot}} = \frac{\varepsilon_0 \varepsilon_{GaAs} \delta F}{e},
\]

where \( \varepsilon_0 \varepsilon_{GaAs} \) is the permittivity of GaAs, \( e \) is the elementary charge. Fig. 2a gives a value of \( \delta V = 0.371 \mathrm{~V} \), from which we find that \( N_{\text{Dot}} \) is \( 6.56 \times 10^{10} \ \mathrm{cm}^{-2} \).

Since the density \( D_{\text{Dot}} \) of InAs dots in our sample is estimated to be \( (5-10) \times 10^{10} \ \mathrm{cm}^{-2} \), the density, \( N_{\text{Dot}} \), of trapped electrons in the dots is very close to \( D_{\text{Dot}} \). This indicates that the dots are charged by one or two electrons.

To clarify trapping mechanisms, we measured the \( N_s-V_g \) characteristics at 4.2 K for three different maximum gate voltages, \( V_{\text{max}} \). Incidentally, measurements were done at 4.2 K but after each measurement, the sample was heated up to room temperature and exposed to a usual room light in order to discharge InAs dots. Fig. 4, \( N_s \), shows the result.

When \( V_g \) is decreased from \( V_{\text{max}} = 0.8 \ \mathrm{V} \) (indicated by \( \bigcirc \)), no shift in \( N_s \) is observed, suggesting that InAs quantum dots capture such a small number of electrons that no potential charging effect is observed. As Fig. 4 shows, \( N_s-V_g \) relation for \( V_{\text{max}} = +1.9 \ \mathrm{V} \) makes a loop and crosses at \( V_g = +1.4 \ \mathrm{V} \). However, \( N_s-V_g \) relation for \( V_{\text{max}} = +1.4 \ \mathrm{V} \) does not agree with that for \( V_{\text{max}} = +1.9 \ \mathrm{V} \). This means that to fully occupy dots with electrons, electrons must be injected by applying excess gate bias. Since the saturation of \( N_s \) observed in a region of \( V_g = +1.0 - +1.9 \ \mathrm{V} \) results from the band bending at InAs dot layer as shown in Fig. 3, one can assume that in this region one or two extra electrons can be trapped at excited levels of InAs dots. Indeed, it has been theoretically and experimentally shown that some electrons can occupy not only the ground state but the excited levels in the dots [14, 23]. When reducing
Fig. 5. The concentrations $N_e$ of electrons in the channel of the dot embedded FET measured as a function of the gate-source voltage at two different temperature 4.2 and 77 K.

$V_{\text{max}}$ electrons in the excited levels may be discharged, since these electrons in the excited levels can be detrapped more easily than those in the ground level even at 4.2 K. When $V_g$ is decreased from $V_{\text{max}} = 1.4$ V, relatively small $N_e$ shift is observed, indicating that not all the InAs dots are occupied. The estimated number of trapped electron is $4.31 \times 10^{10}$ cm$^{-2}$, using Eq. (1) and Eq. (2).

$N_e - V_g$ characteristics were also studied at 77 K. The result is shown in Fig. 5. It is clear that both the slope $dN_e/dV_g$ and the shift of the gate threshold $\delta V_g$ at 77 K are almost the same as those of 4.2 K, suggesting that the number of trapped electrons is almost unchanged. Note that a loop is observed only at 4.2 K, as shown in Fig. 4. Although the enhancement of $V_{\text{th}}$ is attributed to the existence of a barrier at 4.2 K, the details of this charging and discharging processes, including a loop at 4.2 K in this region, will be published elsewhere. Since the barrier around the dot is quite high ($\sim 100$ meV) [15, 19], electrons trapped in the ground level cannot be thermally activated even at 77 K. The rate of tunnel escape process is also expected to be small, since the surrounding GaAs barrier is quite thick. Hence, the reduction of $N_e$ persists even if $V_g$ is biased at $-1.0$ V, at which the density of 2DEG almost vanishes.

4. Conclusions

We have studied MBE growth of InAs dots and the effects of InAs quantum dots inserted in a novel FET structure and demonstrated that the InAs dots work as electron trap centers, in a region where the wave function of 2DEG hardly overlaps the InAs dots or strain region. Although $\mu$ of electrons is not degraded by inserted InAs dots, the electron concentration, $N_e$ at a given gate voltage $V_g$ is found to be reduced once a positive $V_g$ beyond 1.0 V is applied, shifting the threshold voltage of this novel FET device. From the reduction of $N_e$ in the channel, we have shown that each InAs dot captures one or two electrons. We have shown that the number of trapped electrons in InAs dots is dependent on the maximum voltage applied to the gate. This effect was observed at both 77 and 4.2 K, suggesting that electrons captured in the ground state are not detrapped at 77 K. If the structure is optimized, this quantum trap (QT) FET may well function at higher temperatures and serve as a novel memory device.

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References

In situ RHEED control of direct MBE growth of Ge quantum dots on Si (0 0 1)

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Abstract

RHEED studies of the morphological transformation of germanium layers during their MBE growth on the Si(0 0 1) surface were carried out. Zero-streak profiles analysis were used to control the formation of coherent and dislocated germanium islands at the film thickness of more than 5 and 8 monolayers (ML), respectively. The dislocation-free islands formed at the first stage (4–8 ML) of the morphological transformation were employed for designing the structures with the self-organized Ge quantum dots. The effects of charge and size quantization of hole transport, such as the Coulomb blockade and the resonance hole tunneling through the discrete energy levels in the quantum dots were observed in these structures.

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Keywords: Germanium; Silicon; MBE; RHEED; Quantum dots

1. Introduction

In the recent years direct synthesis of low-dimensional GeSi structures has received considerable attention. The self-organization of 3D Ge islands on silicon according to Stranski–Krastanow’s growth mechanism was used for the first time in Refs. [1–3] to produce a layer of quantum dots as small as 10 nm in diameter. A lot of quantum-sized effects were observed in heterostructures containing quantum dots [1–3]. This approach was widely used later to make GeSi dot structures as well as III–V and II–VI ones. A direct growth allows obtaining of smaller quantum dots against those obtained by lithographic methods. Additionally, the direct synthesis makes it possible to overcome numerous technological problems in designing quantum-sized structures. During the synthesis of the quantum dot structures, it is important to know sizes of the 3D germanium islands and to be able to control in situ the process of their formation. In this connection, the present work was aimed at RHEED studies of the morphological transformations in germanium-on-silicon layers and of the
direct growth of the structures containing self-organized quantum dots.

2. Experimental details

Germanium on silicon films were grown using a MBE installation “Katun-C” equipped with two e-beam evaporators for germanium and silicon and two Knudsen cells for doping. Si (0 0 1) wafers of 60 mm diameter were used as substrates. The growth process was controlled in situ using the RHEED technique (20 kV). Diffraction patterns and intensity variations versus time at separate points and along the given profiles were recorded by using a computer-aided TV system “FOTON-4”. In the present work variations in intensity profiles versus time were studied along (0 0) streak in the [1 1 0] azimuth. The shortest recording interval for one intensity profile was 0.5 s. The Ge growth rate was measured through observation of RHEED oscillations of specular beam, it was equal to about 0.01 nm/s.

3. RHEED control of quantum dots formation

The initial stages of germanium growth on the Si (0 0 1)-(2 x 1) surface were studied by observing variations in the diffraction patterns and the intensity profiles. From the onset, there were observed RHEED oscillations (Fig. 1a) each period of which corresponded to filling of one monolayer (ML) of a/4 (0.14 nm) thickness. At the film thickness of 4-6 ML, highly smeared bulk spots appeared in the diffraction pattern (Fig. 2a). In the literature, such a pattern is accounted for by the formation of germanium clusters shaped as a tetrahedral pyramid with the side orientation of {1 0 5} type (the so-called hut-clusters) [4, 5]. According to Refs. [5, 6], there are no misfit dislocations in the clusters, and the strain relaxation in them proceeds due to deformation of the silicon substrate. Fig. 3a presents germanium hut-clusters formed upon growth of the 7 ML film.

The thickness of 5 ± 0.3 ML of the Ge film corresponding to the formation of the smeared bulk spots was determined by analyzing profiles (Fig. 4a, b). The specular beam is divided into two. As the film grows, they transfer towards the bulk diffraction positions. The most spectacular is the transfer observed for the (0 0 6) beam. The nature of this transfer is not clearly understood now; we assign it to the redistribution of intensities of electron beams reflected by the free silicon surface and by the surface of germanium {1 0 5} planes facing the hut-clusters. The bulk beam is a superposition of electron beams reflected by the surfaces of these two types. Two of these reflections are of large half-widths due to the small cluster size and, thus, are merged into one spot in the diffraction pattern. As germanium is deposited, the cluster grows and...
the intensity of the beam reflected by the \{1 0 5\} plane increases. On the other hand, the cluster-free Si\{0 0 1\} surface area decreases, which results in a lower intensity of the specular beam. As a consequence, the bulk spot is shifted from the original position of the specular beam towards the range of large angles.

A similar behavior of bulk spots was observed by us at the growth temperature range of 200–600°C. The beams move more readily at high growth temperatures, probably, due to a larger size of the clusters and a higher rate of their plane growth. The data on profile analysis of RHEED patterns used to elucidate dynamics of the growth of 3D germanium islands will be reported elsewhere.

When the thickness is above 8–10 ML, the diffraction pattern changes gradually (Fig. 2b). According to Refs. [4–6], now it is attributed to the formation of germanium islands with misfit dislocations faced by \{1 1 3\} planes. When the thickness is above 7.5–8 ML, bright and narrow spots, assigned to bulk diffraction on the islands, appear in the intensity profiles close to the bulk diffraction position (Fig. 4). This is clearly seen after splitting the \{0 0 6\} spot into two Gaussians (Fig. 4c and Fig. 4d). The wide spot assigned to reflection from \{1 0 5\} planes disappears rather quickly. For example, it reduces to one-sixth height when the thickness increases by 2 ML only (Fig. 4c and Fig. 4d). The small width of the bulk reflections indicates a markedly larger size of the dislocated islands against the size of the hut-clusters. The latter assumption is supported by the TEM studies of a Ge film of 12 ML thickness (Fig. 3b). The islands with dislocations are formed at a later stage of the morphological reconstruction, and their size (50 nm) is considerably larger than that of the hut-clusters (below 20 nm).

Hence, the germanium hut-cluster films are preferable to design structures with smaller quantum dots. Additionally, the absence of misfit dislocations allows defect-free layers to be grown at the quantum dot surface, which is of importance for development of structures for electric physical measurements. Notice that analysis of intensity profiles in the RHEED pattern makes it possible to detect islands with dislocations before the diffraction pattern changes. Therefore, the formation of large 3D islands incorporating misfit dislocations can be prevented.

4. Direct growth of quantum dot structures

To study tunnel current through the 3D germanium hut-clusters, we grew the structures consisting of the following layers (in order of growth from the p⁺-Si\{0 0 1\} substrate):
- a base electrode p⁺-Ge₀.₇Si₀.₃, 17 nm;
- a tunnel barrier Si, 8 nm;
• a layer of isolated Ge islands, 1 nm (average thickness);
• a tunnel barrier Si, 16 nm;
• a counter electrode p⁺-Ge₀.₅Si₀.₅, 17 nm;
• an amorphous p⁺-Ge contact layer, 500 nm

The layers were processed into mesas of 200–500 μm diameter. Indium ohmic contacts were linked with the substrate and top contact. The differential transversal conductivity \( G = \frac{dI}{dU} \) was measured at 4.2 K and conductivity oscillations were observed. Transversal conductivity versus bias voltage for a sample grown at 400°C is shown in Fig. 5. The observed peaks were explained in terms of resonant tunneling of holes through individual energy levels in quantum dots. A splitting of peaks into doublets is observed due to Coulomb blockade of holes transport [2]. The positions of the doublets calculated in accordance with Ref. [7] are indicated by arrows in Fig. 5. The dot sizes used for calculation were obtained from the TEM image (Fig. 3a) with a diameter of 19 nm and height of 30 nm (on the assumption of cylinder-shaped dots). The general consistency between the theory and experimental data is apparent. Details on studies of electrical and optical properties of the grown GeSi heterostructures were given in our previous papers [1–3, 8].

5. Conclusions

It was shown that in situ analysis of intensity profiles in the RHEED pattern is more effective, compared to visual examination of its variations, for the control of the morphological transformation of germanium films on the Si(0 0 1) surface. The thickness of formation of dislocation-free hut-clusters (5 ML) and plastically relaxed 3D germanium islands (8 ML) was determined.

The structures containing self-organized quantum dots of germanium were grown in a direct manner. Effects of size quantization of the hole energy spectrum and Coulomb blockade of hole transfer were observed for these structures.
Fig. 4. Zero-streak intensity profiles in RHEED pattern during Ge/Si(0 0 1) films growth at 400°C. Film thickness are 4.6(a), 5.3(b), 8.6(c) and 11.1(d) ML.

Fig. 5. Transversal differential conductivity versus bias voltage for a structure (see the text) grown at 400°C [2].

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Growth controlled fabrication and cathodoluminescence study of 3D confined GaAs volumes on non-planar patterned GaAs(001) substrates


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Abstract

We report on (a) the effect of growth interruption on the growth profile evolution in growth on non-planar patterned mesa tops via substrate-encoded size-reducing epitaxy (SESRE) and (b) the optical behavior of isolated 3D-confined GaAs volumes as well as 3D-confined GaAs volumes coupled with 1D-confined quantum wells (QWs) fabricated by SESRE. Steady-state excitation and time-resolved cathodoluminescence (CL) are used for these optical studies.

Keywords: Patterned substrate; Growth interruption; Quantum boxes; Time-resolved cathodoluminescence

1. Introduction

The fabrication of semiconductor structures with electronic states confined in more than one dimension is a subject of much activity in recent years. Of the various approaches to the fabrication of such structures, in situ, one-step growth approaches are promising since the structures thus fabricated are expected to be devoid of the problems of damage and contamination. Substrate-encoded size-reducing epitaxy [1] is one such in situ approach in which the differential growth rates on various facets and inter-facet adatom migration are exploited to reduce the mesa top size from as-patterned length scales (typically 1 μm) down to the nanoscale (< 100 nm) regime. One of the critical issues in this approach is the growth condition dependence of the growth profile evolution. Previous studies [1-3] of the growth condition dependence of the growth profile evolution have been limited to the case of continuous growth. In this paper we report on some findings on the effect of growth interruption on the growth on stripe mesas on GaAs(001).

Additionally, 3D confined GaAs volumes synthesized on <100> oriented square mesas [3] via SESRE are examined via steady state as well as time-resolved CL studies to shed light on carrier dynamics of significance to carrier collection.

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efficiency in isolated or communicating 3D-confined volumes.

2. Experimental procedure

GaAs (001) ± 0.1° substrates were patterned via conventional photolithography followed by wet chemical etching. The effect of growth interruption was studied for growth on stripe mesas oriented along the ⟨100⟩ and [110] directions, with mesa width of ~1 μm and depth of ~0.5 μm. The growth interruption used in these experiments has a duration of ~120 s. GaAs layers of 45 ML thickness followed by 5 ML AlGaAs marker layers were grown to reveal the growth profile evolution. Square mesas for the fabrication of 3D-confined GaAs volumes were patterned along ⟨100⟩ directions. The details of the substrate preparation, MBE growth procedures and the growth profile evolution for 3D-confined GaAs volumes have been given in an earlier paper [3]. For the samples with interacting 3D-confined GaAs volumes and 2D-confined GaAs QWs, the MBE growth consisted of a size-reducing buffer of four periods of [175 ML Alₐ₀.₂₅Ga₀.₇₅As/525 ML GaAs] followed by a 40 period multiple quantum well (MQW) composed of [40 ML Al₀.₂₅Ga₀.₇₅As/20 ML GaAs]. For the sample containing isolated 3D-confined GaAs volumes, the growth consisted a size-reducing buffer composed of 16 period of [5 ML Al₀.₂₅Ga₀.₇₅As/195 ML GaAs] followed by 100 ML Al₀.₂₅Ga₀.₇₅As/20 ML GaAs/200 ML Al₀.₂₅Ga₀.₇₅As and capped by 50 ML GaAs.

The CL studies were performed with a modified JEOL-840A scanning electron microscope (SEM) with a 0.25 m monochromator. Time-resolved CL experiments were performed with the method of delayed coincidence in an inverted single photon counting mode with a time resolution of ~100 ps [4]. Electron beam pulses of 50 ns width and a frequency 1 MHz were used for excitation for the time-resolved studies.

3. Results and discussion

3.1. Effect of growth interruption on the growth profile evolution

In Fig. 1 are presented the transmission electron microscope (TEM) images showing the growth profile evolution on stripe mesas along the [110] (parallel to the As dimer direction; panels (a) and (b)) and the [010] (at 45° to the As dimer direction; panels (d) and (e)) directions for different temperatures. The As₄ pressure is kept constant at 4.8 × 10⁻⁶ Torr and the GaAs growth rate at

![Fig. 1. TEM images of growth on stripe mesas: (a) and (b) along [110] direction, (d) and (e) along [010] direction. Schematic drawings of the growth profile evolution are shown in panels (c) and (f) for the stripe mesas along [110] and [010] directions, respectively.](image)
0.5 ML/s. Only a portion of the mesa top and sidewalls is shown in these images corresponding to the area marked by the dotted rectangle in the schematic drawings in panels (e) and (f). The thin white lines are the AlGaAs markers which delineate the growth profile. The arrows indicate the GaAs layers in which growth was interrupted.

In the case of the growth on stripe mesas along the [1 1 0] direction, layer thickness on the mesa top (sidewall) for the GaAs layers in which the growth was interrupted is found to be higher (lower) than the preceding and following layers. This implies that migration of Ga atoms from the sidewalls to the mesa top continues in the absence of Ga delivery. Careful measurements of these layer thicknesses for the growth at 610°C indicate that more than 1 ML worth of material migrated from the sidewalls. This indicates that during growth interruption it is not only the initial adatoms on the sidewalls but the atoms that were incorporated into the GaAs layers during the growth also migrate to the mesa top. The effect of interruption on the growth profile evolution becomes less significant when the substrate temperature is lowered to 590°C. The flat morphology on the mesa top as evidenced by the AlGaAs markers implies that the migration length of the adatoms on the mesa top, perpendicular to the stripe direction, is larger than half the mesa width (i.e. the migration length is at least ~0.5 μm for the growth condition used).

Unlike the above case, in the case for the growth on stripe mesas along [0 1 0] direction at 610°C the thickness on the mesa top of the GaAs layers for which the growth was interrupted is found to be lower than the preceding and following layers (which is evident for the first two interruptions in panel (d)). This implies that in this case Ga atoms migrate in the opposite direction, i.e. from the mesa top to the sidewalls in the absence of Ga delivery. Moreover, a curved morphology is present at the intersection of the mesa top and sidewalls for the GaAs layers with growth interruption. Again, the effect of interruption on the growth profile evolution becomes less significant when the substrate temperature is lowered to 590°C.

3.2. CL studies of 3D-confined GaAs volumes

3.2.1. Interacting 3D-confined GaAs volumes and 1D-confined quantum wells (QW's)

Fig. 2 shows a TEM image of a typical square mesa containing interacting 3D-confined GaAs volumes (dark regions) on the mesa top below pinch-off and 1D-confined GaAs QWs on the (101) sidewalls. The area-averaged CL spectra, taken from a region ~1 μm x 1 μm centered on the mesa top [5], at various sample temperatures are shown in Fig. 3. The spatial region responsible for the identifiable peaks in Fig. 3 can be inferred from the spatially resolved monochromatic CL images and the SEM image shown in Fig. 4. The luminescence from p1 and p2 is primarily from QWs grown on sidewalls away from mesa top while p3 is localized about the mesa top and hence is from thick 3D-confined GaAs volumes just below the pinch-off region. Peak p4 is similarly identified to be from the underlying bulk-like GaAs in the size-reducing buffer.

The peak intensity decrease of p1 relative to p2 and p3 with increasing temperature seen in Fig. 3 indicates thermal re-emission of carriers from thinner QWs (p1) and their subsequent transport and recombination in the thicker QW (p2) and the 3D-confined volumes (p3). Further insight into these thermalization effects is gained from the time-delayed CL measurements. The intensity of peaks p1–p4 (I_p) as a function of time (t) for both decay and onset stages is shown in Fig. 5a and b, respectively. The onset rate r, defined as Δln(I_p)/Δt,
is the slowest for p4 (see Fig. 5b) reflecting the large distances (∼0.5 μm) that the hot carrier must traverse before recombining in the thick bulk-like GaAs layers in the size-reducing buffer. These relative rates are also observed in Fig. 5c from the peak intensity ratios. The initial decay time τ, measured from the slopes in the ln(Ip) versus time transients shown in Fig. 5a increases in the order p1, p2 and p3 implying the enhanced thermal re-emission of carriers in the QWs (p1 and p2) to feed continuously the larger 3D-confined GaAs volumes represented by p3, as the system proceeds towards equilibrium.

3.2.2. Isolated 3D-confined GaAs volumes

Fig. 6a shows a TEM image of growth of the SQW on the square mesa top. As seen in the figure, the mesa top size at the start of the GaAs well deposition is ∼120 nm. The thickness of the GaAs layer on the mesa top (∼11 nm) is higher compared to the deposition thickness (∼5.7 nm) due to adatom migration from the {1 0 1} sidewalls to the (0 0 1) mesa top. Negligible GaAs growth on the {1 0 1} sidewalls in the vicinity of the 3D-confined volume is observed indicating good lateral confinement.

Results from CL imaging and spectroscopy are shown in Fig. 6 (panels (c) and (d)) and Fig. 7, respectively. The monochromatic CL images (Fig. 6) show that the high energy peak seen in the spectra shown in Fig. 7 is associated with 3D-confined GaAs volume and the low energy peak has its origin in the bulk-like GaAs in the size-reducing buffer layer. The CL spectra from the

![Fig. 3. Constant excitation CL spectra taken at various temperatures. The peak shifts for p1 to p4 with temperature are shown by dashed lines.](image)

![CL images](image)

Fig. 4. Monochromatic CL images at various wavelengths (a)-(d), and (e) SEM image. The arrow tips point to the mesa apex position. CL imaging conditions are indicated in the upper right corner.
Fig. 5. CL transients for peaks p1–p4: (a) the decay curves with linear fits for the initial ∼ 6 ns of decay; (b) the onset curves with linear fits for the first ∼ 2 ns; (c) CL peak intensity ratios p2/p1, p3/p1, and p4/p1.

non-patterned planar region (not shown) shows a peak at λ = 778 nm at 87 K which is associated with the SQW deposition. The red shift of the CL peak from well deposition on the mesa top with respect to that from the planar region is attributed to interfacet adatom migration induced GaAs thickness enhancement on the mesa top. The temperature dependence of the CL spectra (see Fig. 7) shows that the ratio of the peak intensity of the 3D-confined to bulk-like GaAs decreases with increasing temperature (T < 90 K) indicating good carrier confinement in these 3D-confined volumes. The reversal of this trend for temperatures beyond 90 K is under investigation.

Strong evidence for 3D confinement of the high energy peak in the CL spectra comes from the results of excitation current dependence. Fig. 8 shows a plot of the full width at half maximum (FWHM) versus the excitation current for the peaks from the mesa top 3D confined GaAs volume (labeled mesa) and the corresponding GaAs/AlGaAs single QW (SQW) in the planar region (labeled planar) of the same sample. The FWHM of the SQW on the planar region increases from ∼6.7 to ∼9.6 meV as I_s increases from 10 pA to 5 nA, whereas the FWHM of the 3D-confined region increases by a factor of almost 3 for the same increase of I_s. This enhanced peak width broadening reflects rapid phase-space-filling in the 3D-confined GaAs volume.

4. Conclusion

The role of growth interruption during SESRE on patterned stripe mesas is examined for the first time and found to reveal remarkable continued interfacet migration during growth interruption.
Fig. 7. Constant excitation CL spectra taken at various temperatures in $18 \, \text{K} \leq T \leq 200 \, \text{K}$. range.

Temperature-dependent and time-resolved CL experiments on coupled 3D-confined volumes grown by SESRE reveal carrier feeding from sidewall quantum wells to the 3D-confined volumes. Isolated confined 3D volumes show the expected phase-space-filling behavior.

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Fig. 8. Full width at half maximum (FWHM) of CL peaks originating from the 3D-confined GaAs volume and the GaAs/AlGaAs single QW (denoted as mesa and planar, respectively) as a function of excitation current.

References

Manipulating InAs island sizes with chemical beam epitaxy growth on GaAs patterns

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Abstract

We demonstrate the manipulation of InAs island size through growth on patterned GaAs substrates. Using chemical beam epitaxy, we deposited islands on patterns consisting of concentric circular trenches. One sample was left uncapped for atomic force microscopy, and another was capped with GaAs for micro-photoluminescence. Luminescence images taken at particular energies show that at different orientations along the circular arcs, the islands have different luminescence energies and thus different sizes. In correlating the island distributions found by atomic force microscopy images with the luminescence, we conclude that the InAs islands that form in high-density one-dimensional chains are smaller than those that are not in chains.

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1. Introduction

The small defect-free islands that form at the early stages of compressively strained epitaxial growth may be interesting for single-electron or single-hole devices. For the case of InAs on GaAs, the islands that form on top of a thin wetting layer after one to two monolayers of growth are pyramidal shaped, having 12–25 nm wide bases, and 3–6 nm heights [1, 2]. These islands have a deep conduction-band potential well which may contain only a single bound state [4]. Many groups have reported such islands on flat substrates grown with molecular beam epitaxy [3, 5–10], metalorganic chemical vapor deposition [11, 12], and chemical beam epitaxy [13]. When island-containing layers are separated by thin GaAs layers, the islands align vertically [3, 14], and may be placed closely enough together to couple the conduction band states [15–17, 23]. Such stacked islands could be at the heart of electrical devices.

The individual stacks may possibly be contacted because island position can be controlled by growing on lithographically patterned GaAs substrates [18]. The islands form selectively in the patterned features, displaying a clear dependence on local surface orientation. The number of islands in small holes varies approximately as the square of the hole diameter [19]. A selectivity ratio has been defined
as the ratio of the island density in the holes to the density away from the pattern, with a ratio of more than 1000 reported [19]. The minimum center-to-center period reported for chains of islands many microns long was 33 nm [18]. When capped with GaAs the islands on patterns retain their quantum dot luminescence properties. Other reports on island growth in patterns include InAs islands aligned on intentionally bunched step edges [20], InAs islands grown on etched gratings that showed surface orientation and deposition shadowing effects [21], and InP and InAs islands grown on corrugations formed by W-wire overgrowth that showed surface orientation and alignment effects [22].

We report here that for a given set of growth conditions the island size depends on the size and orientation of patterned features. InAs islands were formed on two GaAs substrates patterned with sets of concentric circular trenches. One sample was left uncapped for atomic force microscopy (AFM) and the other was capped with GaAs for microscopic photoluminescence (µPL). The island linear density seen by AFM in the trench varies with angle around the trench. In µPL the island luminescence energy also varies with angle around the circles. This demonstrates that not only can patterns be used to position islands, but may as well be used to engineer the island sizes.

2. Sample growth and experimental details

The chemical beam epitaxy machine, of our own design, uses pressure-regulated sources without a carrier gas. The precursors are triethylgallium, trimethylindium, and tertiary-butylarsine, with the tertiary-butylarsine thermally cracked to As2. The data we present here are from growths 941230A, an uncapped sample, and 941230B, a GaAs-capped sample. Arrays of sets of seven concentric circles were defined in (1 0 0)GaAs substrates with electron beam lithography and wet etched to a depth of 100 nm. The radial pitch of the circles is 1.25 μm. A GaAs buffer layer of 100 nm was grown before depositing approximately 1.5 monolayers of InAs at 493°C. More detailed growth conditions are given in Ref. [18]. The InAs island density outside of the patterned area of the uncapped sample was \(2 \times 10^{7}\) cm\(^{-2}\). The buffer layer growth was very orientation-dependent.

We operated our AFM in a constant force mode. Our transmission-electron microscopy measurement on islands grown under similar condition showed approximately pyramidal islands of 4 nm height and 20 nm base. Our estimate of the AFM tip radius is 80 nm, using the \(\approx 50\) nm apparent island width and assuming a spherical tip and a 4 nm high sharp point for the island. The data we present here have been leveled and flattened, and are displayed to simulate oblique illumination, but are otherwise unprocessed. The PL data were taken at 5 K using the 488 nm line of an Ar\(^+\) laser for excitation. The luminescence signal was either dispersed with a monochromator and detected with a Ge detector for spectra or was directed through interference filters into a CCD camera for imaging. Image integration times were from 1 to 10 s.
Fig. 2. Linear island densities and trench width as functions of arc angle measured at the circle center and with respect to the [0\,1\,1] direction. The center islands are those aligned in the center of the trench, and the wall islands are those that have otherwise formed on the side wall of the trench. The inset depicts the meaning of arc angle.

3. Atomic force microscopy

An AFM micrograph of InAs islands formed in circular arcs on the uncapped sample is given in Fig. 1. Illumination of the data surface is from the left. Here we will refer to the position along the circular arcs by giving the angle measured at the circle center, increasing counterclockwise from the [0\,1\,1] direction. Only angles from the first quadrant are discussed because the circles are symmetric. This geometric definition is depicted schematically in an inset to Fig. 2. While the as-etched circles were formed with anisotropic etching, the buffer layer growth was very orientation-dependent. Where the trenches run in the [0\,1\,1] direction, at 90° arc angle, they are almost closed off, and those in the [0\,1\,1] direction, at 0° arc angle, are very wide and have even contributed GaAs to the (1\,0\,0) ridge top between them. At positions along the arcs from 30 to 60°, the trench side walls have several shallow pockets with characteristic sizes of 0.25 μm.

The InAs islands have formed very selectively in the etched and overgrown trenches. For the narrow portions of the arcs from about 45 to 90°, the islands are concentrated in dense chains in the center of the trenches. A high proportion of the pockets on the side walls, perhaps originating from lithographic irregularities, have single or few islands in them. At angles from 30° down to 0° along the arcs, some InAs islands and roughness can be seen in the trenches. The roughness is perhaps due to partially formed islands. The islands and roughness below 30° are scattered and not clearly associated with the trench center.

We have measured the trench width and counted the island density as functions of angle and plotted the results in Fig. 2. The data are from the next to the outermost circle of the pattern, a portion of which is visible in the upper right corner of Fig. 1. The radius of the circle is 7.5 μm. The islands have been put into two classifications for the counting: those aligned in the trench center and those on the trench side walls, including those in shallow
pockets. The resulting counts are plotted as 1D densities along the arc. The centered island chains are seen at linear densities above 20 μm⁻¹ for trench widths of 600 nm and less. The centered chains essentially begin to disappear for angles below 45°. The islands on the trench side walls are seen over the whole arc length, though have the highest density from 30 to 45°, in the region with the most side wall pockets.

4. Micro-Photoluminescence

The broad area photoluminescence from the circles on the capped sample, given in Fig. 3, shows the typical features of a low- to medium-density InAs island sample. The tall peak at 1.440 eV is from the approximately one-monolayer-thick wetting layer, and the low broad peak at 1.32 eV is from the InAs islands.

Fig. 4 shows six μPL images taken at the energies indicated by the markers (a)-(f) in the spectrum of Fig. 3. The overall nonuniformity in intensity from each image is due to the nonuniformity of the exciting laser spot. The four lower energy images, from 1.240 to 1.348 eV in Fig. 4a–Fig. 4d, have luminescence coming almost exclusively from the sets of concentric circles. At the imaging energy of 1.409 eV, Fig. 4e, the luminescence principally comes from the circles, but a substantial amount of intensity also comes from the unpatterned areas between the sets of circles. The highest energy image, Fig. 4e, taken with the wetting layer luminescence is approximately complementary to the island images.

The angular distribution of the luminescence is different for each of the images in Fig. 4. For the images of island PL, Fig. 4a–Fig. 4d, most of the intensity is above some angle with respect to [011]. In the image taken at the intermediate
energy of 1.409 eV, Fig. 4e, the intensity shows an “X” pattern near 32°, and most of the intensity within the circles comes from lower angle regions. Essentially, all of the wetting layer luminescence from the patterned area, Fig. 4f, comes from regions below an angle of approximately 30°. Here we will characterize each image by the angle of the edge that defines the upper or lower limit of the luminescing region. The energy of the images are plotted in Fig. 5 versus the angles which characterize the edges of the luminescing regions. Two points are plotted for the image of Fig. 4a because two edges are seen. The luminescence edge of a seventh image at 1.319 eV which is not shown in Fig. 4 is also plotted.

5. Discussion

The AFM and μPL images may be combined to draw some conclusions about which islands are luminescing at what energies. From the island density distributions plotted in Fig. 2, it is seen that center island chains are principally at angles higher than 40°, but the side wall islands are distributed to much lower angles, down through 20°. The
luminescence edges in the images taken at energies between 1.26 and 1.35 eV are at angles above 40°, but the luminescence edge of the images taken at 1.24 eV extends down to approximately 20°. We therefore infer that the center-chained islands are smaller and luminesce at the higher energies than the side wall islands which can be larger and luminesce at the lower energies.

The “X” shaped pattern seen at 32° in the image taken at the intermediate energy of 1.409 eV seems to mark the boundary between the center island chain luminescence and the wetting layer luminescence. This is the angle below which the chains thin out and disappear, and may be considered as some sort of phase boundary for the condensation of the InAs islands into 1D chains.

In this study we have not attempted to determine island size from AFM measurements, which are highly complicated by tip-island convolution effects. On a flat substrate, we estimate our vertical resolution to be about 2 InAs monolayers for these islands that are 10–15 monolayers high. Our lateral resolution of the shape differences, though, is no better than 5 nm. Together, these lead to an uncertainty in the volume of around 50%. On the tilted faces of the patterned features, the uncertainties are greater because different islands on different local orientations will be measured with different portions of the tip.

A principal conclusion that we reach using the data of this paper, and especially those of Fig. 5, is that for a given set of growth conditions, the island size can be manipulated by changing the size and orientation of etched and overgrown patterns. Here the changing size and orientation are the trench width and angle. The indication of changing island size is the measured change in luminescence-energy distribution characterized by the angle of the edge of the luminescing region. Thus, in Fig. 5, the independent variable is the pattern and the dependent variable is the size. Unfortunately, while these circular patterns provide an existence proof of
manipulating island size with patterns, these patterns do not lend themselves well to separating the independent influences of trench orientation and width on island formation.

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References

Self-organized quantum dot structures in strained \((\text{GaP})_n(\text{InP})_m\) short period superlattices grown on GaAs \((N\ 1\ 1)\) by gas-source MBE

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Abstract

\((\text{GaP})_n(\text{InP})_m\) short-period superlattices (SLs) are grown on GaAs \((N\ 1\ 1)\) \((N = 2\ 5)\) and \((1\ 0\ 0)\) substrates by gas-source molecular beam epitaxy. Scanning tunneling microscopy and transmission electron microscopy observations show that the SLs grown on GaAs \((N\ 1\ 1)A\) have lateral-composition-modulated dot/columnar structures with a lateral period of about 10–20 nm, while wire structures are formed on GaAs \((1\ 0\ 0)\). Quantum dots formed in \((\text{GaP})_n(\text{InP})_m\) SL/\(\text{In}_{0.45}\text{Ga}_{0.55}\)P multilayers by self-organization exhibit strong 77 K photoluminescence (PL) with a narrow full-width at half-maximum of about 40 meV. Anomalous temperature variation of PL peak energy is also observed in these self-organized structures.

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Keywords: Self-organized quantum dot; GaP/InP superlattice; Gas source MBE; TEM; STM; PL

1. Introduction

Low-dimensional structures such as quantum wires (QWRs) and quantum dots (QDs) are attracting great interest for novel device applications and physical studies. Progress in the fabrication of high-quality QWR/QD structures using self-organized growth method has supplied opportunities for detailed experimental studies on two/three-dimensional quantum confinement structures [1–4].

Recently, we have reported the substrate-orientation dependence of lateral composition modulation in the \((\text{GaP})_n(\text{InP})_m\) superlattices (SLs) grown by gas-source molecular beam epitaxy (MBE) [5]. We have shown that the formed structures of the SLs grown on GaAs \((1\ 0\ 0)\), and \((3\ 1\ 1)A\) and \((4\ 1\ 1)A\) are wire and dot structures, respectively, by transmission electron microscopy (TEM) observation. On the other hand, we have found that the
(GaP)$_n$(InP)$_m$ SLs grown on GaAs (1 1 1) have strong CuPt-type ordering and exhibit a band-gap reduction as large as 321 meV (large red-shift), compared with the band gap of disordered InGaP of the same average composition.

In this paper, we report the scanning tunneling microscopy (STM) and TEM observations on the high density of self-organized structures formed in the (GaP)$_n$(InP)$_m$ SLs grown on GaAs(N 1 1)A by gas source MBE. We also describe the TEM and photoluminescence (PL) results on the self-organized QD structures fabricated by growing (GaP)$_n$(InP)$_m$ SL/In$_{0.49}$Ga$_{0.51}$P multilayers.

2. Experimental procedure

(GaP)$_n$(InP)$_m$ SLs and (GaP)$_n$(InP)$_m$ SL/In$_{0.49}$Ga$_{0.51}$P multilayer QD structures ($n \leq m$) were grown by gas source MBE. Elemental Ga, In and thermally cracked AsH$_3$, PH$_3$ were used as group III and group V sources, respectively. Reflection high-energy electron diffraction (RHEED) intensity oscillations were used to calibrate GaP and InP growth rates (GaP: 3.3 s/ML; InP: 3.7 s/ML). Both the SLs and QD structures were grown at 420–460°C after the growth of 300 nm GaAs buffer layer at 600°C. AsH$_3$ and PH$_3$ flow rates were 1.0 and 1.2 sccm, respectively. Samples for STM observation were doped with Si and the carrier concentrations were 2 x 10$^{18}$ cm$^{-3}$. Formed structures were investigated by STM, TEM and PL measurements. Electrically etched tungsten (W) tips were used as a STM probe in this study. TEM observation was performed to compare with the STM image. Both the 325 nm line of a He–Cd laser (19 mW) and 488 nm line of an Ar$^+$ laser (70 mW) were used in the PL measurement as excitation sources.

3. Results and discussion

3.1. STM and TEM observation of self-organized structures in (GaP)$_n$(InP)$_m$ SLs

TEM images (not shown) for 0.3 µm-thick (GaP)$_n$(InP)$_m$ SLs grown on GaAs (1 0 0) substrates showed lateral wire composition modulation (formation of the Ga-rich and In-rich regions) with a lateral period of about 10–20 nm and a lateral wire length of over 200 nm along the [0 1 1] and [0 1 1] directions, respectively, as observed by Hsieh et al. [1]. The length of the lateral wire structures was increased by the growth on vicinal (1 0 0) substrates. This composition modulation is probably due to the island formation along the [0 1 1] direction during growth, which is primarily determined by the direction of the group V dimer bonds related to the migration direction [1].

Plan-view TEM observations on the 0.3 µm-thick (GaP)$_2$(InP)$_2$ SLs grown on GaAs (N 1 1)A revealed the lateral periodic composition modulation (dot structures) with a period of 10–20 nm, along both the [0 1 1] and [K l m] directions ([0 1 1] equivalent direction on the (1 0 0) plane). Both (0 1 1) and (0 1 1) cross-sectional TEM images showed columnar structures with a lateral period of about 10–20 nm [3]. They are oriented along the [1 0 0] direction for N ≥ 4 or [2 1 1] direction for N ≤ 3. It is noteworthy that on the (1 0 0) surface the lateral periodic composition modulation occurs only along the [0 1 1] direction (wire structures), while on the (N 1 1)A surfaces the lateral periodic composition modulation occurs along the [0 1 1] direction as well as along the [K l m] direction. The dot/columnar structures observed here are not considered to be caused by spinodal decomposition because the structures caused by spinodal decomposition were observed along the [0 1 0] and [0 0 1] equivalent directions [6, 7]. The cross-sectional TEM images for the SLs grown on (1 1 1)A and (1 1 1)B substrates revealed that the lateral composition modulation does not occur and that the quasi-perfect monolayer SL is formed until n ≤ 4. PL peak energies were also greatly dependent on the substrate orientation and monolayer number n [5].

Thirty six periods of (GaP)$_{1.5}$(InP)$_{1.88}$ SLs and (GaP)$_2$(InP)$_{2.5}$ SLs were here grown on GaAs(3 1 1)A and GaAs(4 1 1)A substrates at 460°C, and were studied by STM and TEM. Fig. 1 shows their STM ((a) and (c)) and plan-view TEM ((b) and (d)) images. These images indicate the formation of uniform high-density dot structures along the [K l m] and [0 1 1] directions. The dot periods and the observed structures in STM and
plan-view TEM images are nearly the same. The dot period studied by STM observation increased with increasing $n$ and $m$. Scanning tunneling spectroscopy (STS) measurement showed that the voltage width for the $dI/dV = 0$ at bright region was narrower than that at dark region, suggesting that they correspond to the In-rich region and the Ga-rich region, respectively, and showing the band-gap variation due to lateral composition modulation.

In these SLs the dot (columnar) structure densities were larger than $10^{11}$ cm$^{-2}$. These results indicate the promising potential for the fabrication of high-density quantum dot (QD) structures by growing (GaP)$_n$(InP)$_m$ SL/GaP multilayers, as shown in the next subsection. High density of QD structures are favorable for obtaining high optical gain and therefore low threshold current density in laser diodes.

### 3.2 Quantum dots formed in (GaP)$_n$(InP)$_m$ SL/Ga$_{0.49}$In$_{0.51}$P multilayers

Self-organized QWR and QD structures were grown on GaAs (1 0 0) and (N 1 1)A, respectively. The grown structures consist of (i) a 300 nm-thick In$_{0.49}$Ga$_{0.51}$P buffer layer, (ii) 5 cycles of (GaP)$_n$(InP)$_m$ SL (18 periods, about 20 nm) layers and In$_{0.49}$Ga$_{0.51}$P (20 nm) layers and (iii) a 300 nm-thick In$_{0.49}$Ga$_{0.51}$P top layer, as shown in Fig. 2a.

Both (0 1 1) and (0 1 1) cross-sectional TEM images show lateral composition modulation, indicating QD structure formation in the (GaP)$_{1.5}$(InP)$_{1.88}$ SL layer regions grown on GaAs (4 1 1)A substrate, as shown in Fig. 2b and Fig. 2c. The QWR structures were formed in the SL grown on GaAs (1 0 0) substrates for the same structures.
PL spectra at 77 K for the QDs grown on (N 1 I)A are shown in Fig. 3. The full width at half-maximum (FWHM) for the QDs are about 39–98 meV. For the QDs on (3 1 I)A the best PL intensity and FWHM were obtained. PL peak energy variation with growth temperature at 420–460°C was very small, though PL intensity increased with increasing growth temperature.
By increasing the deviation percentage of the SL periodicity $\Delta T$ [8],

$$\Delta T = (\text{SL period} - l \lambda_0)/\lambda_0,$$

the PL peak energy of the QDs showed a larger red-shift due to larger lateral composition modulation in the (GaP)$_n$(InP)$_m$ SL region, where SL period $= (n + m)\lambda_0/2$, $n = l = \text{integer}$ and the GaAs lattice constant $\lambda_0 = 5.6533 \text{ Å}.$

The PL peak energy for the QDs was higher than that for the QWRs. The PL intensity of the QDs was stronger than that of QWRs. The PL intensity for the QDs formed in (GaP)$_2$(InP)$_{2.5}$ SL/In$_{0.49}$Ga$_{0.51}$P multilayers was three orders of magnitude stronger than that of QWRs. However, the PL intensity of the QWRs for the (GaP)$_{1.5}$(InP)$_{1.88}$ SL was two orders of magnitude stronger than that for the (GaP)$_2$(InP)$_{2.5}$ SL.

3.3. Temperature dependence of PL peak energy

We have observed anomalous temperature dependence of the PL peak energy for these QDs. Fig. 4a shows the temperature variation of the PL spectrum for QDs formed in the (GaP)$_{1.5}$(InP)$_{1.88}$ SL/In$_{0.49}$Ga$_{0.51}$P multilayers grown on GaAs (3 1 1)A. FWHM is less than 40 meV below 80 K, though it decreases with increasing temperature due to tunneling effect between neighbor quantum states up to 45 K. At temperatures over 80 K it increases further, the origin of which is not known at present. Similar temperature variation of

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**Fig. 3.** PL spectra at 77 K for the self-organized quantum dots (QDs) formed in the (GaP)$_{1.5}$(InP)$_{1.88}$ SL and (GaP)$_{1.5}$(InP)$_{1.88}$ SL/In$_{0.49}$Ga$_{0.51}$P multilayers grown on GaAs (3 1 1)A.

**Fig. 4.** Temperature dependences of (a) PL spectra and (b) PL peak energies for the QDs formed in the (GaP)$_{1.5}$(InP)$_{1.88}$ SL and (GaP)$_{2.5}$(InP)$_{2.5}$ SL/In$_{0.49}$Ga$_{0.51}$P multilayers grown on GaAs (3 1 1)A.
FWHM was also observed for the QDs formed in the (GaP)$_{2}(InP)_{2.5}$ SL.

With increasing temperature the PL peak energy for the (GaP)$_{1.5}$(InP)$_{1.88}$ SL decreases rapidly in the low-temperature region, has a minimum at about 100 K, then increases in the range of 100–190 K, and has a relatively small but normal temperature variation over 200 K, as shown by closed circles in Fig. 4b. Much more pronounced temperature variation was also observed for the QDs of the (GaP)$_{2}(InP)_{2.5}$ SL. PL peak energy has a minimum at about 120 K, then increased in the range of 125–175 K, and had a normal temperature variation over 200 K, as shown by quadrangles in Fig. 4b. The temperature variation of PL intensity observed in these QDs is similar to those observed for the InAs QD structures formed on GaAs by the Stranski–Krastanov mode MBE [2]. On the other hand, the PL peak energy for the QWRs on GaAs (1 0 0) showed very small temperature variation up to 120 K. The origin of these anomalous temperature variations is not known at present, but the existence of the strained ordered structures in the (GaP)$_{n}$(InP)$_{m}$ SL regions may play a role. Similar anomalous temperature variation of the PL peak energy to that for the QDs was observed for the ordered alloy of InGaP grown by MOVPE [9, 10]. The anomalous temperature variation of PL peak energy observed here is much larger than those observed for the InGaP ordered alloy [9, 10].

4. Summary

(GaP)$_{n}$(InP)$_{m}$ SLs were grown on GaAs (N 1 1) and (1 1 0) substrates by gas-source MBE. STM and TEM observations showed that the quantum dot/columnar structures were formed in the SLs grown on GaAs (N 1 1)A substrates by the lateral periodic composition modulations both along the [0 1 1] direction and the [0 1 1] equivalent direction with a lateral period of 10–20 nm. The surface densities of these dot structures were larger than $10^{11}$ cm$^{-2}$.

Quantum dots were fabricated by growing (GaP)$_{n}$(InP)$_{m}$ SL/In$_{0.45}$Ga$_{0.55}$P multilayers on GaAs (N 1 1)A substrates by self-organization. Strong 77 K PL with a narrow FWHM of about 40 meV were observed for these QDs grown on GaAs(3 1 1)A. Anomalous temperature variation of the PL peak energy was observed in all of these self-organized structures.

Acknowledgements

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References

Intersubband absorption from In$_{0.26}$Ga$_{0.74}$As/GaAs quantum dot superlattice

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Abstract

We have grown a high-quality 20 period InGaAs/GaAs quantum dot superlattice with a standard structure typically used for quantum well infrared photodetector. Normal incident absorption was observed around 13–15 μm. Potential applications for this work include high-performance quantum dot infrared detectors.

Keywords: Quantum dot; Intersubband absorption; Quantum dot infrared photodetector

1. Introduction

Zero-dimensional (0D) quantum dots (QD) structures have attracted much interest in recent years due to their δ-function-like density of states, strong carrier localization, large exciton binding energies, and enhanced oscillator strength [1–3]. High-quality quantum dot can be self-formed in situ in Stranski–Krastanow growth mode without resorting to any substrate patterning process [4–11].

The growth of quantum dot superlattice has also been of much interest [9, 10]. The formation mechanism of quantum dots needs to be further investigated, such as the complex indium migration dynamics in the process of self-organization growth and the formation of wetting layer. On the other hand, very high detectivity quantum dot infrared photodetector can be achieved by using intersubband transition taking place between bound levels in quantum dot. The discrete levels in quantum dots hinder carrier relaxation toward the ground state and thus a very long carrier lifetime [12–14]. This is called the “phonon bottleneck effect”. Unlike the conventional GaAs/AlGaAs QWIPs that require a coupler (e.g. gratings) to couple the normal incident radiation [15], the intersubband transition in quantum dot can be induced by the normal incident radiation due to the localized state in quantum dots. Hence, the intersubband absorption quantum dot infrared photodetector (QDIP)

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has a great advantage over the conventional GaAs/AlGaAs QWIP.

Quantum dot multilayer is required for enhancing infrared absorption. High-quality InGaAs/GaAs QDs have been difficult to grow due to dislocation occurring with increasing layer thickness. Presently, there have been only a few reports on high-quality InGaAs/GaAs quantum dot superlattice [9, 10]. In Stranski–Krostanow growth mode, initial growth starts two dimensionally in the layer-by-layer mode, but after a certain thickness is reached, islands form spontaneously and a very thin 2D wetting layer is left behind the islands. If the growth is allowed to continue beyond the initial island formation stage, misfit dislocations start forming at the island edges [4, 12]. Therefore, the dislocation-free InGaAs/GaAs quantum dots require that the growth be interrupted immediately after the formation of the islands but before the islands reach a size for which strain relaxation and misfit dislocation occurs [4]. The superlattice is a more complex system than the single layer and dislocation easily occurs in the process of quantum dot growth.

Using lower indium fraction permits the thicker layer without dislocations, and the island formation may be completely controlled by growth condition. In the paper, we have grown a high-quality 20 period InGaAs/GaAs quantum dot superlattice (QDS) with low indium fraction (x = 0.26). The normal incident infrared absorption can be observed at 13–15 μm.

2. Experimental procedure

The InGaAs/GaAs QDS was grown by molecular beam epitaxy (Riber-32P) on semi-insulating GaAs (1 0 0) substrate. The structure of InGaAs/GaAs QDS is very similar to the conventional QWIP. The layers consisted of, a 1.3 μm buffer layer, a 1.0 μm n⁺ contact layer, a 20 period InGaAs/GaAs quantum dot arrays, and a 1.0 μm GaAs and 200 nm AlGaAs (x = 0.04) top contact layer. The InGaAs layers and the contact layers were Si-doped with doping concentrations of 1 × 10¹⁸ cm⁻³, and δ-doping for InGaAs layer. The growth rate for GaAs was 2.83 A/s. The indium fraction in InGaAs is about 0.26–0.28, which is determined by reflection high-energy electron diffraction oscillations and equivalent beam pressure. The quantum dots were self-formed by the coherent relaxation into islands of InGaAs between undoped GaAs layers. The actual amount of indium incorporated in the dots is difficult to measure or calculate due to the complex dynamics of the atoms during island formation.

The cross-sectional transmission electron microscope (TEM), low-temperature photoluminescence and X-ray double-crystal diffraction have been used to measure the structural and optical properties for quantum dot superlattice. The infrared absorption was measured for QDS by using a Fourier transform infrared (FTIR) spectrometer at room temperature, which was carried out under two modes: multipass 45° wave guide and normal incident mode. These measured results are presented in Figs. 1–6.

3. Result and discussion

Fig. 1 shows the cross-sectional TEM micrograph of the quantum dot array. A 20 period defect-free quantum dot array is clearly shown. Using cross-sectional view TEM results, statistics on the QD size can be extracted. Fig. 2 shows these results. It shows that the average size of quantum dots is 28 nm (diameter), 7 nm (height) and 0.26, height to diameter ratio. It is worth noting that the actual size of quantum dot is smaller than the one measured by TEM because the distributions measured by TEM are affected by strain which tends to overestimate the size. Fig. 3 shows the low-temperature (10 K) photoluminescence (PL) spectrum of QDS. A strong luminescence peak around 1.25 eV is observed. The peak PL intensity dependence of excitation power is also presented in Fig. 3 and it is noted that there is no saturation of intensity, which indicates that island-attibuted PL is not due to defects or impurities. The dot peaks have a full-width at half-maximum of about 53 meV, which is due to the size fluctuation of quantum dots. These results are in good agreement with the present reports for quantum dots [2–4]. It is worth noting that the vertical correlation of quantum dot is not
obvious in our sample, which is due to the thicker spacer layer (34 nm). This observation agrees with a report by Xie [10]. Fig. 4 shows the measured X-ray double-crystal rock curve. Over ten satellite peaks are observed, indicative of good structural quality. The period of superlattice is calculated to 34 nm. The quantum dot superlattice structure is shown schematically in Fig. 5.

As measured in the conventional QWIPs, we measure the infrared spectrum of quantum dot array using Brucker IFS-120HR FTIR in a multi-pass 45° wave-guide configuration [15], as seen in Fig. 6(a). Strong absorption peaks are observed at 13–14 μm with a 13 meV FWHM. Fig. 6b shows the measured infrared absorption spectrum under the normal incident radiation. The obvious absorption peaks around 14–15 μm with a 11 mev can be observed. Compared with that measured in a multi-pass 45° wave-guide configuration, the peaks is shifted to the lower energy, which is similar to the

Fig. 1. Cross-sectional view of quantum dot superlattice.

Fig. 2. Histogram of dot (islands) measured from cross-sectional TEM micrograph and fit with a Gaussian curve: (a) the island diameter; (b) the island height; (c) the ratio of the height to diameter.

Fig. 3. Low-temperature (10 K) photoluminescence spectra of sample. The insets show the PL intensity dependence of excitation power and it is noted that there is no saturation of intensity, which indicates that island-attributed PL is not due to defects or impurities.
depolarization effect in normal incident absorption InGaAs/GaAs quantum well [16]. The measured multi-peak structure may be due to the different size of quantum dots, as seen in the PL results.

Since we do not know accurate In content in islands, it is impossible to calculate the energy level for the quantum dot superlattice. The observed intersubband absorption probably results from either quantum dots or quantum well (wetting layer) since the normal incident intersubband transition is possible in InGaAs/GaAs QWs [16, 17]. However, the narrow absorption line width suggests that the absorption takes place between the bound levels in low-dimensional material. Based on the nominal growth parameters, it is impossible to find such the intersubband transition from bound level to bound level in quantum well (the bound-to-continuum instead) [15, 18]. In particular, the normal infrared absorption can be directly observed without using a multipass 45° waveguide and polarization configuration. In the case of quantum well, the normal incident absorption is so weak that it is difficult to observe without using waveguide [15]. Hence, the 13–15 μm infrared absorption results from the intersubband transition in quantum dots. To the author's knowledge, this is the first report that the normal incident infrared absorption can be observed from the QDS. Since the detectivity is proportional to carrier lifetime, the quantum dot infrared photodetector can be predicted to have much higher sensitivity than the quantum well infrared photodetector (QWIPs).

4. Conclusion

We report for the first time, the infrared absorption around the wavelength of 13–15 μm from 20 period dislocation-free InGaAs/GaAs quantum dot superlattice (QDS) with the low In fraction (x = 0.26). It opens a way to high sensitivity quantum dot infrared photodetectors. The QDS has been investigated by the transmission electron micrograph, low-temperature photoluminescence and X-ray double-crystal diffraction. The transmission
electron micrograph reveals that the size distribution of quantum dots: \( \sim 28 \text{ nm} \) in diameter and 7–15 nm in height. A strong luminescence peak is observed around \( \sim 1.25 \text{ eV} \) from 10 K photoluminescence. The superlattice period is 34 nm.

References

Molecular-beam epitaxy of self-assembled InAs quantum dots on non-(1 0 0) oriented GaAs

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Abstract

In this paper we report optical properties of InAs quantum dots (QD) grown by molecular-beam epitaxy on GaAs (2 1 1)A, (n 1 1)A/B, where n is 1, 5 and 7, and on reference (0 0 1) substrates. The photoluminescence (PL) spectra reveal differences in amplitude, integral luminescence, peak position and shape. Temperature dependence indicates an additional lateral confinement on (0 0 1), (n 1 1)B, (2 1 1)A and (1 1 1)A substrates. Our results also show an enhancement of QD onset thermal quenching energy by a factor of ~ 3 for these orientations, when compared with the reference quantum well. PL polarization measurements show strong in-plane dependence caused by the quantum dot’s structural anisotropy. However, the structure grown on (5 1 1)A and (7 1 1)A surfaces does not exhibit QD formation.

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Keywords: Quantum dots; Self-assembled; High-index; MBE

1. Introduction

The growth of strained films has become a field of intense study in recent years. In particular, InGaAs/GaAs and InAs/GaAs systems have been intensively investigated due to the possibility of producing nearly defect-free quantum dot (QD) structures. These structures are a result of the Stranski–Krastanow (SK) growth mode [1]. Such a growth mode starts two-dimensionally; after a certain critical thickness is reached, islands are formed spontaneously, and a thin wetting layer is left below the islands. The main QD parameters, such as size distribution, lateral size, height and form, are dependent on the mechanism of stress relaxation of the two-dimensional layer (2D) to three-dimensional (3D) islands. Such quantum dots have mainly been obtained by molecular-beam epitaxy (MBE) [2, 3] or metalorganic chemical vapor deposition (MOCVD) [4, 5], with (0 0 1)-orientation substrates. Few works studying QD formation on GaAs high-index surfaces have been reported [5, 6]. During MBE growth these high-index planes break up into periodic arrays of nanometer scale facets, thus lowering the surface energy. This microscopic surface morphology represents the

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equilibrium surface structure, and is generally stable during growth. Therefore, the lateral periodicity is fixed for a certain crystallographic orientation [7, 8]. Such periodicity might influence the indium adatom kinetics, resulting in different stress relaxation for each surface. These conditions determine the growth mode for each orientation.

In this work, we used high-index GaAs surfaces to study the formation of self-assembled InAs quantum dots on \((n\ 1\ 1)A/B\) planes. Photoluminescence (PL) results indicate QD formation for all orientations except \((5\ 1\ 1)A\) and \((7\ 1\ 1)A\). PL polarization measurements suggest the presence of developed microfacets on QDs.

2. Experimental procedure

The samples were grown by MBE technique, using a Meca 2000 system, on GaAs reference \((0\ 0\ 1),\ (2\ 1\ 1)A\) and \((n\ 1\ 1)A/B\) (where \(n\) is 1, 5 and 7) substrates. Growth process was monitored in situ by means of reflection high-energy electron diffraction (RHEED) (15 kV). The substrates were degreased and etched ex situ, then soldered side by side with indium on a molybdenum holder and immediately transferred to the sample load-lock system. After transfer to the MBE growth chamber, the surface oxide had been thermally desorbed by heating the substrate wafers to \(\sim 580^\circ\text{C}\) under an arsenic flux. An As\(_4\) background pressure of \(9 \times 10^{-6}\) Pa was used during structure growth. Then, the substrate temperature \(T_s\) was raised to \(620^\circ\text{C}\) and samples were outgassed for 5 min. After this, \(T_s\) was lowered to \(600^\circ\text{C}\) and a 80 nm GaAs/AlAs (2/2 nm) superlattice was grown, followed by a 0.5 μm GaAs buffer layer. During the last 0.1 μm buffer layer, \(T_s\) was decreased to \(500^\circ\text{C}\) to grow a 3 nm In\(_{0.2}\)Ga\(_{0.8}\)As reference quantum well (QW) and the remainder of the structure. The gallium- and indium-cell temperatures were kept constant during \(T_s\) variations. In order to compare optical properties, the QW was grown with the same nominal estimated thickness as the InAs quantum dots [9] and separated from them by a 100 nm GaAs layer. In order to finish the structure, a 50 nm GaAs cap layer was grown. The growth rates of GaAs and InAs determined by the RHEED oscillation technique were, respectively, 0.23 and 0.061 nm/s on GaAs(0 0 1) plane. Since \(T_s\) (500°C) was below critical temperature for starting indium reevaporation, we assumed these growth rates to be the same for all orientations. Interface smoothing was carried out, using growth interruption, during 1 min before and 3 min after InAs deposition. During these growth interruptions, growth was stopped momentarily by closing the gallium and indium cells for a specified duration, while the arsenic cell was always kept open to maintain an excess arsenic pressure condition. The QD nucleation was directly observed by RHEED on (0 0 1) surface, when the pattern changed from streaky to spotty. We observed that the growth mode was 2D during the deposition of the first three monolayers (ML) and later it changed to 3D (SK mode). Afterwards three InAs ML were deposited for QD formation. The surface morphology of the \((1\ 1\ 1)A\) and \((2\ 1\ 1)B\) orientations was hazy due to choice of growth parameters for optimum quality of the \((0\ 0\ 1)\) planes. The other planes showed a mirrorlike surface. The photoluminescence measurements were performed at 16 K in a closed-cycle He cryostat using a 514.5 nm line of Ar\(^+\) laser. The average excitation density was 150 W/cm\(^2\). A photomultiplier mounted on a 0.5 m monochromator was adapted for detection of PL signal in lock-in mode.

3. Results and discussion

Fig. 1 shows PL spectra at 16 K for all orientations. Concerning just the reference quantum well (QW) PL signal, we have observed that the peak position and the full width at half-maximum (FWHM) depend on crystallographic orientation. Such an effect has already been reported in Ref. [10]. For \((0\ 0\ 1),\ (n\ 1\ 1)B,\ (2\ 1\ 1)A\) and \((1\ 1\ 1)A\) surfaces, an additional peak has been detected at around 1.35 eV. This peak is attributed to the quantum dot formation on each orientation. High QD PL efficiency was obtained for \((0\ 0\ 1)\) and \((7\ 1\ 1)B\) surfaces, where QD–QW integrated PL intensity ratio is 3.2 and 1.9, respectively. In the other planes this ratio was one order of magnitude smaller. Peak positions, FWHM, and the integral
plane, fitting is done by two Gaussians. The Gaussian with high-energy peak was attributed to the contribution of wetting layer to PL signal. We also measured QD PL excitation density dependence in the range of $10^{-10}$ W/cm$^2$ ($T = 16$ K) in order to deduce the density of stress-related defects in the QDs. The result shows linear dependence (not shown here), indicating a low defect density on QD interfaces. The inset of Fig. 1b exhibits PL spectra for (5 1 1)A and (7 1 1)A planes. The arrows indicate an unexpected peak of an additional structure, with energy position higher than the QD. We believe that this PL characteristic is related to a two-dimensional layer (QW) formed after the stress relaxation of a critical InAs layer. An enhancement of the FWHM of this PL signal is observed in comparison to reference QW. This increase is probably due to the formation of dislocations or related defects as a result of stress relaxation. Non-QD formation on (5 1 1)A has previously been reported [5, 10].

In order to obtain additional information about the QD properties, measurements of PL temperature dependence were carried out. Fig. 2 displays the (7 1 1)B PL spectra. The QW PL intensity decreases drastically when the temperature changes from 16 to 70 K, contrary to that of QDs, which persists up to ~160 K. This persistence at high temperatures is related to the increase of oscillator strength due to additional lateral confinement [11, 12]. In addition, an unusual large red shift for QD peak position (~0.1 eV) is observed when temperature oversteps ~45 K for all orientations. The rate of energy decrease with temperature is about two or three times higher than InAs band gap decrease rate. In order to explain this red shift and reduction of FWHM (see below), a model involving exciton recombination and thermal activation and transfer has recently been proposed [13]. In our case, this model does not describe completely our experimental data because the Gaussian shape of QD PL peak is maintained as temperature raises [14]. Another possible explanation is related to a temperature dependence of electron effective mass in InAs. Because the quantization energy of an electron–hole pair in the QD is higher by about a factor of 4 than that in the QW, this causes a greater change in the quantization

intensity ratios QD/QW for all samples are listed in Table 1. For each orientation the QD peak can be fitted by one Gaussian shape. In the case of (5 1 1)B

Fig. 1. PL spectra of (a) (1 1 1)A/B, (2 1 1)A, and (b) (5 1 1)B, (7 1 1)B, and (0 0 1) samples (dotted lines). Solid line represents Gaussian fitting for QDs, indicating formation of QDs with one predominant size. The inset in (b) shows the PL spectra for (5 1 1)A and (7 1 1)A surfaces, where grown the structure does not exhibit QD formation.
Table 1
Peak energy (in eV), FWHM (in eV), and QD–QW wavelength integrated PL (IPL) intensity ratio for all orientations at 16 K

<table>
<thead>
<tr>
<th>Orientation</th>
<th>Quantum dot</th>
<th>Quantum well</th>
<th>IPL_qq/IPL_QW</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$F_r$</td>
<td>FWHM</td>
<td>$E_r$</td>
</tr>
<tr>
<td>(0 0 1)</td>
<td>1.338</td>
<td>$\sim$ 123</td>
<td>1.461</td>
</tr>
<tr>
<td>(7 1 1)B</td>
<td>1.344</td>
<td>$\sim$ 102</td>
<td>1.444</td>
</tr>
<tr>
<td>(5 1 1)B</td>
<td>1.330</td>
<td>$\sim$ 58</td>
<td>1.430</td>
</tr>
<tr>
<td>(2 1 1)A</td>
<td>1.381</td>
<td>$\sim$ 128</td>
<td>1.466</td>
</tr>
<tr>
<td>(1 1 1)A</td>
<td>1.392</td>
<td>$\sim$ 136</td>
<td>1.468</td>
</tr>
<tr>
<td>(1 1 1)B</td>
<td>1.387</td>
<td>$\sim$ 132</td>
<td>1.468</td>
</tr>
</tbody>
</table>

Fig. 2. (7 1 1)B and (7 1 1)A (inset) PL temperature dependence. The arrow in the inset indicates presence of an unexpected peak (see text).

Fig. 3. QW and QD peak amplitude thermal energy dependence for (7 1 1)B plane. The QW peak is thermally quenched at $\sim$ 1.5 meV, whereas the QD peak persists up to $\sim$ 4 meV (see Table 2).

Energy of the QD relative to that of the QW when the effective mass is changed [15]. The inset of Fig. 2 shows PL temperature dependence for (7 1 1)A surface. We can see that the behavior of the unexpected peak, indicated with arrow, is different from the (7 1 1)B QD. This dependence also suggests the existence of stress-related defects, and lack of relationship between peak and QD formation.

Fig. 3 exhibits QD and QW PL intensities shown in Fig. 2 as a function of the thermal energy. From similar plots for all orientations, we extracted the value for thermal quenching energy ($E_Q$). Further, by plotting the logarithm of wavelength-integrated PL values versus $1/kT$, we determined activation energy ($E_A$) of thermionic emission.
Table 2
Thermal quenching energy ($E_q$) and electron–hole emission activation energy ($E_a$) values in meV for all orientations

<table>
<thead>
<tr>
<th>Orientation</th>
<th>$E_q$</th>
<th>$E_a$</th>
</tr>
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<tbody>
<tr>
<td></td>
<td>QD</td>
<td>QW</td>
</tr>
<tr>
<td>(0 0 1)</td>
<td>~ 6</td>
<td>~ 2</td>
</tr>
<tr>
<td>(7 1 1)B</td>
<td>~ 4</td>
<td>~ 1.5</td>
</tr>
<tr>
<td>(5 1 1)B</td>
<td>~ 6</td>
<td>~ 2</td>
</tr>
<tr>
<td>(2 1 1)A</td>
<td>~ 6</td>
<td>~ 2</td>
</tr>
<tr>
<td>(1 1 1)A</td>
<td>~ 6</td>
<td>~ 2</td>
</tr>
<tr>
<td>(1 1 1)B</td>
<td>~ 6</td>
<td>~ 2</td>
</tr>
</tbody>
</table>

$E_a$ and $E_q$ values are summarized for all orientations in Table 2. We observed that QD’s $E_q$ is about 3 times larger than QW’s. $E_a$ values observed for QD are about 1.4–2.4 times higher than for QW, suggesting suppression of electron-hole emission from QDs to GaAs barrier, due to a binding energy increase. Activation energy of unexpected peak of (5 1 1)A and (7 1 1)A surfaces was also determined. The $E_a$ values are around 18 and 24 meV for (5 1 1)A and (7 1 1)A orientations, respectively. For (5 1 1)A plane, this value is almost the same as for reference QW (17 meV). In the case of (7 1 1)A, activation energy is 1.3 times higher than that for the reference QW (19 meV). Taking into account temperature dependence, power density dependence of peak position and intensity, as well as activation energy values, we attributed the unexpected peak, already referred to, to the emission of a 2D layer.

Fig. 4 exhibits an unexpected effect present in PL FWHM as a function of temperature for (7 1 1)B, (2 1 1)A, (1 1 1)A/B orientations. In the case of QDs, an unusual decrease of FWHM was detected for Gaussian-shaped emissions from 16 to 85 K for (1 1 1)B, 100 K for (1 1 1)A and (2 1 1)A, and 130 K for (7 1 1)B planes, respectively. This effect was reported in Ref. [14] and described in terms of suppression of non-predominant size QD emissions, and persistence of predominant size emissions present at the center of a Gaussian shaped PL function. In Ref. [13] as noted above, another model was proposed to explain this reduction of FWHM.

QL PL polarization dependence measurements were also carried out to determine probable structural anisotropy of QDs in-plane. Fig. 5 shows the angular dependence of normalized PL peak intensity for (0 0 1), (7 1 1)B and (5 1 1)B orientations. These planes demonstrate a strong polarization effect. A maximum of PL intensity has been set for PL component when polarization angle is zero, occurring when polarization is parallel to the following crystallographic directions: [2 7 7] for (7 1 1) plane, [2 5 5] for (5 1 1)B and [1 1 0] for (0 0 1) plane.
(0 0 1), respectively. Based on our results, we believe that QDs grown on these oriented planes have well-developed microfacet configuration. Similar results have been reported for InAs QD on (0 0 1) [16] and InGaAs QD on high-index surfaces [6].

4. Conclusions

Optical properties of self-organized InAs quantum dots grown on high-index GaAs substrates have been investigated. Such properties are strongly dependent on substrate orientations. Quenching energy for QDs is around three times higher than for reference QW. Dependence of optical characteristics with temperature is related to an increase of exciton binding energy and activation energy for thermionic emission, due to additional lateral confinement. Our results show an unusual decrease of FWHM with temperature, suggesting suppression of non-predominant size QD emissions and persistence of those of predominant size. Non-QD formation was detected on (7 1 1)A and (5 1 1)A orientations. For such surfaces, an unexpected peak related to a two-dimensional layer was observed. These results suggest that another growth mechanism predominates in such A surfaces as a result of stress relaxation.

Acknowledgements

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References

InAs/GaAs self-assembled quantum dots grown by ALMBE and MBE

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Abstract

We have studied the effect of different growth schemes of structures with InAs/GaAs self-assembled dots on the size of dots and, then, on their PL features. Lower confining layers, dots and upper confining layers were grown by different techniques (ALMBE and MBE), at different growth temperatures and by using growth interruptions. Both AFM and PL show that the ALMBE is particularly suited to grow relatively large dots with sharp size distributions at relatively large InAs coverages without incurring dot coalescence. Moreover, the low temperature ALMBE growth of caps reduces the interaction between dots and caps. The control of dot sizes and of dot/cap interaction results in PL emissions shifted towards the spectral windows of photonic interest. In particular, structures with ALMBE dots and with optimized confining layers show a significantly bright emission at 1.26 µm at 300 K.

PACS: 68.55

Keywords: Quantum dots; MBE; ALMBE

1. Introduction

Self-assembled dots originating from deposits of highly strained materials on lattice-mismatched substrates are very simple systems which show low-dimensionality effects that can be exploited both for applications and for fundamental studies on semiconductor physics and crystal growth [1–3]. Among the most important parameters of the dot populations, there are the island size and its distribution; the size determines the luminescence emission energy from the dots, while the size distribution affects the full widths at half maximum (FWHMs) of the transitions. InAs/GaAs dots generally have dimensions such that their emission occurs at ≤ 1 µm at low temperatures; therefore, in order to have room temperature (RT) emissions within the spectral windows of photonic interest, such as the 1.3 µm one, the dot sizes should be substantially increased, in order to reduce the extent of carrier confinement. Emissions at 1.24 µm at RT have been obtained from structures with
almost-coalesced dots and, then, with reduced carrier confinement [4]. However, uncontrolled coalescence as a tool for increasing the dot dimensions should be avoided, since it may result in the formation of dots which include misfit dislocations, that drastically reduce the PL efficiency [5]. The achievement of FHMs lower than the thermal energy at room temperature \((kT \sim 25 \text{ meV})\) is of paramount importance so as not to spoil the advantages of the zero-dimensionality of the structures.

While MBE and MOVPE are widely used for the preparation of dot structures, new growth techniques are investigated [6–8] since they may give interesting benefits. Atomic layer MBE (ALMBE, also termed as migration enhanced epitaxy, MEE [9]) is considered interesting since the alternate supply of anions and cations gives rise to a significant increase of cation surface migration. It is expected that this feature would result in sharper size distributions of larger dots, since it may cause the selective enlargement of existing dots, instead of the continuous nucleation of new small islands. Moreover, the enhanced migration of cations allows the growth of InGaAs alloys at low temperatures where the In segregation towards the growth front is substantially minimized [10]; this should reduce the interaction of quantum dots with the upper confining layers in structures intended for luminescence; such an interaction may reduce the dimensions of dots and substantially modify their shape [11].

The aim of this work is to study: (i) how the sizes and the size distribution of self-assembled InAs/GaAs quantum dots are affected by the growth technique (MBE or ALMBE), (ii) how the growth parameters and procedures of the upper confining layers influence the interaction between the dots and the caps, and then, the dot dimensions and the photoluminescence (PL) emission energies and (iii) which are the design and preparation parameters that give efficient PL at 300 K in the spectral windows of photonic interest.

2. Experimental details

The InAs/GaAs dot structures were grown by MBE and ALMBE under different conditions and with different InAs coverages \(\theta\); they consist: (i) of GaAs buffer layers, (ii) of InAs dots, and (iii), only in the case of the structures intended for PL studies, of GaAs upper confining layers. The buffer layers were generally grown by MBE at 580°C; then the growth was interrupted for 210 s to lower the growth temperature to that required for the dot deposition. The dots were deposited at 460°C either by MBE or ALMBE. The upper confining layers were grown after interruptions of 210 s by ALMBE (at 360°C or 460°C) or by MBE; in the latter case, 5 ML of GaAs were deposited at 360°C so as to limit In segregation [10] and, then, the effects on dot morphology due to dot interaction with caps; the topmost 10 nm of the caps were grown at temperatures increasing up to 580°C.

As\(x\)/Ga and As\(x\)/In beam equivalent pressure ratios of \(\sim 12\) and \(\sim 45\) were used; the In and Ga fluxes were adjusted for InAs and GaAs growth rates of 0.12 and 2.8 ML/s, respectively; during ALMBE, the growth rate was set at 0.2 ML/cycle and the As supply time was chosen so as to give a sharp \((2 \times 4)\) surface reconstruction at the end of the As cycle. The substrates were radiatively heated at growth temperatures \(T_g\) measured by a suitable optical pyrometer for \(T_g \geq 450°C\) and by a thermocouple (TC) not in direct contact with the substrate for \(T_g < 450°C\). For \(T_g < 450°C\), the TC readings were corrected by the difference \((\sim 200°C)\) between the TC and the optical pyrometer values measured at \(T_g > 450°C\). The coverages were determined by using the growth rate measured by observing the 2D–3D growth transition on a calibration substrate just before the preparation of the dots under investigation and by assuming that the transition takes place at an InAs coverage \(\theta_r = 1.6\) ML. This value is consistent with that deduced from the measurement of the growth rate by means of RHEED oscillations; moreover, it is very close to that \((1.57 \text{ ML})\) determined by very careful experiments described in Ref. [12], under slightly different conditions; it should be recalled, however, that the values of \(\theta_r\) are not very sensitive to growth conditions [5].

The PL spectra were taken in the temperature range 10–300 K with a spectral resolution of 0.5 meV, and were excited by the blue line of an Ar+ laser, using power densities as small as 0.5 W/cm².
Topographic images of uncapped dots were taken by contact-mode Atomic Force Microscopy (AFM) under constant probe force conditions. The measurements were performed using silicon nitride cantilevers and tips with curvature radii of \( \sim 20 \) nm. Then, the images were analyzed in order to derive the mean lateral size and the height distributions of dots. In the analysis, the convolution effects of AFM tips were not taken into account, but they are the same for all the lateral size distributions discussed here.

\section*{3. Results and discussion}

Let us compare first dots grown by ALMBE and by MBE at temperatures of 460°C. In Fig. 1a and Fig. 1b we show the AFM images of dots grown by ALMBE and MBE at 460°C, respectively, with InAs coverages of 3.0 ML. It is very interesting to note that for relatively large coverages the dots grown by MBE tend to coalesce more extensively than the ALMBE ones, and that the uncoalesced dots have larger lateral size when they are grown by ALMBE. Further details can be deduced by the inspection of Fig. 2 where we show the distribution of lateral sizes of dots grown with InAs coverages of 2.4 and 3.0 ML. The main results are that: (i) for both coverages the dots grown by ALMBE have mean lateral sizes larger than that of the MBE counterparts (19 versus 14 nm and 21 versus 16 nm, for \( \theta = 2.4 \) and 3.0 ML, respectively) and (ii) the lateral size distributions are always sharper in the case of ALMBE dots. Moreover, also the height distribution (not shown here) are peaked at larger values of \( \theta \) when ALMBE dots are considered (4.2 versus 2.8 nm and 5.0 versus 3.8 nm, for \( \theta = 2.4 \) and 3.0 ML, respectively). The coalescence of MBE dots for \( \theta = 3.0 \) ML can be also deduced from the analysis of Fig. 2 that shows that MBE dots have a lateral size distribution with a tail and a possible small peak at large lateral sizes (\( \sim 50 \) nm); these features are completely absent in the case of ALMBE dots. The data relative to structures with different coverages (not presented here) show that the coalescence of dots begins at coverages of \( \sim 2.4 \) ML and of \( \sim 3.6 \) ML, for MBE and ALMBE, respectively. The value relative to MBE is close to that

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{Fig_1.png}
\caption{AFM micrograph of InAs/GaAs dots grown by ALMBE (panel a) and by MBE (panel b) at 460°C. The InAs coverage is 3.0 ML. The dots have been deposited on a MBE GaAs layer grown at 580°C, after a 210 s interruption. The ALMBE dots have been grown in 15 ALMBE cycles. The image has been obtained by a tip with a 20 nm radius.}
\end{figure}
(2.18 ML) very recently measured by Madhukar and coworkers [12]. For ALMBE dots and $\theta \sim 3.0$ ML, a distribution of tiny dots (with mean lateral size of $\sim 4$ nm) shows up, which is absent at lower coverages and in MBE structures.

The size features discussed so far have implications also on the PL of dot structures; by comparing different sets of samples grown by both techniques at 460°C and capped with GaAs layers grown by ALMBE at 360°C, we observe that the photoluminescence (PL) of ALMBE dots have (Fig. 3): (i) energies smaller and (ii) FWHMs narrower than those of the corresponding MBE ones; these features are consistent with the larger lateral sizes and heights (at the same ML coverages) of ALMBE dots and with their sharper size distributions. This observation is particularly interesting since it suggests that ALMBE is a viable technique for the growth of relatively large dots, suitable for the fabrication of photonic devices.

Let us now turn our attention to the effects of growth conditions of the upper confining layers on the PL properties of self-assembled dots. Therefore, we compare first structures with dots grown by MBE at 460°C with coverages of 2.0 ML and with caps prepared either by MBE (at 360°C for 5 ML and then at 580°C) or by ALMBE at 360°C. While in the former case the PL emission is at $\sim 1.26$ eV at 10 K, the ALMBE capped structures have emissions at energies lower by $\sim 0.1$ eV. This observation suggests that growing caps by MBE results in a substantial modification of dots, even if we grew the first 5 ML of the GaAs cap at sufficiently low temperatures to reduce the effects of In segregation [10], if the low-temperature grown layer was thick enough to cover the dots. On the other hand, the PL FWHMs of ALMBE capped structures are somewhat larger ($\sim 70$ meV) than that of the MBE counterparts ($\sim 50$ meV), thus suggesting that the conditions following the growth of dots affect their size uniformity.

Then, we consider the effect of the growth temperature (360°C versus 460°C) of ALMBE caps on ALMBE dots deposited at 460°C with InAs coverages of 2.4 ML. The results show that the emission energies are 1.04 and 1.25 eV for caps deposited at 360°C and 460°C, respectively, thus suggesting that the ALMBE growth of caps at relatively high
4. Conclusions

We have studied the effect of different growth schemes of structures with InAs/GaAs self-assembled dots on: (i) the size of dots and (ii) their PL features. The results show that ALMBE is very effective in growing relatively large dots at relatively large InAs coverages without incurring dot coalescence, that may result in the formation of misfit dislocations. Moreover, we have shown that the growth of the upper confining layer by ALMBE at low temperatures (360°C) gives rise to a much lesser interaction between dots and caps (and, then, in a lesser decrease of the dot effective size) than the other investigated schemes, such as the growth by ALMBE (at 460°C) or by MBE (5 ML at 360°C and 10 nm at 580°C). The larger dot lateral size and the lower reduction of dot size due to cap–dot interaction results in dot structures emitting at lower energies and, then, more suited for photonic applications. For instance, structures with MBE dots/MBE caps, MBE dots/ALMBE caps and ALMBE dots/ALMBE caps and with InAs coverages of 3 ML have 10 K emissions at 1.01, 1.08 and 1.17 μm, respectively. The last structure has an intense 300 K emission at 1.26 μm. Our results show that the advantage of ALMBE in growing dot structures is twofold since the enhanced cation surface migration (i) results in larger dots with sharper size distributions since it allows the selective enlargement of existing dots, instead of the continuous nucleation of new small islands, and (ii) allows the growth at lower temperatures where the cap–dot interaction that reduces the effective dimensions of the dots and modify their shape is substantially decreased.

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Formation of self-organized In_{0.5}Ga_{0.5}As quantum dots on GaAs by molecular beam epitaxy

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Abstract

Self-organized In_{0.5}Ga_{0.5}As quantum dots have been successfully grown on vicinal GaAs substrates by molecular beam epitaxy. The density of the quantum dots can be changed by nucleating the dots under different As overpressure. Substrate tilt angle of 15° results in much larger dot size and density than that of 4° due to the closely spaced step edges on the surface. Through investigations of the dots grown on In_{0.1}Ga_{0.9}As buffer, the strain energy of the buffer layer is also found to be an important factor that affects the size and density of the quantum dots.

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Keywords: Quantum dots; Step-bunching; Molecular beam epitaxy

1. Introduction

Semiconductor quantum dots is an area of great interest for low-dimensional quantum physics and devices. Since ex situ processes such as lithography, etching, and regrowth cannot avoid the surface effects, process-induced damages, and rough heterostructure interface, which seriously deteriorate the properties of the quantum dot structures, the formation of self-organized quantum dots during epitaxial growth becomes one of the key techniques for the realization of high-quality quantum devices. This technique basically relies on the lattice strain and/or the bunched steps to induce coherent island growth at the edge of the terraces. Both molecular beam epitaxy (MBE) [1, 2] and metalorganic vapor-phase epitaxy (MOVPE) [3, 4] have been successfully used to demonstrate the strain-induced self-organization. Since the quantum dots formed by this method do not suffer from the adverse effects mentioned above, a number of InGaAs and InAs quantum dot lasers have been realized [5, 6]. Nevertheless, further efforts have to be made to lead to the performance as expected. The control of size, density, and arrangement of the quantum dot

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is one of the essential tasks for this purpose. In this work, we have investigated the effects of several growth parameters, such as As overpressure, substrate tilt angle, strained buffer layer, on the formation of In_{0.5}Ga_{0.5}As self-organized quantum dots on GaAs by molecular beam epitaxy. The factors that influence the nucleation of InGaAs quantum dots on GaAs are discussed.

2. Experimental procedure

The samples investigated were grown by solid-source molecular beam epitaxy. Vincinal GaAs substrates (4° and 15° towards (1 1 1)A) were used in this work for comparison. The surface reconstruction during growth was monitored by reflection high-energy electron diffraction (RHEED). After oxide desorption, a 200 nm thick GaAs buffer layer was grown in the step-flow regime, i.e. low V/III flux ratio at 580°C, which led to the (3 × 1) surface reconstruction. In order to enhance the step-bunching effect, which facilitates the formation of self-organized quantum dots, the GaAs buffer was annealed at 580°C for 30 min before the substrate temperature was lowered to 520°C for the deposition of InGaAs quantum dots. The resultant step edges, running parallel to the [1 1 0] direction, were found to be uniformly distributed over a large area as shown in Fig. 1. When the substrate temperature was reduced to 520°C, the RHEED pattern changed from (3 × 1) to (2 × 4). The In_{0.5}Ga_{0.5}As quantum dots were then grown with interruptions corresponding to 0.5 monolayer (ML) of In_{0.5}Ga_{0.5}As and 5 s of As$_4$ irradiation for each period. The growth was stopped when the RHEED pattern transformed from a streaky pattern to a spotty pattern, indicating the presence of three-dimensional islands on the growing surface. These quantum dots were formed after the deposition of 5 ML of In_{0.5}Ga_{0.5}As at a growth rate of 0.5 ML/s. It was found that a thicker layer was needed to form the quantum dots if lower growth rate was used. The In_{0.5}Ga_{0.5}As quantum dots were investigated using atomic force microscopy (AFM). The AFM images were taken in the contact mode with a scan rate of typically 1 Hz.

Fig. 1. Atomic force micrograph of the bunched steps formed on 4°-off (1 0 0) GaAs substrate.

3. Results and discussion

Images taken by AFM show that the self-organized In_{0.5}Ga_{0.5}As quantum dots are formed spontaneously at the edge of the terraces. Shown in Fig. 2a is the image of the quantum dots formed on the 4°-off substrate at 520°C. The central blurry area is caused by successive measurements. The size of these well-organized InGaAs quantum dots is about 25 nm in diameter and 3 nm in height. A lower density of quantum dots, as shown in Fig. 2b, can be obtained by commencing the growth under an As-free condition, i.e. when the As shutter remains closed during the annealing and the cooling steps. The dot density decreased from about 3.6 × 10$^{10}$ to 1.2 × 10$^{10}$ cm$^{-2}$, while the size remains about the same. This is attributed to the longer diffusion length at the growing surface for the group III adatoms under low As background conditions. The effect of the diffusion length was also examined by growing the quantum dots using alternating (In, Ga) and As$_4$ beams. This growth scheme is supposed to enhance the migration of the group III adatoms on the growing surface, and leads to two-dimensional growth. No quantum dots, except for the bunched steps, were observed when the total thickness of the InGaAs was about
Fig. 2. Atomic force micrograph of the MBE grown InGaAs quantum dots on 4°-off (1 0 0) GaAs substrate with (a) As shutter open and (b) As shutter closed before the growth.

7.5 ML. It can be concluded that more deposition is needed to form quantum dots in this growth scheme because the wetting layer is thicker in this case.

However, the dots grown on the 15°-off substrate exhibit significantly different characteristics as compared with the dots on the 4°-off substrate.

Fig. 3. Atomic force micrograph of the MBE grown InGaAs quantum dots on 15°-off (1 0 0) GaAs substrate.

Fig. 3 shows the AFM images of the quantum dots formed on these two substrates side-by-side after the deposition of 3.5 ML of InGaAs at a growth rate of 0.7 ML/s. The size of the dots on the 15°-off substrate is about 190 nm, which is much larger than that of the dots on the 4°-off substrate. It is believed that the smaller distance between the terrace edges on the 15°-off substrate limits the diffusion of the adatoms during the growth and a large number of nucleation sites was provided for the formation of three-dimensional islands. Therefore, the dots thus formed have larger size due to their energetically favorable surface configuration. Since their size is larger than the terrace, the dots overlap with each other and lose their regularity.

We have also studied the effects of surface strain on the formation of quantum dot since the nucleation of quantum dot is closely related to the surface energy which may be modified by lattice strain. Fig. 4 shows the AFM images of the In₀.₅Ga₀.₅As quantum dots formed on a 15 nm thick pseudomorphic In₀.₁Ga₀.₉As buffer and a 0.2-μm-thick strain-relaxed In₀.₁Ga₀.₉As buffer, respectively. These dots were formed by depositing 5 ML of In₀.₅Ga₀.₅As at 520°C. Obviously, the pseudomorphic In₀.₁Ga₀.₉As buffer results in higher dot density, even higher than that for the dots grown on GaAs buffer. This is attributed to the higher
strain energy stored in the pseudomorphic layer, which facilitates three-dimensional growth. This In$_{0.1}$Ga$_{0.9}$As pseudomorphic layer plays the same role as the wetting layer in the Stranski-Krastanov growth mode often observed in the growth of (In, Ga)As quantum dots on GaAs. Using such a strained buffer layer, quantum dots of smaller size and high density can be obtained as compared to the case of direct growth on GaAs. This might be beneficial for the realization of low-threshold quantum dot lasers since both quantum confinement and gain region are increased, although the dots do not seem to be well-organized. On the other hand, the density for the dots grown on the relaxed buffer is slightly lower than that for the direct growth while the dot size is about the same. It implies that the lattice mismatch between In$_{0.1}$Ga$_{0.9}$As and In$_{0.5}$Ga$_{0.5}$As is the dominant factor in the dot nucleation process. More interestingly, it was found that the dot density on the relaxed buffer is modulated by the cross-hatching resultant from strain relaxation. As shown in Fig. 5, higher dot density is observed at the top region of the hatch. This is closely related to the lower surface energy at the top of the hatch [7]. Since most of the adatoms are driven towards the more energetically favorable region, the nucleus of quantum dots, the dot density is enhanced at this region. Similar behavior has also been observed for the growth of vertically stacked InAs quantum dots on GaAs where the dots beneath provide lower surface energy for the successive nucleation of InAs dots [8]. Therefore, perfect vertically aligned quantum dot structure is spontaneously formed.

Fig. 4. Atomic force micrograph of the MBE grown InGaAs quantum dots on (a) 15 nm-thick pseudomorphic In$_{0.1}$Ga$_{0.9}$As and (b) 0.2 μm-thick strain-relaxed In$_{0.1}$Ga$_{0.9}$As buffer layers.

Fig. 5. Atomic force micrograph of the MBE grown InGaAs quantum dots on 0.2 μm-thick strain-relaxed In$_{0.1}$Ga$_{0.9}$As buffer layers showing the modulation of dot density by the cross-hatch patterns.
4. Conclusion

Self-organized In$_{0.5}$Ga$_{0.5}$As quantum dots have been grown on vicinal (1 0 0) GaAs by molecular beam epitaxy. Parameters that affect the configuration of the quantum dots, such as substrate tilt angle, as background pressure, and the strain in the buffer layer, have been investigated. The associated surface strain energy and free energy for the adatoms are the key factors in the formation of the quantum dots. The size and density of the quantum dots can be varied accordingly for various applications.

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MBE growth and magnetotunnelling transport properties of a single GaAs/AlAs/GaAs barrier incorporating InAs quantum dots


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Abstract

We describe the growth and investigation of a novel type of structure in which self-organised InAs quantum dots are incorporated in the AlAs tunnelling barrier of an n-i-n single-barrier GaAs/AlAs/GaAs heterostructure. The low-temperature current–voltage curves exhibit a series of pronounced peaks which are absent in a control sample grown without InAs in the barrier. By studying their behaviour in a magnetic field B, we attribute these peaks to single-electron tunnelling through single discrete zero-dimensional states of individual InAs dots in the barrier.

1. Introduction

The fabrication of low-dimensional structures, such as quantum dots or wires, has been intensively pursued in recent years in order to realize novel devices using confinement effects. One of the promising fabrication methods is to use self-organised three-dimensional structures which are formed during the initial stage of heteroepitaxy growth in lattice-mismatched systems, called the Stranski–Krasnotov (SK) growth mode. In the SK growth mode the mismatched epitaxy is initially accommodated by biaxial compression in a layer-by-layer (2D) growth region, traditionally called the wetting layer. After deposition of a few monolayers the strain energy increases and the development of islands (3D) becomes more favourable than planar growth. In the III–V semiconductor material system, SK growth has been used to grow InAs islands on GaAs and it has been shown that the size fluctuation of dots is relatively small (≤10%) and the small dots and surrounding host matrix are dislocation-free and strained coherently with GaAs. It has been reported that the InAs growth mode changes from 2D to 3D upon the deposition of less than 2 ML of InAs. To date, studies have been limited to the optical properties of self-organised InAs quantum dots on GaAs.

In this paper we describe the growth of a novel type of structure in which the dots are incorporated in the AlAs tunnelling barrier of an n-i-n single
2. Experimental procedure

The epitaxial layers were deposited by molecular beam epitaxy (MBE) using a Varian Gen II system on 2 in (1 0 0) n⁺-GaAs substrate oriented within ± 0.5°. The preparation of the MBE machine followed the method adopted previously to give optimum quality of material [1]. Prior to loading, the substrates were cleaned using a 5 min ultrasonic degrease in Summa-Clean followed by a 5 h soak in Summa-Develop. Both solutions are commercially available choline etches [2]. The wafers were rinsed in deionised water and spun dry before being mounted onto Mo blocks with In. The substrates were then heated in the growth chamber at 600°C for 30 min in order to desorb the thin oxide film on the substrates.

The sample consisted of the following layers, in order of growth from the substrate: (i) a 1 μm thick buffer layer of GaAs doped at $n = 4 \times 10^{18}$ cm$^{-3}$; (ii) 100 nm of GaAs, $n = 2 \times 10^{16}$ cm$^{-3}$; (iii) a 100 nm thick spacer layer of undoped GaAs; (iv) an undoped AlAs barrier of thickness 5.1 nm; (v) 1.8 monolayers of InAs; (vi) an undoped AlAs barrier of thickness 5.1 nm; (vii) a 100 nm thick spacer layer of undoped GaAs; (viii) 100 nm of GaAs, $n = 2 \times 10^{16}$ cm$^{-3}$; and (ix) a 0.5 μm top contact layer of GaAs, $n = 4 \times 10^{18}$ cm$^{-3}$. The growth temperature was 550°C as monitored by a pyrometer, except during the growth of InAs when the temperature was lowered to 520°C. The growth rates are one monolayer/s (ML/s) for GaAs, 0.5 ML/s for AlAs and 0.066 ML/s for InAs. The formation of InAs islands was monitored by RHEED, and the average thickness of InAs deposited is 1.8 ML. A control sample without the InAs layer was also grown. The size and density of InAs quantum dots were estimated by SEM and STM from samples of the same design but with the growth terminated after depositing the InAs layers. The STM samples were capped with a thick arsenic layer before they were transferred to a separate UHV STM chamber.

During growth the GaAs surfaces showed a (2 × 4) reconstruction. The samples were rotated during growth to improve uniformity. After growth, the epitaxial surfaces were examined, using a Nomarski phase-contrast optical microscope and were found to be mirror smooth and nearly defect-free.

The graded doping in the contact layers and the undoped spacer layers adjacent to the barrier are thought to improve the performance of the devices by reducing dopant diffusion into the region close to the barrier during growth, thus reducing the number of bound impurity scatterers.

The epitaxial layers were processed into circular mesas of various diameters ranging from 30 to 400 μm by means of photolithography and wet-etching techniques. Au and AuGe were evaporated and alloyed to form ohmic contacts.

A schematic band diagram of the device under an applied bias $V$ is shown in Fig. 1. Because of the low doping in the regions close to the barrier, an applied bias produces a 2DEG in an accumulation layer adjacent to the emitter barrier.

3. Results and discussion

The current–voltage characteristics $I(V)$, recorded for a 100 μm diameter mesa in the absence of magnetic field, are shown in Fig. 2. Forward and reverse bias correspond to electron flow from and to the substrate, respectively. An $I(V)$ curve for the
control sample is also shown for comparison. Both devices have a very high impedance (∼10¹² Ω) around zero bias and exhibit a monotonically increasing background current. In addition, pronounced, low-current (a few pA) peaks, superimposed on the background current, are observed for the InAs quantum dot device for forward bias above 100 mV at 4.2 K. The peaks are absent in reverse bias at 4.2 K, where there is only indistinct structure. The asymmetry in the \( I(V) \) characteristics in forward and reverse bias as shown in Fig. 2 results from the asymmetry of dot positions in the AlAs layer. The dots are grown on the centre plane of the barrier, but the AlAs layer covering the dots is effectively thinner due to the size and shape of the dots, and thus has a higher tunnelling probability than the AlAs layer on the substrate side. Thus, for reverse bias the tunnelling rate out of the dots is much smaller than the rate of tunnelling into the dots from the accumulation layer. On lowering the temperature to 0.4 K, the structure in reverse bias evolves into a set of distinct steps. The \( I(V) \) curve of the control sample has no structure in either bias direction but the background current is of similar magnitude.

The peaks in \( I(V) \) arise from resonant tunnelling through states in the barrier, and our observations indicate that these states are associated with the incorporation of InAs in the barrier. We have also examined the effect of magnetic field on the tunnelling current in forward bias in order to confirm that the discrete (zero-dimensional) electron states of InAs quantum dots in the barrier are the states through which the resonant tunnelling occurs. For \( B \) applied parallel to the current, the \( I(V) \) curves change qualitatively as shown in Fig. 2. At fields as low as 0.4 T a series of narrow peaks arises in the curves. The peaks diverge in bias and their number falls with increasing \( B \) up to 3–4 T. Increasing \( B \) from 4 to 12 T causes the peaks to shift to lower bias with little change in shape.

Fig. 3a shows examples of \( I(B) \) at constant bias \( V_0 \). If \( V_0 \) is equal or close to the bias at which a peak occurs, there are pronounced oscillations in \( I(B) \). Their maxima and minima shift to smaller \( B \) with increasing \( V_0 \). The \( I(B) \) curves exhibit no structure at \( V_0 \) just below or above a peak in \( I(V) \).

The energy spectrum of the 2DEG in the emitter-accumulation layer splits into a set of discrete Landau levels for \( B \) normal to the sample plane. Therefore, we attribute the sharp peaks in \( I(V) \) to the Landau quantisation of the 2DEG in the emitter which is consistent both with the peak divergence with increasing \( B \) and with their shift to lower \( V \). As to the oscillations in \( I(B) \), these behave quite differently from magneto-oscillations earlier reported in single-barrier tunnelling devices [3], for
which the maxima should shift to higher $B$ with increasing $V_o$. In our case the maxima in $I(B)$ occur when the magnetic field brings an occupied Landau level in the 2DEG in resonance with the energy levels in the barrier. In effect, both $I(V)$ and $I(B)$ probe the density of states (DOS) of the 2DEG in the emitter accumulation layer.

To estimate the size of the resonant states in the barrier, we have used the diamagnetic shift of peaks in $I(V)$ in magnetic fields applied normal to the current, i.e. parallel to the barrier plane. Applying $B \parallel I$ up to 11 T causes no qualitative change in the $I(V)$ characteristics at 4.2 K. Their evolution is shown in Fig. 4a. The peak current values fall with increasing $B$ for the lower-voltage peaks. The peak positions shift to lower voltage quadratically in $B$, as shown in Fig. 4b.

The variation in peak voltage is determined by the relative diamagnetic shifts of the electron state in the 2DEG and the resonant state in the barrier. Within the uncertainty in the values of parameters used, we estimate the size of the resonant barrier state to be $\approx (10 \pm 5)$ nm. This value is fully consistent with the size of InAs quantum dots measured by SEM and STM on samples of the same design but with the growth terminated after depositing the InAs layers. The samples for STM measurements were arsenic-capped in the growth chamber before being transferred to a separate STM chamber where arsenic desorption and imaging took place. From SEM and STM imaging we estimated the density of dots as $\approx 2 \times 10^{11}$ cm$^{-2}$, with a dot size $\approx (10 \times 10)$ nm$^2$. We therefore conclude that the peaks in $I(V)$ originate from the tunnelling through the electron states of the self-organised InAs dots incorporated in the barrier.

The observed decrease of peak amplitude with increasing $B \perp I$ in Fig. 4a is qualitatively similar to that reported recently for magnetotunnelling into donor states [4]. As will be discussed in another paper, this provides a separate estimate for the spatial extent of the dot wave function which is consistent with the above value.

A PL spectrum of our tunnel structure, recorded with Ge detector at He–Ne laser excitation ($\lambda = 6328$ Å) is shown in the inset of Fig. 1. The spectrum exhibits a broad line with a maximum few hundred meV below the GaAs band-gap energy.

![Image](https://via.placeholder.com/150)

Fig. 4. (a) $I(V)$ characteristics at various $B \parallel I$. Curves are offset. (b) Diamagnetic shift of lower voltage peaks in $I(V)$. Solid lines are parabolic fits.

The line corresponds to the emission from the dot ensemble and is similar to that reported by other groups [5–7].

We now discuss why we are able to resolve tunnelling through only a few single dots rather than a broad tunnelling peak due to the ensemble of dots, about $10^7$ dots in a typical mesa, in analogy with the broad lines observed in PL. To understand this, note that the PL spectrum from the sample (Fig. 1) suggests that for the majority of dots the electron ground energy level is below the conduction band edge $E_c$, in agreement with capacitance spectroscopy studies [8]. These levels are unavailable for energy-conserving tunnelling processes. We observe tunnelling through extremal dots with electron level energies above $E_c$. Such dots can arise
due to fluctuations in size (30–40%), shape and strain or possible Al alloying of dots. Our picture implies that a fraction of dots should be charged at zero bias, in order to align the chemical potentials of the dot ensemble and the collector and emitter n-doped contact layers. The accumulation of the 2DEG is followed by dot discharge when the device is biased, which contributes strongly to the leverage factor dependence on voltage.

4. Conclusion

We have fabricated novel resonant tunnelling devices with buried InAs quantum dots and demonstrated resonant tunnelling through single zero-dimensional states of individual InAs dots in the barrier. The MBE technique coupled with SK growth provides us with a novel means of producing 0D tunnel structures in which we can freely vary the tunnel barrier heights and widths, and the alloy composition of the dots. It should also be possible to produce stacked dot structures [9] and dot structures in different index substrate planes.

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[2] Summa-clean and Summa-develop are brand name products of Mallinckrodt Inc. (Science Products Division).
Selective MBE growth of GaAs wire and dot structures using atomic hydrogens and their electronic properties

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Abstract

The selective growth of GaAs wire and dot structures by molecular beam epitaxy in the presence of atomic hydrogen was studied on a GaAs substrate with stripe and window shaped SiN\textsubscript{x} mask patterns. It was found that 20 nm ridges were successfully formed on long stripe patterns, but the ridge width increased substantially when grown on short stripes of less than several \textmu{}m. This is ascribed to the longitudinal migration of Ga flowing from the two ends of a stripe to its central zone. The incorporation of Si dopants into a 0.3 \textmu{}m-wide wire was achieved. Negative magnetoresistance and conductance fluctuations indicative of the one-dimensional nature of electrons were observed. On a 2 \times 2 \textmu{}m square window pattern, a pyramidal structure with two-fold symmetry was formed, on which a GaAs dot with the lateral dimension of 100 nm was grown. Photoluminescence (PL) spectra from this dot exhibited a shoulder structure 15 meV, suggesting the filling of several discrete levels and the slow relaxation of carriers.

1. Introduction

The fabrication of 10–100 nm scale quantum wires (QWIs) and quantum dots (QDs) is important not only for device applications but also for the physics of low dimensional systems. Among various methods, selective area growth of nanostructures using SiN\textsubscript{x} or SiO\textsubscript{2} mask patterns is attractive. Indeed, both metalorganic chemical vapour deposition [1–6] and molecular beam epitaxy (MBE) with the assistance of atomic hydrogen [7] have been used to produce nearly damage-free nanostructures.

The characterization of such selectively grown QWIs has been mostly done by optical methods since transport characterization requires the formation of electrical contacts and the introduction of dopants. Since various facets appear in the selective growth due to the migration of atoms from one plane to another, the shape of nanostructures is strongly influenced by boundary conditions of crystal growth including the presence of end zones for electrical contacts. It is thus indispensable to clarify these facets interaction to form quantum structures suitable for transport studies.
In this work we investigate structural features and electronic properties of n-GaAs wires and dots prepared by the selective MBE growth using thermally cracked atomic hydrogen [8]. Specifically, we discuss the effect of longitudinal diffusion of Ga atoms in short stripe patterns bound by two wide patterns for electrical contacts and also on a GaAs substrate with window patterns.

2. Sample preparation

Patterns were made into a 50 nm thick SiNₓ film deposited on a (1 0 0) semi-insulating GaAs substrate by using electron-beam lithography and wet chemical etching. Fig. 1a and Fig. 1b illustrate patterns for wires and dots, respectively. In Fig. 1a, stripe window is parallel to ⟨0 1 1⟩ and is connected to 40 µm wide windows on both ends. In Fig. 1b, a 2 × 2 µm square window pattern is aligned along ⟨0 1 0⟩ and ⟨0 0 1⟩.

A GaAs substrate with these patterns was first cleaned in MBE chamber by exposing atomic hydrogen for 30 min at the substrate temperature Tₛ of 350°C [8]. Then, a 10 nm GaAs was deposited to smoothen the surface at Tₛ of 590–600°C. We raised Tₛ by 10–20°C and then deposited a 900 nm thick GaAs buffer layer for wire structure or a 300 nm buffer for dot structure. By this time, relatively sharp ridge structures were formed, as schematically shown in Fig. 2. Then, 10 periods of a superlattice (SL) buffer layer were grown consisting of a 5 nm AlGaAs and a 1.5 nm GaAs with 10 s of growth interruption (GI). At this point a 6 nm thick GaAs QW structure was formed which had a 5 nm AlGaAs barrier and 7 nm AlGaAs barrier on the bottom side and top side of the QW, respectively. A 60 nm Si-doped AlGaAs layer was further deposited with the donor density of 5.5 × 10¹⁷ cm⁻³ to supply electrons and the growth was completed by a 10 nm thick GaAs capping layer. During the MBE growth, thermally cracked atomic hydrogens were supplied [9] with the sample heating block rotating at the rate of 15 rpm. The cracking cell was set at about 1600°C and the hydrogen flux was 0.90 cm³. The growth rate was 0.24 µm/h for GaAs and 0.10 µm/h for AlAs. The flux ratio Asₓ/Ga was 5–6. The mole fraction of Al in AlGaAs was 0.29. In this growth condition, Ga atoms impinging onto SiNₓ masks desorbed efficiently with the assistance of atomic hydrogen. However, Al containing materials such as AlGaAs are hard to desorb and, therefore, form polycrystals on masks, as shown in Fig. 2.

3. Fabrication of GaAs wire structures

Fig. 2a–Fig. 2c show SEM images of a sample grown on a 1.6 µm-wide stripe pattern of Fig. 1a. Note that the growth took place selectively in the window region although a little amount of polycrystalline AlGaAs was deposited on SiNₓ mask regions. The central portion of the ridge wire

![Diagram](attachment:image.png)

(a) Wire pattern

(b) Dot pattern

Fig. 1. Schematic illustrations of SiNₓ mask patterns for the fabrication of wire (a) and dot (b) structures. A stripe pattern in the wire pattern is parallel to ⟨0 1 1⟩. The dimensions of the stripe window are 0.8–1.6 µm in width and 1–6 µm in length. A square pattern in the dot pattern is aligned along ⟨0 1 0⟩ and ⟨0 0 1⟩ with a size of 2 µm.
meets with the 40-μm wide lead regions, more complicated facets were found. In the case of long stripes 20 nm scale ridges were successfully formed since the ridge tends to sharpen due to the up-flow of Ga from the two (1 1 1)B side facets to the top (1 0 0) facet [10]. In short stripes, however, the self-narrowing process was found to be suppressed. This is because the diffusion of Ga is strongly modified by the presence of wide regions on both ends of stripes.

We then examined how the final ridge width depends on the stripe length. The ridge width in the central part of wires was measured and shown as a function of L in Fig. 3. Here the original width W of stripes was 1.6 μm. The width was found to widen exponentially as L decreases, indicating that the diffusion of Ga along the stripe direction plays an important role in determining the ridge shape. The ridge width decreases by a factor of 1/e every time L increases by 3 μm. Note that this figure is close to the reported diffusion length on (1 1 1)B plane [10]. We found also that the amount of GaAs incorporated on ridge regions was somewhat greater than the amount deposited, indicating the in-flow of materials. Hence, the dominant process which prevents the sharpening of ridges is the longitudinal flow of materials from the stripe ends to the stripe center.

Next, we studied transport properties of a series of 3 μm long selectively doped GaAs wires at 1.6 K. In Fig. 4, the magnetoconductance of four wire samples A–D are shown for the width W, of wire channel ranging from 0.3 to 1.1 μm. The original stripe width W is also indicated in the figures. For comparison, the data on a 40 μm-wide ordinary two-dimensional electron gas (2DEG) sample E are also shown. Note that the zero-field resistance depends on the wire width. In our samples, electrons accumulate mainly on the top (1 0 0) plane, since the QW on the (1 1 1) facet is too thin to accommodate electrons. Very strong Shubnikov–de Haas (SdH) oscillations seen in Fig. 4 for B > 1 T in all samples originate from the 2DEG flowing in the 40 μm wide lead regions. In contrast, the resistance of samples for B < 1 T depend on the stripe widths W and display distinct negative magnetoresistances (NMR) especially in sample A. This NMR is indicative of a weak localization effect in quasi-1D...
system. In sample A, random oscillatory fluctuations on NMR were reproducibly observed. This may be related to a kind of universal conductance fluctuation (UCF), although its amplitude is far less than that expected from theory. The presence of junctions [11] and/or the geometrical inhomogeneity of wires may reduce the interference of wavefunctions and UCF, though more study is necessary to conclude this.

Next we discuss the quality of sample A. The mobility $\mu$ and the electron concentration $N_e$ of 2DEGs of a 6 nm thick GaAs QW grown simultaneously on a flat substrate were studied by Hall effect and found to be $4.7 \times 10^{11}$ cm$^{-2}$ and $3.4 \times 10^6$cm$^2$/V s at 4.2 K. These values are close to those derived from the SdH oscillations of Fig. 4. On the basis of these values, the resistance of sample E is calculated to be 6 kΩ as the 40 μm-wide 2DEG channel stretches about 250 μm to both ends. This value is close to the zero-field resistance of sample E.

Since the zero-field resistance of sample A is 15.3 kΩ, the resistance of the wire region is estimated to be about 8.5 kΩ. If we assume $N_e$ in the wire region to be comparable to that of 2DEG and take account of the wire geometry (3 μm in length and 0.3 μm in width), $\mu$ is estimated to be about $2 \times 10^4$ cm$^2$/V s. This value is almost half of 2DEG, probably due to the presence of inhomogeneities of the wire geometry. We expect that the quality of wires can be further improved by optimizing the growth conditions.

4. Fabrication of GaAs dot structures

By using a window pattern of Fig. 1b, we grew a pyramidal facet structure and then a crescent shaped GaAs dot on its top. Fig. 5a shows an SEM image taken from the top, and Fig. 5b is the cross-sectional view of the structure seen after cutting along the line A of Fig. 1b. Note that the symmetry of a pyramidal dot structure is not fourfold but twofold although a pyramidal structure tends to exhibit fourfold symmetry as the growth proceeds. In this pyramidal structure, facet planes with indices higher than (1 1 0) are formed. One can also see that crescent shaped GaAs dots with 100 nm width and 30 nm thickness, sandwiched by AlGaAs, are successfully formed.

PL spectra of GaAs dots at 15 K are shown in Fig. 6 at various excitation powers. The main peak
5. Conclusions

In conclusion, we have fabricated GaAs wire and dot structures by MBE with assistance of atomic hydrogen. We have found that the interfacet interaction is modified by the presence of leads and that the longitudinal diffusion suppresses the sharpening process of ridges particularly in short stripe patterns.

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References


Molecular beam epitaxial growth of InGaAs/InGaAsP quantum wires on V-grooved InP substrates with (1 1 1) sidewalls

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Abstract

Bulk InGaAsP and InGaAs/InGaAsP quantum well structures have been grown by gas source molecular beam epitaxy on V-grooved InP substrates having (1 1 1)A and (1 1 1)B sidewalls. The growth of the InGaAsP layer in a (1 1 1)A V-groove results in a flat and wide bottom, which rules out the possibility of the formation of InGaAs/InGaAsP quantum wires (QWRs). However, the growth of the InGaAsP layer in (1 1 1)B V-grooves results in a sharper bottom so that crescent-shaped InGaAs/InGaAsP quantum wells are formed such that QWR behaviour is possible. Transmission electron microscopy shows that the crescent-shaped InGaAs layer has a very large thickness variation at the V-groove bottom and that all layers at the V-groove bottom are free of extended defects. The photoluminescence from different spatial regions of the sample has been identified using a selective etching technique. Two peaks are associated with the QWRs at V-groove bottom, with the higher energy one apparently being from the second lateral subband transition. The energy separation between these two peaks is consistent with values calculated from the measured thickness variation. Consistent with QWR behaviour, the photoluminescence is polarized, with a stronger signal along the wire direction than perpendicular to the wire direction.

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Keywords: Quantum wire; V-groove; MBE; InGaAsP; InGaAs; Polarization

1. Introduction

Growth onto V-grooved substrates has been demonstrated to be one of most promising ways, among various techniques, to make quantum wires (QWR) [1]. QWR lasers with very low threshold currents have been grown onto V-grooved GaAs substrates [1, 2]. However, much less progress has been made on InP based-materials, even though they are very important to telecommunication applications. To our knowledge, neither QWR laser structures nor InGaAs/InGaAsP QWRs have been successfully grown on V-grooved InP substrates [3, 5]. One difficulty arises from a sidewall...
diffusive flux and increased incorporation of group III adatoms in the V-groove bottoms. Consequently, a planarization develops as the growth of buffer and barrier layers proceeds, making QWR formation impossible. A second difficulty arises due to the differential migration of indium over gallium, resulting in strain development and possible relaxation generated defects [1,3–6]. Nevertheless, InGaAs/InP quantum wire formation was recently reported on V-grooves with (1 1 1)A sidewalls [7]. For a laser structure, however, InGaAsP barrier or waveguiding layers with sufficient thicknesses have to be grown. Such InGaAsP layers need to be defect-free and V-groove bottoms must be sharp before the growth of the quantum well layer [3,4]. The subsequent quantum wire layer should form a crescent-shape profile and also be free of defects. In this work, using gas source molecular beam epitaxy (GSMBE) we have grown quaternary and quantum well structures on V-grooves with both (1 1 1)A and (1 1 1)B sidewalls, which are expected to show different growth behavior. Transmission electron microscopy (TEM) has been used to examine layer morphology and to determine the dimensions of the crescent-shaped InGaAs/InGaAsP quantum wells in (1 1 1)B V-grooves. Based on the measured quantum well thickness variation, a simple quantum wire model has been used to estimate lateral subband separations. Optical characterizations, including spatially resolved photoluminescence (PL) and PL polarization, have also been performed on the InGaAs/InGaAsP quantum wires.

2. Experimental procedure

V-grooves with both (1 1 1)A and (1 1 1)B sidewalls were chemically etched in (1 0 0) InP substrates. V-grooves with (1 1 1)A sidewalls were etched using 2HBr : 2H₃PO₄ : 1K₂Cr₂O₇ as an etchant and SiO₂ as a V-groove etching mask, with patterns of 1μm wide openings aligned along the [0 1 1] direction. V-grooves with (1 1 1)B sidewalls were obtained using an etch solution of 5HCl : 1H₃PO₄, with 1μm wide photoresist mask openings aligned along the [0 1 1] direction [8].

A bulk InGaAsP quaternary layer and an InGaAs/InGaAsP quantum well structure were grown by GSMBE on the V-groove etched substrates. The quaternary layer has a nominal composition of In₀.₇₂Ga₀.₂₈As₀.₆₁P₀.₃₉ (with a bandgap wavelength of 1.3 μm) and a nominal thickness of 40 nm. In the quantum well structure, the InGaAs layer is lattice-matched to InP and nominally 1.2 nm thick; the lower and upper InGaAsP barrier layers, also having a composition of In₀.₇₂Ga₀.₂₈As₀.₆₁P₀.₃₉, are nominally 20 nm thick. All InGaAs and InGaAsP layers were grown at a growth rate of 1 μm/h, a substrate temperature of 450°C and a V/III flux ratio of 2.6. Along with the quaternary and ternary layers, InP layers were also grown, under conditions giving sharp V-groove bottoms and smooth morphology [9], to serve as buffer and capping layers in the above structures. Buffer layers were 50 nm thick, while capping layer thicknesses varied from 50 to 500 nm.

Cross-sectional TEM was used to examine layer morphology and any extended defect structures. Conventional low temperature (15 K), top-emission PL measurements were performed on the quantum well structure samples using a 488 nm Ar⁺ laser with a spot diameter of about 250 μm which typically covered 20 grooves. The sample was selectively etched [7] with the planar (1 0 0) regions and upper parts of sidewalls removed, (i.e. all but the bottom ~1.5 μm of sidewalls was removed). The polarization of photoluminescence originating from quantum well in the V-groove bottom region was also investigated, with the backside of the sample mechanically polished and optically excited by a cw YAG laser. The 1.06 μm YAG laser light passed through the InP substrates and selectively pumped the InGaAsP and InGaAs layers. The spectra were calibrated with respect to the polarization response of the PL system.

3. Results and discussion

Fig. 1 is the TEM image of the InGaAsP layer, sandwiched between InP buffer and capping layers, grown on a (1 1 1)A V-groove. It shows no extended defects present at the V-groove bottom, but a dramatic planarization at the V-bottom region is evident. After the growth of the InP buffer layer, the V-groove bottom is faceted with (3 1 1) planes and
is only 20 nm wide. Subsequently, the top surface becomes flat and expands to 80 nm width after the growth of the InGaAsP layer. This result implies that the growth behaviour of an InGaAsP layer in a (111)A V-groove is an obstacle to the formation of QWRs. This is further demonstrated by Fig. 2, which is the TEM image of a 3-quantum well structure grown on a (111)A V-groove. It is seen that all three InGaAs layers are wide, having no significant thickness variation to produce lateral quantum confinement. Thus, InGaAs/InGaAsP QWRs cannot be obtained in a (111)A V-groove, even though InGaAs/InP QWRs have been achieved in such a type of V-groove [7].

In contrast, the InP and InGaAsP layers grown on a (111)B V-groove show different layer morphology, as shown in Fig. 3. The V-groove bottom is flat and small (about 30 nm wide) after the growth of the InP buffer layer. However, the subsequent InGaAsP quaternary layer evolves during growth such that its top surface in the V-groove bottom is sharper, i.e. it develops into a rounded bottom with a small radius of curvature. This feature indicates that the InGaAsP growth behaviour on a (111)B V-groove is different from that on a (111)A V-groove, in a way which could be used for quantum-wire formation. The causes for the different growth behaviour of InGaAsP on (111)A and (111)B V-grooves and the cause for

**Fig. 1.** The cross-sectional TEM image of InP/InGaAsP/InP structure at the bottom of the (111)A V-groove.

**Fig. 2.** The cross-sectional TEM image of InGaAs/InGaAsP 3 quantum well structure at the bottom of the (111)A V-groove.
the different growth behavior of InGaAsP to that of InP are not currently understood.

Fig. 4 is the TEM image of a InGaAs/InGaAsP quantum well structure grown on a (111)B V-groove. As seen in the image, all ternary and quaternary layers at V-groove bottoms are free of any extended defects. Moreover, the InGaAs layer at the V-groove bottom clearly shows a crescent-shape. The thickness is about 7 nm at the center of V-bottom, which is nearly 6 times the nominal thickness on a planar (1 0 0) region and more than 10 times the estimated thickness on the sidewall, indicating a significant group III adatom migration to the bottom from the sidewalls. The physical width of the crescent-shaped InGaAs region is about 50 nm. A big thickness variation in such a small width is an indication of possible quantum wire formation. Furthermore, even the top of upper InGaAsP layer is sharp, which implies promise for the formation of multiple InGaAs/InGaAsP QWRs.

To estimate the lateral quantum confinement for the InGaAs/InGaAsP QWR, a simplified lateral one-dimensional parabolic potential model was used, which neglects band intermixing and is similar to that used for GaAs/AlGaAs QWRs [1, 10, 11] and InGaAs/InP QWRs [7]. This involves determining the InGaAs layer thickness at different positions from the TEM images and calculation of the associated confinement energies with a conventional quantum well calculation. The confinement energy distributions follow parabolic shapes near the center of the wire, for which the lateral subband separations are calculated to be 17, 5.7 and 222 meV for electrons, heavy holes and light holes respectively. The effective widths, $W_{eff}$, of the wire for electrons, heavy holes and light holes are 22, 13.6 and 20 nm, respectively [1, 10, 11]. These subband separations are slightly higher than those reported on InGaAs/InP QWR grown on (1 1 1)A V-grooves [7] and are about the same as those obtained on GaAs/AlGaAs QWRs [1, 11]. The larger light-hole subband separation compared to that of electrons is due to the valence band offset being larger than the conduction band offset. For the InP-based system, the ratio of conduction band offset to valence band offset is taken as 40/60, while the ratio for GaAs/AlGaAs is 60/40. The different band offset ratios result in different
relative subband separations among electrons, heavy holes and light holes between the InGaAs/InGaAsP and GaAs/AlGaAs systems. Therefore, there is less lateral confinement for electrons and larger lateral confinement for holes in the InGaAs/InGaAsP system compared to the GaAs/AlGaAs system [1, 11].

Conventional low temperature photoluminescence was carried out on the InGaAs/InGaAsP QWR sample grown on (1 1 1)B V-grooves to examine the optical properties of the QWRs. Fig. 5a is the spectrum from the patterned region of the as-grown sample, displaying multiple peaks, at 0.756, 0.776, 0.892, 0.946 eV (with a shoulder at 0.93) and 0.982 eV. To resolve the spectrum spatially a selective etching technique, as described in reference [7], was used. Fig. 5b is the PL spectrum obtained with top planar (1 0 0) regions and upper sidewalls removed, showing that the PL peaks at 0.946 and 0.892 eV have been removed. Therefore, it is concluded that these peaks are originating from either the top (1 0 0) surfaces or upper sidewalls where there exists a composition variation and a quantum well thickness variation. The peak at 0.982 eV is assigned to the lower part of the sidewalls, which region has a high energy due to a thinner quantum well thickness than the (1 0 0) regions. The peaks at 0.756 and 0.776 eV are from quantum wires at the V-groove bottoms. Their low energies, compared to that on the planar region, are consistent with the composition (about 70% In) as measured by energy dispersive X-ray analysis. The full width at half maximum of the peak at 0.756 eV is about 15 meV, which is narrower not only than that of the quantum well peak on the non-patterned (1 0 0) region but also than that of InGaAs/InP QWRs grown on (1 1 1)A V-grooves [7]. It seems that a broad peak with a high energy tail which was observed in Ref. [7] is resolved into two peaks in the spectra in the present work. The peak at 0.756 eV is from the QWR ground energy levels (1e-1hh), while the peak at 0.776 eV is assumed to be from the higher QWR subband transition (2e-2hh). The 20 meV separation between these two peaks is consistent with estimated energy separations (22.7 meV) between 1e-1hh and 2e-2hh transitions. The low PL intensity from QWRs, compared to that of quantum wells on the lower part of sidewalls, is due to its small size. The size of the lower part of sidewalls is about 60 times that of QWRs. This suggests that carrier diffusion from sidewalls to the QWRs is not significant.

This study suggests that the appearance of the second QWR subband transition with PL spectra is related to bandfilling, because the relative intensity of the 2e-2hh subband transition with respect to that of the ground level transition increases with pumping power and sample temperature. A similar phenomenon has been observed in GaAs/AlGaAs QWRs where the band-filling argument was experimentally supported by magneto-PL and time-resolved PL measurements [11, 12]. The band-filling effects are believed to be associated with a sharp QWR density of states and small subband separations.

Polarization has also been used to characterize the QWRs in this work. Fig. 6 shows the PL spectra polarized along, and normal to, the wire direction at a given spot. It is seen that the PL intensity is strongly polarized, with the signal along the wire direction being more than twice that perpendicular to the wire direction. This polarization behaviour results from heavy-hole transitions.
of QWRs [13–15]. The InGaAs wire is sufficiently strained and thin that the light-hole band is separated from the heavy-hole band, and the transitions observed in the PL spectra are associated with heavy holes.

4. Conclusions

In summary, the growth of InGaAsP in (1 1 1)A V-grooves in InP results in a significant planarization development and flat V-bottoms; thus, InGaAs/InGaAsP QWRs cannot be obtained in (1 1 1)A V-grooves. In contrast, the growth of an InGaAsP layer in (1 1 1)B V-grooves leads to a sharper bottom, in which InGaAs/InGaAsP quantum wires have been achieved. The QWRs are defect free and crescent-shaped. A significant thickness variation across a small width results in subband separations comparable to those achieved in the GaAs-based system. The existence of lateral subbands has been demonstrated in PL spectra and also supported by polarization measurements. The subband separation observed in PL spectra is consistent with a calculated value based on the wire geometry observed by TEM.

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References

Vertically stacked quantum wires fabricated by an in situ processing technique

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Abstract

We have fabricated vertically stacked structures consisting of two arrays of AlGaAs/GaAs quantum wires using a processing technique conducted under an ultra-high vacuum or in a controlled ambient, called in situ electron beam (EB) lithography. In this technique a thin GaAs oxide layer was selectively formed on a clean GaAs surface by EB-stimulated oxidation in an oxygen atmosphere, and then used as a mask material to define mesa stripes by Cl\textsubscript{2} gas etching. Subsequently, ridge structures were formed on the mesa stripes by the MBE growth of a GaAs layer. The first array of quantum wires was formed on the top of the ridges by the growth of an AlGaAs/GaAs quantum well. Then, these wire structures were buried and the surface was flattened out by the growth of a thick GaAs layer. The second array of quantum wires was successively formed by a second pattern and regrowth process. The successful fabrication of these structures was confirmed by cathodoluminescence measurements at 77 K.

1. Introduction

During the fabrication process of nanostructures, in particular when dealing with III–V compound semiconductors, unintentional contamination of the processed surfaces must be minimized, since this would severely deteriorate the electrical and optical properties of the resulting structures. Several approaches have been proposed in order to synthesize nanostructures, like quantum wires and boxes, with interfaces free of contaminants. One of the most promising methods to fabricate such nanostructures is the epitaxial growth on pre-pattern substrates \cite{1-6}. However, so far, the patterning step has been carried out using conventional lithographic techniques using organic resists and under air-exposure conditions, which yields to unavoidable contamination. Furthermore, this method is presently limited in the sense that the final array is ultimately determined by the initial patterned structure. Great flexibility would be achieved if substrate patterning and regrowth could be realized successively, and as many times as required without exposing the wafer to air using an in situ processing technique \cite{7}.

In this direction, in the last few years we have been developing a technique called “in situ electron-beam (EB) lithography” \cite{8}, in which lateral
patterning and molecular beam epitaxy (MBE) are conducted sequentially, without exposing the sample to air. In this technique an ultrathin GaAs oxide layer is selectively formed on a clean GaAs surface by EB-stimulated oxidation under a controlled oxygen atmosphere [9], and then it is used as a mask material against subsequent Cl₂ gas etching. In this way in-situ patterning of the underlying GaAs layer is achieved, then the oxide mask is thermally removed, and the MBE regrowth step on a clean patterned substrate is carried out. In this work, to show the great flexibility of this technique we have fabricated vertically stacked structures consisting of two arrays of quantum wires. The high quality of these structures is demonstrated by cathodoluminescence measurements.

2. Experimental system

Fig. 1 shows a schematic illustration of the ultrahigh vacuum (UHV) multichamber system used to perform the in situ EB lithography process. Seven chambers, for sample loading, sample exchange, preheating, MBE, surface treatments, analysis, and etching are connected by UHV tunnels, so that samples loaded into the system can be transferred from one chamber to another without exposure to air. The base pressure of the system is better than 1.3 \times 10^{-7} \text{ Pa}, except for the loading chamber. An EB gun column placed on the top of the etching chamber is used for the patterning. The EB gun was designed so as to obtain a small EB diameter at a high current density by using a high-brightness Zr/O/W thermal field emitter [8]. The entire etching chamber is made of iron in order to shield the EB from any fluctuations in the environmental magnetic fields. This chamber is also equipped with a gas introduction system with a thin stainless-steel nozzle in order to supply O₂ gas to the sample surface along with EB irradiation.

3. Fabrication of vertically stacked quantum wires

The process used to fabricate quantum wires is illustrated in Fig. 2. First, in order to obtain a clean GaAs surface a 500 nm thick buffer layer was

![Schematic illustration of the UHV multichamber system](image-url)
grown by MBE on a GaAs(001) substrate mis-oriented 2° towards the [1 1 0] direction (Fig. 2(1)). Under an As pressure the sample was cooled down to room temperature, and then it was transferred to the etching chamber. Then, O₂ was gradually introduced into the etching chamber through the nozzle up to reach an O₂ pressure of 2.66 × 10⁻⁴ Pa. In order to locally enhance the O₂ pressure the nozzle was placed at a distance of 400 μm from the sample surface. Using a pattern generator a 25 kV-EB with a beam current of 4 nA was raster-scanned over the surface at room temperature so as to selectively form a thin GaAs oxide-mask layer by EB-stimulated oxidation with a total dose of 6 × 10¹⁸ electrons/cm² (Fig. 2(2)). Here it is worth to mention that we have used slightly misoriented GaAs(0 0 1) substrates because more resistant oxide masks can be obtained on this kind of substrates [9]. The O₂ gas was evacuated from the etching chamber to obtain a pressure of less than 1.3 × 10⁻⁶ Pa. Then, the substrate temperature was increased to 150°C, and Cl₂ gas was introduced to the chamber to perform the etching step at a Cl₂ gas pressure of 6.6 × 10⁻³ Pa. We have found, in a study using in situ Auger electron spectroscopy and reflection high-energy electron diffraction, that stoichiometric etching is realized and smooth As-stabilized surfaces are obtained under these etching conditions, which are excellent for the regrowth of high-quality quantum wells [10]. The etching time used here was 5 min, in this way mesa-stripe structures oriented along the [1 1 0] direction having a width of 200 nm, and 300 nm depth were defined (Fig. 2(3)). We used the [1 1 0] mesa direction because the Cl₂ etching produces normal-type mesa stripes exhibiting (311)A slanting planes along this direction, which can be easily buried during the regrowth step. The etching chamber was evacuated to about 1.3 × 10⁻⁷ Pa, and then the sample was transferred back to the MBE chamber. In order to remove the oxide mask the substrate temperature was increased to 620°C in an As flux, it was kept at this temperature for 20 min to complete the desorption of the oxide mask (Fig. 2(4)). Then, for the
Quantum wires
-second array-
λ = 804 nm

Quantum wires
-first array-
λ = 790 nm

Fig. 3. Schematic illustration of the vertically stacked structure consisting of two arrays of wires.

Second array of wires
CL image (λ = 804 nm)

![Image](image1)

First array of wires
CL image (λ = 790 nm)

![Image](image2)

Fig. 4. Spatially resolved CL images showing the emission from the first (λ = 790 nm) and second (λ = 804 nm) array of wire structures.

Regrowth step the substrate temperature was lowered to 600°C. The mesa-top width was reduced during the growth of a 200 nm thick GaAs buffer layer, due to the migration of Ga atoms from the (311)A sidewalls to the top (001) surface. In this way narrow ridge structures were formed on the mesas (Fig. 2(5)). Then, a 3.5 nm thick GaAs quantum well with 20 nm thick Al_{0.3}Ga_{0.7}As barriers was grown, quantum wires at the top of the ridges were formed directly in this step because of the slow growth rate on the sidewalls compared to that on the mesa top (Fig. 2(6)). Subsequently, these wire structures were buried, and the surface was flattened out by the growth of a 500 nm thick GaAs layer. Then, the second array of wire structures was successively formed by a second pattern (EB-stimulated oxidation and Cl₂ gas etching), and regrowth process. In order to obtain wire structures emitting at a different wavelength from the first array, a different quantum well width of 5 nm was grown during the formation of the second array of wires.

Fig. 3 shows a schematic illustration of the final structure. Cathodoluminescence (CL) measurements on this structure were performed at 77 K, using a 10 keV electron beam with 1 nA current as an excitation source. We found that the first and second array of wire structures emitted at 790 and 804 nm, respectively. Fig. 4 shows spatially resolved CL images taken at a selected wavelength of 790 nm (bottom image) and 804 nm (upper image), showing the emission from the first and second array of wire structures, respectively. These results show the feasibility of fabricating optically active stacked structures by using in situ EB lithography, thus showing the great potential of this technique.

4. Conclusion

We succeeded in fabricating vertically stacked structures consisting of two arrays of quantum wires by in situ EB lithography. Since the patterning and regrowth can be repeated as many times as required without exposing the wafer to air, this all-UHV processing technique is very promising for fabricating arbitrarily designed arrays of low dimensional structures.

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References

UHV-AFM study of MBE-grown 10 nm scale ridge quantum wires

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Abstract

We evaluate the shape of ridge quantum wires (RQWIs) with nm-scale resolution by using SEM and ultra-high-vacuum (UHV) AFM system which is connected to MBE chamber. This MBE-AFM system provides us a detailed information about the evolution of size and uniformity of ridges grown under various conditions. In this report, we investigate systematically how the growth temperature \(T_g\) and As flux affect the width and morphology of GaAs ridge structure. The results show that a very sharp and uniform GaAs ridge structures \((W < 10 \text{ nm})\) can be obtained. On the basis of this understanding, we fabricated the ridge quantum wire of 10 nm width.

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Keywords: MBE; UHV-AFM; Ridge structure; Quantum wire

1. Introduction

Facet formation by epitaxial growth on a patterned substrate has been explored with great interests as a promising method to organize nanometer size structures [1–5]. We have earlier reported successful growth of 16 nm-wide RQWIs by MBE [6–10]. Their lifetime of photoluminescence has shown a unique temperature dependence indicative of one-dimensional excitons [11]. For their device applications the structural uniformity and the width control of the wire are extremely important.

In this work, we evaluate the shape of RQWIs with nm-scale resolution by using ultra-high-vacuum (UHV) AFM system which is connected to our MBE chamber. This MBE-AFM system provides us systematic and detailed information about the time evolution of the size and uniformity of ridges grown under various growth conditions with nanometer resolution.
Previously, we have shown that the ridge width \( W \) can be controlled by adjusting the growth temperature \( T_s \) [12]. However, variation of \( T_s \) affects not only the width \( W \), but also the uniformity of ridges. For example, the ridge grown at \( T_s \) of 515°C is quite narrow \( (W < 10 \text{ nm}) \), but its morphology is poor. Since a very sharp GaAs ridge with good uniformity is indispensable with a RQWI of 10 nm width, a better growth condition must be explored. In this report, we investigate the effects of not only \( T_s \) but also of As flux to show that a very sharp and uniform ridge structures \( (W < 10 \text{ nm}) \) can be obtained.

2. Experimental procedure

GaAs(0 0 1) wafers were patterned by photo lithography and etched by a reactive ion etcher using SiCl₄. The width and height of mesa were about 1.6 and 2–3 μm, respectively. To form a ridge structure a 2.5 μm thick GaAs layer was grown at the substrate temperatures \( T_s \) of 560°C. Then the GaAs and AlAs layers were alternatively grown at various \( T_s \), under the As₂ beam equivalent pressure ranging from \( 2 \times 10^{-6} \) to \( 3 \times 10^{-5} \) Torr. The growth rates of GaAs and AlAs were 0.25 and 0.10 μm/h, respectively. The width \( W \) of ridge structure was measured by UHV-AFM and cross-sectional SEM [14, 15]. By UHV-AFM we can evaluate the ridge width \( W \) and especially its morphology, without exposing the sample to the air. Hence, we can study effects of growth parameters on a single sample. As AFM is influenced by the finiteness of tip radius, each tip was characterized by SEM and the actual ridge width \( W \) was determined by the deconvolution of the raw AFM data. Then the ridge width \( W \) was also evaluated by SEM using reflected electron mode.

Fig. 1 is the width \( W \) of the GaAs ridges thus determined as a function of \( T_s \). Solid circles are for low As flux, while blank circles are for high As flux. The width \( W \) increases with increasing \( T_s \), but the use of high As flux is effective in reducing \( W \), especially at high \( T_s \) (\( > 530°C \)). Note that \( W \) can be as narrow as 8 nm even at 540°C if high As flux is used. As for the mechanism of increase in width \( W \) at high growth temperature \( T_s \), we previously reported that the frequent dissociation of Ga atoms followed by re-arrangement process at the summit of the ridge plays a key role [8]. Hence, the use of high As flux and/or low \( T_s \) seems effective in suppressing this dissociation process of Ga atoms.

Variation of \( T_s \) influences the morphology of ridge structure. For example, when \( T_s \) is lowered to 520°C or below, the width \( W \) becomes less than 10 nm, but the (1 1 1)B surface and the (1 1 0) sidewall surface which stands vertically along the mesa stripe direction get rough. In such a case, the (1 1 0) side surfaces have some blistered facet structures, which indicates that a substantial amount of Ga will be incorporated there. The surface of (1 1 1)B nearby also has random facet structures. Fig. 2 shows the AFM scan of a sample grown at 515°C. Note that step bunching is clearly observed on the (1 1 1)B surface [13]. Note also that (1 1 1)B surface has an anomalous recess where the ridge width loses its uniformity. In the region where the ridge is uniform, the (1 1 0) surface is quite flat. There the growth rate on the (1 1 0) surface is very low, and the mesa width does not increase so much. Hence the smoothness of the ridge top is strongly correlated with the smoothness of the nearby (1 1 1)B and (1 1 0) surfaces.

Next, we grew another sample at \( T_s = 560°C \), while As flux was increased to \( 1.5 \times 10^{-5} \) for small width \( W \). Fig. 3a shows the AFM image of this sample right after the growth, which indicates that
Fig. 2. (a) AFM image of a GaAs ridge structure grown at 515°C. (b) the shaded area indicates the zone where UHV-AFM measurement was done.

Fig. 3. (a) AFM image of a GaAs ridge structure grown at 560°C. (b) the height distribution of the ridge structure along the ridge top.
a sharp ridge of 15 nm width was uniformly formed. Fig. 3b shows the height distribution along the ridge and indicates that the roughness is quite small within ±1 nm for the entire region (>1 µm²) measured. The planes forming the ridge are almost as flat as the surface of GaAs grown on a (0 0 1) substrate grown under the optimum condition.

Moreover, we have grown at 540°C a stack of three RQWIs consisting of 12 nm GaAs layers and 4 nm thick AlAs barrier layers on top of the ridge structure. Fig. 4 is its cross-sectional SEM micrograph where GaAs layers appear as bright layers and AlAs as dark and thin layers. Note that all three RQWIs have almost the same width (W ~ 10 nm) and height, indicating that the shape of the ridge grown under this condition is stable and reproducible.

Next we discuss the ways to fabricate 10 nm scale RQWI structures. To achieve a strong lateral confinement of carriers in RQWIs, one must form a uniform and narrow ridge structure of AlGaAs or AlAs with the width of less than 30 nm which functions as a bottom barrier. Then one must form on its top a thin GaAs quantum well (QW) in such a way that the resultant GaAs ridge is even narrower than the bottom ridge structure. As described before, the first condition has already been achieved. However, the second requirement is even harder to meet, because this GaAs ridge tends to be broader than that of AlAs ridges if grown under the same condition. Hence, we attempted to control the ridge width W first by varying the growth temperature Tg. The shrinkage of ridge width was achieved as Tg was lowered from 580 to 530°C. Its effect was marginal, due to the limited range of acceptable growth temperature and due to the short growth time of GaAs.

Next we examined the effect of controlling As flux. To change the As flux quickly, we prepared two As sources, a conventional K-cell source A,
and a valved K-cell source B. First we grew a GaAs ridge at 560°C with a low As flux (≈2 × 10⁻⁶ Torr) from source B. After the formation of a broad GaAs ridge, we covered the surface by a thin AlAs layer of 3 nm thickness to preserve the shape. Then we increased the As flux to ≈3 × 10⁻⁵ Torr by opening the shutter of source A and loosening the valve of source B. Under this high As flux, we grew a GaAs QW layer of 8 nm thickness and a 5 nm thick AlAs layer. Fig. 5a is the cross-sectional SEM image of this RQWI structure. Note that a sharp RQWI is sandwiched between two AlAs barrier layers. Note that the GaAs layer on the top of ridge is thicker than that on the (1 1 1)B. Hence, electrons are expected to be confined strongly in the wire of 10 nm width × 8 nm height. Fig. 5b shows the cross-section of a RQWI grown under an As flux of ≈1.5 × 10⁻⁵ Torr. As the thickness of a GaAs layer on (1 1 1)B is about 3 nm, and is smaller than that of Fig. 5b, the quantized energy of electrons in the side QW on (1 1 1)B of Fig. 5a is 220 meV, which is higher than that of Fig. 5b by 100 meV. Hence, electrons in the RQWI of Fig. 5a will be tightly confined.

3. Conclusions

From the morphological study of MBE-grown GaAs ridge structures by UHV-AFM and cross-sectional SEM, we have shown that the increase of As flux is quite effective in reducing the ridge width down to 10 nm even at high substrate temperatures. As a result, the condition to form very narrow GaAs and AlAs ridges with good morphology has been clarified. Very narrow and uniform ridge quantum wire structures, including the stacked wires have been successfully grown by optimizing both the substrate temperature and the As flux.

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References

Improved optical qualities of GaAs/Al$_{0.3}$Ga$_{0.7}$As tilted T-shaped quantum wires fabricated by glancing-angle molecular beam epitaxy

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Abstract

High-quality GaAs/Al$_{0.3}$Ga$_{0.7}$As tilted T-shaped quantum wires (tilted T-QWRs) were fabricated with a two-step growth of molecular beam epitaxy (MBE), which consists of glancing-angle MBE growth of GaAs/Al$_{0.2}$Ga$_{0.8}$As multi-quantum well (MQW) layer (five 6.6 nm thick GaAs wells) on a reverse-mesa etched (1 0 0) GaAs substrate and MBE over growth of GaAs/Al$_{0.3}$Ga$_{0.7}$As/single-quantum well (SQW) (7.3 nm well width) on a (1 1 1)B facet. This new technique enable us to fabricate lots of tilted T-QWRs over a large area of a GaAs substrate surface. Full-width-at-half-maximum (FWHM) of a cathodoluminescence (CL) peak from the tilted T-QWRs was reduced down to 10 meV at 78 K, which is almost comparable with those of GaAs/AlGaAs T-QWRs fabricated by the cleaved-edge overgrowth.

Keywords: GaAs; Al$_{0.3}$Ga$_{0.7}$As; Glancing-angle MBE; T-shaped quantum wire; (1 1 1)B facet; Selective growth

1. Introduction

Recently, low-dimensional semiconductor structures such as quantum wires (QWRs) and quantum dots (QDs) have been intensively investigated because of their novel optical properties [1–3].

Pfeiffer et al. [4] fabricated GaAs/Al$_{0.3}$Ga$_{0.7}$ As T-shaped QWRs (T-QWRs) by the cleaved edge overgrowth (CEO) method. In this method, a GaAs quantum well (QW) layer and an Al$_{0.3}$Ga$_{0.7}$As barrier layer are overgrown on a (1 1 0) cleaved plane of a GaAs/Al$_{0.3}$Ga$_{0.7}$As multi-quantum well (MQW) layer grown on a (1 0 0) GaAs substrate by MBE. Therefore, the cross-sectional dimension of the T-QWRs can be precisely controlled in the atomic scale and high quality T-QWRs have been obtained [4, 5]. These T-QWRs, however, are fabricated only on a very limited area of the cleaved
surface of the MQW layer and it is difficult to apply the T-QWRs to electric or optical devices.

In order to find some solutions of this problem, we have attempted to fabricate GaAs/Al$_{0.5}$Ga$_{0.7}$As tilted T-shaped QWRs (tilted T-QWRs) using (1 1 1)B facet planes, instead of the (1 1 0) cleaved plane, preferentially grown on a reverse-mesa etched GaAs substrate by glancing-angle molecular beam epitaxy (GA-MBE) [6–8]. In this case, many tilted T-QWRs can be easily fabricated on a lot of mesa stripes along [1 1 0] direction on a large area of a (1 0 0) GaAs substrate. During GA-MBE growth (no substrate rotation), the preferentially formed (1 1 1)B facet planes are not directly exposed to both Ga and Al beams due to a self-shadowing effect [6]. After the GA-MBE growth of a GaAs/Al$_{0.5}$Ga$_{0.7}$As MQW layer, a cross-section of the MQW structure appears on the (1 1 1)B facet plane when we choose a growth condition where the inter-facet migration of Ga and Al atoms do not occur from the (1 0 0) surface to the (1 1 1)B facet. GaAs/Al$_{0.5}$Ga$_{0.7}$As tilted T-QWRs are fabricated by overgrowing a GaAs QW layer and an Al$_{0.5}$Ga$_{0.7}$As barrier layer on the (1 1 1)B facet plane by a normal MBE with rotating the substrate just after the GA-MBE growth [6, 7]. In the previous work [6], we observed strong cathodoluminescence (CL) at 78 K from 20 GaAs/Al$_{0.5}$Ga$_{0.7}$As tilted T-QWRs formed with a GaAs/Al$_{0.5}$Ga$_{0.7}$As MQW of the well width of $L_w = 5.4$ nm on the (1 0 0) plane and a GaAs/Al$_{0.5}$Ga$_{0.7}$As QW of $L_w = 4.6$ nm on the (1 1 1)B facet, but the full-width-at-half-maximum (FWHM) of the CL peak from the tilted T-QWRs was as large as 61 meV which is mainly due to not optimized conditions for both GA-MBE growth and overgrowth on (1 1 1)B facets. Recently, we reduced this FWHM value down to 19 meV by using improved growth conditions but the CL peak from the tilted T-QWRs was observed only as a shoulder next to the large CL peak from GaAs/Al$_{0.5}$Ga$_{0.7}$As QWs on the (1 0 0) plane [7]. This FWHM was still about twice as large as those of GaAs/Al$_{0.5}$Ga$_{0.7}$As T-QWRs (10 meV [4], 4–7 meV [5]) fabricated by the CEO method, which is mainly due to smaller size of GaAs/Al$_{0.5}$Ga$_{0.7}$As tilted T-QWRs (4.4 nm-wide MQW on the (1 0 0) plane and 4.5 nm-wide QW on the (1 1 1)B facet). In this paper, we report on much improved CL spectra from GaAs/Al$_{0.5}$Ga$_{0.7}$As tilted T-QWRs with a little bit larger size fabricated on a reverse-mesa etched (1 0 0) GaAs substrate by the two-step growth (GA-MBE/MBE).

2. Fabrication of GaAs/Al$_{0.5}$Ga$_{0.7}$As tilted T-QWRs

The structure of the fabricated sample is shown in Fig. 1a. A GaAs/Al$_{0.5}$Ga$_{0.7}$As (87 nm/4.3 nm, 20 periods) superlattice buffer, an Al$_{0.5}$Ga$_{0.7}$As layer (87 nm) and a GaAs layer (6.6 nm) were grown on a reverse-mesa etched GaAs (1 0 0) substrate by GA-MBE ($T_s = 630^\circ C$, $V_{III} = 25$). Next, a GaAs/Al$_{0.5}$Ga$_{0.7}$As (6.6 nm/17 nm, 4 periods) MQW layer and an Al$_{0.5}$Ga$_{0.7}$As barrier layer (47 nm) were grown by GA-MBE ($T_s = 570^\circ C$, $V_{III} = 25$) to reduce the inter-facet migration of Ga atoms [8]. Finally, a GaAs/Al$_{0.5}$Ga$_{0.7}$As

Fig. 1. Sample structure of GaAs/Al$_{0.5}$Ga$_{0.7}$As tilted T-QWRs (a) and its cross-sectional SEM image (b).
(7.3 nm/51 nm) SQW and a 12 nm-thick GaAs cap layer were overgrown on the (1 1 1)B facet under the optimized overgrowth condition on the (1 1 1)B facet (T_s = 630°C, V/Ii = 10) to obtain high optical quality of a GaAs/Al_{0.3}Ga_{0.7}As SQW with rotating the substrate at 60 rpm [7]. Fig. 1b shows the cross-sectional SEM image of the grown sample. Five tilted T-QWRs were successfully fabricated at the junction region of MQW on the (1 0 0) plane and the SQW on the (1 1 1)B facet plane.

3. CL measurements

A CL spectrum of the GaAs/Al_{0.3}Ga_{0.7}As tilted T-QWRs at 78 K is shown in Fig. 2, which was obtained when a focused electron beam (0.2 µm diameter, 7 keV) excited the (1 1 1)B facet region (the tilted T-QWR region). CL peaks at 787 nm and at 795 nm come from the GaAs/Al_{0.3}Ga_{0.7}As MQW on (1 0 0) plane and from the GaAs/Al_{0.3}Ga_{0.7}As SQW on the (1 1 1)B facet, respectively. These MQW on the (1 0 0) plane and SQW on the (1 1 1)B facet form the tilted T-QWRs at their cross regions. The luminescence from the tilted T-QWRs was observed at 803 nm. This peak energy was 1.544 eV which is 31 meV lower than that of the MQW on (1 0 0) plane and is 15 meV lower than that of the SQW on the (1 1 1)B facet plane. This red-shift is caused by the weaker confinement of carriers of the tilted T-QWRs. A peak at 817 nm comes from the edge of the SQW on the (1 0 0) plane which was formed with a thicker QW due to the surface migration of Ga atoms from (1 1 1)B facet plane to (1 0 0) plane when the GaAs/Al_{0.3}Ga_{0.7}As SQW was overgrown on the (1 1 1)B facet. As can be seen in Fig. 2, all these CL peaks are well resolved.

In order to investigate the optical qualities of the tilted T-QWRs fabricated by the two-step growth of GA-MBE / MBE, we measured CL spectra with changing the intensity of the exciting electron beam as shown in Fig. 3. The CL peaks from the T-QWRs were clearly observed in all cases. In the case of a weak electron beam (1 × 10^{-9} A), the peak from the SQW on the (1 1 1)B facet plane does not become clear, while that from the tilted T-QWRs is still clear, implying that almost all of electrons and holes excited in the (1 1 1)B facet region diffuse into the tilted T-QWR regions and radiatively recombine there. With increasing the exciting

![Fig. 2. CL spectrum (78 K) observed in the GaAs/Al_{0.3}Ga_{0.7}As T-QWRs sample when the (1 1 1)B facet region is excited by the electron beam.](image1)

![Fig. 3. CL spectra (78 K) observed in the GaAs/Al_{0.3}Ga_{0.7}As T-QWRs. Excitation intensities are changed from 1 × 10^{-9} to 1 × 10^{-7} A.](image2)
electron beam, the luminescence from the SQW on
the (1 1 1)B facet become clear and FWHM of the
peak from the T-QWRs become larger. It is partly
due to the band filling effect in the tilted T-QWRs.
In the case of the low excitation intensity
(1 x 10^-9 - 6 x 10^-9 A), FWHM of the peak from
the T-QWRs is 10 meV, which is about half of that
(19 meV) of previous GaAs/Al_{0.3}Ga_{0.7}As tilted T-
QWRs [7], and is almost comparable with those
(10 meV [4], 4 - 7 meV [5]) of GaAs/Al_{0.3}Ga_{0.7}As
T-QWRs fabricated by the CEO method.

4. Calculation of electron and hole states in the
T-QWRs

In order to understand quantitatively the optical
properties of the tilted T-QWRs, we calculated states of
electrons and holes in the GaAs/Al_{0.3}Ga_{0.7}As tilted T-QWRs based on the
finite element method (FEM). The advantage of
FEM is that the eigenenergy and eigenstates of
carriers confined in a volume with an arbitrary
shape can be calculated without any approximation
in principle. We take the direction perpendicular
to the (1 0 0) substrate surface as the z-axis.

The y-axis is taken to be parallel to the tilted T-QWRs
(i.e., [1 1 0] direction) and the x-axis is taken to be
perpendicular to the tilted T-QWRs in (1 0 0) plane,
respectively. In the effective-mass approximation,
the envelope wave function of conduction electrons
in the GaAs/Al_{0.3}Ga_{0.7}As tilted T-QWR is of the
form e^{iS} y(x, z), where S(x, z) satisfies the 2D
Schrödinger equation given by

\[ -\frac{\hbar^2}{2m^*_e}\left(\frac{\partial^2 S}{\partial x^2} + \frac{\partial^2 S}{\partial z^2}\right) + U(x, z)S = E_eS, \]

where \( \hbar \) is the Plank’s constant, \( E_e \) is an
eigenenergy of an electron confined in the tilted
T-QWR. In Eq. (1), \( m^*_e(x, z) \) is a position-dependent
effective mass of an electron. In the GaAs and
Al_{0.3}Ga_{0.7}As regions, we used the effective masses
of \( m^*_e = 0.067m_0 \) and \( m^*_e = 0.071m_0 \), respectively,
where \( m_0 \) is the static mass of an electron.
The conduction-band offset of 225 meV was used for
\( U_e(x, z) \). The Schrödinger equation for holes is
given by

\[ -\frac{\hbar^2}{2m^*_h}\left(\frac{\partial^2 \Psi}{\partial x^2} + \frac{\partial^2 \Psi}{\partial z^2}\right) + U_h \Psi = E_h \Psi. \]

The valence band offset of 131 meV was used for
\( U_h(x, y) \). We used the common hole mass \( m^*_h \) of
0.45m_0 for GaAs and Al_{0.3}Ga_{0.7}As regions. In this
calculation, the electron–hole Coulomb interaction
was neglected.

Fig. 4a shows \( |\Psi|^2 \) of an electron confined in the
tilted T-QWR; the contours are lines of constant
probability \( (|\Psi|^2 = 0.2, 0.4, ..., 0.8) \) for electrons. The contour map of \( |\Psi|^2 \) is given in Fig. 4 b. The top of
calculated \( |\Psi|^2 \) slightly shifts from the crosspoint of
the QW on (1 0 0) plane and the QW on (1 1 1)B
facet due to effectively thinner width of the
Al_{0.3}Ga_{0.7}As barrier on the (1 0 0) QW layer near the
T-shaped region. In this calculation, we obtained a red-shift of 22 meV for recombination
energy in the tilted T-QWRs from the corresponding
MQW transition. The observed red-shift is about
31 meV. The difference (9 meV) between the
observed and calculated red-shifts is considered to

![Fig. 4. Calculated probability density |\Psi|^2 (a) and contour map of |\Psi|^2 (b) for a electron confined in the tilted T-QWRs.](image-url)
equal to an excess part of the exciton binding energy for the one-dimensional structure (the tilted T-QWR). Taking into account of an exciton binding energy of 11 meV for the 6.6 nm wide MQW, we obtain an exciton binding energy of 20 meV for the tilted T-QWRs, which agrees well with the value (17 meV) obtained for a GaAs/AlGaAs T-QWR by Wegscheider et al. [9].

5. Conclusion

GaAs/Al₀.₃Ga₀.₇As tilted T-QWRs (6.6 nm wide MQW on the (1 0 0) plane and 7.3 nm wide SQW on the (1 1 1)B facet plane) were successfully fabricated by the two-step MBE growth (GA-MBE/MBE). Well-resolved CL peaks from the MQW on (1 0 0) plane, the SQW on (1 1 1)B facet plane, and the tilted T-QWRs were observed. FWHM of the CL peak from the tilted T-QWRs was as small as 10 meV at 78 K, which is comparable with those (10 meV [4], 4–7 meV [5]) of GaAs/Al₀.₃Ga₀.₇As T-QWRs fabricated by the CEO method. Electron and hole states in the T-QWR were investigated with the use of the finite element method (FEM). The binding energy of excitons in the GaAs/Al₀.₃Ga₀.₇As tilted T-shaped QWRs is estimated to 19 meV by comparing with calculated recombination energies and the observed CL peak energies for the MQW and the tilted T-QWRs.

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References

High-density GaAs/(GaAs)$_2$(AlAs)$_2$ quantum wires naturally formed on $(7\,7\,5)$B-oriented GaAs substrates by molecular beam epitaxy

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Abstract

GaAs/(GaAs)$_2$(AlAs)$_2$ quantum wires (QWRs) were naturally formed in a thin GaAs/(GaAs)$_2$(AlAs)$_2$ quantum well (QW) with a regularly corrugated AlAs/GaAs upper interface (a period of 12 nm) and a flat GaAs/AlAs lower interface grown on $(7\,7\,5)$B-oriented GaAs substrates by molecular beam epitaxy. The QWRs were formed side by side with an extremely high density of $8 \times 10^9$ QWRs/cm, which is the same to that of the previous GaAs/AlAs QWRs grown on the $(7\,7\,5)$B substrate (the highest density of QWRs ever reported [8]). A photoluminescence from the QWRs formed in the QW with an average well width ($L_w$) of 2.1 nm, which have a cross section of about $12 \times 2$ nm$^2$, showed a strong polarization dependence (the polarization degree $P = [(I_H - I_L)/(I_H + I_L)] = 0.21$). The polarization degree is about twice as large as that of the previous GaAs/AlAs QWRs with an average $L_w$ of 3.3 nm grown on the $(7\,7\,5)$B substrate. This improvement of the polarization degree is mainly due to the further reduced QW width, which results in an improved one-dimensional confinement of carriers in the present QWRs.

1. Introduction

Recently, low-dimensional quantum structures such as quantum wires (QWRs) and quantum dots (QDs) have attracted much interest from the viewpoints of low-dimensional physics and device applications, and many endeavors have been devoted to fabricate these structures. The confinement of carriers in QWRs or QDs gives rise to sharp peaks in the density of states, and it is expected to realize high efficient lasers with a low threshold current and a reduced temperature dependence of threshold current by using QWRs or QDs [1, 2]. Among many fabrication techniques of QWRs, molecular beam epitaxy (MBE) [3-6] and metalorganic chemical vapor deposition [7] on high index planes with multi-atomic steps of GaAs crystals are very promising, because the fabrication techniques are very simple and QWRs can be formed without any fabrication damage. When we studied optical properties of GaAs/AlAs quantum wells (QWs) grown on GaAs substrates with various tilted...
angles from the (111)B plane toward the (110) plane by MBE, we eventually found that the GaAs/AlAs QWRs structure is naturally formed on GaAs substrates with an off-angle of 8.5° from the (111)B plane [(7 7 5)B-oriented GaAs substrates] in 1995. And we have reported the GaAs/AlAs QWRs grown on the (7 7 5)B GaAs substrate by MBE [8]. The GaAs/AlAs QWRs on the (7 7 5)B substrate were naturally formed in a 3.3 nm thick GaAs/AlAs QW with a corrugated AlAs/GaAs upper interface (lateral period 12 nm and vertical amplitude 1.2 nm) and a flat GaAs/AlAs lower interface. Density of the QWRs was very high (8 × 10⁵ QWRs/cm), but the one-dimensional confinement of carriers was rather weak. It is difficult to reduce the GaAs well width further in order to enhance the one-dimensional confinement of carriers for the GaAs/AlAs QWs, because the ground energy level of electrons in the QW approaches the X band edge of the AlAs barrier layer with decreasing Lₔ down to about 3 nm.

In this paper, we study to avoid the X-band edge problem by using (GaAs)₅(AlAs)₅ short-period superlattice (SPS) barriers instead of AlAs barriers and improve the one-dimensional confinement of carriers in the GaAs/(GaAs)₅(AlAs)₅ QWRs grown on (7 7 5)B GaAs substrates by MBE.

2. Fabrication of QWRs

Samples in this study were grown by a Nissin RB-2001G MBE system. V/III pressure ratio was 9 (14) for GaAs (AlAs), and growth rates of GaAs and AlAs were 1.0 μm/h. The (7 7 5)B GaAs substrate was degreased and etched using a sulfuric acid etchant (H₂SO₄ : H₂O₂ : H₂O = 5 : 1 : 1) before loading into the growth chamber, and they were thermally cleaned at a substrate temperature (Tₛ) of 650°C for 15 min in As₄ atmosphere (10⁻⁶ Torr) just before MBE growth.

The surface of GaAs grown on the (7 7 5)B GaAs substrate can be changed between a flat surface and a corrugated surface by controlling the substrate temperature. The surface of GaAs becomes flat at Tₛ = 540–580°C and corrugates above Tₛ = 640°C, while the surface of AlAs is always flat above Tₛ = 540°C.

Fig. 1 shows a schematic illustration of the GaAs/(GaAs)₅(AlAs)₅ QWRs structure grown on the (7 7 5)B GaAs substrate. The GaAs/(GaAs)₅(AlAs)₅ QWRs structure on the (7 7 5)B substrate was fabricated as follows. First, a 25-period (30 nm thick) (GaAs)₅(AlAs)₅ SPS barrier layer was grown on a 800 nm-thick GaAs/AlAs buffer layer at Tₛ = 580°C (Fig. 2a). Second, the
substrate temperature was raised to 640°C, and a GaAs QW (QWRs) layer was grown on the structure at \( T_s = 640°C \) (Fig. 2b). Regular corrugation with very straight step edges along [1 \( \bar{1} 0 \)] direction is formed. Lateral period and vertical amplitude of the corrugation is 12 and 1.2 nm by cross-sectional transmission electron microscopy. Details of the corrugation have been reported previously in Ref. [8]. Third, (GaAs)\(_2\)(AlAs)\(_2\) SPS barrier was grown after reducing the substrate temperature from 640 to 580°C. The surface turned to be flat again [Fig. 2c]. The corrugated interface results in a periodic modulation of the well width in the QW. Therefore, lateral confinement effect of carriers arises and QWRs are naturally formed at the thick parts in the GaAs/(GaAs)\(_2\)(AlAs)\(_2\) QW grown on the (7 7 5)B substrate. After repeating second and third processes with changing QW layer thickness \( L_w \), a 20 nm thick GaAs cap layer was finally grown on the structure. Hence, high-density GaAs/(GaAs)\(_2\)(AlAs)\(_2\) QWRs were fabricated using the interesting MBE growth characteristics of GaAs and AlAs on the (7 7 5)B GaAs substrate. Average well widths \( L_w \) were 2.1, 3.0, 4.2, 6.6 and 12 nm, respectively.

3. Photoluminescence measurements

We measured photoluminescence (PL) at 14 K not only from the GaAs/(GaAs)\(_2\)(AlAs)\(_2\) QWs (including QWRs) grown on the (7 7 5)B GaAs substrate but also from GaAs/(GaAs)\(_2\)(AlAs)\(_2\) QWs grown on a (1 0 0) GaAs substrate at \( T_s = 640°C \) without any growth interruptions. The samples were excited by a He–Cd laser beam with a power of 2 mW and a beam diameter of about 200 \( \mu \)m.

Fig. 3 shows PL spectra at 14 K from the QWs (QWRs) on the (7 7 5)B substrate and the QWs on the (1 0 0) substrate for two different polarization directions. For the sample on the (7 7 5)B substrate, a solid line shows the PL spectrum from the QWs for the polarization parallel to the QWRs ([1 \( \bar{1} 0 \) direction), and a dotted line shows that for the polarization perpendicular to the QWRs ([5 5 14] direction). PL peaks at 670, 707, 741, 776 and 804 nm are from the QWRs with average \( L_w \) of 2.1, 3.0, 4.2, 6.6 and 12 nm on the (7 7 5)B substrate, respectively. For the reference QWs grown on the (1 0 0) substrate, a solid line shows PL spectrum for the polarization parallel to the cleaved surface ([0 1 1] direction), and a dotted line shows that for the polarization perpendicular to the cleaved surface ([0 1 1] direction).

Full-width at half-maximum of the PL peak from the QWRs with an average \( L_w \) of 2.1 nm on the (7 7 5)B substrate was 30 meV, while that from the QW with a \( L_w \) of 2.1 nm on the (1 0 0) substrate was 20 meV. The integrated PL intensity from the QWRs on the (7 7 5)B substrate was almost
the same as that from the QW on the (1 0 0) substrate. These results indicate that the GaAs/(GaAs)$_2$(AlAs)$_2$ QWRs grown on the (7 7 5)B substrate had high optical quality.

The polarization degree, $P = [(I_1 - I_0)/(I_1 + I_0)]$, of PL peaks from the GaAs/(GaAs)$_2$(AlAs)$_2$ QWRs (QWRs) on the (7 7 5)B substrate, the GaAs/(GaAs)$_2$(AlAs)$_2$ QWs on the (1 0 0) substrate and the previous GaAs/AlAs QWs (QWRs) grown on the (7 7 5)B substrate [8] are plotted as a function of $L_w$ in Fig. 4. With decreasing $L_w$ from 12 to 2.1 nm, the polarization anisotropy of PL from the QWs (QWRs) on the (7 7 5)B substrate monotonously increases. The PL peak from the QWRs with the average $L_w$ of 2.1 nm on the (7 7 5)B substrate exhibits the strongest polarization degree ($P = 0.21$). On the other hand, the polarization dependence is constantly almost zero for the QWs on the (1 0 0) substrate. These results are in good agreement with theoretical investigations that PL from a QWR is polarized parallel to the wire direction [9, 10]. The polarization degree of PL from the QWRs with the average $L_w$ of 2.1 nm on the (7 7 5)B substrate was about twice as large as that ($P = 0.11$) from the previous GaAs/AlAs QWRs with an average $L_w$ of 3.3 nm grown on the (7 7 5)B substrate [8]. These results indicate that lateral confinement effect of carriers due to the corrugated interface is actually much enhanced by decreasing $L_w$ down to 2.1 nm, and high-density GaAs/(GaAs)$_2$(AlAs)$_2$ QWRs with a considerably strong one-dimensional confinement of carriers are realized on the (7 7 5)B substrate by MBE.

Using the finite element method (FEM), we also calculated the envelope wave functions $\psi$ for ground energy levels of electrons and heavy holes in the GaAs/Al$_{0.3}$Ga$_{0.7}$As QW with an average $L_w$ of 2.1 nm, which is almost equivalent to the GaAs/(GaAs)$_2$(AlAs)$_2$ QWRs with the average $L_w$ of 2.1 nm grown on the (7 7 5)B substrate. This FEM calculation showed that the envelope wave function of electrons spreads rather laterally beyond the boundary of the QWR, but that of heavy holes is clearly localized within the QWR.
The high-density QWRs are formed in a GaAs/(GaAs)$_2$(AlAs)$_2$ N-multi-layer structure grown on the (7 7 5)B substrate, then they are expected to be applied to semiconductor lasers with extremely high-density QWRs ($8 \times 10^7$ QWRs/cm).

4. Conclusion

GaAs/(GaAs)$_2$(AlAs)$_2$ QWRs were naturally formed in a thin GaAs/(GaAs)$_2$(AlAs)$_2$ QW with a regularly corrugated AlAs/GaAs upper interface (a period of 12 nm) and a flat GaAs/AlAs lower interface grown on (7 7 5)B-oriented GaAs substrates by MBE. The density of the QWRs was as high as $8 \times 10^7$ QWRs/cm. The PL peak at $\lambda = 670$ nm from the QWRs formed in the QW with an average well width of 2.1 nm, which have a cross section of about $12 \times 2$ nm$^2$, showed a strong polarization dependence (the polarization degree $P \equiv (I_1 - I_\perp)/(I_1 + I_\perp) = 0.21$). The observed degree of polarization is about twice as large as that ($P = 0.11$) from the previous GaAs/AlAs QWRs with an average well width of 3.3 nm on the (7 7 5)B substrate [8]. In addition, the integrated PL intensity from the GaAs/(GaAs)$_2$(AlAs)$_2$ QWRs was as large as that from the corresponding 2.1 nm QW grown on the (1 0 0) GaAs substrate. These results imply that the GaAs/(GaAs)$_2$(AlAs)$_2$ QWRs grown on the (7 7 5)B substrate have high potential for applications in optical devices.

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References

Growth of GaInAsP quantum wire heterostructures using the strain-induced lateral-layer ordering process


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Abstract

We present a generalized strain-balance theory on the strain-induced lateral-layer ordering (SILO) process, where various combinations of short-period superlattice (SPS) constituents are used to fabricate novel quantum wire (QWR) heterostructures. Based on the theory, we have fabricated two sets of QWRs in the GaInP and GaInAsP material systems with emission wavelengths of 6200 Å and 1 µm, respectively. Growth of (GaP)_{1/3}(InP)_{1/3} SPS was used to create QWRs on GaAs_{0.6}P_{0.4} based substrates. Also, QWRs were formed using (GaP)_{1/2}(As)_{1/2} SPS on GaAs substrates. In both these cases, the monolayer pairings of the SPS constituents were unequal. Transmission electron microscopy and polarized photoluminescence spectroscopy confirm the formation of QWRs. These results support the validity of the generalized theory of the SILO process.

1. Introduction

The development of semiconductor quantum wires (QWR) has been driven by the theoretically predicted improvements in performance over quantum well (QW) devices, as well as the scientific aspect of exploring new concepts in engineering. While current efforts into creating QWRs include etch and regrowth [1], growth on patterned substrates [2], and cleaved edge overgrowth [3], the strain-induced lateral-layer ordering (SILO) process has proven itself to be a simple and reliable technique for QWR formation [4]. The in situ SILO technique can use standard (1 0 0) on-axis or vicinal substrates [5], creates lateral composition modulation perpendicular to the growth direction, and applies to a wide range of growth conditions. It also avoids damage at interfaces because no pre- or post-growth processing is required. The SILO technique uses localized strain within short-period superlattices (SPS) to drive atomic migration, resulting in lateral quantum wells (LQW) formed in the [1 1 0] direction. Combined with common QW fabrication methods, which confine charge carriers in the (1 0 0) growth direction, this technique creates two-dimensionally confined QWRs.

A key factor of the SILO technique is that the strain balance is maintained through the SPS layer. That is, the layers which comprise the SPS have
 alternating compressive and tensile strain, with respect to the growth substrate. Thus, while the strain within the SPS can be large, the net strain is finite and near zero. This near-zero overall strain is significant so that defects due to lattice relaxation are minimized. Previous efforts utilizing the SILO process studied the formations of QWRs through the growth of (GaP)\textsubscript{m}/(InP)\textsubscript{n} SPS on GaAs [6] and (GaAs)\textsubscript{m}/(InAs)\textsubscript{n} SPS on InP [7]. In order to fulfill the nearly balanced strain requirement in these materials systems, only near-equal monolayer pairings of the SPS (m ≈ n) could be studied. As a consequence, photonic devices fabricated using the SILO process exist in limited wavelength domains.

In this study, we have generalized the strain-balance concept in the SILO process where the monolayer pairings of the SPS are not equal (m ≠ n) and yet still satisfy the near-zero net strain condition. This is demonstrated by the growth of (GaP)\textsubscript{2}/(InP)\textsubscript{1} SPS on a GaAs\textsubscript{0.66}P\textsubscript{0.34} substrate. Additionally, QWR formation is also explored using (GaP)\textsubscript{1.2}/(InAs)\textsubscript{1} SPS grown on GaAs. Polarized emission from these samples was observed near 6200 Å and 1 μm, respectively. Thus, this study provides important evidence about the validity and versatility of the SILO process as it is applied to QWR formation in a wide selection of materials.

2. Theory

Realizing one-dimensionally confined structures in semiconductors has been routinely explored and exploited in the past 20 years. Typically, these fabricated QWs confine carriers in the growth direction. A more difficult challenge has been to create two-dimensionally confined QWRs. For this type of structure, composition variation of the alloy must occur simultaneously in two orthogonal directions.

In an effort to create viable QWRs in semiconductors, the SILO process was developed [8]. This growth technique is an in situ method for forming LQWs in the growth plane through molecular beam epitaxy growth. The basic technique of the SILO process is based on the deposition of alternating thin layers of two different materials to form a SPS, where each of the layers is strained with respect to the growth substrate. This nearly strain balanced structure, however, is tensile for one of the layers and compressive for the other. During the growth of this SPS, the localized finite net strain that exists can trigger spinodal decomposition [9] or induce a morphological instability [10] on the surface. This will cause atoms in the alloy to migrate in a plane perpendicular to the growth direction. As predicted by Gläs, once the composition modulation is initiated, it is stable and propagates as growth continues [11]. In previous studies, this modulation has always been observed to occur in the [110] direction, creating LQWs. When combined with larger band-gap material in the growth direction via standard QW growth methods, somewhat uniform arrays of QWRs are formed. The key factor of the SILO process is that the strain in each of the alternating layers is opposite in sign but nearly equal in magnitude. Thus, the SPS as a whole has minimal strain, and defects due to lattice relaxation or critical thickness boundaries are not induced.

Previous studies have explored the growth of (GaP)\textsubscript{m}/(InP)n SPS on GaAs substrates to create Ga\textsubscript{x}In\textsubscript{1-x}P QWRs [6]. The lattice mismatch between GaP/GaAs and InP/GaAs is approximately equal in magnitude (∼4%), but opposite is sign. By growing this superlattice using the condition that m ≈ n on GaAs, the tensile and compressive strain forces within the SPS region compensate each other to provide a near-zero global strain (ΔS). Similarly, (GaAs)\textsubscript{m}/(InAs)\textsubscript{n} SPS has been grown on InP substrates to create Ga\textsubscript{x}In\textsubscript{1-x}As QWRs [7]. In this case also, the condition m ≈ n was used. Thus, because of the nature and magnitude of the strain that exists in the SPS using the aforementioned material systems, only the case of using nearly equal monolayer pairing (m ≈ n ≈ 2) has been previously studied. While success has been achieved in creating QWRs in these systems, the necessity to fulfill the balanced strain conditions has also limited the wavelength ranges for photonic devices fabricated via the SILO process.

The near strain balanced conditions of the SILO process in general are not limited to the case of m ≈ n. Consider the general SPS consisting of (A)\textsubscript{m}/(B)n grown on a substrate with a lattice
constant $a_0$, where A and B are arbitrary semiconductor alloys. The amount of mismatch between the SPS and substrate can be characterized using the equation

$$\Delta T = \frac{\text{SPS period} - N(a_0/2)}{N(a_0/2)}$$

where SPS period = $(ma_A + na_B)/2$, $N$ is the nearest integer of $(m + n)$, and $a_A$ and $a_B$ are the lattice constants of the SPS constituent layers. In order for strain balance to occur within the SPS, the lattice constant of the substrate ($a_0$) must be between the constants of the constituents ($a_A$ and $a_B$). The global strain can be calculated using the relation

$$\Delta S = |m\Delta a_A + n\Delta a_B|,$$

where $\Delta a_A = (a_A - a_0)/a_0$ and $\Delta a_B = (a_B - a_0)/a_0$. The only necessary constraint of the SILO process is therefore that $\Delta S$ and $\Delta T$ are small but finite (~5%). This condition provides the strain to drive the atomic migration within the alloy, and yet maintains the near lattice match of the SPS as a whole [8]. The number of monolayers used in the SPS need not be equal, provided that the proper substrates and SPS constituents are used.

3. Experimental procedure

To verify the generalized strain-balance theory, two QWR heterostructures were created using the SILO process with $m \neq n$ in the SPS layers. The first was \([(GaP)_{12}/(InP)] \ SPS)/AlGaInP grown on GaAs$_{0.66}P_{0.34}$ substrates. This substrate, commonly employed for light-emitting diode fabrication, uses vapor-phase epitaxy to compositionally grade GaAs to GaAsP on a (100) oriented GaAs substrate [12]. Each of the 5 SPS QW grown consisted of 12 pairs of the binary constituents, with QWs separated by 100 Å Al$_{10.15}$Ga$_{0.53}$In$_{0.32}$P barriers. The second QWR structure was \([(GaP)$_n/(InAs)] \ SPS)/GaAs grown on (100) oriented GaAs. The QWR region consisted of 5 SPS QWs, where 12 pairs of (GaP)$_{2.2}/(InAs)$ SPS made up each well, separated by 75 Å GaAs barriers. Surrounding this QWR region above and below was 2500 Å of lattice-matched Ga$_{0.52}$In$_{0.48}$P. Lateral modulation in the [110] direction within the SPS regions, along with the barrier materials in the [001] direction, create the two-dimensionally confined QWRs.

Samples were grown using a gas-source molecular beam epitaxy system, with elemental In and Ga group III sources, and thermally cracked AsH$_3$ and PH$_3$ group V sources [13]. Both a cryogenic and turbomolecular pump were used to evacuate the growth chamber. Growth rates of individual effusion cells were calibrated using reflection high-energy electron diffraction intensity oscillations.

The structural and optical properties of the QWR samples were characterized using transmission electron microscopy (TEM) and polarized photoluminescence (PPL) spectroscopy, respectively. Excitation for the PPL measurements was performed using the 5145 Å line of an Ar ion laser. The photoluminescence (PL) emission signal was measured using a 0.5 m focal length grating spectrometer and a liquid-nitrogen-cooled Ge detector, using the lock-in technique. Samples were mounted in an evacuated cold finger type cryostat. To determine the polarization of the emission, PPL additionally employed a polarization analyzer. This analyzer was rotated so that its axis could be oriented either parallel or perpendicular to the QWR direction. Also, a depolarizer was employed to remove any anisotropic effects of the spectrometer grating. The polarization value ($p$) for the PPL was calculated using the equation $p = (I_\parallel - I_\perp)/(I_\parallel + I_\perp)$, where $I_\parallel$ is the PL peak intensity measured with the polarizer axis parallel to the QWR direction, and $I_\perp$ is the PL peak intensity measured with the polarizer axis perpendicular to the QWR direction.

4. Results and discussion

In TEM images of both samples, a uniform contrast appears across the SPS region in the [110] cross section (not shown), while strong dark and light fringes are seen in the [110] cross-section. The dark field [110] cross-sectional TEM image
of a QWR structure using a [(GaP)\(_{2.2}/(InAs)\)] SPS]/GaAs is shown in Fig. 1a. Fig. 1b displays the [\(\{1\ 1\ 0\}\)] cross section of the QWRs formed using (GaP)\(_{2}/(InP)\) SPS and Al\(_{0.15}\)Ga\(_{0.85}\)In\(_{0.32}\)P. The light fringes correspond to the higher-energy band-gap material. The 5 QWs existing in the [0 0 1] direction are clearly visible. Lateral modulation occurs in the [1 1 0] direction, the same direction as was observed in previous SILO studies using equal monolayer pairings [6, 7]. The atomic segregation forms wires which lie in the [1 1 0] direction, and the lateral periodicity of the QWR in the [1 1 0] direction is on the order of ~150–200 Å. The interface non-uniformity apparent in the (GaP)\(_{2.2}/(InAs)\) SPS sample is most likely a result of the non-optimal source switching scheme used, where As and P intermixing in the lattice can affect the surface roughness [14]. The general layout of the cross section that the images display is similar to that seen in other instances of the QWRs formed by the SILO process [6, 7].

Fig. 2 displays the PPL spectrum of the [(GaP)\(_{2.2}/(InP)\)] SPS]/AlGaInP QWRs at 77 K. As seen in this figure, evidence of the QWR formation is clearly present due to the highly polarized nature of the PL emission (\(\rho = 0.84\)), with its peak wavelength near 6200 Å. As a reference, the spectra from the surrounding AlGaInP and the GaAsP substrate are shown and are seen to be nominally polarized. For this application of the SILO process, the global strain is \(\Delta S = 0.36\)\% and the mismatch between the SPS and the substrate is \(\Delta T = 0.12\)\%. The room-temperature PPL spectra for the [(GaP)\(_{2.2}/(InAs)\)] SPS]/GaAs based QWRs are shown in Fig. 3. The emission from this structure, near 1.02 μm, is also highly polarized, with a value \(\rho = 0.67\). Using the strain calculations described in a previous section, strain values for this structure are \(\Delta S = 0.7\)\% and the SPS/substrate mismatch is \(\Delta T = 6.6\)\%. The observed luminescence intensity and degree of polarization is comparable to the previous studies of SILO produced QWRs.

It is evident from the periodic fringes in the cross-sectional TEM micrographs and the strong polarization of the PPL spectra that QWRs have been formed in these two heterostructures [15]. These results, in combination with previously reported QWR formation using SPS with equal number monolayer pairings, verify the generalized strain-balance theory and its relation to lateral
composition modulation through the SILO process. Indeed, this method has been employed to generate QWRs across numerous material systems and substrates. Fig. 4 summarizes the distinct emission wavelength regimes from visible to near infrared where the SILO technique has thus far been used to create QWRs [6, 7].

5. Conclusion

Due to the restrictions of certain material systems, previous endeavours into QWR formation by using the SILO process with this technique have used equal monolayer pairings within the SPS. In this paper, we presented a generalized theory of the SILO process where the required number of monolayer pairings within the SPS to maintain strain-balanced conditions has been established. The data gathered for this report provide evidence for the theory's generality by its extension into new material systems (GaInAsP on GaAs and GaInP on GaAsP) as well as new growth schemes (non-equal monolayer SPS pairings). A number of QWR heterostructures with a wide range of PL emissions from visible to near-infrared have been demonstrated. It is therefore seen that the SILO technique is a versatile tool to create functional QWRs in semiconductors.

Acknowledgements

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References


MBE growth of high-power InAsSb/InAlAsSb quantum-well diode lasers emitting at 3.5 μm


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Abstract

Molecular beam epitaxy (MBE) has been employed for the growth of strained quantum-well laser structures on InAs substrates. These lasers consist of compressively strained InAsSb wells, tensile-strained InAlAsSb barriers, and lattice-matched AlAsSb cladding layers. Broadstripe lasers, with emission at wavelengths between 3.2 and 3.55 μm, have exhibited cw power of 215 mW/facet at 80 K, pulsed threshold current density as low as 30 A/cm² at 80 K, characteristic temperatures (T₀) between 30 and 40 K, and maximum pulsed operating temperature of 225 K. Ridge-waveguide lasers have cw threshold current of 12 mA at 100 K, and a maximum cw operating temperature of 175 K. In this paper we will present some of the key issues regarding the MBE growth of such high-power lasers on InAs and discuss future directions for improved device performance.

PACS: 68.55.Bd; 42.55.Px

1. Introduction

A number of different approaches are actively being investigated with the goal of obtaining high-performance mid-infrared (3–5 μm) diode lasers that are capable of high-power cw operation at room temperature. These include laser structures based on conventional electron-hole recombination in quantum-well (QW) and superlattice active regions of either Type I [1] or Type II [2] band alignment, and quantum cascade structures which are based on unipolar intersubband transitions [3]. All of these approaches attempt to overcome limitations, such as nonradiative Auger recombination, that are increasingly important for room-temperature operation of long-wavelength diode lasers. We have previously reported Type I multiple-quantum-well (MQW) diode lasers, grown by molecular beam epitaxy (MBE) on GaSb substrates, consisting of active regions of compressively strained InAsSb wells and tensile-strained InAlAsSb barriers, and lattice-matched AlAsSb cladding layers. Such lasers emitted at 3.9 μm and operated cw up to 128 K [4]. In this paper, we describe our investigation of similar strained MQW diode lasers, based on active regions of InAsSb wells and InAlAsSb barriers, and AlAsSb cladding layers, but grown by MBE on

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InAs substrates. These laser structures are grown with epitaxial compositions adjusted to (a) achieve the desired strains and Type I band alignments in the active region materials and (b) maintain lattice matching of the cladding layers to the InAs substrate. The reasons for investigating such QW structures on InAs substrates will be explained and the resultant performance of high-power, mid-infrared diode lasers will be discussed. We will also describe some of the MBE growth issues which were found to influence the characteristics of these long-wavelength diode lasers.

2. Experimental procedure

The InAsSb QW lasers were grown in a solid-source EPI Gen II 3 in MBE system, equipped with two Sb, sources, an EPI valved As cracking source used to provide As₂ flux, and conventional ion pumps. InAs(100) n-type substrates were prepared by either (a) solvent cleaning and chemical etching in a 4:1:7 H₂SO₄:H₂O₂:H₂O solution, followed by In-bonding for typical substrate material, or (b) directly loaded into a solderless block containing an Al₂O₃ backing plate for 2 in epi-ready substrate material. After careful UHV outgassing and oxide desorption of the substrates in an As₂ flux, the following layers were grown: n⁺-InAs buffer, ~2.0 μm n-AlAs₀.₁₆Sb₀.₈₄ lower cladding, MQW active region consisting of 150 Å compressively strained InAs₀.₉₃₅Sb₀.₀₆₅ wells and 300 Å tensile-strained In₀.₇₅Al₀.₁₅As₀.₉Sb₀.₁ barriers, ~2.0 μm p-AlAs₀.₁₆Sb₀.₈₄ upper cladding, and 500 Å p⁺-GaSb cap layer. These values give approximately 0.45% strain in the wells and 0.27% strain in the barriers, with the strain–thickness product in wells and barriers set to approximately the same value in an attempt to achieve a strain-balanced active region. A cross-section of the laser structure is shown in Fig. 1. Be was used as the p-type dopant and GaTe was the source for Te as an n-type dopant. Hall measurements of nominally undoped InAsSb and InAlAsSb layers grown on semi-insulating GaAs substrates yielded n-type conductivity, as was observed for growth of the related alloys on GaSb. Since the active regions used for this work were undoped, the p–n junction was formed at the active-layer/upper-cladding interface. The two Sb₄ effusion cells allowed some optimization of the III/V ratio for each alloy, when used in conjunction with the valved As₂ source. The buffer and cladding layers were grown at ~0.8–0.9 μm/h, and the MQW active regions were grown at slightly lower growth rate. The substrate temperature was controlled at 520°C for the lower cladding, 430°C for the QW

![Fig. 1. Cross section of InAs QW laser structure.](image-url)
active region, and reduced to 490°C for the upper cladding and GaSb cap layer. As will be shown below, this reduction in upper cladding temperature was necessary to prevent degradation of the active regions. As previously reported [5], the necessity of precise control of composition, strain, and lattice parameter required a number of test growths and calibration procedures.

After dismounting of the In-bonded samples, the In-alloyed region on the back side of the InAs wafer was removed by etching in HNO₃, making sure that the wafer surface was protected during the etching process. Samples grown on the 2 in epiready substrates did not require this additional step, as there was no apparent degradation of the back side of the InAs substrate at the growth temperatures used. The surfaces of the InAs substrates were characterized by atomic force microscopy (AFM), and the epitaxial layers were characterized by photoluminescence spectroscopy, double-crystal X-ray diffraction (DCXRD), and Auger microprobe chemical analysis. The complete MBE-grown laser wafers exhibited excellent surface morphology, and especially for the epiready 2 in substrates, very good yields of laser diodes with consistent characteristics were obtained across the entire wafer.

To evaluate laser performance, broad-stripe lasers 100–250 μm wide were fabricated by using SiO₂ patterning. For both n and p contacts, non-alloyed Ti/Pt/Au was used. Ridge-waveguide lasers, 8 μm wide, were fabricated from some wafers using reactive ion etching in a BCl₃/Ar plasma to define the ridges. Lasers were mounted junction-side up on Cu heat sinks using In, and loaded into a dewar for low-temperature I–V and L–I measurements.

3. Results

Because of the desire to grow complete laser structures on large-area, nonbonded substrates, epiready InAs material was evaluated by AFM. Fig. 2 shows a comparison between conventional InAs material and epiready material from two different vendors, with the respective samples taken directly from the vendor’s packaging to the AFM apparatus. The remarkable smoothness of the epi-ready InAs surface was also confirmed by a comparison of reflection high-energy electron diffraction images obtained before oxide desorption in the growth chamber. In the AFM image of the epi-ready samples there is some evidence of atomic structure on the as-received surface, and this structure was determined to be crystallographically oriented by AFM scanning in different directions.

A key reason for the consideration of the growth of strained InAsSb/InAlAsSb QW active regions on InAs substrates is the presence of a large predicted miscibility gap for the InAlAsSb barrier material [6]. As shown in Fig. 3, changing from GaSb to InAs substrates increases the predicted region of stable growth for InAlAsSb from a maximum Al content of less than 6% to approximately 15%. Increasing the Al content in the InAlAsSb barrier material is important to improve both the conduction and the valence band offset in the active region, for improved carrier confinement and enhanced laser performance [7]. Although these theoretical predictions of regions of stable growth for the InAlAsSb alloy are strictly valid for equilibrium growth conditions, our previous experience with the nonequilibrium, MBE growth of InAsSb/InAlAsSb QWs on GaSb substrates confirmed the practical difficulty of increasing the Al content in this barrier alloy, while still maintaining high-quality QWs.

As an indication of the quality of Inₐ₉₅Sb₀₆₅Tₐ₀₈₅Al₀₁₅As₀₉Sb₀₁ strained MQWs that can be grown on InAs substrates by MBE, Fig. 4 shows a comparison of a 10-well test structure grown on an InAs buffer layer. The excellent agreement between the actual DCXRD data and the simulated diffraction curve shows high structural perfection in the MQW test structure. For these QW structures, the best results were obtained with an As₅₂ interruption at each well/barrier interface. While the use of InAs substrates permits the growth of high-quality MQW active regions, the growth of the Alₐ₀₆Sb₀₈₅ lattice-matched cladding region becomes more difficult, since this alloy now contains ~2X as much arsenic as was the case for the lattice-matched cladding alloy grown on GaSb, and is near a region of
alloy miscibility [8]. In spite of this difficulty, high-quality n- and p-type AlAsSb alloys were grown on InAs. In addition, as was found with the growth on similar structures on GaSb, the transition from Sb-dominated growth in the lower cladding to As-dominated growth in the active regions (and the reverse transition to the upper cladding) was best achieved by enhancing Sb-like interfaces and suppression of As-like interfaces in these transition regions. This observation has also been reported for the growth of InAs/AlSb quantum wells [9].

A more challenging problem with the growth of complete MQW laser structures on InAs was the degradation of the active region caused by the overgrowth of the upper cladding layer at the increased growth temperature necessary for the good-quality material. To investigate this effect, a MQW test structure, similar to that in Fig. 4, was furnace annealed in a sealed ampoule for various times, and DCXRD scans were taken after the annealing. Fig. 5 shows the results of these experiments, where annealing at 530°C for 3 h (a typical growth time for the upper cladding) completely obliterated the satellite structure, while annealing at 510°C for 3 h resulted in less degradation of the satellite structure. A growth temperature of 490°C was chosen as a reasonable compromise between optimizing upper cladding layer electrical and optical quality and active layer degradation.

Broad-stripe diode lasers fabricated from these structures showed emission near 3.4 μm, with pulsed current density as low as 30 A/cm² at 80 K, and characteristic temperatures in the range 30–40 K. By measuring devices with cavity lengths of 500, 1000 and 1500 μm the internal quantum efficiency and internal loss coefficient are estimated to be
Fig. 3. Calculated alloy stability curves for In–Al–As–Sb system (from Ref. 6). The two dotted lines represent alloys matched to GaSb and InAs.

Fig. 4. DCXRD scan and simulation of InAsSb/InAlAsSb QW test structure grown on InAs.
Fig. 5. DCXRD comparison of annealed QW test structures: (a) control sample, (b) after annealing at 510°C for 3 h, and (c) after annealing at 530°C for 3 h.

63% and 9 cm⁻¹, respectively. Fig. 6 shows the cw power versus current for a 1500 μm device at several temperatures. At 80 K the maximum cw output power is 215 mW/facet, limited by junction heating. The maximum output power decreases with temperature, and at 150 K it is ~35 mW/facet. The operating voltage at the maximum power is ~4 V at 80 K. This surprisingly large operating voltage (more than 10 times the photon energy) is caused by the presence of large internal heterostructures, as seen in the band offset diagram, Fig. 7. Substantially higher output power would be expected if these large operating voltages can be reduced by incorporating graded interfaces in the laser structure. Lasers with different active layer QW structures have been evaluated and emission wavelengths at 80 K of 3.21–3.38 μm have been observed. For a laser with 80 K emission at 3.21 μm, emission wavelength versus temperature is

Fig. 6. CW output power versus current for QW broad-area device at various temperatures.
shown in Fig. 8. At low temperatures the wavelength shifts at \( \sim 1 \) nm/K, gradually increasing to \( \sim 2 \) nm/K at 220 K. Ridge-waveguide lasers 8 \( \mu \)m wide exhibited CW threshold current of 12 mA at 100 K and maximum cw operating temperature of 175 K. Additional details on laser performance are reported elsewhere [10].

4. Discussion

Impressive diode laser performance has been obtained at \( \sim 3.5 \) \( \mu \)m with strained InAsSb/InAlAsSb MQW structures grown on InAs. The use of InAs substrates has enabled the growth of InAlAsSb barriers with increased Al content, which in turn
lead to the development of improved QW active regions. The use of InAs substrates, however, further complicated the growth of lattice-matched AlAsSb cladding layers. Although we have not fully optimized the present laser structures, we are confident that by incorporating graded regions to reduce heterobarrier offsets, we can obtain improved performance with reduced operating voltage. With these complex laser structures, careful attention to the growth issues such as strain and lattice matching, thermal effects, and chemistry at interfaces is required for optimum performance.

5. Summary

We have demonstrated that MBE growth on InAs substrates is capable of producing high-performance InAsSb/InAlAsSb QW diode lasers with emission near 3.5 μm. Further improvements in device structures should lead to even higher performance lasers.

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Microstructural properties of InAs/InAs\textsubscript{x}Sb\textsubscript{1-x} superlattices and InAs\textsubscript{x}Sb\textsubscript{1-x} ordered alloys grown by modulated molecular beam epitaxy

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Abstract

Modulated molecular beam epitaxy (MBE) has been demonstrated to be useful in controlling group-V alloys such as InAs\textsubscript{x}Sb\textsubscript{1-x} and AlAs\textsubscript{x}Sb\textsubscript{1-x}. Further studies of the MBE grown InAs\textsubscript{x}Sb\textsubscript{1-x} ordered alloys and InAs/InAs\textsubscript{x}Sb\textsubscript{1-x} superlattices by using cross-sectional scanning tunneling microscopy show well defined interfaces between InAs\textsubscript{x}Sb\textsubscript{1-x} ordered alloys and InAs. Clear composition modulation is observed in the InAs\textsubscript{x}Sb\textsubscript{1-x} ordered alloy layers. Transmission electron diffraction and microscopy, X-ray diffraction, and low temperature photoluminescence experiments show no obvious sign of Cup orderings. These results suggest that MBE provides a possible means to bypass the ordering problem of InAs\textsubscript{x}Sb\textsubscript{1-x} random alloys.

PACS: 81.15.Hi; 61.16; 78.66 - w; 68.65. + g

Keywords: MBE; InAs; InAsSb; InSb; STM; TEM

Control of alloy composition of group-V alloys has been problematic for molecular beam epitaxy (MBE). The problem is even more acute for the epitaxial growth of InAs\textsubscript{x}Sb\textsubscript{1-x} random alloys due to the spontaneous Cup-orderings [1–3]. Such orderings result in substantial bandgap shrinkage [1, 2]. Due to the spontaneous nature of these orderings, it is difficult to accurately and reproducibly control the desired bandgap for optoelectronic device application. Previously, we reported a straightforward and accurate means of controlling the incorporation of As and Sb by rapidly alternating As\textsubscript{2} and Sb fluxes, i.e. modulated MBE (MMBE), for As/Sb group-V alloys [4]. In this paper we will present some latest results of the atomic-scale crystalline properties of the MMBE grown InAs\textsubscript{x}Sb\textsubscript{1-x} ordered alloys and superlattices (SL) using cross-sectional scanning tunneling microscopy (STM), transmission electron diffraction and microscopy (TED and TEM), X-ray
diffraction, and optical spectroscopy. These results show that MMBE provides not only a better control of the composition but also a possible way to bypass the problems of spontaneous CuPt orderings in InAsSb_{1-x} random alloys.

The samples used in the present study were grown by MMBE with a conventional solid-source MBE system equipped with cracking As and Sb cells. The growth rate for all the epilayers was kept around 0.50 μm/h while the growth temperatures were kept around 480–500°C measured by both a calibrated pyrometer and a thermocouple. Different from conventional MBE, MMBE uses a straightforward and accurate means of controlling the incorporation of As and Sb by rapidly modulating As2 and Sb beams during growth. This method will be referred to as MMBE later on. By exploiting this method, the MBE growth yields InAsSb_{1-x} ordered alloys (or InAs/InSb short-period SLs) rather than random alloys. Detailed discussion of MMBE has been published in Ref. [4].

Three types of samples are used for present study. The first sample (Sample A) is a 172 Å InAs/52 Å InAs_{0.76}Sb_{0.24} SL grown on an InAs substrate. Each of the 53 Å InAs_{0.76}Sb_{0.24} layer consists of 4 periods of nominal 7.8 Å InAs/5.2 Å InSb SL structures. The second sample (Sample B) is a laser structure consisting of an InAs/InAs_{0.924}Sb_{0.076} SL active region. The details of the whole laser structure can be found in Ref. [5]. The third type of samples (Sample C and D) has a 3000 Å InAsSb_{1-x} ordered-alloy layer consisting of 250-period 12 Å InAs/InSb short-period SL. This layer is sandwiched by two 200 Å InAs layers grown on GaSb substrates.

The group-V interfaces, such as As/Sb interfaces, have been the topic of extensive study. It has been reported that the interface quality has strong impact on the material structural properties. Since the compositions of ordered alloys are controlled by the shutter duty cycle, it is extremely helpful to understand the detailed atomic structural properties of MMBE-grown samples. Cross-sectional STM can perfectly serve this purpose.

STM experiments for this study focused on Sample A. Fig. 1a shows a constant-current cross-sectional STM image of a single InAs_{0.76}Sb_{0.24} ordered-alloy layer surrounded by InAs layers within a 172 Å InAs/52 Å InAs_{0.76}Sb_{0.24} superlattice. Topographic contrast across the InAs_{0.76}Sb_{0.24} layer, averaged over approximately 200 individual line scans corresponding to this image, is plotted in Fig. 1b. Several features in the structure of the ordered alloy layer and the InAs/InAs_{0.76}Sb_{0.24} interfaces may be noted. Contrast between the InAs and InAs_{0.76}Sb_{0.24} is clearly visible, with the interfaces between the layers appearing abrupt to within ~1–2 monolayers. The four-period ordered structure within the InAs_{0.76}Sb_{0.24} alloy layer arising from the MMBE growth technique is also evident, with the bright regions corresponding to the InSb-like layers.

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Fig. 1. (a) A high-resolution constant-current cross-sectional STM image of a single InAs_{0.76}Sb_{0.24} ordered-alloy layer surrounded by InAs. (b) An averaged topographic line scan of the layer and the surrounding InAs layers.
From the topographic line scan, an increase in contrast, corresponding to increasing Sb content, is observed from the first to the third InSb-like layer. This was seen consistently with different cleaved samples and tips, and suggests that the presence of As cross incorporation is most severe at the start of the InAs$_{0.76}$Sb$_{0.24}$ growth, and decreases as the layer is grown. An asymmetry in interface abruptness is also observed, with the InAs$_{0.76}$Sb$_{0.24}$-on-InAs interface appearing atomically abrupt and the InAs-on-InAs$_{0.76}$Sb$_{0.24}$ interface appearing to be graded over 1–2 monolayers. We believe this grading arises from segregation of Sb on the wafer surface during the InAs$_{0.76}$Sb$_{0.24}$ growth, and incorporation of the segregated Sb into the subsequently grown InAs layer. A detailed discussion of the STM studies may be found in Ref. [6].

TED study of InAs/InAs$_{0.924}$Sb$_{0.076}$ SL active region in a laser structure (Sample B) reveals additional detailed information about the structure properties of the SL. The sample is examined in a [1 1 0] cross-section direction perpendicular to the [0 0 1] growth direction. TED diffraction pattern of the InAs/InAs$_{0.924}$Sb$_{0.076}$ SL active region is shown in Fig. 2. Only zinc-blende reflections appear on the TED image. Special attention is paid to the possible reflections caused by CuPt orderings. No reflection spots at half the distance between (0 0 0) and [1 1 1] reflection spots are visible, indicating that there is no detectable CuPt orderings in the studied InAs/InAs$_{0.924}$Sb$_{0.076}$ SL. The absence of CuPt orderings is further confirmed by photoluminescence study of MMBE grown thick InAs$_{0.9}$Sb$_{0.1}$ ordered-alloy layers almost lattice matched to GaSb substrates. Close lattice match of the epitaxial layer to the substrate allows one to determine the composition more accurately. In this case, no residual or local strains exist, excluding many uncertainties such as the determination of compositions and residual strains.

InAs$_{0.9}$Sb$_{0.1}$ ordered-alloy layers lattice matched to GaSb substrates are used for PL experiments. The average compositions of these layers are determined by X-ray diffraction measurements. As an example, the X-ray diffraction spectrum of Sample D is plotted in Fig. 3a. A distinct sharp X-ray diffraction peak is observed for the 3000 Å InAs$_{0.931}$Sb$_{0.069}$ layer. The small peak on the large angle side is the diffraction from the two 200 Å InAs layers. The average strain in the InAs$_{0.931}$Sb$_{0.069}$ layer is 3.0E-3. Taking into account the layer thickness of only 3000 Å, the epitaxial layer is coherently strained based on the model given in Ref. [7].

The low temperature PL spectrum for the same sample (Sample D) is plotted in Fig. 3b. The bandgaps of free standing InAs$_{1-x}$Sb$_{x}$ random alloys as a function of temperature and alloy composition are given by [8]

$$E_g(\text{Cal}) = 0.411 - \frac{3.4 \times 10^{-4} T^2}{210 + T} - 0.876(1 - x)$$

$$+ 0.70(1 - x)^2 + 3.4 \times 10^{-4}(1 - x)x T^2.$$ 

By taking into account the strain effects [9], the calculated bandgaps [$E_g(\text{Cal})$] of samples C and D are listed in Table 1 together with sample parameters and PL peak positions. FTIR photoluminescence excitation study of similar samples has revealed that there is a very small Stokes
(5–7 meV) shift between PL peaks and the bandgaps for similar samples at low temperatures (~10 K) [10]. Such Stokes shift for low-temperature PL peak is mainly due to localization of photogenerated carriers. At room temperature the photogenerated carriers are delocalized. Therefore, the room temperature PL peak reveals more directly the bandgap. One can see from Table 1 that the agreement between the measured bandgaps with the calculated ones are very good, confirming the absence of CuPt orderings in the As rich InAs$_x$Sb$_{1-x}$ (x ~ 0.9) ordered alloys.

It has been evidenced by TEM and optical spectroscopy studies that there is strong CuPt orderings in InAs$_x$Sb$_{1-x}$ random alloys. These orderings can be clearly observed for even As-rich alloys (x > 0.9) [3] and reaches maximum when x = 0.5 [11]. In comparison with these facts, we believe that the absence of CuPt orderings in the ordered alloys is mainly due to the following reasons: First, the ordered alloys consist mainly of binary–binary short-period InAs/InSb SL. As shown in the STM images the interfaces are well defined although some atomic intermixing exists. These MMBE introduced artificial ordering along growth direction probably hinders the formation of spontaneous CuPt orderings. Second, CuPt ordering is maximum at lower growth temperature (<400°C) [11]. At higher growth temperatures (>400°C), MBE grown InAs$_x$Sb$_{1-x}$ alloy layers show uniform compositions. The samples used in the present study are grown at 480°–500°C. At such high growth temperatures, it is not expected to have CuPt orderings even for MBE grown InAs$_x$Sb$_{1-x}$ random alloys.

Table 1

<table>
<thead>
<tr>
<th>Sample</th>
<th>x</th>
<th>PL peak (meV)</th>
<th>Stocks shifts (meV)</th>
<th>$E_g$(Exp) (meV)</th>
<th>$E_g$(Cal) (meV)</th>
<th>$\Delta E_g$ (meV)</th>
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<tr>
<td>C</td>
<td>0.934</td>
<td>341 (10 K)</td>
<td>6</td>
<td>347</td>
<td>355</td>
<td>8</td>
</tr>
<tr>
<td></td>
<td></td>
<td>301 (300 K)</td>
<td>–</td>
<td>301</td>
<td>301</td>
<td>0</td>
</tr>
<tr>
<td>D</td>
<td>0.931</td>
<td>342 (10 K)</td>
<td>6</td>
<td>348</td>
<td>353</td>
<td>5</td>
</tr>
<tr>
<td></td>
<td></td>
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<td>–</td>
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</tbody>
</table>
In summary, STM and TED studies have demonstrated excellent atomic-scale structural properties of modulated MBE grown InAs$_x$Sb$_{1-x}$ ordered alloys and superlattices. These alloys ($x \sim 0.9$) do not show any sign of CuPt orderings in TED images. No obvious bandgap shrinkage beyond experiment errors has been observed. MMBE technique thus does provide a way to avoid CuPt orderings in As-rich InAs$_x$Sb$_{1-x}$ alloys.

References

Sb-surface segregation and the control of compositional abruptness at the GaAsSb/GaAs interface

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Abstract

Quantitative measurements of Sb surface segregation in pseudomorphic GaAs$_{0.8}$Sb$_{0.2}$ layers grown on GaAs (0 0 1) using molecular beam epitaxy (MBE) were performed in situ using the line-of-sight mass spectrometry (MS) technique. It was observed that substantial surface accumulation of Sb occurs under normal growth conditions, giving rise to compositional broadening of the GaAsSb on GaAs interface. This compositional grading is found to strongly depend on the growth temperature as well as the presence of Sb at the GaAs film surface prior to the initiation of GaAsSb layer growth. We report the effect to substrate temperature and shutter sequence on the evolution of Sb surface accumulation near the interface and demonstrate that a compositionally abrupt interface can be formed by populating the GaAs surface with an amount of Sb equivalent to that expected on the GaAsSb surface during growth at steady state. Additionally, compositional abruptness at the GaAs on GaAsSb interface can be achieved by a “chemical flashoff” of the surface Sb population during exposure to an incident As$_2$ flux.

PACS: 07.75.+h; 68.55.Bd; 68.10.Gw

Keywords: Surface segregation; Interfacial broadening; Mass spectroscopy

1. Introduction

Much of the success of III–V semiconductor devices containing antimony, such as infrared photodetectors [1] and lasers [2], depends on the ability to grow precisely tailored heterostructures. A critical component of this is the formation of compositionally abrupt interfaces, especially between arsenide and Sb-containing layers.

The interface between a ternary and a binary III–V alloy is often compositionally broadened due to the surface segregation of the more weakly bound species occupying the shared sublattice of the ternary alloy [3]. Compositional broadening occurs as a result of the gradual buildup or decay of surface accumulation of this species near the intended interface. For example, in pseudomorphic InGaAs/GaAs heterostructures grown by molecular beam epitaxy (MBE), the transient buildup of
the surface segregated “In floating layer” on InGaAs occurs at the expense of In incorporation into the film near the InGaAs on GaAs interface [4–10]. This results in an In-deficient graded InGaAs region which can be extensive at normal growth conditions. Similarly, the GaAs on InGaAs interface is broadened by the incorporation of the “In floating layer” in the InGaAs layer into the GaAs overlayer [5, 8]. Studies performed to date on interfacial broadening in III–V heterostructures have almost exclusively addressed group-III atom segregation (e.g., Ga on AlGaAs, In on InGaAs, In on InGaP), while little is known about Sb surface segregation and compositional broadening at the GaAs/GaAsSb interface. However, Sb surface segregation during GaAsSb growth can be expected since Sb is considerably more volatile than As [11]. For example, Sb surface segregation on InAs has been observed [12]. Very few in situ techniques allow quantitative measurement of surface accumulation or broadening. We have taken advantage of the fact that a static GaAs0.8Sb0.2 film surface containing a surface segregated population of Sb is unstable when exposed to an incident arsenic flux [13]. Under these conditions, an efficient As/Sb exchange reaction results in the removal of the Sb surface population by desorption. We have performed precise desorption measurements to quantify the “Sb floating layer” using the line-of-sight mass spectrometry (MS) technique and monitored its evolution near the GaAs0.8Sb0.2 on GaAs interface. The data demonstrate that the compositional grading at this interface can be eliminated when the Sb population at the GaAs starting is properly controlled. We also propose to take advantage of the As/Sb exchange to eliminate the “Sb floating layer” in order to form a more abrupt GaAs on GaAsSb interface.

2. Experimental procedure

Film growth and in situ measurements were performed in a Varian Gen-II MBE chamber equipped with a quadrupole mass spectrometer (QMS) mounted onto the center port in the source flange, directly facing the wafer as described in detail elsewhere [14]. The line-of-sight of the ionizing region of the QMS was limited to approximately the central ½ of the radiatively heated 2 in diameter wafer. A variety of GaAs0.8Sb0.2 surfaces were carefully prepared by pseudomorphic growth on GaAs (001) using cracked As2 and Sb2 beams at substrate temperatures Tₜ ranging between 325°C and 515°C. The GaAs0.8Sb0.2 growth rate was 0.75 ML/s, and the entire incident antimony flux (J(Sb2) = 0.15 ML/s) was incorporated (i.e., unity Sb sticking coefficient throughout the Tₜ range) to give the nominal Sb-mole fraction, x(Sb) = 0.2.

The surface accumulated population of antimony ΘSb was measured using the QMS by quantifying the amount of antimony purged from the surface as a result of As/Sb surface exchange. It was previously shown that the QMS Sb signal from the monomer (mass/q = 121 or 123 amu/e) is proportional to the total Sb-flux leaving the film surface within a wide range of temperatures and surface composition [14]. The QMS Sb signal versus time profile shown in Fig. 1 describes a typical measurement of ΘSb following the deposition of a thin GaAs0.8Sb0.2 layer. Initially, both the As2 and Sb2 beams are coincident on the static GaAs surface, and a constant QMS Sb signal is observed. At this stage, the QMS Sb signal is proportional to the total antimony desorption rate which is equivalent to the total incident Sb2 flux of 0.15 ML/s. Based on this measurement, the QMS Sb signal during

![Fig. 1. Evolution of QMS Sb signal as GaAs0.8Sb0.2 layer growth is initiated, interrupted, and As/Sb exchange occurs.](image-url)
the remainder of the experiment can be calibrated with the desorption rate. As shown in Fig. 1, the Sb desorption signal is observed to drop to zero when the Ga cell shutter is opened and alloy growth is initiated, indicating that all of the incident Sb$_2$ is consumed and the Sb$_2$ sticking coefficient is indeed unity. In the case shown in Fig. 1, both the Ga and Sb$_2$ fluxes are interrupted after 35 ML of GaAs$_{0.8}$Sb$_{0.2}$ growth at $T = 450^\circ$C, and the film surface remains exposed only to the incident As$_2$ beam. Immediately following the growth interruption, a sharp rise in the QMS Sb signal is observed which is generally followed by a gradual decay. This indicates that antimony is purged from the surface of the film and that the surface becomes slowly depleted of Sb in the presence of the As$_2$ beam. The area under this feature can be quantified based on the calibration described above, and was measured to be $\sim 0.62 \pm 0.05$ ML for the case shown in Fig. 1. This is considerably larger than the expected 0.2 ML based on x(Sb), verifying substantial Sb surface accumulation in GaAs$_{0.8}$Sb$_{0.2}$ at these growth conditions. Assuming steady-state conditions are reached after 35 ML of growth, $\Theta^{\text{Sb}}$ is comparable to $\sim 0.79$ ML of In measured to accumulate at the surface of a 90 Å thick In$_{0.22}$Ga$_{0.78}$As layer deposited on GaAs at the same temperature in a previous study [10].

3. Results

3.1. Evolution of the surface segregated layer

When $\Theta^{\text{Sb}}$ reaches a steady-state value during film growth, the nominal GaAsSb alloy composition can be attained because the presence of the “Sb floating layer” does not alter the Sb sticking coefficient. However, it is expected that a transient buildup of surface Sb population occurs near the GaAsSb on GaAs interface where $\Theta^{\text{Sb}}$ is initially zero. In order to measure the evolution of $\Theta^{\text{Sb}}$ as a function of the distance from the interface, a series of GaAs$_{0.8}$Sb$_{0.2}$ layers of varying thickness were deposited on GaAs at $T = 450^\circ$C. In each case an antimony desorption measurement was performed during As/Sb exchange following growth interruption in a manner similar to that described above. In this series, the GaAs static surface was bathed only in As$_2$, while Sb$_2$ was introduced together with Ga to initiate alloy growth. Fig. 2 shows the measured values of $\Theta^{\text{Sb}}$ as a function of nominal GaAs$_{0.8}$Sb$_{0.2}$ layer thickness. It is observed that $\Theta^{\text{Sb}}$ gradually increases from 0.31 $\pm$ 0.05 ML after 3 ML of GaAs$_{0.8}$Sb$_{0.2}$ deposition to 0.62 $\pm$ 0.05 ML after 10 ML of GaAs$_{0.8}$Sb$_{0.2}$ deposition where a steady-state value $\Theta^{\text{Sb}}(ss)$ of 0.62 $\pm$ 0.05 ML is approached. The observed evolution of $\Theta^{\text{Sb}}$ suggests that the nominal value of x(Sb) = 0.2 is not reached until nearly 30 Å from the interface, and that the graded GaAsSb layer at the interface is deficient in Sb. Based on the measured value of $\Theta^{\text{Sb}}$ after 3 ML of GaAs$_{0.8}$Sb$_{0.2}$ growth, the average Sb mole fraction in the first 3 ML of the alloy is $\sim 0.10$, significantly lower than the intended value of 0.2.

The GaAsSb on GaAs interface can also be formed using an alternative shutter sequence: the GaAs static surface is bathed in both As$_2$ and Sb$_2$ prior to opening the Ga shutter to initiate GaAsSb as described in Fig. 1. In this case, a steady-state surface population of Sb is already present at the GaAs surface and can be expected to influence the evolution of $\Theta^{\text{Sb}}$ when growth is initiated. A series of $\Theta^{\text{Sb}}$ values measured over incrementally thicker GaAs$_{0.8}$Sb$_{0.2}$ layers deposited on GaAs in this
manner is also shown in Fig. 2. It was observed that \( Sb \) is initially large (\( Sb = 0.90 \pm 0.05 \) ML after \( \sim 1 \) ML deposition of GaAs\(_{0.8}Sb_{0.2}\)) and decays to the steady-state value of \( \sim 0.62 \) ML after nearly 10 ML of alloy deposition. The lack of Sb desorption signal during the growth of the GaAs\(_{0.8}Sb_{0.2}\) layer implies that the excess Sb at the film surface near the interface is incorporated into the film. This results in a compositionally graded GaAsSb layer at the interface which contains excess Sb in contrast to the result of the previously described shutter sequence. Assuming that the observed reduction in \( Sb \) within the first 3 ML of GaAs\(_{0.8}Sb_{0.2}\) is homogeneously incorporated into the alloy, the Sb mole fraction adjacent to the interface is \( \sim 0.25 \), significantly higher than the intended value of 0.2.

3.2. Control of compositional abruptness

The evolution of \( Sb \), as demonstrated above, depends strongly on the amount of Sb at the initial surface prior to the deposition of the GaAsSb layer. It has been previously demonstrated that if a surface accumulated population equivalent in size to the steady-state value on the ternary alloy surface can be maintained at the initial static surface, then the transient behavior of \( Sb \) near the interface, thus compositional grading, can be eliminated [9, 10]. The fact that we can populate the starting surface in excess of \( Sb \) as shown in Fig. 2 suggests that the surface Sb population on GaAs, \( Sb \) (GaAs), can be matched to \( Sb \) (GaAs) under appropriate conditions.

To search for growth conditions where \( Sb \) = \( Sb \) (GaAs), the effect of \( Sb \) on \( Sb \) (GaAs) and \( Sb \) (GaAs) was studied. A series of GaAs\(_{0.8}Sb_{0.2}\) layers were deposited on GaAs at \( T_s < 515^\circ C \) using the same incident fluxes, and \( Sb \) (GaAs) was measured. The layer thickness in each case was 35 ML in order to ensure that steady-state conditions were achieved. It was observed that \( Sb \) (GaAs) is as large as \( 1.20 \pm 0.05 \) ML at \( T_s = 515^\circ C \), and is reduced to \( 0.36 \pm 0.05 \) ML at \( T_s = 325^\circ C \), as plotted in Fig. 3. Also shown in Fig. 3 are the measured values of \( Sb \) (GaAs) for a range of \( T_s \).

In this case, \( Sb \) (GaAs) was measured by blocking off the incident Sb flux on the GaAs static surface irradiated by both As\(_2\) and Sb\(_2\), and attributing the decay of the QMS Sb signal to the surface buildup of antimony. We observe the general trend that \( Sb \) increases with \( T_s \) while \( Sb \) (GaAs) decreases with \( T_s \), and that a crossover point exists where \( Sb \) (GaAs) = \( Sb \) (GaAs).

Data shown in Fig. 3 indicate that, given the incident fluxes used, a crossover point where \( Sb \) (GaAs) = \( Sb \) (GaAs) exists near \( T_s = 485^\circ C \). As is generally observed for surface segregating species, \( Sb \) (GaAs) increases with increasing \( T_s \); however, \( Sb \) (GaAs) has an inverse relationship with \( T_s \). \( Sb \) (GaAs) is observed to decrease from \( 0.90 \pm 0.05 \) ML at \( T_s = 450^\circ C \) to \( 0.71 \pm 0.05 \) ML at \( T_s = 520^\circ C \). Both of these trends are expected since surface segregation is kinetically limited [7, 15] while the stoichiometry on a static surface is not.

To measure the evolution of \( Sb \) on GaAs\(_{0.8}Sb_{0.2}\) layers near the interface when \( Sb \) (GaAs) is matched to \( Sb \) (GaAs), a set of GaAs\(_{0.8}Sb_{0.2}\) surfaces were prepared at \( T_s = 485^\circ C \). In situ measurements of \( Sb \) using line-of-sight mass spectrometry as described earlier is shown in Fig. 4. It is observed that \( Sb \) remains constant at a value of \( 0.82 \pm 0.05 \) ML throughout the region near the GaAs\(_{0.8}Sb_{0.2}\) on GaAs interface. The lack of a transient in \( Sb \) indicates that Sb is incorporated into the film at the steady-state level and the Sb-deficient region near the interface is eliminated.
The formation of a compositionally abrupt GaAs on GaAsSb interface, on the other hand, necessitates the elimination of the “Sb-floating layer” prior to the deposition of GaAs. As demonstrated in Fig. 1, we propose to achieve this by interrupting the growth at this interface to expose the surface to the incident As$_2$ flux. This will ensure that the Sb population at the GaAsSb surface will be purged so that GaAs growth can proceed without surface riding and incorporation of Sb.

4. Discussion

Following the kinetic model of surface segregation developed by Muraki [7] in which only the exchange between the top GaAsSb monolayer and the “Sb floating layer” is allowed, a segregation ratio $R$ can be assigned. $R$ is defined as the fraction of Sb atoms at the top layer which segregate to the floating layer after each new monolayer of growth, and is given by $1 - [x(Sb)/\Theta_{Sb}^{SS}]$. $R$ is measured to be 0.76 at $T_s = 485^\circ C$ in these experiments, and is comparable to the values reported for In in InGaAs (0.78 < $R$ < 0.85) at similar growth temperatures [6, 10].

The growth temperature determined to be necessary to eliminate compositional grading near the GaAs$_{0.8}$Sb$_{0.2}$ on GaAs interface is specific to the incident fluxes and growth rate used, as well as nominal x(Sb) in the alloy. Based on the results of previous investigations of In surface segregation on InGaAs [5, 10, 16], it is expected that $\Theta_{Sb}^{SS}$ will increase as x(Sb) is increased. However, it is not clear whether $T_s$ will have to decrease to yield $\Theta_{Sb}^{SS}(GaAs) = \Theta_{Sb}^{SS}(ss)$ for larger x(Sb). This is because $\Theta_{Sb}^{SS}(GaAs)$ is also expected to increase as a larger incident Sb$_2$ flux is employed to give a larger x(Sb). It is clear, however, that arbitrary choices of growth parameters can result in interfacial broadening for which the magnitude and the direction of the compositional grading is not known. Heterostructures for devices which depend on compositionally abrupt interfaces can therefore benefit from such experiments to fine-tune the growth parameters.

These qualitative effects of segregation are expected in other alloy systems; thus compositional grading in InAsSb, AlAsSb and quaternary III–II–AsSb layers near the binary III–As interface is expected.

5. Conclusion

Line-of sight mass spectrometry was used to investigate Sb surface segregation on GaAs$_{0.8}$Sb$_{0.2}$ during MBE growth by quantifying the desorbed antimony as a result of As/Sb surface exchange. It was observed that size of the surface segregated Sb population at steady state increases with $T_s$, and that its evolution near the GaAs$_{0.8}$Sb$_{0.2}$ on GaAs interface results in compositional broadening. We have developed a growth approach based on controlling the surface Sb population which is expected to result in compositionally more abrupt interfaces than can be obtained by standard MBE growth. Our data suggests that the compositional grading can be eliminated when the shutter sequencing and $T_s$ is precisely chosen so that $\Theta_{Sb}^{SS}(GaAs)$ is well matched to $\Theta_{Sb}^{SS}(ss)$.

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References

MBE growth of Si-doped InAlAsSb layers
lattice-matched with InAs

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Abstract

The two-dimensional electron gas concentration in InAs channel modulation-doped heterostructures can be controlled by using an In0.5Al0.5As0.56Sb0.44 quaternary alloy – doped with Si as a donor – as an electron-supplying layer. When lattice-matched with InAs, an electron concentration of more than $4 \times 10^{18}$ cm$^{-3}$ was obtained in the quaternary layer. The two-dimensional electron gas concentration of an undoped-In0.5Al0.5As0.56Sb0.44/undoped-InAs heterostructure was $7 \times 10^{11}$ cm$^{-2}$ and that of a Si-doped In0.5Al0.5As0.56Sb0.44/undoped-InAs heterostructure was $1.24 \times 10^{12}$ cm$^{-2}$. The electron mobility was low, though, due to the roughness of the heterointerface. If the interface can be made smooth, however, In0.5Al0.5As0.56Sb0.44 will be a useful material for forming the electron-supplying layer of InAs channel modulation-doped heterostructures.

PACS: 68.55.Bd; 73.61.Ey

Keywords: Two-dimensional electron gas; Modulation-doped heterostructures; InAs channel; Electron-supplying layer

1. Introduction

Al(Ga)(As)Sb/InAs heterostructures have been widely studied for future high-frequency field effect transistor (FET) applications because of their very high electron mobility which exceeds 20000 cm$^2$/V·s at room temperature (RT) [1–6]. In this material system, two-dimensional electron gas (2DEG) concentrations of the order of $10^{13}$ cm$^{-2}$ are known to accumulate in the InAs layer even though no layers are intentionally doped. In this heterostructure, the 2DEG concentration can be changed by changing the top barrier thickness [7, 8]. However, it is hard to control the concentration without changing the top barrier thickness, because n-type doping of Al(Ga)Sb is difficult. The Si, which behaves only as a donor for (Al)GaAs, turns into partially compensated acceptors in GaSb and AlSb [9]. Although column VII materials, such as S, Se, and Te, become donors for GaSb and AlSb [9], the vapor pressure of these materials is too high for accurate control of the doping concentration in MBE. Chow et al. reported that Si-modulation-doped InAs/AlSb
superlattices yield controllable n-type carrier concentration up to $1 \times 10^{18}$ cm$^{-3}$ [10]. However, for FET application, more than $3 \times 10^{18}$ cm$^{-3}$ of carrier concentration is thought to be required.

To overcome these problems, we propose the use of a Si-doped In$_{0.5}$Al$_{0.5}$As$_{0.56}$Sb$_{0.44}$ layer as an electron-supplying layer lattice-matched on the InAs channel in this paper. We have found that Si behaves as a donor in In$_{0.5}$Al$_{0.5}$As$_{0.56}$Sb$_{0.44}$ through our investigation of the electronic properties of Si-doped In$_{0.5}$Al$_{0.5}$As$_{1-x}$Sb$_x$ ($x < 0.5$). Finally, we demonstrate here that the 2DEG concentration in the InAs layer can be controlled by changing the Si-doping concentration in the In$_{0.5}$Al$_{0.5}$As$_{0.56}$Sb$_{0.44}$ electron-supplying layer.

2. Predicted site occupation of Si in InAlAsSb

According to the prediction by Pindoria et al., Si occupies the site by replacing the largest atom in III–V compounds under normal growth conditions (i.e., usually a group-V flux rich condition) [11]. The most direct confirmation of this size effect comes from the observed gradual shift from n-type to p-type in Si-doped GaAs$_{1-x}$Sb$_x$ [12]. When the Sb content is below 0.2, the Si-doped GaAs$_{1-x}$Sb$_x$ layers are n-type. When the Sb content is above 0.25, on the other hand, the Si-doped GaAs$_{1-x}$Sb$_x$ layers become p-type. The experimental site occupation, which depends on the V/III atomic size ratio (the Sb content), for GaAs$_{1-x}$Sb$_x$ is shown in Fig. 1. The tetrahedral covalent radius of the element is used as the atomic size [13]. As shown, the transition from n-type to p-type occurs at a V/III atomic size ratio of about 1.03. The predicted site occupation, which depends on the Sb content of the Si-doped In$_{0.5}$Al$_{0.5}$As$_{1-x}$Sb$_x$, is also shown in Fig. 1. If we assume that the n/p-type transition of Si-doped In$_{0.5}$Al$_{0.5}$As$_{1-x}$Sb$_x$ occurs at the same V/III atomic size ratio as GaAs$_{1-x}$Sb$_x$, we can predict that Si-doped In$_{0.5}$Al$_{0.5}$As$_{1-x}$Sb$_x$ will be n-type when the Sb content is below 0.7. Since the Sb content of In$_{0.5}$Al$_{0.5}$As$_{0.56}$Sb$_{0.44}$ is 0.44, Si-doped In$_{0.5}$Al$_{0.5}$As$_{0.56}$Sb$_{0.44}$ will be n-type and can be used as an electron-supplying layer for InAs channel modulation-doped heterostructures.

Therefore, we have decided to concentrate on studying a Si-doped In$_{0.5}$Al$_{0.5}$As$_{1-x}$Sb$_x$ alloy ($x < 0.5$) for use as electron-supplying layers of InAs channel modulation-doped heterostructures.

3. Material growth and characterization

All samples for this study were grown with a conventional solid-source MBE system. After thermal cleaning of semi-insulating (1 0 0) GaAs substrates at 600°C in an As$_4$ flux, we successively grew an undoped In$_{0.5}$Al$_{0.5}$As layer (300 nm), an undoped In$_{0.5}$Al$_{0.5}$As$_{1-x}$Sb$_x$ layer (1 μm), a Si-doped In$_{0.5}$Al$_{0.5}$As$_{1-x}$Sb$_x$ layer (110 nm, Si concentration: $7.3 \times 10^{18}$ cm$^{-3}$), and an undoped In$_{0.5}$Al$_{0.5}$As$_{1-x}$Sb$_x$ layer (110 nm). The substrate temperature was fixed at 350°C. The growth rate of the In$_{0.5}$Al$_{0.5}$As(Sb) layer was about 1.2 μm/h. The As$_4$ pressure at the sample position was fixed at $1 \times 10^{-5}$ Torr, and the Sb$_4$ pressure was varied from 0 to $9.6 \times 10^{-7}$ Torr. The Al and In content was calibrated by double-crystal X-ray diffraction (DXRD) using an In$_{0.52}$Al$_{0.48}$As layer grown on an InP substrate. Two In$_{0.5}$Al$_{0.5}$As$_{0.56}$Sb$_{0.44}$/InAs heterostructures were also grown, one modulation-doped and one undoped (see Fig. 2). An AlSb/InAs heterostructure was also grown as a reference.
The mobility and free electron concentration of the samples were evaluated by the conventional van der Pauw method at RT (297 ± 3 K).

The $\text{In}_{0.5}\text{Al}_{0.5}\text{As}_{0.5}\text{Sb}_{0.44}$/InAs heterointerface quality was investigated directly by transmission electron microscopy (TEM).

4. Results and discussion

The lattice constant and Sb content of $\text{In}_{0.5}\text{Al}_{0.5}\text{As}_{1-x}\text{Sb}_x$ as a function of the Sb$_x$ beam pressure is shown in Fig. 3. The lattice-matching with InAs is given at an Sb beam pressure of about $9 \times 10^{-7}$ Torr.

The free electron concentration and mobility of Si-doped $\text{In}_{0.5}\text{Al}_{0.5}\text{As}_{1-x}\text{Sb}_x$ as a function of the Sb content is shown in Fig. 4. The Si-doping concentration was fixed at $7.3 \times 10^{18}$ cm$^{-3}$. All layers showed n-type conduction as we predicted in Section 2. The free electron concentration decreases from $5.76 \times 10^{18}$ to $4.21 \times 10^{18}$ cm$^{-3}$ as the Sb content increases from 0 to 0.47. The discrepancy between the doping concentration and the free electron concentration at the Sb content of 0 is due to dislocations in the epitaxial layer, because there is a large mismatch in the lattice constants of the
GaAs substrate and the In$_{0.5}$Al$_{0.5}$As epitaxial layer. When we grew Si-doped In$_{0.52}$Al$_{0.48}$As on an InP substrate as a reference, an electron concentration of $6.9 \times 10^{18}$ cm$^{-3}$ was obtained. By improving the buffer layer quality, larger electron concentration can be obtained on GaAs substrates [14].

The mobilities, on the other hand, were almost constant and remained between 356 and 383 cm$^2$/V·s. These values are comparable with that of the reference In$_{0.52}$Al$_{0.48}$As grown on an InP substrate at the same doping concentration (360 cm$^2$/V·s).

These results indicate that In$_{0.5}$Al$_{0.5}$As/AlSb heterostructures (Fig. 2). The InAlAsSb layer of sample #1 was intentionally left undoped and the InAlAsSb layer of sample #2 was Si-doped at a concentration of $1 \times 10^{18}$ cm$^{-3}$. An AlSb/InAs heterostructure was also grown as a reference. The result of the Hall measurement is listed in Table 1. The 2DEG concentrations of the undoped and Si-doped modulation-doped heterostructures were $7 \times 10^{11}$ cm$^{-2}$ and $1.24 \times 10^{12}$ cm$^{-2}$, respectively. This is the first time that the 2DEG concentration in InAs channel modulation-doped heterostructures has been controlled by Si-doping using a lattice-matched electron-supplying layer. The mobility, however, was very small compared with that of the AlSb/InAs heterostructure. We believe this is due to the rough heterointerface between the InAlAsSb and the InAs.

To investigate the roughness of the InAlAsSb/InAs heterointerface, we used TEM (Fig. 5). As you can see in the TEM micrograph, the InAlAsSb/InAs heterointerface was very rough and many dislocations were observed. This suggests that the surface smoothing effect of InAlAsSb is weaker than that of an AlGaAsSb material system. When an AlGaAsSb material system was used as a buffer layer in an InAs channel heterostructure, very high mobility (over 20 000 cm$^2$/V·s) was obtained even without using a superlattice as a smoothing layer [5]. So we must improve the buffer layer quality of InAlAsSb/InAs material systems to obtain a smooth interface and high mobility.

5. Conclusions

The 2DEG concentration in InAs channel heterostructures can be controlled by using a Si-doped In$_{0.5}$Al$_{0.5}$As$_{0.56}$Sb$_{0.44}$ quaternary alloy as an electron-supplying layer. Under growth conditions suitable for lattice-matching with InAs, an electron concentration of more than $4 \times 10^{18}$ cm$^{-3}$ was obtained in the Si-doped In$_{0.5}$Al$_{0.5}$As$_{0.56}$Sb$_{0.44}$ quaternary layer. The 2DEG concentration of the
undoped-In_{0.5}Al_{0.5}As_{0.5}Sb_{0.44}/undoped-InAs heterostructure was $7 \times 10^{11}$ cm$^{-2}$ and that of the Si-doped In_{0.5}Al_{0.5}As_{0.5}Sb_{0.44}/undoped-InAs heterostructure was $1.24 \times 10^{12}$ cm$^{-2}$. This is the first time that the 2DEG concentration in InAs channel heterostructures has been controlled by Si-doping. The mobility, however, was lower than that of the AlSb/InAs heterostructure due to the rough heterointerface. This indicates that In_{0.5}Al_{0.5}As_{0.5}Sb_{0.44} will be a suitable material for use as the electron-supplying layer for InAs channel modulation-doped heterostructures as soon as the buffer layer quality can be improved.

References

Molecular beam epitaxial growth of GaInSbBi for infrared detector applications

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Abstract

The growth of a new quaternary semiconductor alloy GaInSbBi suitable for 8–12 µm infrared detector applications by molecular beam epitaxy is reported for the first time. By the introduction of Ga into the InSbBi alloy, the following have been achieved: (1) enhancement of Bi incorporation, (2) improved lattice-matching to InSb, and (3) suppression of multiple-phase growth and improved surface morphology. The dependence of Bi content and surface morphology on growth temperature and substrate orientation has been investigated. Significant improvement in surface morphology is achieved for films grown on large angle off-axis substrates including (3 1 1)A and (5 1 1)A. Infrared absorption measurements performed at 77 K indicate that a cutoff wavelength as long as 10.7 µm was achieved. The principles demonstrated in this work are quite general and can be applied to other quaternary material systems containing N, P, As, and Sb such as GaInTIsb and GaInAsN to cover a wide range of band gaps.

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Keywords: Molecular beam epitaxy; Fourier transform infrared spectroscopy; Infrared detector

The InSb$_{1-x}$Bi$_x$ material system has undergone much investigation in recent years due to its potential application in 8–12 µm infrared detectors as the band gap can be varied from that of InSb (~0.23 eV at $T = 0$ K) to a metallic phase with increasing $x$. Joukoff and Jean-Louis first reported the growth of bulk InSb$_{1-x}$Bi$_x$ using the Czochralski technique [1]. However, a solid solubility limit of 2.6 mol% InBi was found due to the different crystal structures of InSb (zincblende) and InBi (tetragonal). Subsequently, nonequilibrium growth techniques such as multi-target sputtering [2–4] and molecular beam epitaxy (MBE) [5, 6], were employed in order to increase Bi incorporation. Although Bi content appeared to have been enhanced from secondary ion mass spectroscopy (SIMS) measurements, poor growth morphology and the formation of multiple phases were shown to degrade the material quality. All previous results [5–9] showed no indication of an extension of a clear band-edge absorption to a longer wavelength. Most likely, the Bi measured by SIMS was not incorporated in the crystal lattice.

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Given the covalent atomic radii of In, Sb and Bi (1.44, 1.40, and 1.46 Å, respectively), it is clear that the lattice constant of the epilayer is increased with increased Bi content, thus increasing the lattice mismatch to the InSb substrates. As a result, Bi, the largest atom, is squeezed out of its lattice site in the lattice-mismatched layer; therefore, inducing the formation of a second phase and limiting the Bi content.

In this work, we report the growth and optical properties of Ga$_{0.04}$In$_{0.96}$Sb$_{0.97}$Bi$_{0.03}$ films grown by MBE on InP substrates and demonstrate that increased Bi content in high-quality crystals can be achieved by the incorporation of Ga into the InSbBi alloy. In addition, (3 1 1)A and (5 1 1)A-oriented growth is shown to increase the incorporation of Bi and improve surface morphology further.

In our experiments, Ga was incorporated into the InSbBi alloy in order to reduce the lattice constant and improve the lattice match to InSb (the covalent atomic radius of Ga is 1.26 Å, which is much smaller than that of Sb and Bi). Although the introduction of Ga to InSb increases the band gap, the increase is minimal for a low Ga content due to the band gap narrowing near the InSb end of the ternary alloy (the band gap for Ga$_{x}$In$_{1-x}$Sb is given as 0.172 + 0.139x + 0.415x$^2$ at 300 K). In addition, film composition can be easily controlled by MBE as GaInSb is a group III alloy with only one group V element. This technique of “lattice compensation” is widely applicable and may also be applied to other narrow gap compounds, such as adding Ga to InTISb and InAsN.

The dependence of surface morphology on substrate orientation is best understood when one considers the surface bonding structure of various substrate orientations [10]. The atoms on the (1 0 0) surface each has one double-dangling bond, while each (1 1 1)A surface atom has one single-dangling bond. The (5 1 1)A surface (16° off (1 0 0) toward (1 1 1)A), as shown in Fig. 1, is composed of both single- and double-dangling bond sites as the flat (1 0 0) terrace consists of two terrace atoms, each with double-dangling bonds, and one step-edge atom with one single-dangling bond. For (n 1 1)A-oriented substrates, the step-edge group III atom has a very stable configuration [10], providing a favorable bond-site for Sb and Bi. We have previously demonstrated that AlGaAs grown on the (n 1 1)A-oriented substrates exhibited superior electrical and optical properties [10].

In our experiments, we observed that the large angle off-axis substrates (i.e., (3 1 1)A and (5 1 1)A) were critical for the incorporation of Bi in the films and for sustaining two-dimensional crystal growth. Although Bi content did not increase significantly for the (3 1 1)A and (5 1 1)A-oriented films as compared to the (1 0 0)-oriented films, the surface morphology for (3 1 1)A and (5 1 1)A is significantly superior, most likely due to the absence of multiple phases. Most importantly, all GaInSbBi films grown on the (1 0 0)-orientation did not show an extension of the cutoff wavelength. Only films grown on (3 1 1)A and (5 1 1)A exhibited a band-edge that was clearly extended to longer wavelengths.

Ga$_{x}$In$_{1-x}$Sb$_{1-y}$Bi$_{y}$ films were grown in a Varian Gen II MBE system on (1 0 0), (3 1 1)A, and (5 1 1)A InP substrates using Sb$_{3}$ and Bi sources. The InP substrates were chemically etched with a H$_2$SO$_4$ : H$_2$O$_2$ : H$_2$O (5 : 1 : 1) solution, mounted on Mo-blocks and then loaded into the chamber. In order to accommodate the large lattice mismatch between the GaInSbBi film and the InP substrate, a 1 µm AlSb intermediate buffer layer was grown at 500°C, followed by a 1 µm InSb epilayer grown at 400°C. The substrate temperature was measured by an infrared pyrometer. Then the Ga$_{0.04}$In$_{0.96}$Sb$_{1-x}$Bi$_{x}$ films were grown at a growth rate of 0.5 µm/h. The Sb/In flux ratio was maintained near unity in order to produce a mirror-smooth InSb buffer layer surface [11], while the Ga/In flux ratio was used to control the Ga...
content. Reflection high-energy electron diffraction (RHEED) was employed in order to provide in situ monitoring, enabling the maintenance of a slightly Sb-rich condition which is necessary to prevent the formation of metal droplets.

The dependence of the surface morphology and optical absorption of GaInSbBi films on growth temperature was investigated for the substrate temperature range 300 and 380°C. Within this range of substrate temperature investigated, a higher growth temperature was found to result in a superior surface morphology, which was also evidenced from the streaky RHEED patterns during crystal growth indicating a two-dimensional nucleation. Lower substrate temperatures often resulted in more spotty RHEED patterns possibly due to the lower surface mobility of adatoms at lower temperatures. However, optical absorption measurements using Fourier transform infrared spectroscopy (FTIR) indicated that the GaInSbBi films grown at lower substrate temperatures exhibited an absorption edge at longer wavelengths (corresponding to an increased Bi content). This indicated that the sticking coefficient of Bi is higher at lower substrate temperatures, which is consistent with previous reported results [5, 6].

The Sb flux was varied during the growth of the GaInSbBi films in order to investigate the nucleation competition between Sb and Bi. Our results indicated that an Sb-rich condition resulted in a nearly-zero Bi sticking coefficient, while an Sb-deficient condition (i.e. Sb/(Ga + In) ratio slightly less than unity) enhanced the Bi incorporation. Again, this is consistent with previously reported results [5, 6].

The FTIR spectrum at 77 K (Fig. 2) indicated an extended absorption wavelength, 10.7 μm, for Ga_{0.04}In_{0.96}Sb_{1 - x}Bi_x films grown under Sb-deficient conditions. In addition, two absorption regions were observed, corresponding to the InSb and Ga_{0.04}In_{0.96}Sb_{1 - x}Bi_x films. Based on a 36 meV decrease in energy gap per atomic percentage of Bi increase [12], Bi content was estimated to be roughly 3% (Fig. 3).

![Graph](image)

**Fig. 2.** The 77 K absorption spectrum of a Ga_{0.04}In_{0.96}Sb_{0.95}Bi_{0.05}(511)A film.
In conclusion, Ga$_{0.04}$In$_{0.96}$Sb$_{0.97}$Bi$_{0.03}$ suitable for 8–12 μm infrared detector applications has been grown by MBE on (1 0 0), (3 1 1)A and (5 1 1)A InP substrates. The FTIR results indicate that a cutoff wavelength as long as 10.7 μm has been achieved for films grown on the (3 1 1)A and (5 1 1)A orientation but not for (1 0 0). Also, significant improvement in surface morphology was observed for films grown on (3 1 1)A and (5 1 1)A. The principles demonstrated in this work are quite general and can be applied to other quaternary material systems containing N, P, As, and Sb such as GaInTISb and GaInAsN to cover a wide range of tunable band gaps.

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References

Large mismatch heteroepitaxy of InSb on Si(111) substrates using CaF$_2$ buffer layers

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Abstract

The results of initial attempts to grow large lattice-mismatched InSb/Si structures using CaF$_2$ buffer layers are reported. Substrate temperatures in the range of 300–400 °C were used and MBE growth was initiated by opening the In and Sb shutters simultaneously, producing In-terminated InSb(111)-A surfaces on Si(111) substrates. High structural quality was confirmed by reflection high-energy electron diffraction, electron channeling and high-resolution X-ray diffraction. Electron mobility as high as 65 000 cm$^2$/V s for a density of $\sim 2 \times 10^{16}$ cm$^{-3}$ was measured at room temperature for an 8 μm-thick InSb layer grown on CaF$_2$/Si(111).

1. Introduction

The maturity of integrated circuit technology and the availability of low-cost, rugged, large-area wafers have made Si the most attractive substrate material for semiconductor thin film growth. With advancements in molecular beam epitaxy (MBE) technology, heteroepitaxy has been achieved on Si substrates with a lattice mismatch as large as 20% [1-12]. InSb has the smallest bandgap, highest intrinsic electron mobility and lowest electron effective mass of all binary III-V compounds. These characteristics make InSb-based structures candidates for infrared devices, high speed transistors and magnetic field sensors. Monolithic integration of InSb devices on Si substrates thus offers advantages in microelectronics and infrared detector array fabrication.

Compared to heteroepitaxy of GaAs on Si ($\sim 4\%$ mismatch) and CdTe on Si ($\sim 19\%$ mismatch), very little has been reported on heteroepitaxy of InSb on Si [8–13]. The main challenges in the growth of InSb on Si stem from the large lattice mismatch ($> 19\%$), the different thermal expansion coefficients ($\alpha_{\text{InSb}} \approx 2 \alpha_{\text{Si}}$ at 300 K) and antiphase domain (APD) formation (polar on non-polar growth for (0 0 1) and (1 1 1) orientations). Tilted substrates [9–11], In pre-deposition [8] and the insertion of various buffer layers [8, 10–14] have been used to alleviate some of these problems.

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Group IIa fluoride layers have been widely used as buffers for the growth on Si of CdTe [1–3] (whose lattice constant is close to that of InSb) and PbSe [7, 15, 16]. As evident from Fig. 1, the ~19% lattice mismatch between InSb and Si can be reduced through graded layers of CaF₂ and BaF₂. Recently, thin InSb films have been grown on Si substrates using a BaF₂/InSb/BaF₂ sandwich-type buffer [10].

MBE growth of high quality CaF₂ and BaF₂ films on Si substrates has been demonstrated by several groups [16]. The CaF₂/Si(1 1 1) interface was found to be primarily Ca–Si bonds with a neutral overlayer containing complete F⁻–Ca₂⁺ + F⁻ triple layers [18, 19]. The CaF₂(0 0 1) surface has a perpendicular dipole moment and the repeat unit consists of Ca⁺–F⁻ double layers [17]. It may therefore be possible to grow APD-free InSb on these polar CaF₂ surfaces. Moreover, fluoride layers are also known to be effective in relieving thermal mismatch stress in heteroepitaxial systems through gliding dislocations [7, 16, 17, 20]. Their low elastic stiffness coefficients, particularly that of BaF₂, can also provide additional strain relief via plastic deformation [20–22].

Although the fluorides have much larger thermal expansion coefficients than Si, fluoride films cooled to room temperature after high temperature growth on Si generally do not contain cracks [17]. For thick CdTe/CaF₂/Si structures, near vanishing strain was measured in the CdTe layer and the CaF₂/Si interface is believed to be responsible for the strain relief of the entire structure [7]. The low substrate temperatures used for InSb growth should also help to minimize thermal mismatch problems in InSb/CaF₂/Si structures.

In this article, we explore the relatively new use of group IIa fluoride buffer layers in the MBE growth of InSb on Si(1 1 1) substrates. Detailed reflection high-energy electron diffraction (RHEED) and scanning electron microscopy (SEM) studies on the initial growth of this heteroepitaxial system are presented. The interfacial properties are investigated using X-ray photoelectron spectroscopy (XPS), the crystalline quality is confirmed using high-resolution X-ray diffraction and the electrical properties of the InSb films are characterized using Hall-effect measurements.

2. Experimental procedure

MBE growth of InSb/fluoride/Si structures was carried out in an Intevac Gen II system with an analysis chamber and two growth chambers: one for fluoride growth and the other for InSb growth. Three-inch diameter Si(1 1 1) substrates (Silicon Sense, Inc.) were cleaned using the Shiraki method [23] and the passivating oxide was thermally desorbed in the fluoride growth chamber at 1100°C, producing well-defined Si(1 1 1)(7 × 7) RHEED patterns. Substrate temperatures (T sub) were measured by a thermocouple located at the center of the substrate heater and could differ from real sample temperatures by as much as 100°C.

A high-purity polycrystalline CaF₂ source was evaporated from a graphite-coated PBN crucible while a background pressure of ~ 10⁻¹⁰ Torr was maintained. The Si substrate was rotated at 10 rpm and heated to 700°C. Typically, 1000 Å of CaF₂ was deposited at ~ 10 Å/min. This growth rate corresponded to a beam equivalent pressure of ~ 7 × 10⁻¹⁰ Torr. Details of fluoride growth on Si substrates have been reported elsewhere [24].

Following fluoride deposition, the samples were cooled to room temperature and transferred under
ultra-high vacuum to another growth chamber for subsequent InSb deposition using solid elemental sources of In (7 N, Nikko Kyodo) and Sb (6.5 N, Dowa). In most cases, a thin nucleation layer (2000 Å thick) was first grown at a low growth rate (~ 0.1 ML/s) and a low temperature (~ 300°C) to reduce the concentration of crystal defects at the interface [25, 26]. A $T_{\text{sub}}$ from 350°C to 400°C, a growth rate of ~ 0.65 ML/s and an Sb$_4$ to In beam equivalent pressure ratio of ~ 3 were used for growth of the active layer. Growth rates were inferred from RHEED intensity oscillations observed during InSb growth on GaAs(0 0 1) substrates. Growth was interrupted occasionally in the early stages to improve crystallinity and surface smoothness.

A Varian electron gun was operated at 9.5 kV and an angle of incidence between 1° and 3° to monitor the RHEED patterns. Digitized images of these patterns were obtained using a CCD camera and a data acquisition system developed by k-Space Associates, Inc. We have previously demonstrated that RHEED characterization, in spite of the known effects of electron-beam-induced fluorine desorption, can be used to study fluoride surfaces [24]. Electron beam irradiation time on the fluoride layers was kept to a minimum and RHEED patterns recorded at the initial InSb growth stages were taken from previously unexposed areas. These precautions prevent modification of the starting surface by the electron beam and ensure that the observed diffraction features reflect actual growth morphology.

An adjoining analysis chamber was used to conduct XPS experiments at room temperature, thereby eliminating complications associated with the use of capping layers and exposure to atmosphere. The analysis chamber was equipped with a VG100AX hemispherical analyzer and an XR3E2 dual-anode X-ray source used in conjunction with a VGX900 data acquisition system (Fisons Instruments). The base pressure of this chamber was kept below 10$^{-9}$ Torr during the experiments. XPS spectra arising from In 3d, Sb 3d, Ca 2p and F 1s transitions were obtained by running the Al K$_\alpha_{1,2}$ source ($h\nu = 1486.6$ eV) at a 15 keV potential and a 10 mA emission current. The constant analyzer energy mode was used with a pass energy of 20 eV.

SEM micrographs were obtained using a JEOL JSM880 system (15 kV potential, ~ 10 nA emission current) and electron channeling patterns were taken at a 20° scan angle and a 25 kV potential. X-ray diffraction data (0-20 scans) were obtained using a Philips Materials research diffractometer operated with a 4-bounce Ge(2 2 0) incident-beam monochromator.

### 3. Results and discussion

Fig. 2 illustrates the evolution of the RHEED patterns during the growth of InSb/CaF$_2$/Si(1 1 1). The smooth transition from the Si(1 1 1)-(7 × 7) to the CaF$_2$(1 1 1)-(1 × 1) surface reconstruction (Fig. 2a and Fig. 2b) provides evidence that CaF$_2$ growth proceeds two-dimensionally. Parallel epitaxy is confirmed by comparing the electron channeling pattern of the bare Si substrate with one taken after growth of the CaF$_2$ film (Fig. 3a and Fig. 3b). The six-fold symmetry characteristic of (1 1 1)-oriented layers is found in both patterns. Fewer details are observed in the CaF$_2$ pattern as a result of a higher defect density in the CaF$_2$ layer than in the Si substrate.

Subsequent growth of InSb was initiated by exposing the CaF$_2$ layer to In and Sb fluxes simultaneously. The emergence of more closely spaced InSb-related spots in the RHEED pattern (Fig. 2c) indicated that nucleation took place via the formation of 3D epitaxial islands, as expected from the large lattice mismatch. The cloudy appearance of the sample’s surface arose from the high density of dislocations at the interface. As growth proceeded, CaF$_2$ diffraction streaks weakened until disappearing after deposition of ~ 10 ML of InSb (Fig. 2d). Meanwhile, the InSb diffraction features became brighter and 1/2-order spots began emerging in both the[1 1 0] and [1 1 2] azimuths (Fig. 2d). At an equivalent layer thickness of ~ 25 ML, 1/2-order streaks could be clearly seen (Fig. 2e). The reflectance of the surface also improved at this stage but was not uniform across the 3 in. wafer. The coalescence of differently-sized islands with differing degrees of strain relaxation, and hence different lattice constants [27], may explain these observations. After depositing 1000 Å of InSb,
Fig. 2. Evolution of RHEED patterns during the growth of InSb/CaF$_2$/Si(1 1 1): (a) Si(1 1 1)-(4×4); (b) 1000 Å CaF$_2$(1 1 1)-(1×1); initial growth of InSb: (c) ~ 2 ML, (d) ~ 10 ML, (e) ~ 25 ML; after the growth of a 0.4 μm film showing (f) well-defined InSb(1 1 1)-(2×2) and (g) static InSb(1 1 1)-(2×6) surface reconstructions.
growth was interrupted while the substrate temperature was raised to 375°C before further growth resumed. Fig. 2f shows a well-defined \((2 \times 2)\) RHEED pattern for a 4000 Å InSb film. It is worth noting that InSb films grown without the low-temperature, low-growth-rate buffer appeared less smooth but more uniform. Increasing the initial growth temperature above 375°C also roughened the surface. When growth was terminated and the substrate temperature lowered under an Sb flux, a change from \((2 \times 2)\) In-stabilized to \((2 \times 6)\) Sb-stabilized surface reconstruction was observed (Fig. 2g). This transition was reversible and is characteristic of an In-terminated InSb(1 1 1)A surface [28, 29]. This behavior differs from that of GaAs grown on CaF\(_2\)/Si(1 1 1), where an As-terminated (1 1 1)B surface is produced [30].

XPS spectra of In 3d, Sb 3d, Ca 2p and F 1s signals were taken at various stages of growth. Within the resolution limit of our XPS system (\(\sim 1\) eV), no interfacial products were detected. The normalized photoemission intensities of the substrate peaks decrease and the epilayer peaks increase exponentially with increasing InSb coverage. The composition of the InSb film appears stoichiometric throughout the growth.

The crystalline quality, epitaxial orientation and surface morphology of the epilayer were further investigated using SEM and optical microscopy. Non-uniformity in film thickness is detected under SEM by the presence of curved growth steps on an otherwise fairly smooth surface. Optical micrographs taken under Nomarski interference contrast revealed the presence of features related to stacking faults (Fig. 4). Their density was found to increase dramatically with growth temperature. The initial low-temperature, low-growth-rate buffer also helped reduce the density of these defects. Occasionally, extended cracks along the \(<1 1 0>\) directions were observed hundreds of microns apart in films thicker than several microns. Also, sawtooth-shaped microcracks were observed to be most pronounced around defects and can be attributed to the pinning of strain-relieving misfit dislocations [17]. The sharpness and six-fold symmetry of the electron channeling pattern for an 8 µm InSb film (Fig. 3c) is similar to that of the underlying CaF\(_2\)/Si layer (Fig. 3b). This indicates that \{1 1 1\} parallel
epitaxy is achieved and that the InSb layer is not twinned. The high crystalline quality indicated by the channeling pattern was confirmed by X-ray diffraction measurements where the full width at half maximum for the InSb layer was 125 arcsec. This value is comparable to that reported for material systems with similar lattice mismatch such as CdTe/Si [2].

To assess the electrical properties of the InSb layers, Hall-effect measurements were carried out at room temperature and 77 K. The results for InSb layers with different growth parameters are summarized in Table 1. Note that variations in the electrical properties were detected across each wafer due to the non-uniform epitaxial growth expected in a system with such a large lattice mismatch. In general, smoother areas yielded higher electron mobilities. All samples were found to be n-type with room-temperature mobilities in the $10^5$ cm$^2$/Vs range. Electron mobility as high as 65 000 cm$^2$/Vs ($n \approx 2 \times 10^{16}$ cm$^{-3}$) was obtained in an 8 µm film grown with a 0.3 µm low-temperature, low-growth-rate buffer. This compares favorably to the best room-temperature values reported to date for heteroepitaxial InSb films [8, 11, 12, 31, 32]. The 77 K mobilities, however, were at least an order of magnitude lower than the room temperature values. Similar behavior has been reported for other InSb-on-Si structures and is attributed to electron scattering from dislocations arising from both lattice and thermal strains [8, 10]. A conversion to p-type conduction at low temperature, previously reported for InSb films grown on BaF$_2$ substrates and attributed to thermal strain [33], was not observed.

It has been reported that only a very narrow growth window exists for optimal GaAs/CaF$_2$ films grown on on-axis Si(1 1 1) substrates [30, 34, 35]. The stringent requirement for optimal growth, however, can be relaxed by growing on off-axis substrates [34, 35]. In addition, the initial nucleation of GaAs can be improved by the formation of an As layer on the CaF$_2$/Si(1 1 1) surface [30, 36, 37]. The epitaxial quality of InSb/CaF$_2$/Si(1 1 1) may also be improved by applying an Sb soak prior to growth on off-axis substrates.

![Nomarski micrograph of an 8 µm thick InSb/CaF$_2$/Si(1 1 1) film revealing the presence of stacking-fault-related features.](image)

### Table 1

<table>
<thead>
<tr>
<th>Sample</th>
<th>Structure</th>
<th>Growth parameters$^a$</th>
<th>$\mu_{RT}$ (cm$^2$/Vs), $n_{RT}$ (cm$^{-3}$)</th>
<th>$\rho_{77K}$ (cm$^2$/Vs), $n_{77K}$ (cm$^{-3}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>S139</td>
<td>InSb/CaF$_2$/Si(1 1 1)</td>
<td>$d_s = 2$ µm, $d_L = 0.1$ µm, $T_{sub} = 375$°C</td>
<td>24 500–28 500; $n = 2.0 \times 10^{-14}$ cm$^{-3}$</td>
<td>–</td>
</tr>
<tr>
<td>S140</td>
<td>InSb/CaF$_2$/Si(1 1 1)</td>
<td>$d_s = 3$ µm, $d_L = 0$ µm, $T_{sub} = 375$°C</td>
<td>13 000; $n = 1.9 \times 10^{-16}$ cm$^{-3}$</td>
<td>–</td>
</tr>
<tr>
<td>S141</td>
<td>InSb/CaF$_2$/Si(1 1 1)</td>
<td>$d_s = 5$ µm, $d_L = 0.2$ µm, $T_{sub} = 350$°C</td>
<td>29 020; $n = 1.7 \times 10^{-16}$ cm$^{-3}$</td>
<td>1686; $n = 2.3 \times 10^{-15}$ cm$^{-3}$</td>
</tr>
<tr>
<td>S144</td>
<td>InSb/CaF$_2$/Si(1 1 1)</td>
<td>$d_s = 8$ µm, $d_L = 0.3$ µm, $T_{sub} = 350$°C</td>
<td>48 647–65 518; $n \approx 2.0 \times 10^{-16}$ cm$^{-3}$</td>
<td>4715; $n = 1.6 \times 10^{-15}$ cm$^{-3}$</td>
</tr>
</tbody>
</table>

$^a$ $d_s$: thickness of active InSb layer, $d_L$: thickness of low-temperature, low-growth rate InSb buffer.
4. Conclusions

Epitaxial growth of InSb on large lattice-mismatch Si(1 1 1) substrates was achieved by MBE using CaF$_2$ buffer layers. Structural and room-temperature electrical data indicate that Si(1 1 1) can be used as an alternative substrate for InSb growth. However, detailed study of the influence of growth parameters such as substrate orientation, shutter opening sequence, flux ratio, growth temperature and buffer layer thickness on the crystalline quality need to be done before the prospect of growing vertically stacked single-crystal narrow-gap compound semiconductor/insulator/Si structures can be realized.

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References

Indium antimonide doped with manganese grown by molecular beam epitaxy

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Abstract

Indium antimonide is of interest for infrared detecting and emitting devices and for magnetic field sensors. In this study, indium antimonide doped with manganese and grown by molecular beam epitaxy was investigated. Secondary ion mass spectroscopy (SIMS) was used to show that the incorporation of manganese is near unity over a wide range of manganese concentrations. Manganese is observed to be an acceptor with a dopant efficiency which follows a power law in which the hole density is proportional to the manganese concentration raised to the power $\alpha$. The power $\alpha$ depends on the growth temperature; at 300°C, $\alpha = 0.86$ and at 360°C, $\alpha = 0.78$. Lightly manganese doped samples have transport dominated by electrons at low temperatures due to hole freeze out, followed by holes at intermediate temperatures and finally by intrinsic electrons at high temperatures. Additional SIMS studies showed that manganese diffuses relatively slowly in indium antimonide.

Keywords: Dopant; Diffusion; Magnetoresistor

1. Introduction

Indium antimonide is of interest for magnetic position sensors, transistors, infrared detectors and infrared emitters. InSb has a narrow energy band gap of 0.18 eV and a very large phonon limited mobility of 78 000 cm$^2$ V$^{-1}$ s$^{-1}$ at 300 K. In general, these devices require well-controlled donor and acceptor impurities. Studies of acceptors in epitaxial films have included the p-type dopants Zn [1], Cd [1], Be [2-4], and to some extent C [5]. Of these dopants, Zn and Cd have generally been the preferred dopants for InSb films grown by metal organic chemical vapor deposition (MOCVD), partially because of environmental concerns about volatile beryllium compounds. Be has been the p-type dopant most frequently used in InSb films grown by molecular beam epitaxy (MBE) despite some problems such as a doping efficiency of about 0.5 or less and a tendency to rapidly migrate toward the growing InSb film surface. Carbon produces highly compensated material which may be p-type at cryogenic temperatures [5].
Manganese has not been previously studied in epitaxial InSb films to our knowledge. However, bulk InSb crystals doped with manganese have been investigated [6, 7]. It was found to be a monovalent acceptor with an ionization energy which varied from 9.5 meV at low Mn concentrations to zero at a Mn concentration of about $1 \times 10^{17}$ cm$^{-3}$. The solubility was about $5 \times 10^{18}$–$1 \times 10^{19}$ cm$^{-3}$. Thus, it was the aim of the current work to examine the properties of Mn doped InSb films grown by molecular beam epitaxy.

2. Experimental techniques

We have previously reported on the growth of InSb by MBE [8, 9]. The equipment and growth techniques used were similar in the present case. The MBE source ovens contained indium, antimony, and manganese, all of 99.9999% purity or higher. The antimony source was not cracked, and thus consisted mostly of Sb$_4$. The vacuum during deposition was $\sim 1 \times 10^{-9}$ Torr. A temperature stabilized quartz crystal deposition monitor could be moved into the growth position before or after growth to measure the fluxes of In, Sb, and Mn. The Sb/In flux ratio (corrected to equivalent atomic fluxes) was 1.3 during these growths. Secondary ion mass spectroscopy (SIMS) studies were done at the University of Western Ontario using a Cameca IMS-3f instrument.

3. Experimental results: growth issue

The flux of manganese at the sample position vs. the inverse Mn oven temperature was very linear for Mn fluxes corresponding to $\sim 1 \times 10^{18}$–$1 \times 10^{20}$ Mn cm$^{-3}$ (at an InSb growth rate of 1 μm/h) and Mn fluxes at lower oven temperatures were found by extrapolation. Such data allows absolute impinging flux calculations. Measurements of the In and Sb fluxes before and after each growth are much easier since their fluxes are several orders of magnitude larger. A sequence of Mn-doped InSb films was grown at a substrate temperature of 300°C or 360°C and with various Mn fluxes. These films were analyzed with SIMS to determine the ratio $^{55}$Mn/$^{121}$Sb. These data are shown in Fig. 1. The straight line is a fit that assumes a linear relationship between the impinging Mn flux and the Mn incorporated into the film. This appears to be a good assumption for all except the highest Mn fluxes in films grown at the higher (360°C) substrate growth temperature. Since the solid solubility of Mn in bulk InSb crystals was found to be about $5 \times 10^{18}$–$1 \times 10^{19}$ cm$^{-3}$ [6], the fact that incorporation of Mn at 360°C is no longer linear with Mn flux above these levels suggests that a bulk solubility limit is being reached. However, at 300°C the Mn fully incorporates in the bulk of the film. This implies that a metastable solid solution is attained or possibly that an Mn-rich phase has precipitated in the bulk of the films. However, there is a “missing mass” problem for the heavily doped films grown at 360°C. Presumably, the missing Mn either reevaporated from the InSb surface during growth or

![Fig. 1. The isotope ratio $^{55}$Mn/$^{121}$Sb determined for several Mn-doped InSb films grown at two different substrate growth temperatures.](image-url)
surface segregated on the growing surface. This latter possibility was examined immediately after growth by transferring a film under vacuum to an analytical chamber which had an Auger electron spectrometer. The Auger spectrum of a film grown at 360°C and doped with Mn at a concentration $1.5 \times 10^{20} \text{ cm}^{-3}$ showed about 0.1 monolayer of Mn on the surface. While this is a non-negligible amount, it is far less than the amount that appears to be missing from the bulk of the film as shown in Fig. 1. This suggests that a significant amount of Mn reevaporates from the InSb surface at a substrate temperature of 360°C and at large Mn concentrations. This reevaporation may involve rejection of some of the Mn from the bulk because of solid solubility limitations resulting in a buildup of Mn on the surface, enhancing the probability of reevaporation of an Mn-rich specie.

An important criterion for a dopant is that it should have an acceptably low solid state diffusion coefficient at the sample growth temperature. To study Mn diffusion, an InSb film was grown at 360°C which consisted of 1000 nm of undoped material followed by 100 nm doped at $N_{\text{Mn}} = 2.4 \times 10^{18} \text{ cm}^{-3}$.

Fig. 2. SIMS depth profile of Mn in InSb grown at 300°C or at 360°C. Only the region between 200 and 300 nm was intentionally doped.
which in turn was followed by 200 nm of undoped material. A nominally identical film was grown at 300°C for comparison. SIMS depth profiles of $^{55}$Mn in these films are shown in Fig. 2. In general, a very well-defined Mn-doped layer is observed in each film, as expected. A small Mn peak is also observed at the InSb/substrate interface. This may be caused by leakage of Mn flux around the Mn oven shutter before growth. Ion bombardment effects may play a role in broadening the profiles, but the fact that a much broader peak is seen for the sample grown at 360°C at low Mn concentration implies a solid state diffusion mechanism. Further data is needed to quantitatively study the diffusion mechanism. The diffusion profiles appear to have approximately symmetrical shapes down to low Mn concentrations. This is much different from the case of Be diffusion in InSb in which anomalously large diffusion toward the surface is seen at high concentrations at the higher growth temperatures. Thus Mn is relatively easy to control and it incorporates well in InSb except at the highest concentrations. Itdiffuses slowly except at low concentration at 360°C.

4. Experimental results: transport studies

Hall effect and magnetoresistivity data were acquired from 2 to 473 K on all samples. The samples were photolithographically defined mesas of the epitaxial films shaped as six-probe Hall bars. The DC Hall resistivity and magnetoresistivity were calculated as the averages over the two polarities of magnetic field and current. In the lowest-doped samples, electrons dominate transport at the lowest temperatures, holes at intermediate temperatures, and intrinsic electrons at high temperatures. Evidence for two and even three-carrier conduction is thus visible in the field dependence of the Hall resistivity and magnetoresistivity near the n-to-p and p-to-n transition temperatures. The

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Fig. 3. The carrier concentration of two manganese-doped films as a function of temperature inferred from the inverse of the Hall coefficient (from Fig. 5). Positive values refer to holes and negative values to electrons.

Fig. 4. The mobility of carriers for two InSb films both doped with manganese at a concentration of $5 \times 10^{16} \text{ cm}^{-3}$. 
highest-doped samples exhibit negative magnetoresistance because of magnetic scattering. This further complicates the analysis of the data in terms of multi-carrier conduction. For this report, we choose to use only the low-field data ($\mu B \ll 1$, where $\mu$ is the mobility of the carrier with the highest mobility). Thus, we used 0.065 T $\geq$ B for $T < 70$ K, and 0.036 T $\geq$ B for 70 K $<$ T $<$ 473 K. From these low-field data, we compute the density and mobility of these high-mobility carriers, which dominate the electrical transport.

The approximate density of the high-mobility carrier ($> 0$ for holes, $< 0$ for electrons), obtained by simply using the inverse of the Hall coefficient, is shown in Fig. 3 for two InSb films both doped with manganese at a concentration $5 \times 10^{16}$ cm$^{-3}$. The divergences near the n-to-p and p-to-n transitions are unphysical, because multi-carrier conduction dominates in that regime. Conduction is only dominated by holes in an intermediate temperature range for the sample grown at 360°C. Holes freeze out allowing electrons to dominate conduction below 33 K in this sample because the Mn acceptor level is above the valence band edge [6]. These electrons apparently are caused by residual donors as unintentional background impurities. Electrons

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**Fig. 5.** The hole density versus manganese concentration measured at 64 K for two film growth temperatures.
again dominate conduction above 142 K because of the high density of intrinsic carriers. A somewhat similar behavior is observed for the sample grown at 300°C, except that the holes only partially freeze out at low temperatures. The mobility of carriers for these same two samples is shown in Fig. 4. One noteworthy feature is that the high temperature electron mobilities approach 30 000 cm² V⁻¹ s⁻¹ at 470 K, which is close to the value 34 600 cm² V⁻¹ s⁻¹ which we attain in undoped InSb films at this temperature. This implies that this concentration of Mn (5 × 10¹⁶ cm⁻³) does not scatter electrons very strongly at this temperature.

The low-temperature n-to-p transition does not occur in samples with higher Mn concentrations because the ionization energy of the Mn level decreases with Mn concentration [6]. It can be seen in Fig. 3 that at an intermediate temperature, near 64 K, the hole density is reasonably temperature-independent (the curve for T₉ = 360°C in Fig. 3 is the worst case). Therefore, we assume that the hole concentration at 64 K represents the saturation hole density induced by the Mn-doping.

The hole densities and mobilities of Mn-doped InSb at 64 K are shown in Figs. 5 and 6, respectively. We observe a power-law relationship, \( p \sim [\text{Mn}]^n \).

![Graph showing hole mobility versus hole density measured at 64 K for two film growth temperatures.](image)

Fig. 6. The hole mobility versus hole density measured at 64 K for two film growth temperatures.
which is closer to linear for films grown at 300°C ($\alpha = 0.86$) than for films grown at 360°C ($\alpha = 0.78$). Our films had a linear relationship between Mn incorporation and the Mn flux during growth except at Mn levels near $10^{19}$ cm$^{-3}$ (see Fig. 1). Thus, except at the highest Mn levels, the sub-linear relationship in Fig. 5 cannot be explained by problems with Mn incorporation in the InSb film. Since manganese was previously found to be a monovalent acceptor in InSb bulk crystals, our results suggest that the lattice site at which manganese incorporates is dependent upon growth parameters. Films grown at 360°C had a less linear relationship between Mn and hole densities. This result is rather similar to one recently reported for Mn incorporation in InAs [10]. In that study, it was reported that MBE-grown InAs films had Mn on the In lattice sites for low growth temperatures (near 200°C) or at low concentrations (below 1 at%). Thus, we speculate that the increasingly sub-linear dependence of hole density on Mn concentration in InSb at higher film growth temperature may imply that an increasing fraction of Mn atoms are not occupying the normal In lattice sites where Mn acts as a normal acceptor impurity in InSb. The hole mobility at 64 K is shown as a function of the hole density at that temperature in Fig. 6.

![Graph](image_url)

**Fig. 7.** Transition temperature above which conduction is dominated by electrons versus manganese concentration. The intrinsic carrier density (determined from an undoped InSb film) is also shown versus temperature on the same axes.
The above discussion was based upon transport measurements at 64 K. Transport measurements were made over the range 2–473 K. As the temperature was increased, the dominant carrier type changed from holes to electrons. The transition temperature at which this occurred is shown in Fig. 7 as a function of Mn concentration and film growth temperature. This transition is caused by the dependence of the intrinsic electron density on temperature and by the fact that electrons have about two orders of magnitude higher mobility than holes in InSb. Thus, when the intrinsic electron density is over an order of magnitude lower than the Mn concentration, the intrinsic electrons will begin to dominate conduction. We determined the intrinsic electron density from an InSb undoped film which was grown under conditions similar to those used in the current study. It was 2.8 μm thick, and had a low temperature background electron density of about $1 \times 10^{15} \text{cm}^{-3}$. Over the temperature range for which data are shown in Fig. 7 (271–471 K), the electron mobility in our undoped InSb film decreased from 66000 to 34600 $\text{cm}^2 \text{V}^{-1} \text{s}^{-1}$. These electron mobilities are roughly two orders of magnitude higher than the hole mobilities generally measured for Mn-doped InSb in this temperature range.

Thus, manganese has been shown to be a useful, well-controlled acceptor dopant for InSb grown by molecular beam epitaxy. While only films grown by molecular beam epitaxy have been studied so far, it appears to be feasible to grow InSb films by metal organic chemical vapor deposition (MOCVD) which are doped with manganese using a compound such as tricarbonyl manganese. This has not yet been studied.

References

Molecular beam epitaxial growth of high electron mobility InAs/AlGaAsSb deep quantum well structures

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Abstract

We investigated InAs deep quantum well structures made from InAs/AlGaAsSb materials on GaAs substrates by molecular beam epitaxy. By accurately controlling Asx/Sbx and Asx/In beam flux ratios, we were able to grow the high-quality crystals of AlGaAsSb and InAs. Very high electron mobility of more than 32,000 cm²/Vs were obtained at room temperature. It was found that the optimum beam-equivalent-pressure (BEP) ratio of Asx/In for growing an InAs quantum well layer is 110 to get the highest electron mobility. Moreover, a strong photoluminescence (PL) peak of AlGaAsSb band-edge emission at 10 K was observed around 1.2 eV. We determined that the optimum Asx/Sbx BEP ratio is 6.7 for AlGaAsSb from the full width at half-maximum of PL peaks and electrical characteristics. We also observed the blue shifted PL of AlGaAsSb as the Asx/Sbx BEP ratio decreased.

1. Introduction

The InAs/AlGaAsSb material system is of interest for its high electron mobility and large conduction band offset of ~1.3 eV at the InAs/AlGaAsSb heterojunction. Fig. 1 shows a schematic cross section and energy band diagram of InAs deep quantum well structures (DQWs). In recent years, the InAs/AlSb quantum well (QW) system has been studied by several groups [1–5]. Bolognesi et al. [5] achieved high electron mobilities of 33,000 cm²/Vs in the InAs/AlSb system, using a nucleation layer, buffer layers as thick as a few microns of AlSb, smoothing superlattices, and InSb-like interface controls. However, InAs/AlSb materials have a disadvantage for use in practical devices because AlSb easily becomes oxidized during the device fabrication process.

We proposed new InAs DQWs, made from InAs/AlGaAsSb materials, which are reliable enough for practical use [6, 7]. In our InAs DQWs, the InAs QW layer is sandwiched between quaternary AlGaAsSb buffer/barrier layers that are lattice-matched to InAs because of the introduction of As. Furthermore, Ga is incorporated into the buffer/barrier layers to suppress the oxidation of
the antimonide layers. Our InAs DQWs consist of five simple layers, including a submicron-thick AlGaAsSb layer. Although neither superlattices nor interface controls are employed during the growth of InAs DQWs, very high electron mobilities of more than 30000 cm$^2$/Vs have been obtained [8]. Moreover, we have applied InAs DQWs to Hall sensors as a practical device and have obtained high sensitivities, small dependence of input resistance on temperature, and good reliability [7].

In this paper, we report on the As$_4$/In beam-equivalent-pressure (BEP) ratio dependence of electrical characteristics of InAs DQWs. We also discuss the optimum As$_4$/Sb$_4$ BEP ratio during the growth of AlGaAsSb buffer/barrier layers from the full width at the half-maximum (FWHM) of the photoluminescence (PL) peak of AlGaAsSb band-edge emission at 10 K.

2. Experimental procedure

All samples were grown on semi-insulating GaAs (1 0 0) substrates in a specially modified VG Semicron V100 MBE system, equipped with elemental group III and group V solid sources, the latter producing As$_4$ and Sb$_4$ beams.

The growth was initiated with a 150 nm GaAs buffer layer at a substrate temperature of 580$^\circ$C. The substrate temperature was measured by an optical pyrometer during the entire time of the InAs DQWs growth. Following the growth of a 600 nm Al$_{0.5}$Ga$_{0.5}$Sb$_{0.5}$As$_{0.5}$Sb$_{0.5}$ buffer/barrier layer at 540$^\circ$C, 15 nm of not-intentionally doped InAs, a 10 nm of Al$_{0.5}$Ga$_{0.5}$Sb$_{0.5}$Sb$_{0.5}$, and a 5 nm of GaAsSb cap layer were grown. Al$_{0.5}$Ga$_{0.5}$Sb$_{0.5}$Sb$_{0.5}$ and GaAsSb cap layer were not intentionally doped. The As content $y$ in Al$_{0.5}$Ga$_{0.5}$As$_{1-y}$Sb$_y$ lattice-matched to InAs is 0.12. The energy gap of Al$_{0.5}$Ga$_{0.5}$As$_{0.1}$Sb$_{0.88}$ is about 1.36 eV. The substrate temperature was gradually decreased to 450$^\circ$C prior to the start of the InAs growth and held there for the remainder of the growth [7]. The Al/Ga ratio was calibrated by measuring the reflection high-energy electron diffraction (RHEED) oscillation of AlAs and GaAs. The BEPs for In and Sb$_4$ were measured just under the substrate holder using the movable ion gauge before raising the arsenic temperature. The BEP of Sb$_4$ was the average of measurements before and after the growth.

The growth rate of AlGaAsSb was 800 nm/h. The growth rate of InAs changed from 400 to 1000 nm/h when the V/III ratio was investigated by changing In temperature.

The morphology of the InAs surface grown on AlGaAsSb buffer/barrier layer was analyzed by high-resolution scanning electron microscopy (HRSEM). The electron mobilities at room temperature were determined by van der Pauw measurement. PL spectra were measured at 10 K using a Ge detector cooled to 77 K. A 488.0 nm line of an Ar$^+$ laser was used as a pump source with an excitation density of 5 W/cm$^2$.

3. Results and discussion

The surface structure of the grown epilayers was analyzed in situ by RHEED patterns during the entire time of the InAs DQWs growth. As soon as the AlGaAsSb growth was initiated on the GaAs surface, the streaked (2 × 4) RHEED pattern became spotty instantly because of a 7% lattice mismatch between the GaAs and AlGaAsSb. However, the strain relaxation of AlGaAsSb was very fast and the spot RHEED pattern turned into a strongly streaked (1 × 3) pattern after growing only a 35 nm thick of AlGaAsSb layer [8]. After 600 nm of AlGaAsSb growth, the InAs layer was grown without growth interruption and interface control. Just after the InAs layer growth started, the RHEED pattern changed from a streaked (1 × 3) to a weakly
streaked \((1 \times 1)\) pattern. This indicates that the InAs layer two-dimensionally grew on the AlGaAsSb buffer layer. We also confirmed that the InAs layer on AlGaAsSb grew two-dimensionally by atomic force microscopy, as reported in Ref. [8]. The strongly streaked \((1 \times 3)\) pattern recovered as soon as the top AlGaAsSb barrier layer started to grow. This streaked \((1 \times 3)\) pattern did not change until the end of the GaAsSb cap layer growth.

We studied the \(\text{As} \text{Sb}/\text{In} \) BEP ratio dependence of electron mobility during the InAs QW layer growth. The \(\text{As} \text{Sb}/\text{In} \) BEP ratio was changed from 85 to 220 by changing the In cell temperature. In Fig. 2 the electron mobility at room temperature is shown as a function of \(\text{As} \text{Sb}/\text{In} \) BEP ratio. The highest electron mobility of 27,000 cm\(^2\)/Vs was obtained by using the \(\text{As} \text{Sb}/\text{In} \) BEP ratio of 110 at an In flux of about \(J_{\text{In}} = 1.0 \times 10^{14}\) atoms/cm\(^2\)/s. We have to control \(\text{As} \text{Sb}/\text{In} \) BEP between 95 and 120 to get a higher electron mobility than 25,000 cm\(^2\)/Vs. The surface morphology was observed by HRSEM after 3 ML growth of InAs on the AlGaAsSb surface. Fig. 3a and Fig. 3b show the HRSEM images of samples grown at \(\text{As} \text{Sb}/\text{In} \) BEP ratios of 110 and 220, respectively. The morphology of the InAs grown at the ratio of 110 was smoother than that at the ratio of 220. We speculate that In atoms may be able to migrate more easily under relatively lower arsenic pressure. If the \(\text{As} \) flux increases, In atoms may be covered with As atoms before In atoms migrate well on the surface. In contrast, if the \(\text{As} \) flux is lower than the optimum BEP ratio of 110, In atoms may start to coalesce and form In droplets.

The \(\text{As} \text{Sb}/\text{Sb} \) BEP ratio during the AlGaAsSb growth was also investigated. The \(\text{As} \text{Sb}/\text{Sb} \) BEP ratio was changed from 6 to 12 by changing Sb cell temperature. The \(\text{As} \) BEP was kept at 1.80 \times 10^{-5} \text{Torr during the growth of the InAs DQWs layers. Fig. 4 shows the FWHM of the PL peak from AlGaAsSb as a function of the \(\text{As} \text{Sb}/\text{Sb} \) BEP ratio. The PL spectra from InAs DQWs were dominated by a single strong peak around 1.2 eV, corresponding to the band-edge emission of the Al\(_{0.5}\)Ga\(_{0.5}\)AsSb layer at 10 K, as shown in Fig. 4b. It was found that there is a minimum FWHM of the sample grown by using an \(\text{As} \text{Sb}/\text{Sb} \) BEP.
ratio of 6.7 at an Sb$_4$ flux of about $J_{Sb} = 8.9 \times 10^{14}$ atoms/cm$^2$ s.

Moreover, we also investigated the PL peak shift of AlGaAsSb band-edge emission as a function of the As$_4$/Sb$_4$ BEP ratio. As the Sb$_4$ flux increased, the As content y in Al$_{x}$Ga$_{1-x}$As$_{1-y}$Sb$_{y}$ decreased and y was almost zero at the As$_4$/Sb$_4$ ratio of 5.5, which was confirmed by X-ray diffraction. If the Sb ratio increases in AlGaAsSb at a constant Al/Ga ratio, the energy band gap should decrease. However, the photon energy increased from 1.20 to 1.24 eV at 10 K as the As$_4$/Sb$_4$ BEP ratio decreased from 12 to 5.5, as shown in Fig. 5a. This indicates that the Al$_{x}$Ga$_{1-x}$As$_y$Sb energy gap increased because of the change in the Al/Ga ratio as the Sb$_4$ beam flux increased. In order to estimate the change in the Al/Ga ratio in AlGaAsSb, we grew calibration samples changing x in Al$_{x}$Ga$_{1-x}$As$_y$Sb from 0 to 0.8 at a constant As$_4$/Sb$_4$ BEP ratio. Fig. 5b shows the energy band gap of Al$_{x}$Ga$_{1-x}$As$_y$Sb as a function of x. We can estimate that the Al content x increases from 0.44 to 0.48 as the As$_4$/Sb$_4$ BEP ratio decreases from 12 to 5.5. The reason the x changes as the As$_4$/Sb$_4$ BEP ratio changes may be explained as follows. The heats of formation ($-\Delta H_f$, spectroscopy) of AlSb, GaSb, AlAs, and GaAs are, respectively, 20.3, 9.5, 17.3, and 16.3 kcal/mol [9]. We speculate that AlSb is thermodynamically more stable than GaSb and the stability of AlAs is almost the same as GaAs. Therefore, AlSb may be formed more easily than GaSb during the growth of AlGaAsSb as the Sb flux increases. Further studies are required to clear the dependence of electron mobility on Al/Ga ratio in AlGaAsSb of InAs DQWs.

Furthermore, we evaluated the crystalline quality of AlGaAsSb by the X-ray diffraction rocking curves. The minimum FWHM of 600 arcsec was also observed at the As$_4$/Sb$_4$ BEP ratio of 6.7. Since we reduced the substrate temperature from 540°C to 450°C before growing the InAs QW layer, the FWHM of AlGaAsSb was suppressed to 600 arcsec.

Finally, we measured the Hall mobility of these samples at room temperature. We found that the mobility curve had a maximum of 32,000 cm$^2$/V s at the As$_4$/Sb$_4$ BEP ratio of 6.7, as shown in Fig. 6.
For a 6.7 As$_4$/Sb$_4$ BEP ratio growth, an electron mobility of 90 000 cm$^2$/V·s at 77 K was obtained. The typical carrier densities and sheet resistances at room temperature were in the ranges of 0.9—1.1 $\times$ 10$^{12}$ cm$^{-2}$ and 185–250 $\Omega$, respectively. The not-intentionally doped InAs DQWs have remarkably high electron sheet concentrations of the order of 10$^{12}$ cm$^{-2}$. We can reproducibly grow the InAs DQWs with high electron mobility of more than 30 000 cm$^2$/V·s by maintaining the As$_4$/Sb$_4$ BEP ratio at 6.7 and As$_4$/In BEP ratio at 110.

4. Conclusion

To improve the electron mobility of InAs DQWs, we have to grow high-quality crystals of AlGaAsSb buffer/barrier layers and the InAs QW layer. By accurately controlling As$_4$/Sb$_4$ for the AlGaAsSb layer growth and As$_4$/In for the InAs QW layer growth in InAs DQWs, we have obtained very high electron mobility of more than 32 000 cm$^2$/V·s at room temperature. The optimum BEP ratios of As$_4$/Sb$_4$ and As$_4$/In are 6.7 and 110, respectively, from the electrical and optical characterizations of InAs DQWs. We believe that InAs/AlGaAsSb DQWs, with their high electron mobility, are promising materials for high-speed electronic devices and magnetic sensor applications.

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References

MBE growth and characterization of high-quality GaInAsSb/AlGaAsSb strained multiple quantum well structures


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Abstract

Molecular beam epitaxial growth and characterization of GaInAsSb/AlGaAsSb strained multiple quantum wells (MQWs) have been investigated. Optimization for the growth have been studied by characterizing the MQWs with double-crystal X-ray rocking curve, photoluminescence (PL) and absorption measurements and evaluating laser diode performance. Up to the fifth-order the satellite peak is resolved in rocking curves, and asymmetric distribution of satellite peaks is found which indicates large strain in quantum well structures. PL and absorption measurements at different temperatures show the high quality of GaInAsSb/AlGaAsSb multiple quantum wells grown under optimized conditions. The full-width of half-maximum of a PL peak as narrow as 9.0 meV is achieved at 4 K, and room-temperature PL and well-resolved excitonic absorption peaks are observed in the MQW structures. By using the GaInAsSb/AlGaAsSb MQWs as active layers in laser diodes, pulsed operation at room temperature with emission at 2.05 μm has been obtained.

1. Introduction

The GaInAsSb/AlGaAsSb quaternary systems have recently been receiving much attention, due to their potential application as semiconductor diode lasers with emission wavelength in the range 2–4 μm [1–6]. High-performance GaInAsSb/Al- GaAsSb quantum well (QW) lasers grown by molecular beam epitaxy (MBE) has been demonstrated [4–6]. The recombination of photoexcited carriers in GaInAsSb/AlGaAsSb QW is a process of vital importance for the optoelectronic devices based on those quaternary QW structures. Photoluminescence (PL) has so far been an important characterization tool for the recombination process as well as for assessing the QW quality. We previously reported the PL characteristics in GaInAsSb/AlGaAsSb QW system [7–9]. On the
other hand, the employment of strained QW in devices extends the choice of compatible materials and greatly increases the ability to control their optical and electronic properties. Recent investigations show that the strained multiple-QW (MQW) laser diodes may be far superior to those with conventional unstrained MQW structures in operational characteristics [10]. PL, though may detect the strain, cannot extract the strain straightforward, for the energy of PL peak depends on well width, barrier height as well as strain. Instead, double-crystal X-ray rocking curve (DCXRC) analysis has been proven to be a sensitive characterization method for investigating period thickness, alloy composition and strain in MQWs [11]. In this paper we report MBE growth, DCXRC, PL and absorption characterization of GaInAsSb/AlGaAsSb strained MQW structures and demonstration of GaInAsSb/AlGaAsSb MQW laser diodes.

2. Experiments procedure

The GaInAsSb/AlGaAsSb MQW structures were grown in a home-made conventional solid-source MBE system. Uncracked As$_4$ and Sb$_4$ were used as the As and Sb sources. The substrate temperature was measured by a thermocouple. The compositions of epilayers were determined by electron microprobe analysis, while layer thicknesses were estimated from the growth rate. They were also confirmed further by DCXRC measurements.

DCXRC measurements were performed on a computer-controlled high-resolution X-ray double-crystal diffractometer. (1 0 0)-oriented GaAs crystal was used as the first crystal to diffract Cu K$_{α1}$ X-radiation from a normal focus Cu tube run at 35 kV, 20 mA in spot-focus mode.

PL and absorption measurements were performed on a Nicolet 800 Fourier transform infrared spectrometer over the temperature range of 4 K to room temperature (290 K). PL measurements have been carried out using an Ar-ion (514.5 nm) laser for excitation and a liquid-nitrogen-cooled InSb detector, while the absorption spectra were also detected by an InSb photodiode with a tungsten lamp focused onto the sample. The optical measurements were made with a resolution of 4 cm$^{-1}$.

3. Results and discussion

3.1. MBE growth and DCXRC characterization

The GaInAsSb/AlGaAsSb MQW structures were grown on (1 0 0) Te-doped GaSb substrates. Prior to the growth of the GaInAsSb/AlGaAsSb MQW, a 1.0 μm-thick n$^+$-GaSb buffer layer was firstly grown at a calibrated real temperature of 500°C. After 10 min interruption, GaInAsSb/AlGaAsSb MQW was grown at 500–520°C under As, Sb-rich condition with the growth rate ranging from 0.7 to 1.0 μm/h. The growth process was controlled by a computer and in-situ monitored by reflection high-energy electron diffraction (RHEED). Several MQW samples grown at different temperatures and with different growth rates were characterized by DCXRC measurements.

Fig. 1 shows the (4 0 0) rocking curve of a 20-period Ga$_{0.75}$In$_{0.25}$As$_{0.02}$Sb$_{0.98}$ (10 nm)/Al$_{0.16}$Ga$_{0.84}$As$_{0.02}$Sb$_{0.98}$ (30 nm) MQW, which was grown at 510°C with growth rate of 0.8 μm/h. The solid line is the measured rocking curve while the solid line with dots is the theoretical one simulated by Speriosu's model [11]. In the measured curve, in addition to the GaSb substrate peak, seven satellite peaks could be clearly seen, indicating high quality of the GaInAsSb/AlGaAsSb MQW structure. The satellite peak spacing yields an average quantum well period thickness.

![Fig. 1](image_url)

Fig. 1. Experimental and simulated (400) rocking curves for a 20-period Ga$_{0.75}$In$_{0.25}$As$_{0.02}$Sb$_{0.98}$ (10 nm)/Al$_{0.16}$Ga$_{0.84}$As$_{0.02}$Sb$_{0.98}$ (30 nm) MQW.
41.9 ± 0.2 nm, in good agreement with the nominal value 40.0 nm. Sparrow's simulation identifies each satellite peak and up to the fifth-order satellite peak of the MQW can be resolved in the measured rocking curve. GaInAsSb/AlGaAsSb MQWs grown at temperatures higher than 520°C or lower than 500°C or with growth rates higher than 0.9 μm/h show fewer and weaker satellite peaks in rocking curves. Notice the asymmetry of the envelope of these satellite peaks, which indicates the existence of considerable strain in the QW. To investigate the strain further, (1 1 5) DCXRC measurements have been carried out, for perpendicular strain ε∥ will be more sensitive in asymmetric diffractometry than symmetric one. Fig. 2 shows the (1 1 5) rocking curve for the same GaInAsSb/AlGaAsSb MQW. Besides the asymmetry of the envelope of the satellite peaks, the figure even shows the disappearance of the zeroth-order peak in the rocking curve, which strongly indicates large strain in the MQW. With detailed simulation, the biaxial strain parallel to and uniaxial strain perpendicular to the plane of the interface is determined as ε∥ = 3.31 × 10⁻³ and ε⊥ = 3.14 × 10⁻³, respectively. These values are less than lattice mismatch Δa/a (~ 1%), which could be attributed to the distribution of compressive and tensile strain in the well and barrier. And the coherence factor is also determined to be as high as 0.97, indicating that the mismatch in the QW is accommodated mainly by the strain.

3.2. Photoluminescence and absorption measurements

Fig. 3 shows the PL spectrum at 4 K for that GaInAsSb/AlGaAsSb MQW sample grown under optimized conditions. From the figure, a single PL peak is observed, corresponding to the recombination between the first electron subband and the first heavy-hole subband (E₁-HH₁). The full-width at half-maximum (FWHM) of 9.0 meV (4 K) and 13.0 meV (10 K), the best result yet reported in this quaternary system, indicates the good optical quality of the MQW. Fig. 4 shows the PL and absorption spectra at different temperatures for another sample, 20-period Ga₀.₇₂In₀.₂₈As₀.₀₂Sb₀.₉₈ (10 nm)/Al₀.₁₆Ga₀.₈₄As₀.₀₂Sb₀.₉₈ (30 nm) MQW.

In PL spectra a single PL peak, corresponding to E₁–HH₁ recombination, is observed throughout the temperature region of the measurements (4–290 K), with FWHM of 12.1 meV at 4 K and 22.1 meV at 290 K. While in the absorption spectra, additional excitonic absorption structures (E₁–LH₁ and E₂–HH₂) appear at lower temperatures due to the increase of the overlap integral of electron and hole wave functions. The sharp rise in the highest energy portion of the absorption spectra is due to absorption in GaSb substrate. Furthermore, small Stokes shift between the PL and absorption peaks is found at any temperature. This fact, along with strong luminescence and well-resolved excitonic absorption peaks observable at room temperature, undoubtedly demonstrates the high quality of the MQW structures. The detailed analysis of the optical characteristics of the GaInAsSb/AlGaAsSb MQWs can be found elsewhere [7, 8]. By using the high-quality
GaInAsSb/AlGaAsSb MQWs as active layers, GaInAsSb/AlGaAsSb MQW laser diodes have been fabricated through broad area stripe laser processing. Fig. 5 shows the emission spectrum for a GaInAsSb/AlGaAsSb MQW laser diode. Room-temperature pulsed operation with stable threshold current density lower than 900 A/cm² has been achieved. The peak wavelength of lasing is 2.05 µm and the FWHM of the peak is 0.6 nm.

4. Conclusion

High-quality GaInAsSb/AlGaAsSb strained MQWs have been successfully grown by MBE. Both structural and optical qualities of those MQWs have been characterized by DCXRC, PL and absorption measurements. Up to the fifth-order, satellite peak of MQW is observed in a rocking curve, and large strain exists in the QW structure. The FWHM of a PL peak as narrow as 9.0 meV is achieved at 4 K, and room-temperature PL and well-resolved excitonic absorption peaks are observed in the MQW structures. By using the high-quality GaInAsSb/AlGaAsSb MQWs as active layers, room-temperature pulsed operation laser diodes with emission at 2.05 µm and threshold current density lower than 900 A/cm² have been fabricated.

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References

MBE growth of GaInAsSb p/n junction diodes for thermophotovoltaic applications

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Abstract

This paper reports recent progress in the development of quaternary III–V thermophotovoltaic (TPV) devices based on MBE grown Ga\textsubscript{x}In\textsubscript{1-x}As\textsubscript{y}Sb\textsubscript{1-y}. TPV is of great interest for a variety of applications (1st and 2nd NREL Conf. on Thermophotovoltaic Generation of Electricity, AIP Conf. Proc. 321 (1994), 358 (1995)). The objective of this work is to develop a TPV cell which is “tunable” to the emission spectrum of a heated blackbody, at temperatures in the range of 1200–1473 K. One aspect of this “tuning” is to match the band gap, $E_{\text{gap}}$, of the photovoltaic device to the peak output of the heat source. An advantage of the quaternary III–V semiconductor systems is that devices can be fabricated by molecular beam epitaxy on a suitable binary substrate, such as GaSb or InAs, and the band gap and lattice constant can be adjusted more or less independently, to match requirements. Quaternary cells, with band-gaps in the 0.5–0.72 eV range, have been fabricated and tested. For 0.54 eV devices we obtained open circuit voltage $V_{\text{oc}} = 0.3$ V and short circuit current $I_{\text{sc}} = 1.5$ A/cm$^2$ under infrared illumination of a 1200 K blackbody. Under high illumination levels the $V_{\text{oc}}$ and $I_{\text{sc}}$ ranged from 0.5 V at 3 A/cm$^2$ for 0.72 eV devices to 0.31 V at 1.2 A/cm$^2$ for 0.5 eV devices, indicating good photovoltaic device characteristics over the range of bandgaps. The diode ideality factor for 0.54 eV devices ranged from 2.45 at low illumination indicating tunneling-dominated dark current, to 1.7 at high illumination intensity indicating recombination-generation dominated dark currents.

1. Introduction

Thermophotovoltaic (TPV) devices have a wide variety of applications [1, 2]. The elements of a TPV system include an emitter which is a heat source coupled to a blackbody radiator, a spectral control element (band-pass filter) and a photovoltaic converter which is matched to the blackbody and spectral control element. A schematic for a TPV system is shown in Fig. 1.

Fig. 2 compares the power emitted by a blackbody at various temperatures and compares the
power at wavelengths shorter than the cut-off wavelengths for GaSb, and two lower bandgaps. The three lower curves represent the maximum power available for conversion by cells with those bandgaps without taking into consideration any factors limiting their conversion efficiency. It must be pointed out that a high efficiency PV diode is only one aspect of demonstrating an efficient TPV system, an effective spectral control device is critical to obtain high overall system efficiency. As an example, for a 1200 K blackbody, almost 70% of the blackbody power density lies below the bandgap of a 0.5 eV bandgap PV cell. This requires an efficient band-pass filter which reflects most of the radiation below the semiconductor bandgap out to 10 μm back to the blackbody to recycle the photons and allows maximum transmittance above the semiconductor bandgap for the PV cell to convert.

Among the unique advantages of the quaternary GaInAsSb system are that the bandgap and the lattice constant can be adjusted to match the requirements. Fig. 3 shows the alloy composition and the bandgap range of interest lattice matched to GaSb.

In this paper we present recent progress on obtaining large area (1 cm × 1 cm) variable bandgap GaInAsSb PV cell lattice matched to GaSb substrates with bandgaps between 0.72 eV and 0.5 eV.
2. Material growth

The material used for the fabrication of the GaInAsSb TPV devices was grown using a Varian Gen II MBE system. 2 in n-type (1–2 × 10^{17} \text{cm}^{-3}) exact (0 0 1) oriented GaSb substrates were used. The oxide on these wafers was desorbed in situ in the MBE system. The wafers were heated to about 400°C, then the Sb₄ shutter was opened, the oxide was observed to come off at temperatures in the 525–535°C range as monitored by the pyrometer. An n⁺ (≈ 5 × 10^{18} \text{cm}^{-3}) 100 nm GaSb buffer layer was grown to smooth the surface and improve the RHEED pattern. During the GaSb growth the substrate temperature was dropped to 500°C and growth of GaInAsSb was initiated. This formed the base region of the TPV device and had a thickness of 5 μm and an n-type doping density of 2 × 10^{17} \text{cm}^{-3}. Next the p-type emitter and p-type GaSb (300 Å) window layer were grown with a doping density of 1–2 × 10^{18} \text{cm}^{-3}. Fig. 4 shows a typical device structure.

In order to obtain the correct bandgap, Ga and In ratios were adjusted. To obtain the lattice constant match with the substrate, As and Sb ratios were adjusted. The GaInAsSb devices were lattice matched to within 0.05% with the substrate. When the mismatch exceeded 0.1% the wafer was observed to bend. The wafers bent in a convex shape with a plus (larger than the substrate-excess Sb) mismatch. The wafers were observed to bend in a concave shape with a negative (smaller than the substrate-excess As) mismatch. This bending of wafers was much worse in the case of InAs substrates, because InAs is a softer material than GaSb. To relieve the lattice mismatch strain quaternary GaInAsSb appears to deform the substrate instead of relieving strain by the formation of dislocations. This is presumably due to the alloy hardening effect of the second group V in the films. Due to this alloy hardening effect, the layer finds it energetically difficult to produce and propagate dislocations and hence bends to relieve the strain. This bending of the wafers took place at the growth temperature and could be observed through the view port, this rules out the bending due to any coefficient of thermal expansion mismatch. To further investigate the possibility of lattice mismatch related bending peculiar to the GaInSb material system we grew the ternary GaInSb (with the same Ga and In ratio as in the quaternary) on GaSb, we did not observe any bending despite a much larger mismatch ( > 0.5%). This indicates that in the mixed arsenide–antimonide quaternaries strain relaxation proceeds differently as compared to the ternaries, and can have important device implications. Further investigations using TEM are needed on the GaInAsSb quaternary system.

3. Device fabrication and characterization

1 cm × 1 cm mesa diodes were fabricated using chemical etching, the mesa diodes had a single 1 mm wide central busbar for making electrical contact to the devices. The busbar was connected to grid lines 0.45 cm long 100 μm apart and 10 μm wide. Front metallization was done by evaporating Ti and Pt in an e-beam evaporator and then depositing a 5 μm thick Au or Al to obtain a low resistance contact for high current densities. For bottom Ohmic contact a thick Sn/Pt or Sn/Au was used. A 0.2 μm Silicon Nitride layer on top was used as an antireflection coating.
Fig. 5. External quantum efficiency as a function of wavelength for a typical GaSb and a GaInAsSb device.

The diodes were tested under dark and under blackbody illuminated conditions. Fig. 5 shows an external quantum efficiency versus wavelength for a typical device. The figure shows a comparison for a typical GaSb and GaInAsSb device. The fall off at short wavelengths indicates high surface recombination, pointing to the need for an effective front surface passivation or a high bandgap window layer. A softer roll off in quantum efficiency at wavelengths near the band edge indicates the absence of an effective back surface field. Adding these two layers to the devices will considerably improve their performance.

Fig. 6 shows comparison of a typical $I-V$ characteristics from a GaInAsSb and GaSb PV diodes under similar low light level illumination conditions. As is to be expected the short circuit current is higher for the lower bandgap device. The $I-V$ characteristics point to the need for better ohmic contacts and this is an area which is under current investigation. Some of series resistance in the GaInAsSb devices could be due to a heterostructure barrier between GaInAsSb and GaSb [3]. The conduction and valence band discontinuities between GaSb and GaInAsSb are 0.24 and $-0.1$ eV, respectively. This is particularly troublesome and needs to be reduced by degenerately doping the junction or by bandgap grading.

Fig. 6. $I-V$ characteristics for a GaSb and a 0.55 eV GaInAsSb device.

Fig. 7 shows the PV short circuit current density versus the blackbody temperature. It can be observed that for low temperature (1200 K) TPV applications the narrow gap (0.54 eV) cells are the answer because they offer more than three times the current density as compared to the GaSb (0.73 eV) cells.

Fig. 8 plots the measured short circuit current density vs the open circuit voltage. Assuming the ideal diode relation

$$I_{SC} = I_0\exp\left(\frac{qV_{OC}}{nkT}\right) - 1,$$

where $I_{SC}$ is the short circuit current, $I_0$ is the dark current, $V_{OC}$ is the open circuit voltage, $n$ is the diode ideality factor, $k$ is the Boltzmann constant and $T$ is the temperature.
There are four potential dark current mechanisms: diffusion ($n = 1$), bulk recombination/regeneration ($n = 2$), surface recombination ($n = 2$) and band to band tunneling ($n > 2$). In Fig. 8 we plotted the $I_{SC}$ versus $V_{OC}$ and fitted the ideal diode equation assuming a single dominant dark current mechanism. We found it useful to compare the data for our MBE grown GaInAsSb and GaSb diodes and JX Crystal LPE grown cells. At low current injection conditions MBE grown devices show a diode ideality factor of 2.43, indicating band to band tunneling. LPE devices on the other showed a combination of diffusion and generation/recombination dark current. This difference could be due to a lower base doping for the LPE grown cells. Lower base doping leads to a lower $V_{OC}$, MBE cells show a higher $V_{OC}$ at a lower $I_{SC}$ (this is very desirable) as compared to LPE cells. This data leads us to conclude that at the high current levels required for TPV device operation, higher base doping is advantageous even though higher ideality factors are observed. Thus GaInAsSb diodes are more efficient energy converters at high illumination levels. Some of the dark current could also be due to a lack of surface passivation. Use of an effective passivant or a high bandgap window layer will also reduce the dark current.

Fig. 9 plots $V_{OC}$ versus $T$ for MBE grown GaInAsSb cells. This is done for various $I_{SC}$ values. This data also helps determine the differences in the dominant dark current mechanisms. From low to high illumination levels the $V_{OC}$ decreases linearly with temperature. The difference in the slope $dV_{OC}/dT$ between very low injection levels and higher injection levels indicate that at lower current levels the current is a combination of tunneling and some other mechanism, for higher injection levels once the tunneling barrier has been overcome the dark current mechanism is almost entirely due to tunneling [4], this is indicated by a relatively small change in the $dV_{OC}/dT$ slope. It will be interesting to measure $dV_{OC}/dT$ for even higher injection levels to see if other dark current mechanisms become dominant.
4. Conclusions

We have presented data on the current status of lattice matched GaInAsSb TPV devices and pointed out the areas in which improvements or device design changes are needed. Quaternary lattice matched TPV devices are very promising for thermal power conversion and offer superior performance due to potentially higher long term reliability and higher open circuit voltages and currents than lattice mismatched InGaAs/InP devices.

References

Molecular-beam-epitaxial growth of Ga$_{1-x}$In$_x$Sb on GaAs substrates

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Abstract

Molecular beam epitaxy has been used to grow Ga$_{1-x}$In$_x$Sb in the entire compositional range. Good Hall mobility values were obtained, but the surfaces were rough and exhibited square defects. The X-ray diffraction Bragg peaks in the middle of the compositional range were broadened due to the disorder inherent in the alloy. A small inclination of the epitaxial layers with respect to the substrate was also found in this region.

1. Introduction

Presently there is much interest in antimony-containing III–V semiconductors. For example, Ga$_{1-x}$In$_x$Sb has been used in the active region of mid-infrared diode lasers [1, 2], in far-infrared strained-superlattice detectors [3] and resonant tunnelling devices [4].

This paper reports molecular-beam-epitaxial growth and characterization of Ga$_{1-x}$In$_x$Sb alloy layers. Although it is clearly related to In$_x$Ga$_{1-x}$As, this alloy has not been as extensively studied. Low growth temperatures are of particular interest since InAs/Ga$_{1-x}$In$_x$Sb superlattices are grown with the highest structural quality at 390-410°C [5].

The first epitaxial growths of Ga$_{1-x}$In$_x$Sb were made by liquid phase epitaxy [6, 7]. Since then the alloy has been grown by molecular beam epitaxy (MBE) [8, 9], metalorganic vapour-phase epitaxy (MOVPE) [10, 11] and metalorganic molecular beam epitaxy (MOMBE) [12, 13]. Much emphasis has been put on the optical properties, but also on structural characterization, mainly for lower In concentrations.

2. Experimental procedure

2.1. Growth

A series consisting of more than thirty 4-μm-thick Ga$_{1-x}$In$_x$Sb layers, spanning the entire compositional range, was grown on (0 0 1)-oriented semi-insulating GaAs substrates. Both unintentionally doped and Si-doped samples were grown in a modified Varian 360 system with arsenic and antimony cracking cells and standard effusion cells for gallium and indium. The cracking temperatures of the V-sources were kept at 1200°C. In situ
cleaning began by pre-heating the substrates in ultra-high vacuum. This was followed by thermal desorption of the native oxide in the growth chamber. The normal procedure during this step is to stabilize the surface under an arsenic flux. Recent studies have shown, however, that arsenic is easily incorporated in antimonides in non-negligible amounts [14, 15]. Therefore the possibility to desorb the oxygen under an antimony flux was tested. As compared to desorption under arsenic, the process seemed more sudden and it was also noted that it seemed to occur at 20–30°C lower temperature. This could be due to a bombardment effect caused by the greater mass of antimony. However, with no further independent means to measure the temperature, it was difficult to establish this point with certainty, and hence all growth temperatures reported in this paper assume that the desorption occurred at the regular temperature of 583°C.

Since most of the growth temperatures were below 500°C, where the optical pyrometer loses its accuracy, substrate temperatures were estimated assuming a linear relationship between the actual temperature and the reading of a thermocouple mounted in non-contact position on the back side of the substrate holder.

Growth rates were calculated from reflection high-energy electron diffraction (RHEED) oscillations prior to growth. The In molar fraction, x, of the alloy was determined in advance by calculating

\[ x = \left[ 1 + \left( \frac{a_{\text{InAs}}}{a_{\text{GaSb}}} \right)^3 \frac{v_{\text{GaSb}}}{v_{\text{InAs}}} \right]^{-1}, \]  \hspace{1cm} (1)

where \( a_{\text{InAs}} \) is the lattice constant of InAs and \( v_{\text{InAs}} \) is the growth rate it has at the chosen In source temperature, and analogously for \( a_{\text{GaSb}} \) and \( v_{\text{GaSb}} \). In most cases, this gave excellent agreement with the values determined from X-ray diffraction. The surface properties were continuously monitored by RHEED. The \( \text{Ga}_{1-x}\text{In}_x\text{Sb} \) grew under a \((1 \times 3)\) reconstruction for \( x < 0.4 \) and a \((2 \times 3)\) for \( x > 0.4 \).

2.2. Analysis

X-ray rocking curves were measured in a high-resolution X-ray diffractometer equipped with a Cu K\( \alpha_1 \) X-ray source and a four Ge-crystal mono-

chromator set in the \((2 \ 2 \ 0)\) position. Rocking curves were obtained by scanning through the \((0 \ 0 \ 4)\) reflection of the sample. Information on the lattice constants parallel and perpendicular to the surface were obtained by measuring the angles of incidence giving the Bragg peak for the \((1 \ 1 \ 5)\) and \((1 \ 1 \ 5)\) reflections. From this, the equilibrium lattice constant and the residual strain can be found [16].

Surface analysis was performed by optical microscopy, surface profiling in a Tencor Alpha-Step 500 and scanning electron microscopy (SEM).

Electrical properties were assessed by measuring the resistivity, \( \rho \), and the Hall constant, \( R_H \), in the square van der Pauw arrangement on \( 3 \times 3 \) mm\(^2 \) samples in a magnetic field of 0.4 T. The Hall carrier concentration was obtained from \( n_H = 1/eR_H \) and the Hall mobility from \( \mu_H = R_H/\rho \) [17].

Optical measurements were made by Fourier transform photoluminescence (FTPL). The excitation source was a 514.5 nm Ar-ion laser. The sample was kept in a He cryostat and the luminescence was analysed with a Fourier transform spectrometer equipped with an InSb detector.

3. Results

All of the X-ray rocking curves exhibited the substrate's Bragg peak and a wider peak originating from the 4-\( \mu \)m-thick overlayer. In some cases the position of the latter depended on the incident azimuth of the X-ray, indicating that the epitaxial layer is inclined with respect to the substrate. This inclination was hardly noticeable for alloy compositions close to the binaries but increased to about 300 arcsec for Ga\(_{0.6}\)In\(_{0.4}\)Sb. The width of the Bragg peak associated with the Ga\(_{1-x}\)In\(_x\)Sb layer increases in the middle of the compositional range. There was also a strong anisotropy revealing itself in different full widths at half maximum (FWHM) for X-rays incident in different azimuths. Fig. 1 shows the FWHM of the epilayer peak with X-rays incident along the [1 1 0] azimuth for samples grown at 430°C. The peaks are broader than for \( \text{In}_x\text{Ga}_{1-x}\text{As} \) samples grown at near-group-III stabilized conditions. The residual strain in the Ga\(_{1-x}\)In\(_x\)Sb layers was of the order 0.1%, which is comparable to \( \text{In}_x\text{Ga}_{1-x}\text{As} \) of similar thickness [18].
Hall effect measurement due to the large ratio of electron-to-hole effective masses, although the conductivity is still dominated by the holes.

Fig. 3 shows a typical optical micrograph of the surface morphology of Ga_{0.4}In_{0.6}Sb. The surface analysis showed that the surfaces particularly in the middle of the compositional range were very rough and covered with square defects. The sides of the squares were aligned along the <110> directions. Many samples also exhibited pyramid-like defects protruding 100–200 nm from the surface. Fig. 4 shows two kinds of pyramid-shaped defects. The sample in Fig. 4a had been excessively heated during the oxide desorption and was the only sample which showed this type of fully grown pyramid. Most other samples exhibited the collapsed pyramid structure in Fig. 4b. It is also noteworthy that the sample in Fig. 4a had a much more narrow Bragg peak than other samples in its compositional range.

There was a strong correlation between alloy composition and surface morphology. Close to the binaries the surfaces were very smooth with tall, sharp pyramids of high surface density. In the middle of the compositional range the surfaces were rougher, and the pyramids were always located on top of a square plateau. The pyramids were also flatter here with larger lateral size and of lower density.

Only samples that had been grown at temperatures below 440°C for low In concentrations,
where the surface reconstruction changes from $(1 \times 3)$ to $(1 \times 5)$ as the group-III flux is removed, showed any detectable photoluminescence.

Other III–V antimonides are known to have surface morphologies and FWHMs that are highly sensitive to V/III-ratio and substrate temperature [19, 20]. Fig. 5 shows the measured FWHM for a series of Ga$_{0.4}$In$_{0.6}$Sb grown at different temperatures. For low temperatures FWHM is larger for X-rays incident along the [1 1 0] azimuth. At approximately 500°C there is a point where the FWHM is isotropic. For higher growth temperatures the [1 1 0] azimuth gives the lowest FWHM. Growth temperatures higher than 530°C resulted in a dim RHEED pattern during growth and extremely wide Bragg peaks. Back-scattered electron imaging in SEM indicated that these samples are of non-uniform composition. The best FWHMs occurred for growth temperatures in the range 480–500°C, but these peaks are still much broader than normally observed for comparable In$_{0.4}$Ga$_{0.6}$As.

4. Discussion

The disorder in a III–V solid solution can be divided into two different categories. The first kind of disorder is due to the fact that some Ga atoms have been replaced with In atoms, with different scattering powers for X-rays. The second kind of disorder comes from the difference in bond length between Ga–Sb bonds and In–Sb bonds. Using extended X-ray absorption fine structure (EXAFS) results, Mikkelsen and Boyce discovered that the Ga–As and In–As bond lengths in In$_{x}$Ga$_{1-x}$As depend on the alloy composition, but in a fashion that allows the average bond length to obey Vegard’s law [21]. These two effects will maximize the disorder and broaden the X-ray FWHM in the middle of the compositional range as seen in Fig. 1.

The XRD anisotropy has also been observed in In$_{x}$Ga$_{1-x}$As [22]. It is believed that it is caused by
different probabilities to create dislocations in different directions.

The defects on the surface seem to be related to the start of the growth. We also compared samples where the native oxide had been desorbed under As flux. The density of the pyramid-like defect was lower but not zero. However no improvement in surface roughness could be seen.

Defects aligned along the crystallographic directions have been reported for Ga$_{1-x}$In$_x$Sb in other studies. Kodama and Parker et al. have noted similar defects which were explained as stacking-fault polyhedra [23, 24].

5. Conclusions

We can conclude that Ga$_{1-x}$In$_x$Sb grows with a rough surface and has broad XRD Bragg peaks in the investigated temperature region, 400–500°C. The structural quality was seen to improve slightly for Ga$_{0.6}$In$_{0.4}$Sb by increasing the growth temperature from 430°C to 480°C. This not unexpected since GaSb is grown with the best quality above 500°C [25].

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References

Stranski–Krastanov growth of InSb, GaSb, and AlSb on GaAs: structure of the wetting layers

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Abstract

Thin layers of InSb, GaSb and AlSb were grown on GaAs(001) by molecular beam epitaxy and characterized in situ with scanning tunneling microscopy. All three materials exhibit a Stranski–Krastanov growth mode. Distinct wetting layers and self-assembled quantum dots are present after deposition of one to four monolayers of (In,Ga,Al)Sb. The wetting layers consist of anisotropic, ribbon-like structures oriented along the [1 1 0] direction, with characteristic separations of 40–50 Å. The initial GaAs surface reconstruction affects both the wetting layer structure and the quantum dot density.

1. Introduction

The Stranski–Krastanov (SK) growth mode in strained-layer heteroepitaxy has been recognized for decades [1–3]. In this mode, the deposited layer initially grows as a two-dimensional (2D) wetting layer for at least one monolayer. Then, three-dimensional (3D) islands form on the wetting layer. Recently, there has been considerable interest in using SK growth to deposit coherent islands, known as self-assembled quantum dots (QDs). These QDs have been reported in semiconductor systems including InAs/GaAs, InP/InGaP, and GaSb/GaAs [4–8]. Although QDs with size uniformities on the order of 10% are routinely achieved, many potential applications such as solid-state lasers require more uniform ensembles of QDs. Theoretical work suggests that the uniformity of QDs is related to the structure of the wetting layer, [9–11] but few experimental studies have been reported [12–15]. In this work, we apply in-situ scanning tunneling microscopy (STM) to investigate the structure of wetting layers of GaSb, AlSb and InSb on GaAs (mismatches of 7.8%, 8.5% and 14.6%, respectively) grown by molecular beam epitaxy (MBE). In all three cases, we observe wetting layers composed of 2D islands with anisotropic ribbon-like structure and characteristic inter-island separations of 40–50 Å.

2. Experimental procedure

Experiments were carried out in an interconnected multi-chamber ultra-high vacuum (UHV)
facility that includes a III–V solid-source MBE and a surface analysis chamber with STM. All growths were performed on n⁺GaAs, oriented to within 0.1 of (0 0 1). Growth temperatures were determined by GaAs band-edge transmission thermometry [16]. First, a GaAs buffer layer 0.5–1.0 μm thick was grown at 580°C (with interrupts) at a growth rate of 1.0 monolayer (ML)/s with the growth monitored by reflection high-energy electron diffraction (RHEED). During the GaAs buffer growth, the RHEED pattern was a streaky (2 × 4) reconstruction. Before the growth of each antimonide layer, a 3–5 min growth interrupt was performed at 580°C under an As₂ flux, resulting in the appearance of sharp diffraction spots along each streak. A well-ordered (2 × 4) or (4 × 4) reconstruction was then prepared by alternately lowering the substrate temperature and As flux until the point at which the As valve could be closed without degradation of the RHEED pattern. This technique produces GaAs surfaces with ~5000 Å-wide terraces separated by monolayer-height (3 Å) steps [17]. The antimonide layer was grown by migration-enhanced epitaxy with a cation deposition rate of 0.10 ML/s and a V:III flux ratio of approximately 2:1. For example, to grow 1.5 ML of InSb, the shutter sequence was: 5 s In, 20 s Sb, 5 s In, 20 s Sb, 5 s In, 20 s Sb. After deposition of the III-Sb monolayers, the sample was held at the growth temperature under an Sb₄ flux for 120 s before cooling. Finally, the sample was transferred under UHV to the STM chamber and imaged at room temperature. All STM images displayed here were acquired in constant-current mode with currents between 0.1 and 1.0 nA and sample biases ranging from −2.0 to −3.2 V. The images are displayed in gray-scale without corrections for thermal drift.

3. Results

Well-ordered surfaces are observed for GaAs(0 0 1)-(2 × 4) and -(4 × 4), in contrast to the disordered surface revealed by both RHEED and STM after the deposition of 1–4 ML of GaSb, AlSb or InSb. After growth of 1.0 ML of GaSb on GaAs(0 0 1)-(4 × 4) at 490°C, RHEED exhibited primary and half-order streaks along with weak transmission spots in the [1 1 0] direction, and chevrons in the [1 1 0] direction [18, 19]. The resulting surface morphology, shown in Fig. 1a, consists of a disordered distribution of 2D islands. The islands are somewhat elongated, with ≤1 ML-deep gaps between them [20]. Note that quantum dots (QDs) have not yet formed at this coverage. After deposition of a total of 3.5 ML GaSb, strong RHEED transmission spots were visible in both the [1 1 0] and [1 1 0] directions, indicating 3D growth. The distance between the spots was clearly smaller than the distance between the diffraction streaks, implying that the QDs had formed with a larger in-plane lattice constant than the GaAs substrate. Because ex situ TEM images of QDs on similarly prepared samples do not exhibit Moiré fringes [21], however, this change in lattice constant must not result from the formation of dislocations but rather must arise from the elastic relaxation of the GaSb bonds within the QD. The presence of QDs has been confirmed on similar samples by ex situ AFM, with typical densities of 1 × 10⁸/cm², heights of 80 Å, and apparent diameters of 400 Å [21]. An STM image of a typical area between QDs is shown in Fig. 1b. (We try to avoid imaging QDs during studies of the wetting layer in order to avoid damage to the STM tip.) At this GaSb coverage, the wetting layer structure exhibits highly anisotropic ribbon-like structures oriented along the [1 1 0] direction. The characteristic ribbon separation, computed from the autocorrelation function for this image, is 50 Å in the [1 1 0] direction.

Ribbon-like wetting layers also result from the growth of AlSb on GaAs. We examined 2.0 ML of AlSb deposited on a (2 × 4) reconstructed GaAs surface at 500°C. After AlSb growth, the RHEED pattern revealed transmission spots superimposed on a (1 × 3) streak pattern. The resulting surface, shown in Fig. 2, is qualitatively similar to that following the growth of 3.5 ML of GaSb (Fig. 1b), with ribbon-like structures running along the [1 1 0] direction. The autocorrelation function yields a characteristic separation of 40 Å. Although not shown in this image, QDs are also present on the surface. Raman spectroscopy measurements of AlSb QDs on GaAs showed a two-mode
behavior indicating the formation of an AlGaSb alloy due to Ga segregation [22]. Hence, the surface shown in Fig. 2 is more appropriately described as Al\textsubscript{x}Ga\textsubscript{1-x}Sb.

Using ex situ AFM, we previously observed QDs after as little as 1.5 ML of InSb on GaAs [6]. In this study, we examined 1.5 and 2.0 ML of InSb on GaAs(0 0 1)-c(4 × 4) and 1.5 ML of InSb on GaAs(0 0 1)-(2 × 4) with STM. The substrate temperature during InSb growth was 410°C. In all three cases, RHEED revealed weak transmission spots in

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Fig. 1. Filled-state STM images of GaSb on GaAs(001)-c(4 × 4). GaSb coverages are: (a) 1.0 ML and (b) 3.5 ML. The gray-scale height ranges are approximately (a) 8 Å and (b) 10 Å. Quantum dots (not shown) are present on sample (b) but not on (a).

Fig. 2. Filled-state STM image of 2.0 ML of AlSb on GaAs(001)-(2 × 4). The gray-scale height range is approximately 12 Å.
the [1 1 0] direction and chevrons in the [1 1 0] direction; STM and AFM confirmed the presence of QDs. In addition to coherent QDs, large dislocated islands were observed for each sample [23]. In Fig. 3a and Fig. 3b we show the STM images acquired between the QDs for 1.5 ML of InSb on GaAs-c(4 × 4) and GaAs-(2 × 4), respectively. Anisotropic ribbon-like structures are present for both initial substrate reconstructions. The characteristic ribbon separations from autocorrelation analyses are 50 Å for growth on GaAs-c(4 × 4) and 40 Å for growth on GaAs-(2 × 4). The characteristic length of the islands in the [1 1 0] direction is somewhat shorter for InSb deposition on the GaAs-c(4 × 4) reconstruction. Fig. 3c displays a higher magnification image of 1.5 ML of InSb on GaAs-(2 × 4), revealing atomic-scale structure. The surface appears to have a multilayer structure terminated by dimers (most likely Sb dimers).

4. Discussion

The traditional view of SK growth is that the initial deposition occurs as a continuous wetting layer. Our results, however, suggest a more complicated structure with a discontinuous morphology which may arise from a combination of effects. Both the relative surface energies of the absorbates and substrate, and the mismatch-related strain which may be relieved by the formation of vacancy lines, may play a role. A simple calculation, assuming complete strain relaxation by vacancy arrays, gives a spacing of 12.5 atomic planes or 54 Å for an 8% mismatch (GaSb or AlSb) and 6.8 atomic planes or 31 Å for InSb. These values are comparable to the characteristic separations of the ribbon-like structures found for all three materials.

A few groups have investigated the initial stages of InAs growth on GaAs by in situ STM.

Fig. 3. Filled-state STM images of 1.5 ML of InSb on GaAs: (a) GaAs(0 0 1)-c(4 × 4) starting surface, (b) GaAs(0 0 1)-(2 × 4) starting surface, and (c) higher magnification image of surface in (b). The gray-scale height ranges are approximately (a) 12 Å, (b) 12 Å, and (c) 8 Å. Note the monolayer-height (3 Å) steps visible in (a) and (b).
Bressler-Hill et al. examined submonolayer coverages of InAs grown on GaAs(0 0 1)-(2 × 4) surfaces [12]. They observed 2D InAs islands elongated along the [1 1 0] direction—the same direction found in this work. Based upon an examination of the scaling properties of the island size, they concluded that growth along the [1 1 0] direction behaves as if it is unstrained while growth in the [1 1 0] direction is quenched by strain. In contrast, Cirlin et al. studied InAs growth on GaAs(0 0 1)-c(4 × 4) and found wire-like structures oriented along the [1 1 0] direction (i.e. 45° from [1 1 0]) [15]. Because of the different symmetries of the (2 × 4) and c(4 × 4) reconstructions it is not surprising that for low coverages the features of the wetting layer would be oriented in different directions. We observe some evidence of this behavior in our data. Surfaces with 1.0 ML of GaSb (Fig. 1a) and 1.5 ML of InSb (Fig. 3a) on GaAs-c(4 × 4) are more isotropic than 1.5 ML of InSb (Fig. 3b) and 2.0 ML of AlSb (Fig. 2) on GaAs-(2 × 4), consistent with the more isotropic structure of the c(4 × 4) reconstruction. As indicated in Fig. 1b, however, after additional GaSb is deposited (3.5 ML total), the remnant character of the symmetry of the GaAs-c(4 × 4) surface reconstruction is eliminated and the wetting layer becomes highly anisotropic. The anisotropy of this structure does not appear to arise from kinetic limitations but, rather, is attributed to the direction-dependent strain associated with the dimer-based surface reconstructions of the wetting layers.

In addition to the impact of the reconstruction symmetry on the morphology of the wetting layer, the differences in the amount of As on the surface are also expected to influence the formation of antimonide quantum dots. For example, because the As coverage on a c(4 × 4) reconstruction is 1.75 ML [24], compared to 0.5 ML for a (2 × 4) surface [25], we expect the initial formation of AlAs (GaAs) when Al (Ga) is incident on a GaAs-c(4 × 4) surface. Because this material would not be under strain, the coverage required to form quantum dots could be expected to be larger for the GaSb and AlSb dots grown on the c(4 × 4) reconstruction. This is in agreement with our earlier studies of AlSb QD formation on the c(4 × 4) reconstruction which found that QDs did not form until over 3 ML AlSb had been deposited [6, 22]. This should be contrasted with the current study which shows that QDs form after only 2 ML of AlSb is deposited on the (2 × 4) reconstruction. This issue becomes even more complex for the deposition of InSb QDs on the c(4 × 4) reconstruction, where the initial deposition of In would be expected to form InAs. (The InSb/GaAs mismatch is twice as large as the InAs/GaAs mismatch.) AFM measurements on the samples of Fig. 3 revealed a higher density of both QDs and dislocated islands for the (2 × 4) reconstruction compared to the c(4 × 4) reconstruction.

5. Summary

Epilayers of GaSb, AlSb and InSb, with mismatches of 8–15%, exhibit a Stranski–Krastanov growth mode on GaAs(0 0 1). In situ STM measurements reveal that the wetting layers are not uniform, but consist of anisotropic, ribbon-like structures oriented along the [1 1 0] direction, with characteristic separations of 40–50 Å and heights of a few angstroms. These wetting layers coexist with self-assembled quantum dots. The initial GaAs surface reconstruction affects both the wetting layer structure and the QD density.

Acknowledgements

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References


Buffer-dependent mobility and morphology of InAs/(Al,Ga)Sb quantum wells

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Abstract

We find that the mobility and morphology of InAs/(Al,Ga)Sb quantum wells grown by molecular beam epitaxy on lattice-mismatched GaAs depends strongly on the composition of the buffer layers. Using atomic force microscopy we show that GaSb buffers provide atomically flat interfaces on the scale of the electron Fermi wavelength for the quantum wells. In contrast, AlSb buffers generate a very rough interface on the same scale. Van der Pauw measurements show that this results in a greatly reduced low-temperature mobility for InAs quantum wells grown on AlSb buffers. We propose that the Ga adatoms tend to be more mobile than Al adatoms and therefore smoothen out the roughness generated in the nucleation layer. Using a GaSb buffer layer, we have achieved new record mobilities in InAs quantum wells.

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InAs quantum wells are promising for many device applications, including high-speed field effect transistors [1, 2], resonant tunneling diodes [3] or superconducting Josephson field effect transistors [4]. Bulk InAs has an inherently higher mobility than bulk GaAs [5]. One would therefore expect InAs quantum wells (QWs) to achieve higher mobilities than GaAs QWs. However, this has not been the case. Initial problems with growing InAs by MBE included the lattice mismatch with commonly available substrates (Si has a lattice constant \( a = 5.43 \text{ Å} \), GaAs has \( a = 5.65 \text{ Å} \) and InP has \( a = 5.87 \text{ Å} \) whereas InAs has \( a = 6.06 \text{ Å} \)). This was partially overcome by growing a thick buffer layer of GaSb or AlSb on GaAs [6]. The (Al,Ga)Sb is nearly lattice matched to InAs and the buffer layer serves to attenuate the dislocations due to the lattice mismatch with GaAs (7%). Another problem with growing high mobility InAs QWs has been the interface with (Al,Ga)Sb barriers. Since both the group-III and group-V elements change at the interface, there are two possible types of interface
[7]: InSb-like and AlAs-like. It has been found that the InSb-like interface is superior [7]. Even with these improvements, however, the mobility of the InAs QWs does not match that of GaAs/(Al,Ga)As QWs. In earlier work, Nguyen et al. [8], found that a GaSb buffer provided better mobilities in the InAs QW than a mixed GaSb/AlSb buffer. However, they could not provide an explanation for the difference, and did not look at samples with only AlSb in the buffer. In this work, we have further improved the mobility in InAs QWs by optimizing the buffer layer growth. We find that a GaSb buffer provides mobilities in the InAs QW nearly an order of magnitude greater than with an AlSb buffer. We have investigated this effect in detail and we show that it is directly related to the morphology induced by the different compositions of the buffer layer.

We begin all growths on semi-insulating GaAs substrates. To grow the 6.1 Å materials (AlSb, GaSb and InAs), we first provide a nucleation layer of ~1 nm AlAs and then grow approximately 80 nm of AlSb. This sequence has been shown [6] to quickly nucleate two-dimensional growth of the 6.1 Å materials. These nucleation layers are grown at 570°C. The buffer layer is then started and the temperature is then lowered, to 530°C. The buffer layer is either GaSb or AlSb of greater than 1 μm thickness. Since the goal of this work is to determine the effect of this buffer layer on the morphology and mobility of InAs QWs, this is the only parameter that is varied. After the buffer layer is grown, the bottom barrier of the QW is grown as an Al_{0.5}Ga_{0.5}Sb digital alloy superlattice, typically 100 nm thick. The digital alloy consists of 2.5 nm GaSb followed by 2.5 nm AlSb, periodically repeated to the necessary thickness. If high electron concentrations are desired in the QW, Te delta-doped layers are inserted in the barriers. 15 nm of InAs is then grown with an InSb-like interface. The samples for AFM measurements are complete at this point. The samples for mobility measurements have a top superlattice barrier on the QW and a surface GaSb cap to prevent Al oxidation.

Immediately after unloading the samples from the MBE system we scan the surface with a Digital Instruments Nanoscope III atomic force microscope. The samples have also been measured after being exposed for several days in the air and show no significant differences in the following results. In Fig. 1a and Fig. 1b we show the surface of the InAs as scanned by AFM. Fig. 1a shows the sample grown on a GaSb buffer, while Fig. 1b shows

![Fig. 1](https://example.com/fig1.jpg)

(a) 8 μm × 8 μm AFM scan of 15 nm InAs surface grown on 1.4 μm GaSb buffer. Slightly anisotropic mounds are visible with lateral dimensions ~ 1 × 1.5 μm and with height ~ 4 nm. (b) 8 μm × 8 μm scan of 15 nm InAs surface on 1.4 μm AlSb buffer showing slightly smaller (~ 0.5 × 0.5 μm) mounds but with the same vertical scale.
a sample grown on an AlSb buffer. On the scale shown, there is very little qualitative difference between the two surfaces. Both show characteristic features about 4 nm in height and with lateral dimensions about 1 \times 1.5 \mu m. These features are associated with screw dislocations for lattice-mismatched heteroepitaxy [9]. Since the mismatch between buffer and substrate is 7%, dislocations are nucleated in the first few monolayers of growth. However, by growing a thick enough buffer layer, the dislocations can be reduced by several orders of magnitude, to a steady state of \( \sim 10^8 \text{ cm}^{-2} \).

However, on a much smaller scale, we observe a significant difference between the two samples. In Fig. 2a and Fig. 2b, we see the same samples, but scanned at a resolution of 800 \times 800 nm. Fig. 2a, which shows the InAs grown on GaSb buffer, exhibits monolayer steps of InAs (height \( \sim 0.4 \text{ nm} \)) with a lateral width \( \sim 40 \text{ nm} \). In contrast, Fig. 2b shows the sample with the AlSb buffer, and shows trenches about 4 nm deep, but also with a characteristic lateral scale of about 40 \times 150 \text{ nm}.

Since the electrons that contribute to conductivity and hence mobility are all at the Fermi surface, only scattering potentials that vary on the order of the Fermi wavelength are important. For our samples, the Fermi wavelength \( \lambda_f = (2\pi/\bar{n})^{1/2} \), varies from 35 to 15 nm (\( \bar{n} \) from 5.5 \times 10^{11} to 4.3 \times 10^{12} \text{ cm}^{-2} \) in the lowest occupied subband). It is only scattering potentials on this scale that will significantly affect the mobility. Thus the variations shown in Fig. 1a and Fig. 1b do not affect the mobility, since the height fluctuations occur over 500 to 1500 nm, which is more than 10 times \( \lambda_f \). On the other hand, the variations shown in Fig. 2 should strongly affect the mobility. For the GaSb buffer, the variations are only monolayer steps, and this should correspond to only a small interface scattering potential in the InAs QW. However, for the AlSb buffer, the variations are an order of magnitude greater in height (but still on the same lateral scale) which is a significant fraction of the QW thickness, and so will generate a large scattering potential. Hence, the mobility of the samples with an AlSb buffer should be much less than those with a GaSb buffer because the AlSb grows much rougher.

![Fig. 2. (a) 800 nm \times 800 nm AFM scan of 15 nm InAs grown on GaSb showing monolayer terraces. The average terrace width is 40 nm. (b) 800 nm \times 800 nm scan of 15 nm InAs on 1.4 \mu m AlSb showing very deep trenches ( \( \sim 4 \text{ nm} \)) on a similar lateral scale as in (a).](image)

To confirm this, we also grew samples for measuring the mobility of the InAs QWs. These samples were identical to those grown for AFM measurements, except that a top barrier super-lattice and GaSb cap were grown on the InAs. Van
under Pauw measurements from room temperature to 12 K were performed to determine the mobility and carrier concentration of these samples. The samples with a GaSb buffer had consistently higher mobilities at low temperature than the samples with AlSb. This effect occurred over more than an order of magnitude difference in carrier concentration. The largest difference in mobility occurred for the lowest concentration, $n_s = 5.5 \times 10^{11} \text{cm}^{-2}$. In this case, the mobility for the sample with GaSb buffer (240 000 cm$^2$/V s) is almost seven times greater than the sample with the AlSb buffer (35 000 cm$^2$/V s). The Fermi wavelength for this sample is also most comparable to the lateral scale of the roughness. For a concentration of $1.3 \times 10^{12} \text{cm}^{-2}$, we achieved a record mobility of 944 000 cm$^2$/V s at 12 K with the GaSb buffer. In contrast, the same concentration but with an AlSb buffer only achieved 244 000 cm$^2$/V s at 12 K. Similar results were obtained for very high electron concentrations, e.g., $n_s = 7 \times 10^{12} \text{cm}^{-2}$. In this case, the low-temperature mobilities for GaSb and AlSb buffers were 236 000 and 91 000 cm$^2$/V s, respectively.

The preceding results have been consistently reproduced over a large number of MBE-grown samples. We can rule out impurity scattering from the buffer layer since this layer is 100 nm from the InAs QW. Furthermore, these same results have been obtained with separate charges of Al in the Al source furnace. The effect is also robust with respect to variations in the buffer layer growth temperature over nearly 100°C, with variations in GaSb cap thickness, and with the presence or absence of delta doping sheets.

The large mobilities we can achieve at low temperatures may be attributed primarily to the growth dynamics of AlSb versus GaSb. The AFM scans suggest that GaSb smooths out the roughness generated due to the lattice mismatch of GaAs and (Al,Ga)Sb. This is possible if the Ga adatoms have a high enough mobility to move around the surface during growth. In contrast, the lower mobility of Al adatoms causes them to stick approximately where they land. Thus AlSb tends to preserve the morphology of the underlying epilayers. This hypothesis is supported by other experiments [9, 10]. The mobilities we have obtained are competitive with the best mobilities achieved in GaAs QWs [11].

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References

Molecular beam epitaxy of vertically compact 
Al\textsubscript{x}Ga\textsubscript{1-x}As/GaAs laser-HEMT structures
for monolithic integration

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Abstract

Vertically compact Al\textsubscript{x}Ga\textsubscript{1-x}As/GaAs laser structures with Al\textsubscript{0.3}Ga\textsubscript{0.7}As/AlAs SPSL-cladding layers were grown on top of HEMT structures by MBE and investigated with regard to series resistance. We found an exponential dependence of series resistance $R_s$ with AlAs and Al\textsubscript{0.3}Ga\textsubscript{0.7}As SPSL-layer thicknesses at ambient temperature. Laser-HEMT structures for monolithic integration were grown and lasers were processed. For $3 \times 200 \, \mu\text{m}^2$ 3-QW-lasers, suitable for high-frequency performance, threshold currents of 20 mA and series resistances below 12 $\Omega$ were obtained. The slope of the linear regression of the $I_s$ values as a function of mesa width yielded a low threshold current density $j_{th}$ of 480 A/cm$^2$. Reduction of the p-cladding thickness from 600 nm down to 450 nm shows no increase in threshold current density if the p-dopants are kept from diffusing into the active region. This clearly demonstrates that laser structures for monolithic integration can be designed very compact without loosing performance.

1. Introduction

The monolithic integration of laser and electronic circuits represents a promising device technology to meet the growing demand for future information data transfer systems. An ideal semiconductor for short-range optical communication is the GaAs/Al\textsubscript{x}Ga\textsubscript{1-x}As system since even sophisticated vertical structures can be routinely grown by molecular beam epitaxy (MBE). However, in the case of monolithic integration, an increased complexity of the layer structure may lead to restrictions for parameters like growth temperatures, doping profiles and layer thicknesses which can result in lower device performance.

Growth of the low-temperature HEMT structure on top of the highly doped cap of the laser structure is excluded since it results in poor high-frequency performance. A processed laser structure overgrown in an second epitaxial run is also excluded due to poor crystal quality of the overgrown layer at the mesa edges [1]. Therefore, we grow the laser structure on top of the HEMT structure. This is done during one epitaxial run. A scheme of the

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vertical layer structure is given in Fig. 1. In order to obtain an easy to manufacture HEMT process, a vertically compact design of the laser structure is necessary. This can be achieved by a moderate thickness of the cladding layers with high aluminum content realized by short period Al$_{0.3}$Ga$_{0.7}$As/AlAs superlattices (SPLSs).

In this paper three points will be addressed: Firstly, the influence of the layer sequence of the SPLSs on the series resistance of the laser diode is studied. Secondly, DC data of lasers grown on top of the HEMT structures are reported. Finally, a successful reduction of p-cladding layer thickness is reported.

2. Growth of heterostructures and results

All samples were grown by MBE on 2 in GaAs substrates in a Varian Modular Gen II. The aluminum content in the ternary compound was $x = 0.3$. Layers with Al mole fractions between 0 and 0.3 were realized by GaAs/Al$_{0.3}$Ga$_{0.7}$As short-period superlattices (SPLSs) and those with $x > 0.3$ by Al$_{0.3}$Ga$_{0.7}$As/AlAs SPLSs. The growth rate for GaAs was 1.25 µm/h. It was controlled by reflection high-energy diffraction (RHEED) before and after the growth. The actual composition of the ternary compound, and by that, the aluminum contents and the growth rates of all other compositions were determined by additional RHEED measurement of the Al$_{0.3}$Ga$_{0.7}$As compound. The knowledge of the actual thicknesses of the various layers is important for the wet etching step during laser processing. The As$_4$/Ga beam equivalent pressure was below 20. Si and Be are used as n- and p-dopants, respectively. Doping concentrations are related to GaAs. Calibration of dopants was done with Hall effect measurements of GaAs layers.

For studying the influence of the AlAs and Al$_{0.3}$Ga$_{0.7}$As layer thicknesses in the cladding layers on the series resistance of the laser diode, samples with different SPLS-layer thicknesses were grown. The layer sequence of the test samples was the same as the structure displayed in Fig. 1. The mean Al mole fraction of the cladding layers was 80%. AlAs- and Al$_{0.3}$Ga$_{0.7}$As-layer thicknesses varied from 2 to 6 nm and 0.8 to 2.4 nm, respectively. The total thicknesses of the cladding layer was 600 nm. The Be concentration in the p-cladding layer was chosen to $1 \times 10^{18}$ cm$^{-3}$ to emphasize the effect of the short period layer thicknesses on $R_s$. Diode mesa with a cavity length of 200 µm and widths from 3 to 32 µm were fabricated by means of wet etching. Electrical contacts were obtained by evaporation of ohmic metals. Series resistance was determined by recording I–V curves of fabricated diodes at ambient temperature. $R_s$ is the differential resistance for diode voltages above 2 V. Fig. 2 shows the measured $R_s$ data for cavity widths of 3 µm (full diamonds) and 16 µm (full triangles). The y-axis is chosen in logarithmic scale. We found an exponential dependence of series resistance $R_s$ on AlAs and Al$_{0.3}$Ga$_{0.7}$As SPLS-layer thicknesses. This functional relation can be understood by taking into account tunneling processes of holes through the AlAs barriers. (The vertical hole transport in the cladding layer should be the main contribution to series resistance since vertical electron transport contributes much less to $R_s$, due to the much higher mobility.) Thus, if the AlAs barrier thickness increases the tunneling probability

![Diagram](image-url)

should decrease exponentially, e.g., the series resistance should increase similarly. However, at room temperature not only tunneling processes occur but also vertical transport in the p-doped superlattice driven by thermionic emission is present. With increasing AlAs barrier thickness, the Al\textsubscript{0.3}Ga\textsubscript{0.7}As quantum well width increases, too, in order to keep the mean Al mole fraction of 80%. This results in a higher energy difference between QW level and valence band edge. Thus, this second contribution also leads to higher series resistance with thicker AlAs layers since the number of carriers crossing the barriers gets smaller. However, the details of the vertical transport can only be discussed based on results for the temperature dependence of \( R_s \) and lies beyond the scope of this paper. As a result, acceptable \( R_s \) values can be obtained even with low Be doping of the cladding layers if the thicknesses of the SPSS layers are sufficiently thin.

Laser-HEMT structures for monolithic integration were grown as follows (see Fig. 1): First, the two-sided \( \delta \)-doped QW HEMT structure was grown at a substrate temperature \( T_s = 520^\circ C \) measured by pyrometer. This growth temperature is sufficiently low to suppress Si segregation into the two-dimensional electron gas (2DEG) channel. After growth of a thin etchstop layer the laser structure was grown. The \( T_s \) was raised to 660°C for the growth of the n-graded layer, the n-cladding layer and the separate confinement layers with the laser active region. This active region consists of three 7.4 nm thick quantum wells which are separated by 7.2 nm thick Al\textsubscript{0.25}Ga\textsubscript{0.75}As barriers. At the beginning of the lower confinement layer, \( T_s \) was lowered for the growth of a few monolayers to prevent Si from segregating into the quantum wells. The p-cladding layer and p-graded layer were grown at slightly reduced temperatures of 625°C and 590°C, respectively. Finally, the heavily-doped GaAs cap layer was grown at \( T_s = 520^\circ C \).

For the AlAs and Al\textsubscript{0.3}Ga\textsubscript{0.7}As layers in the SPSS-claddings of the actual laser-HEMT structure we choose a thickness of 3 nm and 1.2 nm, respectively. These values seem reasonable with regard to a limitation of shuttering times and frequency. The Si-doping concentration of the n-buffer, n-graded layer, and n-cladding layer was \( 4 \times 10^{18} \text{ cm}^{-3} \).

Resistivities for higher p-doping concentrations were estimated from Fig. 2 on the supposition that the mobility is only weakly dependent on \( N_{\text{be}} \) in the considered doping regime and that the p-cladding layer and the p-graded layer contribute most to \( R_s \). In order to obtain a series resistance of 15 \( \Omega \) the doping concentration of the p-cladding was set to \( 1 \times 10^{18} \text{ cm}^{-3} \) for the first 300 nm and raised to \( 2 \times 10^{18} \text{ cm}^{-3} \) for the growth of the remaining 300 nm. This doping profile leads to an effective lowering of Be diffusion into the active region for two reasons. First, the Be concentration is held below the solubility limit of \( 2 \times 10^{18} \text{ cm}^{-3} \) in Al\textsubscript{0.8}Ga\textsubscript{0.2}As. Solubility limits of Be in Al\textsubscript{1-x}Ga\textsubscript{x}As as a function of Al mole fraction were found in Be doped samples by depth profiling with Secondary Ion Mass Spectrometry [2]. Our investigations have clearly demonstrated the ability of Be to diffuse through the complete p-cladding layer if the intended doping concentration in the cladding layer is beyond these upper limits. It was also shown that in this case this diffusion process cannot be suppressed by using low growth temperatures. As a second point, if the Be concentration is below the solubility limit like in the laser-HEMT structure reported here, Be diffusion can be reduced by lowering the growth temperature \( T_s \). For that
reason \( T_b \) values of 625°C and 590°C were chosen for the p-cladding and the p-graded layer, respectively. The p-graded layer was doped with \( 8 \times 10^{18} \text{ cm}^{-3} \). Laser diodes with fixed cavity length and various widths were processed by means of wet etching and evaporation of ohmic metals. Details of the process technology can be found in Ref. [3]. Finally, laser mirrors were fabricated by either reactive ion etching (RIE) or chemical assisted ion-beam etching (CAIBE). These two methods have proved to be highly reproducible and have yielded Al\(_x\)Ga\(_{1-x}\)As/GaAs lasers with almost the same threshold currents as lasers with cleaved facets.

\( I_{th} \) and \( R_s \) were determined by on-wafer measurements. Here, \( R_s \) is the differential resistance for a diode current 5 mA above threshold. The lasing wavelength was 850 nm. The quality and homogeneity of the laser mirrors were tested by measuring 40 identical lasers which were located on a diameter of the wafer. We found a scattering of the \( I_{th} \) data points of less than 10%. Values for \( I_{th} \) and \( R_s \) for mesa widths between 3 and 32 \( \mu \text{m} \) are shown in Fig. 3. For 3 \( \times \) 200 \( \mu \text{m}^2 \) 3 QW-lasers we found threshold currents \( I_{th} \) below 20 mA and series resistances below 12 \( \Omega \). The slope of the linear regression of the \( I_{th} \) values yielded a threshold current density \( j_{th} \) of 480 A/cm\(^2\). This is almost the same value as obtained for high-speed Al\(_x\)Ga\(_{1-x}\)As/GaAs lasers grown in our laboratory on top of a 1 \( \mu \text{m} \) buffer under optimized growth conditions [4]. The DC performance of our laser grown on the HEMT structure can be compared with data from literature and with earlier results. Nichols et al. [5] obtained a threshold current density of 700 A/cm\(^2\) and a series resistance of 50 \( \Omega \) for a strained GaAs/AlGaAs/GaInAs MQW laser structure grown on top of a MESFET by MOCVD although much thicker cladding layers were used. The circuits operated with bandwidths as high as 3.4 GHz. A similar bandwidth of a MBE grown GaInAs/GaAs MODFET/laser device was reached by Offsey et al. [6]. Typical threshold currents were 80 mA for the \( 20 \times 800 \mu \text{m}^2 \) lasers. Earlier data of monolithic integrated laser-HEMT devices from our laboratory were reported by Hornung et al. in Ref. [7]. Typical results were a threshold current of 28 mA and a series resistance of 22 \( \Omega \) for a \( 4 \times 360 \mu \text{m}^2 \) laser diode grown on top of a HEMT structure. Laser diodes and driver circuits showed performance up to data rates of 7.4 Gbit/s in a short-range optical data transfer setup. Hence, taking the actual vertical layer structure as a starting point it should be possible to obtain a fully integrated laser-HEMT device exceeding the bandwidth reported in Ref. [7].

Finally, attempts were made to make the laser structure even more compact. Laser-HEMT structures with different p-cladding thicknesses but identical doping profiles were grown. Cladding layers

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**Fig. 3.** Threshold currents and series resistances of 3-QW-lasers grown on top of a HEMT structure with mesa widths between 3 and 32 \( \mu \text{m} \) and a cavity length of 200 \( \mu \text{m} \).

**Fig. 4.** Threshold currents of 3-QW-lasers grown on top of a HEMT structure with 450 and 600 nm p-cladding thicknesses.
were doped as reported above with the exception of the p-graded and the cap layers which were doped throughout with $2 \times 10^{18}$ cm$^{-3}$. This was done to keep Be diffusion into the active region to a minimum. Fig. 4 shows the results for laser diodes with a cavity length of 200 μm and widths between 3 and 32 μm. Threshold current densities below 485 A/cm$^2$ were found for both p-cladding thicknesses of 600 and 450 nm, respectively. $R_s$ were above 30 Ω as expected. Nevertheless, no increase in $j_{th}$ was observed upon reducing the cladding layer thickness by 150 nm.

3. Conclusion

Vertically compact Al$_{x}$Ga$_{1-x}$As/GaAs laser structures grown on top of HEMT structures with Al$_{0.3}$Ga$_{0.7}$As/AlAs SPSL-cladding layers were grown by MBE. We found an exponential dependence of series resistance $R_s$ on AlAs and Al$_{0.3}$Ga$_{0.7}$As SPSL-layer thickness at ambient temperature. Laser-HEMT structures suitable for high-frequency performance were grown and processed. A threshold current density of 480 A/cm$^2$ was reached for a 3-QW-laser grown on top of a low-temperature grown HEMT. Laser-HEMT structures grown with different p-cladding thicknesses showed no increase in threshold current density upon reducing the cladding layer by 150 nm. This clearly demonstrates that laser structures for monolithic integration can be designed even more compact when the p-doping profile is chosen in a way that p-dopants are kept from diffusing into the active region.

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References

MBE growth of quaternary InGaAlAs layers in InGaAs/InAlAs HBTs to improve device performance


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Abstract

InAlAs/InGaAs HBTs grown on InP substrates can achieve excellent high-frequency performance but suffer from low breakdown and low current gain due to the small InGaAs bandgap. By replacing the InGaAs collector and/or base with a quaternary InGaAlAs layer the device characteristics are improved. Using an InGaAlAs collector the breakdown voltage is found to increase the breakdown voltage of the device from 15 to 23 V. Using such a layer in both the collector and base has the added benefit of increasing the device current gain from 40 to greater than 60. Current gain can be increased even more dramatically to 88 by using a graded InGaAlAs layer in the base. However, the use of InGaAlAs layers was found to increase the device $V_{BE}$ voltage and degrade the RF properties somewhat. These parameters can be traded off and optimized for a given application.

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Keywords: MBE; InGaAs; InGaAlAs; Heterojunction bipolar transistor

1. Introduction

InAlAs/InGaAs HBTs grown on InP substrates can provide high-frequency performance useful in many applications. [1, 2]. However, these devices tend to have low breakdown voltage and low current gain. These limitations can be improved by adding a larger bandgap material to either the collector and/or the base of the device [3, 4]. In this paper we report the effects of using InGaAlAs layers for this purpose. The use of this quaternary allows the bandgap to be varied while keeping the layer lattice matched to the InP substrate. When an InGaAlAs layer is used only in the collector the breakdown voltage is increased but complex graded quaternary layers must be grown surrounding the collector layer and a thin, pulse-doped layer must be added to suppress current blocking [5]. This collector structure has been shown to achieve a combination of high-breakdown and high-frequency performance [6]. Alternatively, both the collector and base layers can be grown at a fixed InGaAlAs concentration. In addition to increasing the breakdown voltage, this structure also increases the device current gain, probably due to a reduction...
in Auger recombination in the base. In this paper we compare these two approaches, and also the use of a graded composition InGaAlAs layer in the base to reduce the base transit time.

2. HBT device structure and growth

The device wafers for this study were grown in a Gen II modular MBE system using nonIndium-bonded 2-in (1 0 0)InP substrates. All the layers in the structures were grown nominally lattice-matched to the InP substrate with a growth rate of 15 nm/min. The InGaAlAs graded layers were graded in composition by continuously varying the Ga and Al concentrations while maintaining a lattice constant equal to that of the InP substrate. Substrate temperatures in the vicinity of 500°C were used for the growth of the different layers. Growth conditions were optimized to reduce any Be diffusion from the base into the emitter. All the layers except the base were doped n-type using a Si doping source.

The wafers grown for this study consisted of variations on two basic profiles shown in Figs. 1 and 2. In Fig. 1 the structure for a pulse-doped InGaAlAs collector with graded InGaAlAs layers and an InGaAs base is shown. This profile consists of an n⁺ InGaAs subcollector layer followed by an n⁺ InGaAlAs layer graded in composition to the desired InGaAlAs collector composition. The collector layers consist of an n⁺ InGaAlAs collector layer followed by a 10 nm InGaAlAs layer doped in the mid-10¹⁷ range to compensate the quasi-field generated by the subsequent InGaAlAs layer graded in composition to the InGaAs base. The base region consisted of an 80 nm InGaAs base doped with Be to a level of 3×10¹⁸ cm⁻³ and an InGaAs spacer layer. The emitter consisted of a graded InGaAlAs layer, followed by an InAlAs emitter and a heavily Si-doped InGaAs contact layer. The quaternary collector of this device is used to increase the breakdown voltage of the device while the pulse-doping serves to reduce current blocking effects as we described in an earlier publication [5]. With this structure we used Al compositions of In₀.₅₃(Ga₀.₅₅Al₀.₄₅)₀.₄₅As, In₀.₅₃(Ga₀.₇₅Al₀.₂₅)₀.₄₅As, and In₀.₅₂Al₀.₄₈As.

In Fig. 2 is shown the alternate structure which is formed by growing a relatively low Al-composition InGaAlAs layer for both the collector and base, avoiding the need for both the graded-composition layer at the base collector junction and the pulse-doped layer. The aluminum composition of the

![Fig. 1. Schematic cross-section of InP-based, InGaAlAs pulse-doped collector HBT.](image-url)
collector–base layers was chosen to give bandgap energy of 0, 50, 100, and 200 meV greater than InGaAs. This corresponds to Al compositions of $x = 0, 0.07, 0.15, \text{and } 0.29$ for In$_{0.53}$Ga$_{0.47}$As [7]. Except for the $x = 0.29$ wafer, the remainder of the structure is the same as the pulse-doped structure of Fig. 1. For the $x = 0.29$ wafer the Al composition of the initial emitter layer was increased to preserve the step in Al composition at the base–emitter junction.

An additional wafer was grown using the pulse-doped collector layers shown in Fig. 1 but with a graded-composition InGaAlAs base. For this sample the base was graded from In$_{0.53}$Ga$_{0.47}$As at the base–collector junction to In$_{0.53}$Ga$_{0.83}$Al$_{0.17}$As at the base–emitter junction. This results in a built-in quasi-electric field across the base of $\sim 14.5 \text{ kV/cm}$ to reduce the electron transit time.

3. Material characterization

After growth, the samples were analyzed with double-crystal X-ray diffraction (DXRD) and Photoreflectance (PR). Representative DXRD spectra for some of these samples is shown in Fig. 3. These spectra indicate very close lattice-matching of all layers, typically to within 100 arcsec of the InP substrate, which corresponds to a difference in composition equivalent to that between In$_{0.53}$Ga$_{0.47}$As and In$_{0.525}$Ga$_{0.475}$As. This extremely accurate and precise control over composition and lattice-constant is made possible by MBE.

Representative PR spectra taken from some of these samples are shown in Fig. 4. As discussed in a previous publication [6], these spectra provide information on both the compositions of the various layers and the built-in electric fields. For the samples with a constant composition base and collector, the spectral region between 0.7 and 1 eV displays an oscillation related to the base-collector junction. This oscillation corresponds to an energy gap in the base of 0.82, 0.86, and 0.98 eV which gives bandgap shifts referenced to In$_{0.53}$Ga$_{0.47}$As of approximately 70, 110, and 230 meV, respectively. The remaining oscillations in the spectra arise from the base-emitter layers of the device [8]. The oscillations for Al compositions of 0, 0.07, and 0.15 are all very similar as can be seen in Fig. 4. This is in contrast to the sample with the Al concentration of 0.29 which displays differences due to the alternate emitter grading described above.
The PR spectra for the graded-composition base device was found to be very similar to the pulse-doped PR spectra. It displayed a high-field oscillation near 0.8 eV associated with the graded base and also displayed collector and emitter oscillations identical to that of a In$_{0.53}$(Ga$_{0.71}$Al$_{0.29}$)$_{0.47}$As collector sample with an InGaAs base.

4. Device performance

The use of InGaAlAs layers in either the collector or in base and collector dramatically improved the DC characteristics of the HBT devices as can be seen in Table 1. The most noticeable improvement is an increase in device breakdown voltage $BV_{CEO}$. For the pulse-doped collector $BV_{CEO}$ can be increased from 15 V for an InGaAs collector to over 23 V for an InAlAs collector. The devices of Fig. 2 with both a quaternary base and collector showed an even larger increase in breakdown voltage with Al composition as can be seen in Fig. 5. Here $BV_{CEO}$ is again seen to increase to 23 V, but at a composition of only In$_{0.53}$(Ga$_{0.71}$Al$_{0.29}$)$_{0.47}$As. This is due to the higher composition at the base-collector junction in these structures compared to the pulse-doped samples.

An additional benefit to be gained by using the InGaAlAs base and collector profile of Fig. 2 is the increase in current gain, from $\beta = 40$ for an InGaAs base to greater than $\beta = 60$ for the In$_{0.53}$(Ga$_{0.71}$Al$_{0.29}$)$_{0.47}$As base plus collector device. We attribute this increase in current gain to a reduction of Auger recombination in the larger bandgap base. In addition, for the In$_{0.53}$(Ga$_{0.71}$Al$_{0.29}$)$_{0.47}$As sample the alternate emitter grading may also be contributing to higher beta. To increase the current gain beyond these levels, the base was compositionally graded from In$_{0.53}$Ga$_{0.47}$As to In$_{0.53}$(Ga$_{0.83}$Al$_{0.17}$)$_{0.47}$As over the 80 nm base. The resulting built-in field substantially reduced the base transient time as can
Fig. 4. PR spectra for samples with base and collector compositions of (a) $\text{In}_{0.55}\text{Ga}_{0.45}\text{As}$, (b) $\text{In}_{0.55}\text{Ga}_{0.93}\text{Al}_{0.07}\text{As}$, (c) $\text{In}_{0.55}\text{Ga}_{0.85}\text{Al}_{0.15}\text{As}$, and (d) $\text{In}_{0.55}\text{Ga}_{0.71}\text{Al}_{0.29}\text{As}$. Dashed line highlights the oscillation from the base region.

<table>
<thead>
<tr>
<th>Profile</th>
<th>Peak $\beta$</th>
<th>$BV_{\text{be}}$ (V)</th>
<th>Base sheet ($\Omega$/sq)</th>
<th>$V_{\text{be}}$ at 1 $\mu$A (V)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\text{In}<em>{0.55}\text{Ga}</em>{0.45}\text{As}$ collector + base</td>
<td>40</td>
<td>15</td>
<td>460</td>
<td>0.320</td>
</tr>
<tr>
<td>$\text{In}<em>{0.55}\text{Ga}</em>{0.93}\text{Al}_{0.07}\text{As}$ collector + base</td>
<td>43</td>
<td>19</td>
<td>490</td>
<td>0.363</td>
</tr>
<tr>
<td>$\text{In}<em>{0.55}\text{Ga}</em>{0.85}\text{Al}_{0.15}\text{As}$ collector + base</td>
<td>47</td>
<td>22</td>
<td>545</td>
<td>0.411</td>
</tr>
<tr>
<td>$\text{In}<em>{0.55}\text{Ga}</em>{0.71}\text{Al}_{0.29}\text{As}$ collector + base</td>
<td>62</td>
<td>23</td>
<td>627</td>
<td>0.509</td>
</tr>
<tr>
<td>$\text{In}<em>{0.55}\text{Ga}</em>{0.50}\text{Al}_{0.50}\text{As}$ collector</td>
<td>38</td>
<td>20</td>
<td>460</td>
<td>0.320</td>
</tr>
<tr>
<td>$\text{In}<em>{0.52}\text{Al}</em>{0.48}\text{As}$ collector</td>
<td>41</td>
<td>21</td>
<td>460</td>
<td>0.343</td>
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<tr>
<td>$\text{In}<em>{0.55}\text{Ga}</em>{0.50}\text{Al}_{0.50}\text{As}$ collector + graded</td>
<td>40</td>
<td>23.5</td>
<td>460</td>
<td>0.338</td>
</tr>
<tr>
<td>$\text{In}<em>{0.55}\text{Ga}</em>{0.83}\text{Al}_{0.17}\text{As}$ base</td>
<td>88</td>
<td>16</td>
<td>480</td>
<td>0.405</td>
</tr>
</tbody>
</table>

be seen from the large increase in current gain to 88.

The breakdown and current gain benefits to be gained from the insertion of the InGaAlAs collector and base layers must be traded off against an increase in the device $V_{\text{be}}$, base sheet resistance, and also a reduction in the RF properties. As can be seen again from Table 1, the addition of Al to the InGaAs base results in an increase in the $V_{\text{be}}$ of the devices which scales directly with the increase in bandgap at the base–emitter junction. For the quaternary base and collector devices we see increases in $V_{\text{be}}$ of 43, 91, and 189 meV compared to the measured increases in bandgap using PR of 70,
with close to twice the current gain of the comparable In_{0.53}(Ga_{0.85}Al_{0.15})_{0.47}As base and smaller increases in $V_{be}$ and base sheet resistance.

A major concern in the use of the quaternary InGaAlAs in the collector of the device is the effects of current blocking, which limits the RF performance of the device. Small (1 $\times$ 10 $\mu$m) dual emitter devices were fabricated to look at the impact of the quaternary collector on RF performance. The resulting performance is shown in Table 2. In Table 2 it can be seen that the addition of any amount of Al results in a drop in both $f_T$ and $f_{max}$. For the InGaAlAs base and collector samples $f_T$ drops off to 50 GHz compared to 80 GHz for a pure InGaAs base and collector. At the same time the current density at which the $f_T$ curve rolls over also drops substantially from 100 to 35 kA/cm$^2$. Similar reductions in $f_T$, $f_{max}$, and $J_{max}$ can be seen in Table 2 for the InGaAlAs collector only samples.

From these results it is clear that the addition of Al to both the collector and base results in a relatively large degradation of device performance, particularly in decreasing the current at which the $f_T$ curve rolls over. Presumably a lower electron saturation velocity in InGaAlAs compared to InGaAs is the cause of this effect. The plot of $f_T$ with Al concentration shown in Fig. 6 indicates the severity of the decrease in $f_T$ particularly for the quaternary base plus collector structures. Note also that the graded base device contained an In_{0.53}(Ga_{0.59}Al_{0.41})_{0.47}As collector which consequently limited its RF performance. The combination of such a graded base with an InGaAs

![Graph showing variation in BV_{be} with aluminum concentration for a 75 $\times$ 75 $\mu$m HBT device. Closed circles represent InGaAlAs base and collector devices while open circles represent InGaAlAs collector only devices.](image)

110, and 230 meV. For the graded base structure we see an increase in $V_{be}$ of 85 meV for a nominal increase in bandgap of 116 meV at the base-emitter junction. The base sheet resistance also increased from 460 $\Omega$/sq for an InGaAs base to 627 $\Omega$/sq for an In_{0.53}(Ga_{0.71}Al_{0.29})_{0.47}As. In terms of base-emitter properties the graded base is clearly superior

**Table 2**

<table>
<thead>
<tr>
<th>Profile</th>
<th>Peak $f_T$ (GHz)</th>
<th>Peak $f_{max}$ (GHz)</th>
<th>$J_{max}$ at 1.5 V (kA/cm$^2$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>In_{0.53}Ga_{0.47}As collector + base</td>
<td>80</td>
<td>180</td>
<td>100</td>
</tr>
<tr>
<td>In_{0.53}(Ga_{0.02}Al_{0.98})_{0.47}As collector + base</td>
<td>64</td>
<td>130</td>
<td>50</td>
</tr>
<tr>
<td>In_{0.53}(Ga_{0.85}Al_{0.15})_{0.47}As collector + base</td>
<td>50</td>
<td>105</td>
<td>35</td>
</tr>
<tr>
<td>In_{0.53}(Ga_{0.71}Al_{0.29})_{0.47}As collector + base</td>
<td>50</td>
<td>94</td>
<td>35</td>
</tr>
<tr>
<td>In_{0.53}(Ga_{0.56}Al_{0.54})_{0.47}As collector</td>
<td>57</td>
<td>140</td>
<td>45</td>
</tr>
<tr>
<td>In_{0.53}(Ga_{0.23}Al_{0.78})_{0.47}As collector</td>
<td>49</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>In_{0.53}Al_{0.47}As collector</td>
<td>40</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>In_{0.53}Ga_{0.50}Al_{0.50}_{0.47}As collector + graded</td>
<td>56</td>
<td>126</td>
<td>40</td>
</tr>
<tr>
<td>In_{0.53}(Ga_{0.85}Al_{0.15})_{0.47}As base</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>
in both the base and collector was found to substantially increase the breakdown voltage of the device. Use of a quaternary InGaAlAs layer in the base was also found to increase the device current gain compared to an InGaAlAs base, while a more substantial increase in the gain was achieved by grading the Al composition of the base. These performance benefits were found to come at a cost of an increase in device $V_{bc}$ and a degradation of RF performance. This presents the opportunity of tailoring the use of these layers for applications where some tradeoff in $V_{bc}$ and RF performance is acceptable and a large advantage is to be gained from higher breakdown and larger current gain.

**Acknowledgements**

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**References**


Improved hole transport properties of highly strained In$_{0.35}$Ga$_{0.65}$As channel double-modulation-doped structures grown by MBE on GaAs

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Abstract

Hole transport properties have been improved by using highly strained In$_{0.35}$Ga$_{0.65}$As channel double-modulation-doped heterostructures grown by molecular beam epitaxy. This structure provided both a high mobility of 354 cm$^2$/V s and a high sheet hole concentration of $1.23 \times 10^{13}$ cm$^{-2}$ at room temperature. Double-modulation-doped field effect transistors with a 0.4-μm gate length and a 20-μm gate width were fabricated. Transconductance of 118 mS/mm, which is about 1.5 times higher than that of single-modulation-doped field-effect transistors, was obtained at room temperature.

1. Introduction

To obtain high performance complementary GaAs-based circuits, which have the potential to provide low power-consumption and high-efficiency operation, improved transconductance in the p-channel field-effect transistors (FETs) is needed. The hole transport properties of GaAs-based p-channel FETs can be improved by using a strained InGaAs layer as a channel due to the strain-induced splitting of the valence band [1]. This splitting decreases the average hole effective mass whose momentum vector is parallel to the channel, and increases the hole mobility. Thus, greater splitting is desired so that a larger concentration of light holes can act as carriers.

To increase the strain-induced separation between light- and heavy-hole bands, the In content of the InGaAs layers needs to be increased. However, growing high quality In$_x$Ga$_{1-x}$As layers ($x \geq 0.3$) on GaAs substrates has been difficult due to the surface segregation of In atoms, even when the layer thickness is less than the critical thickness [2].

Recently, it was reported that the In content of InGaAs layers pseudomorphically grown on GaAs substrates could be raised above 0.3 by optimizing the growth condition [3, 4]. These highly strained layers improved the electron transport properties; mobility of over $6500$ cm$^2$/V s and sheet electron concentration of $2.94 \times 10^{12}$ cm$^{-2}$ were obtained.

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from highly strained AlGaAs/InGaAs modulation-doped structures [3].

We previously showed that hole transport properties are also improved by using this highly strained layer as a channel of a p-type double-modulation-doped (D-MOD) Al_{0.35}Ga_{0.65}As/ GaAs/In_{0.35}Ga_{0.65}As heterostructure [5]. We obtained a mobility of 284 cm²/V s and a sheet hole concentration of $1.84 \times 10^{12}$ cm⁻².

In this paper, we show that the hole mobility can be further improved by optimizing the growth conditions. We also discuss our results from testing fabricated p-channel double-modulation-doped field-effect transistors (D-MODFETs) that use the highly strained In_{0.35}Ga_{0.65}As layer as a channel.

2. Experimental details

All samples for this study were grown with an ANELVA 850 V conventional solid-source molecular beam epitaxy (MBE) system. Before growing the FET structures, we grew three types of samples to check the quality of the epitaxial layers, whose structures are shown in Fig. 1. The thermal cleaning temperature of the semi-insulating (1 0 0) GaAs substrates was 600°C in an As₄ flux of 1 × 10⁻³ Torr. The substrate temperature for the growth of the 200-nm-thick undoped GaAs and 100-nm-thick undoped AlGaAs (Al content: 0.39) buffer layers was 580°C. This was reduced to 500°C after growing the buffer layers to suppress the Be surface segregation. After growing a 5-nm-thick p-AlGaAs, a $d_{sp}$-nm-thick undoped AlGaAs and a 2-nm-thick undoped GaAs layer, the substrate temperature was further reduced to 420°C to suppress the In surface segregation. After growing a 6-nm-thick InGaAs (In content: 0.35) layer and 2-mm-thick undoped GaAs layer, the growth temperature was increased to 500°C and a $d_{sp}$-nm-thick undoped AlGaAs, a 6-mm-thick p-AlGaAs, and a 10-mm-thick undoped AlGaAs layer, plus a 10-mm-thick p-GaAs cap, were grown at 500°C. The growth temperature was changed with a 2-5-min growth interruption under As₄ flux. The growth temperature of 420°C is optimum for the growth of pseudomorphic In_{0.35}Ga_{0.65}As layers on GaAs with a high In content of 0.35 [6], and 500°C is the upper limit for the growth of a Be-doped AlGaAs layer in which the segregation and/or diffusion of the Be atom is suppressed to the order of 3-4 monolayers [7]. The Be concentration of the carrier-supplying layer below the channel was $1.5 \times 10^{18}$ cm⁻³, while the Be concentration of the carrier-supplying layer above the channel was either $5 \times 10^{16}$ cm⁻³ (sample 1) or $3 \times 10^{18}$ cm⁻³ (sample 2 and 3). The thickness of the AlGaAs spacer was 4 nm (sample 1 and 2) or 8 nm (sample 3). The GaAs growth rate was fixed at 1.0 μm/h. The In content was calibrated by photoluminescence (PL) measurements at 77 K using an InGaAs/GaAs single quantum well structure [8] and the Al content was calibrated in the same way using a thick AlGaAs layer.

The mobility and sheet hole concentration of these samples were evaluated by the conventional van der Pauw method at room temperature (RT).

The layer structure (from the substrate) of the p-channel D-MODFET consists of 200-nm-thick
undoped GaAs, 100-nm-thick undoped AlGaAs (Al content: 0.39), 4-nm-thick p-AlGaAs (sample 4: $2.1 \times 10^{18}$ cm$^{-3}$, sample 5: $4.2 \times 10^{18}$ cm$^{-3}$). Notice that doping concentration was increased to cancel the process damage while the gate recess), 3-nm-thick undoped AlGaAs, 2-nm-thick undoped GaAs, 6-nm-thick undoped InGaAs (In content: 0.35), 2-nm-thick undoped GaAs, 3-nm-thick undoped AlGaAs, 4-nm-thick p-AlGaAs ($1.5 \times 10^{19}$ cm$^{-3}$), 11-nm-thick undoped AlGaAs, 20-nm-thick undoped GaAs, 3-nm-thick undoped AlGaAs and 160-nm-thick p-GaAs ($4 \times 10^{19}$ cm$^{-3}$).

A single modulation-doped field effect transistor (S-MODFET) was also fabricated as a reference (sample 6). The layer structure was the same as that of the D-MODFET except that the GaAs buffer layer was 300 nm thick, and the 100-nm-thick undoped AlGaAs and the 4-nm-thick p-AlGaAs layers were not grown.

Conventional mesa isolation, nonalloyed ohmic contacts for source and drain, and a double recessed gate process were used in the device fabrication. The gate lengths were 0.4 or 1.75 μm, and the gate width was 20 μm.

### 3. Results and discussion

The mobility of the D-MOD structure as a function of sheet hole concentration is shown in Fig. 2. The results of previously reported p-channel modulation-doped heterostructures [5, 9, 10] are also plotted for comparison. Very high mobility of 354 cm$^2$/(V s) was obtained at a sheet hole concentration of $1.23 \times 10^{12}$ cm$^{-2}$ (sample 3). At a higher concentration of $2.88 \times 10^{12}$ cm$^{-2}$, still higher mobility of 276 cm$^2$/(V s) was obtained (sample 1). As far as we know, these mobilities are the highest yet reported at these hole concentrations. These high mobilities were achieved by individually optimizing the growth temperatures of each layer. This confirms that the Be segregation and/or the diffusion length at 500°C was smaller than the spacer thickness of 4 nm. The discrepancy in the mobility between this work and our previous report [5] is due to the different growth temperature used for the carrier-supplying layer. In our previous report, the growth temperature for the carrier-supplying layer (420°C) appears to have been too low to maintain good crystalline quality in the AlGaAs layers.

The typical current–voltage characteristic of the highly strained InGaAs channel D-MODFET and the extrinsic transconductance and drain current as a function of gate bias at room temperature are shown in Fig. 3. These figures show that this D-MODFET offers good FET characteristics. At a gate length of 0.4 μm, the maximum extrinsic transconductance exceeded 100 mS/mm constantly in the 3-in. wafer used to fabricate the FETs. The highest value of extrinsic transconductance reached in this work was 118 mS/mm at a gate length of 0.4 μm (sample 5). The source resistance was about 5 Ω mm.

The relation between transconductance and gate length is shown in Fig. 4. The results from the S-MODFETs and other p-channel FETs [1, 9–14] are also plotted for comparison. To our knowledge, the transconductance of 118 mS/mm is the highest yet reported. This shows that highly strained D-MODFET structures can be used to improve the performance of p-channel FETs.

Transconductance of 82 mS/mm was obtained for the S-MODFET (sample 6) at a gate length of 0.4 μm. This difference in the transconductances
between the D-MODFET and the S-MODFET was due to the difference in the maximum carrier concentration that could be accumulated in the channel while maintaining high mobility. It is difficult for S-MOD structures to maintain high mobility (above 260 cm²/(V·s)) at sheet hole concentrations exceeding $2 \times 10^{12}$ cm⁻² [5]. The improved carrier confinement in the channel also improved the transconductance of the D-MODFETs, while S-MODFETs do not have a potential barrier below the channel (only a GaAs buffer layer exists between the channel and the substrate).

Fig. 4. The relation between transconductance and gate length at RT. The curve is a guide for the eyes.

4. Conclusions

A highly strained In₀.₃Ga₀.₆₅As layer can be used as a channel in p-type AlGaAs/InGaAs D-MOD heterostructures grown on GaAs substrates by MBE. Having grown such a structure, we obtained a high mobility of 354 cm²/(V·s) and a high sheet hole concentration of $1.23 \times 10^{12}$ cm⁻² at room temperature. At a higher concentration of $2.88 \times 10^{12}$ cm⁻², a still higher mobility of 276 cm²/(V·s) was obtained. These high mobilities were achieved by individually optimizing the growth temperatures used for each layer.

The transconductance of a D-MODFET with a 0.4-μm gate length and a 20-μm gate width reached 118 mS/mm. To our knowledge, this is the highest transconductance yet reported.

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References


MBE growth of double-sided doped InAlAs/InGaAs HEMTs with an InAs layer inserted in the channel

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Abstract

InAlAs/InGaAs high electron mobility transistors (HEMTs) with an InAs layer inserted in the In$_{0.53}$Ga$_{0.47}$As channel were grown on InP-substrates by solid source molecular beam epitaxy (MBE). By optimizing position, thickness and growth temperature of the InAs layer, the mobility at 300 K has been increased by 47%. HEMTs with a gate length of 0.15 μm showed very high transconductances and cut-off frequencies $f_T$ in excess of 1270 mS/mm and 235 GHz, respectively.

PACS: 81.05.Ea; 81.15.Hi

Keywords: MBE; HEMT; InGaAs; InAlAs; InAs

1. Introduction

InP-based InAlAs/InGaAs high electron mobility transistors (HEMTs) have demonstrated excellent high-frequency and low-noise performance. These are explained by the high electron mobility, saturation drift velocity and sheet carrier density of the InAlAs/InGaAs two-dimensional electron gas (2DEG) system. In this work we investigate InAlAs/InGaAs HEMTs with a thin InAs layer inserted into the InGaAs channel. This structure improves the electron transport properties. We optimized the structure by changing the position, thickness and growth temperature of the InAs layer.

In conventional HEMT structures high contact and sheet resistances in the access regions between alloyed contact and gate electrode can cause large parasitic source and drain resistances. This is effected by the large conduction band discontinuity between the InGaAs cap layer and InAlAs supply layer, forming a barrier in the current flow between these layers. We demonstrate that an additional Si-$\delta$-doping at the interface of the cap and supply layer reduces the barrier heights and lowers the parasitic resistances, leading to excellent DC and RF characteristics.
2. MBE growth of InAs

The heterostructures were grown by solid source MBE on Fe-doped semi-insulating (1 0 0) InP epilayer substrates with growth rates of the lattice matched InAlAs and InGaAs layers around 1 μm/h, and a V/III ratio of 27 (beam equivalent pressure). Recently [1], we have demonstrated that a channel thickness of 12 nm and double sided δ-doping are essential for low output conductance and kink-free I/V characteristics of lattice-matched HEMTs. Therefore, the HEMT structure consists of a 12 nm wide InGaAs channel, in which a strained InAs layer was embedded, Fig. 1a. Si-δ-doping was used in both InAlAs barriers, with a spacer thickness of 5 nm and a donor concentration of 5 x 10^{12} and 1.5 x 10^{13} cm^{-2} in the upper and the lower barrier.

![Layer structure](image)

Fig. 1. (a) Layer structure of the conventional double-sided δ-doped HEMT. Sample 1 was grown without InAs for comparison. In sample 2, InAs with 4 nm was symmetrically centered in the InGaAs channel. (b) Modified layer structure for HEMT application: increasing δ-doping concentrations and Al-content in the barriers, introduction of an additional Si-δ-doping at 1.5 nm into the top InAlAs.

The position z of the inserted InAs is the distance between the InAlAs spacer layer and the center of the InAs, ranging from 3 to 9 nm. At two different growth temperatures, 470°C and 420°C, the thickness d of the InAs layer was raised from 2 nm up to 5 nm, keeping the total channel thickness constant.

Hall measurements at 300 and 77 K confirmed the enhanced mobility of the InAs inserted channel HEMT. Fig. 2 shows the correlation between the position z of 2 nm InAs, grown at a substrate temperature of 470°C, and the mobility. The optimized mobility increase was obtained for symmetrically centered InAs layers. Solid lines are plotted to guide the eye, with the assumption that mobility is attributed to scattering due to the ionized impurities and interface roughness of both δ-dopings and interfaces of the channel

\[
\frac{1}{\mu} \sim \frac{1}{z} + \frac{1}{t-z}
\]

Compared to the reference sample without InAs, the room temperature mobility is increased from 8500 cm² V^{-1} s^{-1} to 10 500 cm² V^{-1} s^{-1} by inserting only 2 nm InAs. Fig. 3 shows the correlation between InAs thickness d, mobility and substrate temperature, where the position is fixed at the center of the channel. For a substrate temperature of 470°C, the largest mobility is achieved for a thickness of 2 nm. It is well known [2] that the critical thickness increases for lower growth temperatures. Therefore, the substrate temperature for
the growth of the InGaAs/InAs/InGaAs channel is changed to 420°C. Obviously, the critical thickness is increased, thus allowing successful use of 4 nm InAs resulting in very high mobilities of 38 400 cm² V⁻¹ s⁻¹ and 12 500 cm² V⁻¹ s⁻¹ at 77 K and room temperature, respectively. This mobility is 47% higher than that of the conventional heterostructure without InAs. All these structures had a 2DEG carrier density of around 3.5 × 10¹² cm⁻².

3. High electron mobility transistor

Thus, having established that, under our growth conditions, 4 nm thick inserted InAs layers give optimized transport properties, we modified the structure for device applications by (i) increasing the δ-doping concentrations to 5.5 × 10¹² and 2.75 × 10¹² cm⁻², (ii) increasing the Al-content in the barriers to 0.6 and (iii) introducing an additional Si-δ-doping at the interface of the cap and supply layer, Fig. 1b. This resulted in (i) 2DEG-densities of 5.6 × 10¹² cm⁻², (ii) improved carrier confinement and breakdown voltages and (iii) reduced source and drain resistances.

The control of the different compositions of the strained layers was verified by X-ray diffraction measurements of the HEMT structure, shown in Fig. 4. Simulation of this spectrum using dynamical scattering theory fits the measured curve very well.

The appearance of interference fringes and the exact position of the pseudomorphic layers confirm the high material quality.

Shubnikov-de Haas measurements at 4.2 K show that two subbands are occupied with a carrier density of 4.0 × 10¹² cm⁻² in the first and 1.6 × 10¹² cm⁻² in the second subband. Even with this high carrier density, the mobility at 4.2 K is 48 000 cm² V⁻¹ s⁻¹.

HEMTs were fabricated from the conventional heterostructure without InAs, the conventional heterostructure with 4 nm InAs and the modified structure with 4 nm InAs inserted. After wet chemical mesa etching for device isolation, ohmic contacts were formed by evaporation of Ge/Ni/Au/Ge/Ni/Au and annealed at 320°C. Using electron-beam lithography, 0.15 μm T-shaped gates were defined in two layer resist. Gate recess was performed in two steps: first by selective sucinic acid etch for the lateral recess definition, second by a nonselective phosphoric acid etch to remove the top Si-δ-doping. The conventional heterostructures are processed in the same way for comparison.

Fig. 5 shows the drain I–V characteristics of the modified structure with an output conductance of 160 mS/mm (at V_DS = 1 V) and a maximum drain current of 1250 mA/mm. Pinch-off behavior is excellent. However, the drain-source breakdown voltage at high current is rather low (1.5 V). Higher breakdown voltages can be achieved with lower doping levels. This device has a very high current
gain cut off frequency $f_T$ of 235 GHz, the maximum oscillatory frequency $f_{\text{max}}$ is measured with 220 GHz, due to the output conductance [1]. The very low source resistance of 0.27 $\Omega$ mm causes an extrinsic transconductance of 1270 mS/mm, shown in Fig. 6. These excellent results are achieved by (i) inserting 4 nm InAs and (ii) introducing an additional Si-$\delta$-doping at the interface of the cap and supply layer. By inserting 4 nm InAs in the conventional heterostructures, the extrinsic transconductance is increased from 735 to 945 mS/mm (Fig. 6), the source resistance is reduced from 0.52 to 0.41 $\Omega$ mm. The higher mobility in the InAs channel reduces the sheet resistance in the access region between the alloyed contact and gate electrode. The additional Si-$\delta$-doping reduces the barrier heights between InGaAs cap layer and InAlAs supply layer, so a tunneling current through the barrier is possible [3] and the source resistance is further reduced to 0.27 $\Omega$ mm. The effective electron velocity, estimated from the intrinsic transconductance with $V_{DS} = 1$ V, of the InAs-channel heterostructure is $2.50 \times 10^7$ cm/s, compared to $1.95 \times 10^7$ cm/s without InAs. The HEMT results are summarized in Table 1.

4. Summary

We have demonstrated that symmetrically centered 4 nm InAs layers in a double-sided $\delta$-doped InAlAs/InGaAs HEMT improve the electron transport properties. The mobility is increased by 47% and the effective electron velocity is increased by 30% at 300 K. By introducing an additional Si-$\delta$-doping at the interface of the cap and supply layer, the source resistance is significantly reduced to 0.27 $\Omega$ mm, resulting in a very high transconductance of 1270 mS/mm and high cut-off frequencies $f_T$ of 235 GHz.

References

Resonant tunnelling of holes in double barrier heterostructures grown by MBE on (1 1 0) oriented GaAs substrates


Abstract

Hole tunnelling in Be-doped GaAs/AlAs double barrier quantum well structures, grown by molecular beam epitaxy on the (1 1 0) GaAs surface, have been investigated. Fewer resonances are observed in the current–voltage characteristic of the device than in the characteristics of similar devices grown on (1 0 0) and (3 1 1)A oriented substrates. High magnetic fields are used to examine the anisotropic in-plane energy dispersion of the QW states of the (1 1 0) diode. The dispersion of one of the resonances is found to be much greater than has been observed in (1 0 0) and (3 1 1)A oriented devices.

1. Introduction

There has recently been renewed interest in quantum well (QW) devices in the GaAs/AlAs system grown on substrate orientations other than (1 0 0). However, the (1 1 0) orientation has received much less attention than other orientations, both experimentally [1] and theoretically [2–7]. This is due in part to the difficulties of epitaxial growth on the amphoteric (1 1 0) GaAs surface. There have been several calculations of the valence band structure in such devices [2–6], and they show the anisotropy of the quasi-two-dimensional subbands to be much larger in (1 1 0) oriented systems than in those with (1 0 0) and (1 1 1) orientations.

To date, there have been no experiments to test these theoretical predictions of the valence band structure in the (1 1 0) plane. In this contribution, resonant tunnelling of holes through a (1 1 0) oriented QW is reported, and magnetotunneling spectroscopy is used to study the valence subband structure. The experimental results are compared with recent theoretical work.

2. Experiment

The structure was grown on a semi-insulating (1 1 0) GaAs substrate misoriented 6° towards the (1 1 1)A face using an Intevac Gen-II molecular beam epitaxy system. The growth was performed at a temperature of 560°C with growth rates for GaAs and AlAs of 1 ML s⁻¹ and 0.5 ML s⁻¹ respectively,
as measured by reflection high-energy electron diffraction. The As/(Ga, Al) beam-equivalent pressure ratio determined by an ionization gauge was approximately 12. On exactly oriented (1 1 0) surfaces, good electrical and optical properties can only be obtained by using low growth temperature (450–500°C), doubling the arsenic overpressure, and halving the gallium flux to 0.5 ML s⁻¹. A misoriented substrate was selected because good surface morphology could be achieved [8] and because the misorientation allows the use of a much higher growth temperature, and standard arsenic overpressure and gallium growth rate, without compromising the quality of the resonant tunnelling devices. Previous devices on (1 0 0) and (3 1 1)A oriented substrates were grown at 630°C [9].

The layer structure of the device is shown in Fig. 1. It consists of a 4.2 nm QW between 5.1 nm AlAs barriers. The upper and lower contact layers are Be-doped and are separated from the barriers by spacer layers. The QW, tunnel barriers and spacer layers are not intentionally doped. The material was processed into circular mesas of 200 μm diameter and contacts formed, using standard wet-etching, photolithography and metallization techniques. Ohmic contacts were made to the top p⁺-layer and the p⁺-layer immediately above the semi-insulating substrate.

![Layer structure of the device considered in the text. The resonant tunnelling structure is formed by the not-intentionally-doped (NID) layers. The substrate is misoriented from the (1 1 0) plane by 6° towards the (1 1 1)A plane.](image)

Fig. 2 shows the I(V) characteristics, at 4.2 K, in various magnetic fields B between 0 and 38 T. Measurements were made with the magnetic field applied parallel to the layer planes along the [1 1 0] direction, Fig. 2(a), and along the [0 0 1] direction, Fig. 2(b). In the absence of a magnetic field, only one resonance is directly visible in the I(V) characteristic, at 858 mV. In the presence of a magnetic field, this resonance moves to higher bias and weakens markedly; it is only visible as a weak feature in the differential conductance, dI/dV, for fields above 30 T. In zero magnetic field, there is also a resonant feature at 122 mV, but is only visible as a weak feature in dI/dV. This resonance also moves to higher bias with increasing magnetic field, but increases in strength, so that it is just discernible in the I(V) characteristic at around 300 mV, for B > 22 T when B is parallel to [1 1 0], and for B > 31 T when B is parallel to [0 0 1].

The dependence of the amplitudes of the resonances with magnetic field has been studied previously and has been shown to be different for different peaks [10, 11]. From Fig. 2 it can be seen that the increase in amplitude of the lower bias resonance with magnetic field is particularly marked for B parallel to [1 1 0]. When B is applied parallel to the [0 0 1] direction, the amplitude also increases, but less strongly. It can also be seen that this resonance becomes much broader with increasing magnetic field for B parallel to [0 0 1] than for B parallel to [1 1 0]. For the higher bias resonance, in both magnetic field orientations, the amplitude decreases rapidly with magnetic field. These effects are typical of the quantum mechanical admixing of light and heavy hole states, and of the complexities of hole resonant tunnelling in high magnetic fields [10, 12, 13].

The complicated dispersion curves of energy versus in-plane wave vector, α(k||), of the valence band QW have recently been investigated on similar structures grown on non-(1 0 0) GaAs planes using magnetotunnelling spectroscopy. The correspondence between magnetic field B and k||, and that between applied bias V and the energy ε at which holes are injected into the quantum well can be used to examine the in-plane energy wave vector dispersion curves, α(k||), for the QW subbands corresponding to the observed resonances [11]. It has
been demonstrated theoretically that for a device with quantum well width $w = 4.2$ nm, magnetic fields $B < 40$ T produce only a small variation from the $\delta k_\parallel$ dispersion when $B = 0$ [12]. Thus plots of resonant peak voltage position, $V_p$, as a function of magnetic field $B$, such as those shown in Fig. 3, should qualitatively represent the $\delta k_\parallel$ curves for these two subbands [10, 12].

The observed anisotropy of the resonances, shown in Fig. 3, is small at low values of $B$ and is
only apparent for $B > 15$ T. For some quantum well widths, the anisotropy is expected to be apparent even at very low magnetic fields, for example, for $w < 3.0$ nm [2] and for $w = 10$ nm [6]. However, this anisotropy is sensitive to the device parameters and may not be so significant for intermediate well widths [4].

For both resonances, the anisotropy indicated by the magnetotunnelling measurements presented here is in the reverse sense to that calculated for $w < 3.0$ nm [2], but in the same sense as calculated for $w \geq 5$ nm [3–6]. From the $V_p(B)$ curves, it can be seen that both the subbands in this device are higher in hole energy for $k_\parallel$ along $[\bar{1} 1 0]$ than for $k_\parallel$ along $[0 0 1]$.

One of the most noticeable features of Fig. 3, in comparison with previously studied p-type devices on the GaAs/AlAs system, is the very large dispersion of the higher bias resonance. This resonance shifts to higher bias by almost 1.2 V for $B$ parallel to $[1 \bar{1} 0]$, corresponding to $k_\parallel$ along $[0 0 1]$, and by more than 1.6 V for $B$ parallel to $[0 0 1]$, corresponding to $k_\parallel$ along $[\bar{1} 1 0]$. Theoretically, the subband with the greatest dispersion is expected to be LH1 [4], suggesting that the higher bias resonance corresponds to tunnelling into LH1. However, the observed dispersion is anomalously large, even for LH1.

For this well width, the HH2 and LH1 resonances are expected to be very close at $k_\parallel = 0$. The first resonance in the $I(V)$ characteristic, which is at a much lower bias than the LH1 resonance, is therefore attributed to tunnelling into the HH1 subband of the quantum well. As is the case here, the HH1 resonance is often very weak in the $I(V)$ characteristics of p-type double barrier diodes. Coupling of the quantum well HH1 subband to the light-hole states of the tunnel barriers is known to be poor [14]. Also, at low biases, there is insufficient charge accumulation in the emitter subband, which is mainly for HH1 character, for states with larger $k_\parallel$, and therefore greater light-hole character, to be occupied. This also leads to poor coupling with the light-hole states of the emitter tunnel barrier. Most of the tunnel current passes through light-hole states in the barrier [15], so the combination of these effects gives a very low resonant current density.

The reasons for the absence of a peak corresponding to tunnelling into the HH2 subband, and for the very large dispersion of the higher bias resonance, are unclear. The observation of only two resonances contrasts with the much larger number seen in devices grown on $[1 0 0]$ and $[3 1 1]A$ oriented substrates which were otherwise identical to the one considered here [9]. In those devices, six and seven resonances, respectively, were observed in the $I(V)$ characteristics [10]. Theoretically, a $(1 1 0)$ oriented quantum well is expected to have more bound states, because the heavy-hole quantization effective mass is large [4], see Table 1. As each resonance in the $I(V)$ characteristic corresponds to tunnelling from an emitter accumulation layer into a subband of the quantum well, a larger number of resonances would be expected. In contrast, the light-hole quantization effective mass is
Table 1
Quantization effective masses of holes in GaAs

<table>
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<th>&lt;1 0 0&gt;</th>
<th>&lt;3 1 1&gt;</th>
<th>&lt;1 1 0&gt;</th>
<th>&lt;1 1 1&gt;</th>
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<td>(m^* (m_e))</td>
<td>0.38</td>
<td>0.55</td>
<td>0.71</td>
<td>0.95</td>
</tr>
<tr>
<td>(m^* (m_h))</td>
<td>0.09</td>
<td>0.08</td>
<td>0.08</td>
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almost isotropic in GaAs. Thus, in principle, the first light-hole resonance should be at similar bias in devices that have different orientation but are otherwise identical. However, differences will occur due to unavoidable variations during device fabrication, such as in dopant density and ohmic contact resistance. The LH1 resonances in the devices presented in Ref. [9] are at 318 mV in the <1 0 0> oriented device and at 310 mV in the <3 1 1>A oriented device. In zero magnetic field, the LH1 resonance of the <1 1 0> device is at 858 mV. This suggests either that the residence of the contact regions is higher in this <1 1 0> device, or that there is greater charge accumulation in the QW of the device. Both of these effects increase the scaling between \(V\) and \(\varepsilon\), and may explain why tunnelling into the HH3 subband is also not observed: the increased scaling between \(V\) and \(\varepsilon\) pushes the resonance beyond the available bias range. Note that in p-type resonant tunnelling devices, charge build-up in the quantum well does not necessarily produce resonances with sharp cut-offs [16], in contrast to the effect in n-type devices. The very flat dispersion expected for the HH2 and HH3 subbands [4] would produce very broad weak resonances [8] and, combined with the poor coupling of heavy hole states in the quantum well to light hole states in the barrier, may be another reason why tunnelling into these subbands is not observed.

3. Summary

Resonant tunnelling of holes into the first heavy hole and light hole subbands of a <1 1 0> oriented quantum well has been observed. The current-voltage characteristics show fewer resonances than are expected from calculations of the valence band states in quantum wells of this orientation. The valence subbands of the quantum well appear to be less anisotropic than expected from some previous theoretical studies.

Acknowledgements

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References

GaAs/AlAs resonant tunneling diodes with super-flat interfaces grown on (4 1 1)A GaAs substrates by MBE

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Abstract

GaAs/AlAs resonant tunneling diodes (RTDs) with atomically flat GaAs/AlAs interfaces over an entire device area (super-flat interfaces) have been successfully fabricated on a (4 1 1)A GaAs substrate by molecular beam epitaxy (MBE). The (4 1 1)A GaAs/AlAs RTDs exhibited a superior peak-to-valley current ratio ($J_p/J_v = 11.8$ at $80 \, \text{K}$) to that ($7.5$) of RTDs simultaneously grown on a conventional (1 0 0)GaAs substrate. The improved $J_p/J_v$ ratio of the (4 1 1)A RTD is believed to be due to the super-flat GaAs/AlAs interfaces, which significantly reduce the valley current density.

Keywords: (4 1 1)A GaAs substrates; Super-flat interface; GaAs/AlAs RTD

1. Introduction

Formation of atomically flat GaAs/AlGaAs interfaces over a macroscopic area is very important for applications to quantum devices such as GaAs/AlGaAs resonant-tunneling hot-electron transistors (RHETs). GaAs/AlGaAs quantum wells (QWs) grown on (4 1 1)A-oriented GaAs substrates by molecular beam epitaxy (MBE) have been reported to show extremely flat interfaces over a large area (“super-flat interfaces”) [1–4]. GaAs/Al$_{0.3}$Ga$_{0.7}$As QWs with well width of 2.4 nm grown on the (4 1 1)A substrate show a very narrow photoluminescence (PL) peak (FWHM = 5.5 meV at $\lambda = 709.2 \, \text{nm}$) at 4.2 K [1] whose line width is the same as that for growth-interrupted QWs grown on (1 0 0) GaAs substrates [5]. There always exist two or three PL peaks from a single QW on a conventional (1 0 0) substrate due to the lateral variation of well thickness by $\pm 1$ ML [5, 6], while the GaAs/Al$_{0.3}$Ga$_{0.7}$As QWs grown on (4 1 1)A substrates exhibit only one sharp PL peak from each QW over an extraordinarily large area ($\sim 1 \, \text{cm}^2$), indicating the formation of effectively atomically flat (4 1 1)A GaAs/AlGaAs interfaces over a wafer-size area [2]. These super-flat interfaces
have been applied to GaAs/Al_{0.3}Ga_{0.7}As resonant tunneling diodes (RTDs) which exhibited a better peak-to-valley current ratio ($J_p/J_v = 5.0$ at 77 K) than that (4.1) of a RTD simultaneously grown on (1 0 0) GaAs substrate, but both devices yielded figures of merit inferior to state-of-the-art RTDs because of their thick barrier layers (10 nm) and low barrier height (Al_{0.3}Ga_{0.7}As barriers) [7]. In this paper, we report the successful fabrication of GaAs/AlAs RTDs on (4 1 1)A GaAs substrates by MBE with much improved $I$–$V$ characteristics due to super-flat interfaces and an improved RTD structure.

2. Experimental procedure

$n^+$-GaAs substrates with (4 1 1)A and (1 0 0) orientations were degreased and chemically etched by sulfuric acid etchant solution ($H_2SO_4 : H_2O : H_2O = 5 : 1 : 1$) prior to MBE growth. GaAs/AlAs double-barrier resonant tunneling (DBRT) structures were simultaneously grown on these substrates in a Nissin RB-2001G MBE system. The substrate temperature was monitored by a pyrometer, which was calibrated by the melting point of aluminum (660°C). The substrates were rotated at 30 rpm during MBE growth. A 400 nm-thick highly Si-doped n-GaAs layer (1.0 x 10^{18} cm^{-3}) and a 100 nm-thick n-GaAs layer (n = 5 x 10^{17} cm^{-3}) were grown at 580°C and V/III ratio of 14 (in pressure) to realize n-GaAs on the (4 1 1)A GaAs substrate which exhibits a low compensation ratio similar to a Si-doped GaAs grown on a (1 0 0) substrate [8]. An undoped AlAs ($L_A = 3$ nm)/GaAs ($L_v = 7$ nm)/AlAs ($L_A = 3$ nm) DBRT structure with 10 nm-thick undoped GaAs spacer layers was grown at 640°C and V/III ratio of 14 in order to form the super-flat (4 1 1)A GaAs/AlAs interfaces [4]. Then 100 nm-thick n-GaAs (n = 5 x 10^{17} cm^{-3}) and 500 nm-thick n-GaAs (n = 1.0 x 10^{18} cm^{-3}) were grown at 580°C and V/III ratio of 14. Finally, a heavily Si-doped n-In_xGa_{1-x}As (x = 0-0.5, n = 3.0 x 10^{19} cm^{-3}) contact layer was grown at 520°C to form the non-alloy ohmic contact with AuGe : Ni. Growth rates of GaAs and AlAs were 1.0 μm/h. Mesa structures for RTDs were made by photolithography and wet etching. A SiO_2 film was used for electric isolation and passivation. After opening windows in the SiO_2 film on the top of the mesa structure, nonalloy ohmic contacts of AuGe : Ni were made there.

3. Results and discussion

Fig. 1 shows $I$–$V$ characteristics (80 K) of the GaAs/AlAs RTDs with a contact area of 120 μm diameter fabricated on the (4 1 1)A (a) and the (1 0 0) GaAs substrates (b). The (4 1 1)A RTD shows a peak current density of 200.2 A cm^{-2} (149.6 A cm^{-2}) at a forward (backward) bias of 0.46 V (–0.36 V), while the (1 0 0) RTD shows the peak current density of 355.7 A cm^{-2} (356.6 A cm^{-2}) at 0.39 V (–0.39 V). The valley current density of the (4 1 1)A RTD is as small as 17.0 A cm^{-2} (13.6 A cm^{-2}) in the forward (backward) bias which is 36% (28%) of that [47.5 A cm^{-2}, (48.3 A cm^{-2})] of the (1 0 0) RTD. Then, the peak-to-valley current ratio ($J_p/J_v$) becomes 11.8 (11.0) in the forward (backward) bias for the (4 1 1)A RTD which is 57% (49%) larger than that [75 (74)] of the (1 0 0) RTD. These improved behaviors of peak and valley current densities of (4 1 1)A RTDs with respect to the (1 0 0) RTDs are similar to the case of GaAs/Al_{0.3}Ga_{0.7}As RTDs previously reported [7], but higher peak current density ($J_p$) and an extremely high peak-to-valley current ratio ($J_p/J_v$) were achieved for the present (4 1 1)A GaAs/AlAs RTD due to its thin (3 nm) and high (AlAs) barriers. The peak current density (the peak-to-valley current ratio) of the GaAs/AlAs RTD became 5 times (2.4 times) as large as that of the previous GaAs/Al_{0.3}Ga_{0.7}As RTD [7]. The observed peak voltage is 0.46 V for the (4 1 1)A RTD, which is 3.5 times larger than the calculated value (0.13 V) of $2E/e$, where $E$ is the resonant energy in the double-barrier structure at zero bias. This is due to a parasitic resistance at the ohmic contacts in both electrodes.

Fig. 2a shows the peak-to-valley current ratios ($J_p/J_v$) of the (4 1 1)A and (1 0 0) RTDs as a function of temperature. At 300 K, a value of $J_p/J_v$ (1.4) of the (4 1 1)A RTD is almost the same as that of the (1 0 0) RTD, while it increases more than that of
Fig. 1. Current–voltage characteristics of GaAs/AlAs double-barrier resonant-tunneling diodes on the (4 1 1)A GaAs substrate (a) and on the (1 0 0)GaAs substrate (b) with a common contact area of 120 μm diameter.

Fig. 2. Temperature dependence of peak-to-valley current ratios ($J_p/J_v$) (a) and peak current density ($J_p$) and valley current density ($J_v$) (b) of GaAs/AlAs double-barrier diodes on the (4 1 1)A and (1 0 0)GaAs substrates.
the (1 0 0) RTD with decreasing temperature and becomes almost constant (~15) below 50 K. This constant value of $J_p/J_s$ is 67% larger than that (~9) of the (1 0 0) RTD. Numerical values of the peak and valley current densities ($J_p$, $J_s$) and the peak-to-valley current ratio ($J_p/J_s$) at 300, 80 and 14 K are also listed in Table 1. The peak and valley current densities of the (4 1 1)A and (1 0 0) RTDs are plotted as a function of the temperature in Fig. 2b. The peak current density of the (1 0 0) RTD is almost constant over the whole range of temperature from 14 to 300 K, and it is almost 1.5 times larger than that of the (4 1 1)A RTD. Decrease of the barrier thickness results in increase in the resonance width due to the enhanced transmission coefficient in the off-resonant voltage range which results in the increase of the resonant current. Tsuchiya and his co-workers reported that the resonant current increases by 1.7 times while decreasing the barrier thickness by only 1 ML for GaAs/AlAs RTD with the well width of 7 nm [9]. Since the (4 1 1)A and the (1 0 0) RTDs were grown simultaneously, the average thicknesses of the AlAs barrier layers of both RTDs were the same within the error limit of 1%. The electron concentration of both sides of the double-barrier structure should affect the peak current density, but the electron concentration of n-GaAs layers on (4 1 1)A substrate is almost the same as that of the n-GaAs layers grown on (1 0 0) substrate because the optimized growth condition was used for growing n-GaAs layers with a low compensation ratio on (4 1 1)A substrate [8]. Therefore, the large peak current density of the (1 0 0) RTD is mainly due to the tunneling through the locally thinner regions of the AlAs barriers with the lateral variation of the barrier thickness of ±1 ML. The valley current density ($J_v$) of the (4 1 1)A RTD is much smaller than that of the (1 0 0) RTD over the whole range of the temperature from 14 to 300 K and it is almost constant below 50 K due to the extinction of the thermal current over the barriers. The forward valley current density is 14.5 A cm$^{-2}$ for the (4 1 1)A RTD at 14 K, which is as small as 35% of that (41.3 A cm$^{-2}$) of the (1 0 0) RTD. We believe that this drastic decrease of $J_v$ for the (4 1 1)A RTD should be due to its super-flat interfaces. The interface roughness deteriorates the lateral momentum conservation of electron tunneling through the double-barrier structure. The lateral potential fluctuation caused by the interface roughness scatters electrons to other states with different lateral and longitudinal momentums. Hence, the interface roughness leads to the increase of off-resonant transmissivity, i.e., increase of $J_v$.

Fig. 3 shows $J_p/J_s$ observed at 77 K as a function of $J_p$ for GaAs/AlGaAs RTDs. An open square and an open circle show $J_p/J_s$ values obtained for the (4 1 1)A and the (1 0 0) RTDs in this study, respectively. $J_p/J_s$ values reported so far for GaAs/AlGaAs RTDs grown on the conventional (1 0 0) GaAs substrates by MBE are also shown (solid squares [9–11], solid triangles [12], solid circles

<table>
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<th>Temperature (K)</th>
<th>(4 1 1)A RTD</th>
<th>(1 0 0) RTD</th>
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<tr>
<td></td>
<td>$J_p$ (A cm$^{-2}$)</td>
<td>$J_s$ (A cm$^{-2}$)</td>
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<td>300</td>
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</table>
4. Conclusions

GaAs/AlAs double-barrier resonant-tunneling diodes (DBRTDs) with 3 nm-thick barrier layers and a 7 nm-thick well layer have been simultaneously grown on the (4 1 1)A and (1 0 0) GaAs substrates by MBE. Much improved peak-to-valley current ratio \( J_p/J_v \) was obtained for the (4 1 1)A GaAs/AlAs RTD. This \( J_p/J_v \) value achieved for the (4 1 1)A RTD is not only much higher than that of the RTD simultaneously grown on (1 0 0) substrate, but also one of the best data reported for GaAs/AlGaAs RTDs. This result is believed to be due to the super-flat interfaces realized in the (4 1 1)A RTD.

Acknowledgements

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References

Photoluminescence characterization of MBE grown AlGaAs/InGaAs/GaAs pseudomorphic HEMTs

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Abstract

We compare the experimental and theoretical photoluminescence (PL) spectra of power and low-noise pseudomorphic AlGaAs/InGaAs high electron mobility transistor (HEMT) profiles. The modeling approach uses a self-consistent solution between Schrödinger equation and Poisson’s equation to calculate the electron and hole energy levels, envelope functions, and concentrations. Results are in good agreement with 4.2 K photoluminescence measurements. PL spectra of the HEMTs show two peaks due to transitions between the e1–hh1 and e2–hh1 channel states. For the low-noise profile, the intensity of the e2–hh1 transition is greater than the e1–hh1 transition intensity while for the power profile, the e1–hh1 transition intensity is greater. The difference in intensities between the low-noise and power profiles is due to differences in the doping of the profiles.

PACS: 85.30.Dc; 78.55.Cr; 71.25.Tn

Keywords: Photoluminescence; Pseudomorphic; HEMT; InGaAs; pHEMT; MBE

1. Introduction

The need for AlGaAs/InGaAs/GaAs high electron mobility transistor (HEMT) devices for low noise and power microwave circuits continues to increase. In order to reduce costs and ensure a stable supply of epitaxial material, the HEMT device structure must be monitored and screened as close to the growth process as possible. Low temperature photoluminescence (PL) provides a very useful technique to monitor the InGaAs channel material properties, providing the relation between the spectral features and the InGaAs channel material characteristics is understood. In this study we have correlated the 4.2 K PL spectral features from the InGaAs channel region of both a low-noise and a power HEMT profile with their material profile characteristics. The low-noise profile has a single AlGaAs–InGaAs interface whereas the power profile has AlGaAs on both sides of the channel providing two AlGaAs–InGaAs interfaces. For convenience, we will call the low-noise profile a single heterostructure profile and the power

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profile a double heterostructure profile. Correlating the PL spectral features with channel material characteristics allows us to use PL for monitoring the MBE grown HEMT material to provide consistent material for low-noise and power applications.

Much work has been done on using PL to extract data from HEMT structures. In particular, estimates of the channel interface roughness can be calculated from the full width at half maximum (FWHM) of the 4.2 K PL spectra [1–3] while at higher temperatures the FWHM can give a measure of the channel sheet charge concentration [2, 4, 5]. Also, at 4.2 K, the PL spectral features are very sensitive to channel composition as well as channel width. At higher temperatures, thermal broadening reduces the PL sensitivity to channel conditions. Since the channel conditions are extremely critical to device performance, 4.2 K PL measurements are necessary to ensure material quality. However, 4.2 K PL can only be measured on witness wafer, not actual device wafers. This is not a problem since one witness sample per day per MBE machine is sufficient for us to monitor the material quality once the growth process has stabilized. If the MBE machine has not stabilized then only witness samples are grown. In this paper we compare the theoretical and measured PL spectra of both the single heterostructure and double heterostructure profiles. The differences between the spectra of the two profiles will be identified and explained. We show that the origins of the PL spectral features are the same for both profiles but that differences in the profiles lead to shifts in the energies and intensities of the spectral features.

2. Material profiles and theory

The structures used in this study were grown at TRW by molecular beam epitaxy using Intevac GenII Modular equipment on 3-in (1 0 0) undoped GaAs substrates. The material profiles are shown in Fig. 1. Sheet doping for the double heterostructure profile is \( N_c = 3.5 \times 10^{12} \) cm\(^{-2}\) at 300 K and \( N_v = 3.3 \times 10^{12} \) cm\(^{-2}\) at 77 K as derived from Hall measurements. For the single heterostructure, the sheet doping is \( N_c = 2.5 \times 10^{12} \) cm\(^{-2}\) at 300 K and \( N_v = 2.3 \times 10^{12} \) cm\(^{-2}\) at 77 K. PL measurements were made using the 514.2 nm line from an Ar\(^+\) laser at an incident power density of \( \approx 50 \) mW/cm\(^2\), a 1-m computer controlled SPEX monochromator, and a Ge detector cooled to 77 K. A lock-in technique was used to reduce noise.

A self-consistent numerical solution of Schrödinger’s equation and Poisson’s equation was used to solve for the electron and hole energy levels, envelope function, and densities [6]. The solution is based on a finite element analysis that assumes a minimum energy is defined along the \( z\)-direction for both the heavy-hole and light-hole valence bands and the conduction band. Spatial variation of the effective mass and dielectric constant are incorporated in the finite element matrix. Excitonic recombination was not considered due to screening and band filling effects. This assumption is confirmed by a lack of exciton luminescence in both the single and double heterostructure spectra. A quasi-static assumption between e–h pair generation and recombination was used to calculate the quasi-Fermi levels and occupational probability. Exchange terms and excitonic effects were neglected due to the high channel carrier concentration [7].

Strain effects were accounted for by appropriately modifying the InGaAs band energy using \( c_{11} = 11.24 \) dyne/cm and \( c_{12} = 5.196 \) dyne/cm [8]. The conduction band energy discontinuity is given by \( \Delta E_C = 0.70 \Delta E_G \). Table 1 lists the material
Table 1
Material constants used to calculate the quantum well energy levels and wavefunctions

<table>
<thead>
<tr>
<th>Material</th>
<th>$E_G$</th>
<th>$m^*_e$</th>
<th>$m^*_h$</th>
<th>$m^*_s$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Al$<em>{0.25}$Ga$</em>{0.75}$As</td>
<td>1.697</td>
<td>0.0763</td>
<td>0.562</td>
<td>0.1010</td>
</tr>
<tr>
<td>Al$<em>{0.25}$Ga$</em>{0.75}$As</td>
<td>1.734</td>
<td>0.0778</td>
<td>0.569</td>
<td>0.1030</td>
</tr>
<tr>
<td>GaAs</td>
<td>1.422</td>
<td>0.0670</td>
<td>0.520</td>
<td>0.0870</td>
</tr>
<tr>
<td>In$<em>{0.21}$Ga$</em>{0.79}$As</td>
<td>1.256</td>
<td>0.0583</td>
<td>0.573</td>
<td>0.0747</td>
</tr>
</tbody>
</table>

*This value includes a 44 meV increase in bandgap energy due to strain energy.

The parameters used for each layer. We assume midgap Fermi level pinning at the free GaAs surface [9, 10]. Doping segregation effects are modeled by using 20 Å uniformly doped regions with the same total dose [10, 11]. Energy state broadening, an excess e–h pair distribution, and impurity scattering were used to calculate the spectral broadening.

3. Spectral analysis

Fig. 2 shows the energies and wavefunctions for the $n = 1$ and 2 electron states ($e_1$, $e_2$) as well as the $n = 1$ heavy hole state (hh1) for the single heterostructure profile. Fig. 3 shows the energies for the $n = 1$, 2, and 3 electron states ($e_1$, $e_2$, $e_3$) and the wavefunctions for the $n = 1$ and 2 electron states as well as the $n = 1$ heavy hole state (hh1) for the double heterostructure profile. The $e_3$ state is not truly a channel state since the wavefunction is localized in the notch formed by the AlGaAs/GaAs buffer interface. The $e_3$ energy level is shown only for reference since the Fermi energy lies just below this level. Only the $n = 1$ heavy hole state is considered because the excess holes would thermalize to the top of valence band prior to recombining.

Fig. 4 shows representative measured and calculated PL spectra for both profiles. The solid lines show the measured spectra while the dashed lines show the calculated spectra. Both measured and calculated spectra are normalized to the peak amplitude. Good agreement is obtained between the measured and calculated spectra. Two peaks are observed in both structures. The lower energy peak corresponds to the $e_1$–hh1 transition while the higher energy peak corresponds to the $e_2$–hh1 transition. The $e_2$–hh1 peak broadening is less than the $E_F - E_2$ separation, therefore no Fermi edge singularity is seen [12]. In both profiles, the $e_1$ electron state is strongly localized at the front surface side of the well while the $e_2$ state is more weakly localized at the buffer side of the well. In the single heterostructure profile, the ionized donors on the surface side of the channel attract the electrons and repel the holes which reduces the intensity of the $e_1$–hh1 transition relative to the intensity of the $e_2$–hh1 transition. In the double heterostructure profile, the ionized donors on the buffer side of the channel increase the localization of the $e_2$ state to the buffer side of the well, decrease the localization of the $e_1$ state at the front surface side of the well, and repel the holes from buffer side of the well. The effect on the double heterostructure PL spectra
Fig. 3. Conduction band minima, valence band maxima, electron energy states (top) and the wavefunctions for each energy state (bottom) plotted as a function of position for the double heterostructure device. The dashed vertical lines in the wavefunction plots denote the well boundaries. In this structure, the holes are localized near the center of the well which results in a stronger overlap between the e1 and hh1 wavefunctions than between the e2 and hh1 wavefunctions.

is to decrease the e2–hh1 transition intensity and increase the e1–hh1 transition intensity relative to the single heterostructure profile. The additional donors in the double heterostructure also reduce the energy of the e2 state and increase the energy of the hh1 state. This has the net affect of increasing the e1–hh1 transition energy and decreasing the e2–hh1 transition energy.

Broadening of the PL lineshapes is very complicated but can be generally accounted for by considering the natural spread in the energy states, thermal broadening of the hole distribution, impurity-assisted broadening, and interface roughness. At low excitation power densities and at low temperatures, the spread in energy states and thermal broadening of the holes is small which confines the holes to a very narrow range of momentum (\(\hbar k\)) values near the bottom of the valence band. Therefore, for direct \(k\) conserving transitions, the recombining electrons can only come from a narrow range of energies near the bottom of the electron band, emitting photons over a narrow range of energies near the minimum of the electron-to-hole state energies.

Impurity and interface roughness broadening are very similar processes. In these processes, the \(k\) conservation is broken by allowing electrons to scatter into intermediate, virtual, states either directly before or after photon emission. High-energy electrons with energies up to the Fermi energy can recombine with holes, emitting a significant number of photons above the minimum electron-to-hole state energies. The transition probability of these higher energy electrons decreases as electron energy increases due to the energy term in the denominator of the transition probability matrices. However, because the density of states increases with energy, the net affect of impurity and interface roughness scattering is to shift the emitted photon distribution toward higher energies [13]. This is seen in the PL spectra as a broadening of the high energy side of the spectral features and an increase in the energy of the peak PL intensities.
In the single heterostructure profile, the PL spectra FWHM is 29.0 meV for the e1–hh1 transition and 27.4 meV for the e2–hh1 transition. The e1–hh1 transition is expected to have a greater FWHM than the e2–hh1 transition because impurity and interface roughness have a greater affect on the e1–hh1 transition than the e2–hh1 transition due to localization of the e1 state at the front AlGaAs–InGaAs interface near the front doping layer. This is also the reason for the high-energy tail in the PL spectra of the e1–hh1 transition. In the double heterostructure profile, the PL spectra FWHM is 33.5 meV for the e1–hh1 transition and 33.0 meV for the e2–hh1 transition. The broadening of the e1–hh1 and e2–hh1 transitions are expected to be similar in the double heterostructure profile since both the e1 and e2 states are localized at an AlGaAs–InGaAs interface near a doping plane.

4. Summary and conclusions

In conclusion, the PL spectra of double and single heterostructure AlGaAs/InGaAs/GaAs HEMT profiles were measured at 4.2 K and compared with the PL spectra calculated using a self-consistent solution of Poisson’s and Schrödinger’s equations. Two peaks are identified corresponding to transitions between the e1 and e2 conduction band states and the hh1 valence band state confined in the well. In both profiles, the e1 electron state is confined at the front surface side of the well while the e2 state is confined at the buffer side of the well. For the single heterostructure profile, the holes are localized near the buffer side of the well, resulting in a stronger overlap integral between the e2 and hh1 states than between the e1 and hh1 states. However, for the double heterostructure profile, the holes are localized closer to the center of the well due to repulsion by the ionized donors in the buffer. This results in a stronger overlap between the e1 and hh1 states than between the e2 and hh1 states. This shift in the relative intensities is seen in the measured and calculated PL spectra. Now that we understand the origin of the spectral features, we monitor the transition energies and intensities to ensure the uniformity of MBE grown HEMT wafers delivered for processing.

References

Multispectral InGaAs/GaAs/AlGaAs laser arrays by MBE growth on patterned substrates

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Abstract

Multispectral semiconductor laser arrays on single chip is demonstrated by molecular beam epitaxial (MBE) growth of In$_{0.25}$Ga$_{0.75}$As/GaAs quantum well lasers on GaAs (1 0 0) substrates patterned by dry etching. No regrowth is needed for simple edge emitting lasers. It was observed that the laser characteristics are not degraded by the patterned growth. The shift in the emission wavelength obtained by this method can be controlled by varying the width of the pre-patterned ridges as well as by selecting the regions with different number of vertical sidewalls on both sides. We have also shown that multispectral vertical cavity surface emitting laser (VCSEL) arrays can be made by this technique with a single regrowth.

PACS: 68.55.Bd

Keywords: Molecular beam epitaxy; Patterned growth; Adatom migration; Reactive ion etching; Laser

1. Introduction

Semiconductor laser arrays with spectrally separated emission wavelengths in close spatial proximity are of interest in applications such as wavelength division multiplexing and multi-wavelength optical recording. Two different techniques have been tried with molecular beam epitaxy (MBE) to obtain dual and multiple wavelength laser arrays. In the first approach [1], the laser structure is grown in two steps. In the first step, lower clad and multiple quantum well active layer with different composition and well widths are grown. By selectively etching the quantum wells (QW), regions with different emission wavelengths are defined. The upper clad is then regrown to complete the laser structure. Although laser arrays with multiple emission wavelengths can be obtained by a single regrowth step in this technique, it has a major disadvantage that the regrown interface is in the active region. In the second approach [2], impurity induced intermixing from a doped layer adjacent to the quantum well by post-growth annealing is utilized to realize the change in band gap of the quantum well. This has the advantage

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that no regrowth is involved. However, only two different wavelengths can be obtained by this method, one without intermixing and one with intermixing. Moreover, the high-temperature anneal used for intermixing might seriously degrade the laser performance. In this paper we present a new technique wherein multiple wavelength laser arrays with close spatial proximity are obtained in a single step growth on patterned substrates.

When an alloy such as InGaAs is grown on a patterned substrate, lateral composition modulation is obtained due to the large difference in surface migration lengths of In (~ 25 μm) [3] and Ga(1–2 μm) [4]. However, most of the earlier work [3–7] was done on patterned substrates with sloping (n 1) walls, which are hard to control accurately. We have recently shown that when InGaAs is grown on GaAs (1 0 0) substrates patterned with vertical sidewalls as shown in Fig. 1a, there is a blue shift in the band gap of the material grown on top of the patterns and in the grooves, when the dimensions of the patterns are of the order of migration length of In on GaAs [8, 9]. The measured blue shift in the photoluminescence of In0.2Ga0.8As/GaAs quantum wells grown on ridges of different widths is shown in Fig. 1b. This blue shift results from changes in alloy composition and thickness, which in turn are due to the dual effects of growth on the sidewalls, and a large density of kink sites provided by the edges of the pattern to the migrating adatoms. As a result InGaAs with increasing blue shift in the bandgap can be obtained by reducing the pattern dimensions. Moreover, the photoluminescence spectra of quantum wells grown on top of the ridges show linewidths as narrow as those on unpatterned substrates [8]. Therefore, laser structures grown on such patterned substrates are expected to exhibit similar threshold characteristics as those on unpatterned substrates.

Here we present the results of our study on In0.2Ga0.8As/GaAs laser arrays with varying emission wavelengths obtained by MBE growth on GaAs substrates patterned by reactive ion etching (RIE). We also explore the possibility of integrating multispectral vertical cavity surface emitting laser (VCSEL) arrays.

2. Experimental procedure

Pseudomorphic In0.2Ga0.8As/GaAs single quantum well separate confinement heterostructure (SQW-SCH) lasers, as shown in Fig. 2, were grown

![Fig. 1. (a) Schematic illustrating the In adatom migration on a patterned substrate with vertical sidewalls obtained by dry etching (b) dependence of pattern dimensions on the blue shift in the band gap of In0.2Ga0.8As/GaAs quantum well grown on ridges.](image)

![Fig. 2. Structure parameters of the SCH laser grown on GaAs (1 0 0) substrates patterned into 2 μm deep ridges with vertical sidewalls. The layers marked inner clad are the carrier confinement regions.](image)
in a Varian GEN II MBE system. The n'-GaAs (1 0 0) substrates were patterned into 2 μm deep ridges (width 10–60 μm) having vertical sidewalls, obtained by RIE. The growth rate of GaAs was 0.72 μm/h and the V/III flux ratio ~ 5. The GaAs and Al0.3Ga0.7As layers were grown at 620°C and the In0.2Ga0.8As layers were grown at 540°C. Silicon and beryllium were used as n- and p-type dopants, respectively. Single mode ridge waveguide lasers were fabricated with alloyed Pd/Zn/Pd/Au and Ni/Ge/Au/Ti/Au for p-ohmic and n-ohmic contacts, respectively, and with wet and dry etching.

For the demonstration of multispectral VCSELs, the structures were grown in two growth steps. The bottom distributed Bragg reflector (DBR) mirror with 20 periods of GaAs/AlAs was grown in the first step. Small circular patterns with 12–45 μm diameters were masked by photoresist and the samples were etched by RIE to reach the substrate. A wet etch with H2SO4/H2O2/H2O (1 : 8 : 800) was performed before introducing the sample in the MBE system for regrowth. One more period of bottom DBR followed by the active region consisting of single In0.2Ga0.8As QW surrounded by 0.1 μm of GaAs on each side were regrown by MBE. Fig. 3 shows the scanning electronic microscope (SEM) image of such pillars before and after regrowth.

Photoluminescence measurements were made on all the samples at 18 K with 488 nm Ar + laser excitation source, 0.75 m scanning spectrometer, liquid nitrogen cooled photomultiplier, and lock-in amplification. The excitation laser was focused to a spot size (~ 15 μm) of the order of the size of the patterns.

3. Results and discussion

Single mode lasers (3 μm wide) fabricated from the unpatterned region has a threshold current of 12 mA for a cavity length of 500 μm. The L–I characteristics of the lasers from unpatterned and patterned regions of the same laser bar are shown in Fig. 4. There is no significant change in either threshold current or the slope above threshold which indicate that the quality of the material grown on top of the ridges is as good as that on the plane substrates. This corroborates our earlier report that there is no significant difference in the intensity or the line width of the PL emission from the material on unpatterned region and on top of the ridge [8].
The emission spectra obtained from lasers placed at three different regions of the same laser bar are shown in Fig. 5. Laser L0 is located farthest – greater than one migration length of indium adatoms – from the patterns so that there is no contribution from the surface discontinuity at the sidewalls. L1 is close to the patterned ridge and hence has sidewalls on one side. L2 and L3 are from a 20 and 10 µm ridges, respectively, and have sidewalls on both sides. As is evident from Fig. 5b, there is a blue shift in the emission energy of the lasers on top of the patterns compared to laser L0. The blue shift is more for 10 µm ridge compared to 20 µm ridge in agreement with the PL measurements [8]. Also, the amount of blue shift in laser emission wavelength for lasers having 1 sidewall is different from that of 10 and 20 µm ridges. Thus, with a suitable combination of the ridge width and the placing of the lasers, a large number of lasers with closely spaced emission wavelength can be integrated in close spatial proximity. We have observed a maximum shift of 11 nm for a minimum ridge width of 10 µm used in this study.

One of the major benefits of using this technique of patterned growth for obtaining multi-spectral laser arrays is that conventional single mode laser processing is used and no changes in the fabrication steps are necessary. However, its limitation lies in the fact that the maximum wavelength difference that can be obtained is limited. Therefore, we can conclude that this technique is useful in applications where a large number of lasers with relatively small wavelength shift is needed with precise control over wavelength spacing.

Another area where this technique of varying emission wavelengths by pre-patternning the substrates is applicable is in the fabrication of multi-spectral VCSELs. Multispectral VCSELs have been tried earlier by differential heating across the wafer wherein the growth rate of the DBR stack and therefore the reflection maximum is varied. However, this has not proved to be a very controllable process. Here, we present the possibility of obtaining multispectral VCSELs by growth on patterned substrates. This is done in two growth steps. After growing the bottom DBR stack, circular patterns are formed by RIE and then the VCSEL structure is completed by a regrowth step. The problems associated with regrowth can be partially eliminated by growing one period of the bottom DBR before growing the active layers. This will put the regrown interface in the DBR region rather than in the active region. In order to demonstrate the possibility of obtaining multiple emission wavelengths, PL samples were prepared by a regrowth of single quantum well on the DBR pillars as explained in Section 2. The PL spectra obtained from plane region and from the top of the patterns of two different diameters are shown in Fig. 6. There is a shift of 11 and 46 meV in the emission peak from the QW on top of pillars of diameters 35 and 12 µm, respectively. These spectra were taken after etching off the QW from rest of the region.
4. Conclusion

We have demonstrated multispectral laser arrays by MBE growth on non-planar patterned substrates, without using regrowth. The blue shift obtained in the band gap of the material on top of ridges having vertical sidewalls is utilized in integrating lasers with different emission wavelengths. It is found that the characteristics of the lasers on top of the patterns are similar to those on unpatterned substrates indicating that there is no degradation in the behavior of the lasers due to patterned growth. Laser emission spectra show that the blue shift obtained depends on the ridge width of the pre-patterns as well as the number of sidewalls. We have also demonstrated that multispectral VCSEL arrays can possibly be made with a single regrowth step.

Acknowledgements

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References

MBE growth of high-quality InP for GaInAs/InP heterostructures using incongruent evaporation of GaP

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Abstract

The incongruent evaporation of GaP source material using a conventional effusion cell equipped with a PBN scavenger at the orifice is demonstrated as a simple and cost-effective way to generate a pure P_2 molecular beam, suitable for the growth of InP layers in a solid-source MBE system. Low residual carrier concentrations in combination with high mobilities as well as narrow and intense photoluminescence spectra were achieved. While some arsenic incorporation from the growth environment was observed, gallium incorporation is limited to below 0.1% of the group-III lattice sites. n- and p-doping using Si and Be has been investigated. Successful growth of InP/GaInAs heterostructures indicates that the use of the GaP source technique is a viable way for growing AlGaInAs/InP device structures by solid source MBE.

1. Introduction

Long wavelength (1.3–1.6 μm) quantum well lasers based on the (Al,Ga)_{1-x}In_{x}As system have gained increasing attention recently as they promise superior laser performance in terms of high-temperature operation, high output power and speed as well as low chirp [1–3] due to the larger conduction band offset in these alloys. Concerning the cladding layer material, the ternary Al_{0.47}In_{0.53}As would be the consequent choice, however, due to its better thermal and electrical conductivity, InP is clearly preferable.

This contribution specifically reports on the MBE growth of InP employing a conventional ion-pumped growth system. As high precision control of the phosphorus flux, as offered by valved cracker cells [4], is not necessary when only the binary InP is required, the incongruent evaporation of GaP from a conventional effusion cell equipped with a scavenger at the orifice appears to be a very simple and cost-effective way to generate an extremely pure P_2 molecular beam [5]. Using this technique, the growth of high-quality InP in terms of morphology, electrical and optical characteristics will be presented. Contamination of InP by Ga from the GaP source as well as by As from the growth environment has been analysed. Results on the incorporation of the conventional dopants Si and Be will be described. Subsequently, the growth of abrupt GaInAs/InP heterostructures will be demonstrated, proving the suitability of GaP.

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source MBE of InP for InP/(Al)GaInAs device structures.

2. Experimental procedure

The InP layers were grown in a conventional MBE system which is exclusively equipped with an ion pump. Except for arsenic, As, conventional effusion cells were used for the group-III and the dopant elements. For As a valved cracker cell was employed (with only the As₄-mode used) to facilitate the rapid switching of the As₄ beam within GaInAs/InP heterostructures. The GaP source material (polycrystalline 6 N) was also evaporated from a conventional cell. The use of the P₂ beam originating from the GaP did not cause any difficulty with the MBE system. Even during warm up of the growth chamber the pressure rise did not exceed 10⁻⁵ mbar. In addition, opening of the growth chamber did not provoke any firing even without pre-baking. Reduction of the residual Ga beam generated during the GaP evaporation was achieved by the implementation of a multiple diaphragm approach [6]. A custom designed PBN device similar to Ref. [5] was applied. The GaP cell was operated without a shutter, i.e. the P₂ beam was switched on and off simply by variation of the source temperature. Consequently, growth interruptions of up to 5 min with an As₄ stabilising beam impinging on the surface were required at InP/GaInAs and GaInAs/InP interfaces. InP layers were deposited onto epiready (100) InP substrates without applying any pre-treatment. The substrate surface was inspected by RHEED during oxide desorption to calibrate the thermocouple reading of the substrate temperature, Tₑ. Growth was initiated at Tₑ = 465°C, and the sample temperature was subsequently ramped to its final value.

The properties of the MBE-grown InP were assessed with conventional characterisation tools: interference microscopy, 10 K photoluminescence, PL, 300/77 K Hall measurements, double-crystal X-ray diffraction, DCXRD, and secondary-ion mass spectroscopy, SIMS, using an oxygen primary beam.

3. Results

The temperature of the GaP source, Tₑ, to deliver an adequate P₂ beam density was determined from an Arrhenius type plot of the P₂ pressure (flux gauge) versus 1/Tₑ. Variation of the cell temperature between 800°C and 865°C results in an increase of the P₂ beam pressure from 1.0 × 10⁻⁶ mbar to 8.0 × 10⁻⁶ mbar. The chamber pressure increases correspondingly from 0.55 × 10⁻⁷ mbar to 1.8 × 10⁻⁷ mbar.

3.1. Material characteristics

InP layers were deposited at temperatures between 400°C and 500°C. Inspection by interference contrast microscopy confirmed smooth and featureless surfaces at minimum BEP(P₂/In) ratios between 2 and 10, depending on Tₑ. The effectiveness of suppressing the residual Ga concentration, originating from the GaP source, by means of the PBN scavenger in front of the cell orifice was studied by SIMS profiling of ⁷¹Ga. In Fig. 1, the SIMS profile of a 0.8 μm thick InP layer grown at 420°C shows a strong ⁷¹Ga interface peak due to Ga deposition occurring during P₂ stabilisation of the InP substrate during oxide desorption, as well as a low and constant amount of ⁷³Ga incorporated during growth. For the particular V/III ratio applied, a ⁷¹Ga concentration of 2 × 10¹⁹ cm⁻³ was determined, which corresponds to an overall

Fig. 1. SIMS depth profiles of residual ⁷³As and ⁷¹Ga in an MBE InP layer grown from a GaP source.
amount of $^{71}$Ga plus $^{69}$Ga of $5 \times 10^{18}$ cm$^{-3}$. Thus, only a fraction of $5 \times 10^{-4}$ of the overall group-III lattice sites is occupied by gallium which is a factor of two lower than previously reported [5].

The dominant contaminant in the InP layer depicted was found to be arsenic. Its concentration exceeds that of residual gallium by roughly two orders of magnitude, thus amounting to several percent. Apart from the accumulation at the layer/substrate interface, related to the formation of an intermediate In(P, As) layer during the oxide desorption process, As incorporation exponentially decreases during growth, which indicates that it is released from any background source. This is not surprising since the MBE system had been extensively used for the growth of AlGaInAs layers in the past. However, by extensive baking of the MBE system As incorporation was dramatically reduced to a level below 0.1%, which lies in the same range as reported for MBE InP growth with valved cracker cells [7, 8]. The high amount of residual As present in Fig. 1 results in some lattice distortion amounting to values even in excess of 1000 ppm with the occurrence of cross-hatching indicating strain relaxation. Successful reduction of residual As by extensive baking gives rise to an X-ray diffraction pattern composed of only one InP peak (FWHM: 21 arcsec) revealing a lattice distortion below 50 ppm.

High $T_x$ values and correspondingly lower V/III ratios during deposition were found to favour the reduction of the residual carrier concentration. This is demonstrated in Fig. 2, in which 300 and 77 K Hall data are shown in dependence of the V/III ratio for samples grown at $T_x = 420^\circ$C. In agreement with the decrease of residual carrier concentration down to $1.6/0.6 \times 10^{15}$ cm$^{-3}$ (300/77 K), when reducing the V/III ratio from 6 to 3, a monotonous increase of the 77 K mobility from 20,000 to 50,000 cm$^2$/V·s was observed as well. At a lower V/III ratio the 300 K carrier concentration drops below $10^{15}$ cm$^{-3}$, however, a degradation of mobility as well as surface roughness due to group-V deficient growth are observed in that case.

High optical quality of the MBE-grown InP was obtained, as depicted in Fig. 3 which shows 10 K PL spectra of a layer grown at 420$^\circ$C in comparison with state-of-the-art MOVPE-grown InP ($T_x = 650^\circ$C). For both epitaxy techniques, the observed dominance of the excitonic near band edge emission indicates high optical quality. The impurity related transitions between 1.40 and 1.37 eV are close to the detection limit. While the linewidth of the MOVPE excitonic emission is as narrow as 2 meV, the MBE grown InP shows a larger line width of 5 meV due to the higher residual carrier concentration of the particular MBE layer investigated ($5 \times 10^{14}$ cm$^{-3}$ (MOVPE) versus $7 \times 10^{15}$ cm$^{-3}$ (MBE)). Interestingly, despite the strong blue shift of 12 meV of the emission from MBE InP indicating a relatively high degree of As incorporation (roughly 2% in this case), the integral intensity of the MBE InP is fairly comparable to that of the MOVPE-grown material.

![Fig. 2. 300/77 K Hall carrier concentrations of undoped MBE InP in dependence of the P$_2$/In ratio.](image)

![Fig. 3. Comparison of 10 K PL spectra of InP layers grown by MBE using a GaP source and by low-pressure MOVPE.](image)
3.2. Si and Be doping

Studies on n- and p-type doping were performed using the conventional dopants Si and Be and comparing their incorporation behaviour with that in GaInAs. Doping levels between $1 \times 10^{17}$ and $1 \times 10^{19}$ cm$^{-3}$ were investigated. Hall carrier concentrations were found to be identical in InP and GaInAs for n- and p-doping over the whole doping level range. Particularly for Be incorporation, the observed Arrhenius-type $p$ versus $T^{-1}$ behaviour in combination with the Be concentration, as measured by SIMS, indicate that up to $1 \times 10^{19}$ cm$^{-3}$ complete activation occurs. The agreement with the GaInAs data suggests 100% Be incorporation in InP. The electrical quality of the n- and p-doped layers compares well with published mobility values obtained in material grown with a valved crater. At a doping concentration of $1 \times 10^{18}$ cm$^{-3}$, 300 K mobilities of 1750 and 55 cm$^2$/V·s were obtained for Si- and Be-doping, respectively.

Although Be atoms are known to be prone to acceptor diffusion during growth [10], controlled Be incorporation in InP could be achieved at a reduced $T_g$ of 420°C. This is demonstrated in Fig. 4 by comparing SIMS $^6$Be profiles of 1.3 µm thick p-InP homogeneously doped to $1.3 \times 10^{19}$ cm$^{-3}$ (highest Be concentration applied throughout this study) and a p$^+/p$-doped structure ($5 \times 10^{18}$ cm$^{-3}$/2 $\times 10^{17}$ cm$^{-3}$) designed for the p-cladding layer of a laser. Even for the highly doped layer only minor movement of Be was detected. The corresponding SIMS profile (a) shows that homogeneous incorporation occurs apart from some depletion observable near the substrate/layer interface, apparently associated with accumulation at the interface, and a constant low-level incorporation in the substrate. The latter appears to be strongly dependent on doping concentration, as demonstrated by comparison with profile (b) in Fig. 4. Reduction of the doping concentration to $5 \times 10^{18}$ cm$^{-3}$ obviously eliminates Be movement allowing for a very abrupt change in doping concentration to a level of $2 \times 10^{17}$ cm$^{-3}$. In addition, no beryllium could be detected in the substrate in this case, hence ruling out any significant Be movement into an active laser region if implemented in a real laser structure.

3.3. GaInAs/InP heterostructures

To further assess the applicability of InP grown from a GaP source to heterostructure devices, we have deposited InP/GaInAs/InP test layers similar to those implemented in optoelectronic devices. SIMS-profiles, presented in Fig. 5, give an impression of the switching ability from As to P and vice versa, confirming that quite abrupt switching is possible. A P concentration drop of nearly three orders of magnitude to a constant low level can be obtained which is determined by the idling temperature of the GaP source operated without a shutter. The As level can also be abruptly changed at the
InP/GaInAs interface, however, as already pointed out, an exponential decay of the As level in InP is observed, ranging from 0.03 in the lower towards 0.01 in the upper InP layer for the specific sample shown. In addition, no substantial P incorporation in GaInAs is observed.

4. Summary

In summary, it has been demonstrated that incongruent evaporation of GaP from a conventional effusion cell equipped with a multiple diaphragm device is a simple and cost effective way to generate a very pure P$_2$ beam. InP layers of high quality and well-controllable incorporation of dopants, particularly Be, were deposited. Abrupt switching from InP to GaInAs was successfully achieved with no substantial P incorporation in GaInAs. While residual incorporation of Ga in InP was demonstrated to remain at a negligible level below 0.1%, As incorporation from the growth environment is of concern. Nevertheless, levels down to 0.1% are attainable.

Acknowledgements

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References

Gas-source molecular beam epitaxial growth of low threshold current 1.3 \, \mu m \, InAsP/InGaAsP lasers

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Abstract

Highly strained InAsP/InGaAsP MQW lasers emitting at 1.3 \, \mu m grown by gas-source molecular beam epitaxy with threshold currents of 1.1 mA at 20°C are reported. Lasers with anti-reflection/high-reflection coated facets exhibited slope efficiencies as high as 0.48 mW/mA, output powers of 65 mW, and wall plug efficiencies of 30% at 10 mW.

1. Introduction

InP-based semiconductor lasers operating at 1.3 \, \mu m have received much attention due to their increasing role in applications such as fiber-optic communication and optical interconnect systems. These applications require that the semiconductor lasers maintain low threshold currents and high output powers over a wide temperature range. For semiconductor lasers operating at 1.3 \, \mu m, separate confinement heterostructure multiple quantum well (SCH-MQW) lasers using compressively strained InAsP wells with either lattice-matched or strained InGaAsP barriers grown on InP have been demonstrated to have superior high-temperature and output power performance over the commonly used InGaAsP/InGaAsP SCH-MQW lasers [1, 2]. We had recently reported on the optimization of growth conditions for high-quality 1.3 \, \mu m InAsP/InGaAsP SCH-MQW lasers by gas-source molecular beam epitaxy (GSMBE) [3]. Here we report on InAsP/InGaAsP SCH-MQW lasers for 1.3 \, \mu m emission grown by GSMBE with very low threshold currents of 1.1 mA at 20°C and 6.1 mA at 100°C. To our knowledge, these are the lowest threshold currents yet reported for any InGaAsP lasers grown by GSMBE [4–6].

2. Experimental procedure

The InAsP/InGaAsP heterostructures were grown on (100) n-type InP substrates. Conventional effusion cells were used to provide the In and Ga atomic fluxes, with growth rates of 0.6–1.0 \, \mu m/h. AsH₃ and PH₃ were thermally cracked at 800°C to obtain the As₂ and P₂ molecular beams, with flow rates of 2–10 sccm.

A schematic of the epitaxial layer structure used for the InAsP/InGaAsP SCH-MQW laser is
shown in Fig. 1. The laser structure consisted of a 1 μm n-type InP cladding layer (1 × 10^{18} \text{ cm}^{-3}), the active region, a 1.5 μm p-type InP cladding layer (7 × 10^{17} \text{ cm}^{-3}), and finally a 0.2 μm p-type InGaAs contact layer (2 × 10^{19} \text{ cm}^{-3}). The active region consisted of six wells of InAs_{0.47}P_{0.53} 4.5 nm thick, with 15 nm InGaAsP barriers with band gap emission of 1.15 μm (denoted as 1.15Q), and 110 nm InGaAsP (1.15Q) separate confinement layers on either side of the MQW region. The InP cladding layers were grown at 460–480°C, while the active region was grown at 430°C, since substrate temperatures between 410 and 430°C were previously determined to be optimum for GSMBE growth of the InAsP/InGaAsP MQWs [3]. Buried heterostructure (BH) lasers were fabricated from the GSMBE wafers with an InP p-n-i-n current blocking structure using MOCVD [7]. For high-reflection coatings alternating layers of SiO_{2} and Si were evaporated on the laser facets; for anti-reflection coatings a single layer of SiO_{2} was used.

3. Results

The light output versus drive current (L–I) characteristics of BH lasers with a cavity length \( L \) of 125 μm and different facet reflectivities \( R_1 \) and \( R_2 \) are shown in Fig. 2. For lasers with uncoated facets (i.e., \( R_1/R_2 = 29%/29\% \)), a minimum threshold current of 6.5 mA was measured. With a high-reflection coating on the rear facet (\( R_2 = 95\% \)), the threshold current reduced to 3.0 mA. A low thres-

![Fig. 2. Light output versus drive current characteristics for 125 μm long buried heterostructure lasers with different facet reflectivities (\( R_1 \) and \( R_2 \)). \( R_1 \) is the front facet reflectivity and \( R_2 \) is the reflectivity of the rear facet.](image)

shold current of 1.1 mA was measured for the 125 μm long lasers with high-reflection coatings on both facets (\( R_1/R_2 = 85%/95\% \)).

The \( L–I \) characteristics as a function of temperature for a BH laser with a cavity length of 125 μm and \( R_1/R_2 = 85%/95\% \) was measured. The threshold current increased from 1.1 mA at 20°C to only 6.1 mA at 100°C.

The output power of the lasers was increased by using anti-reflection coatings for the front facet (\( R_1 = 10\% \)) and high-reflection coating for the rear facet (\( R_2 = 95\% \)). The \( L–I \) characteristics of a 250 μm long and a 750 μm long laser with anti-reflection/high-reflection coated facets are shown in Fig. 3. A maximum output power of 49 and 65 mW were measured for the 250 μm and 750 μm long lasers, respectively, at 20°C. Very high slope efficiencies of 0.48 mW/mA and wall plug efficiencies of 30% at 10mW were observed for devices of 250 μm in length at 20°C.

The maximum output power as a function of device temperature is shown in Fig. 4. The maximum output power dropped from 49 mW at 20°C to 4 mW at 90°C for the 250 μm long laser and from 65 mW at 20°C to 15 mW at 90°C for the 750 μm long laser. Even at these high operating powers and elevated temperatures no catastrophic failure of the lasers were observed in our preliminary evaluation.
Fig. 3. Light output versus drive current characteristics for a 250 μm long laser and a 750 μm long laser with anti-reflection/high reflection coated facets ($R_1/R_2 = 10%$/95%).

Fig. 4. Maximum output power as a function of device temperature for 250 μm and 750 μm long buried heterostructure lasers with $R_1/R_2 = 10%$/95%.

4. Summary

In summary, we report low threshold current InAsP/InGaAsP SCH-MQW lasers for 1.3 μm emission grown by GSMBE. Threshold currents as low as 1.1 mA at 20°C and 6.1 mA at 100°C were measured for buried heterostructure lasers with high-reflection coatings on the laser facets. To our knowledge, these are the lowest threshold currents yet reported for InGaAsP lasers grown by GSMBE. Furthermore, external slope efficiencies as high as 0.48 mW/mA with wall plug efficiencies of 30% at 10mW were observed, and a maximum output power of 65 mW at 20°C was achieved.

Acknowledgements

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References

High temperature (Ga)InAsP/high band gap GaInAsP barriers 1.3 µm SL-MQW lasers grown by gas source MBE

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Abstract

We report on an experimental investigation of the temperature sensitivity of (Ga)InAsP compressively strained multi-quantum well (MQW) 1.3 µm broad area lasers with respect to the amount of strain and barrier band gap. The composition of wells are adjusted to let the strains range from 0.6% to 1.4%, whereas barrier band gap scales from 1.13 to 1.3 eV. Threshold current densities per well for infinite length as low as 60 to 135 A cm$^{-2}$ at 20°C assesses for the high quality of the structures grown by gas source MBE. Characteristic temperatures of 80 to 110 K, depending on the barrier band gap, are measured between 20°C and 80°C for 1200 µm long devices.

Keywords: 1.3 µm lasers; Strain-compensated multi-quantum wells; Characteristic temperature

1. Introduction

Low cost, low drive current, wide temperature range [$-40^\circ$C, $+80^\circ$C] 1.3 µm lasers are key components for large volume fibre to the home applications. In the GaInAsP/InP system, Auger recombination and/or carrier overflow in the barrier and SCH layers, resulting in low characteristic temperature $T_0$ (40–50 K for short cavity length), have been identified to limit high temperature performances of the devices [1, 2].

The introduction of strain in quantum well lasers is well known to improve threshold current densities through a reduction of both densities of state and nonradiative recombination processes (Auger, inter valence band absorption) [3]. Highly strained (≈1.4%) InAsP 1.3 µm MQWs have been shown to be of interest for low threshold lasers [4, 5]. However, the effect of strain on the temperature sensitivity, as seen from many experimental results, is controversial [3]. As a matter of fact, temperature sensitivity strongly depends on the lasing cavity design parameters, i.e. vertical and lateral confinements, length, facets coating, leading to a large discrepancy of results.

More recently, the benefit of high band gap barriers on the temperature characteristics of devices
has been evidenced. Lasing at temperatures in excess of 130°C has been reported with structures including 1.05 μm InGaAsP barriers and SCH layers grown by chemical beam epitaxy (CBE) [6]. To's as high as 117 K between 20°C and 60°C have been shown on 10 InAsP strained-layer multi-quantum well (SL-MQW) with InGaP barriers devices grown by atmospheric pressure metal organic vapor phase epitaxy (AP-MOVPE) [7]. It has to be pointed out that To is not the most relevant parameter regarding high temperature laser operation. From this point of view, threshold current density and external quantum efficiency at 80°C are best suited to characterize the potential of a device structure.

We report on the experimental investigation of both compressive strain in the wells and barrier (lattice-matched or tensile strained) band gap on the temperature characteristics of SL-MQW broad area (BA) lasers grown by gas source MBE (GSMBE).

2. Structure design and growth

The strain in the In(Ga)AsP QWs increases as their Ga concentration (Xw) is reduced. For a wavelength of 1.3 μm and a well width of 8 nm, compressive strain values of 0.6, 1% and 1.4% can be reached with $X_w \approx 0.19$, 0.11 and 0.00, respectively.

The compositions of the barriers have been calculated to meet the following requirements: (i) to let the electron escape energy $E_{be}$ range in a large interval, $E_{be}$ being defined as the difference between the well/barrier conduction bands offset $\Delta E_c$ and the 1st quantized electron level $e_1$, (ii) to compensate with tensile strain the highly compressive InAsP wells. Tensile strain of about −0.8% is achievable by GS-MBE. Moreover, tensile strain in barriers improves slightly the conduction bands offsets, and thus further increases $E_{be}$. From those conditions, a maximum $E_{be} \approx 200$ meV is calculated for Ga$_{0.11}$InP (−0.5%) barriers and InAs$_{0.43}$P wells. Such values compare favorably with those obtained in the AlGaInAs/InP system [8].

The structures consist of six QWs and seven barriers, both 8 nm thick embedded in thick 1.05 μm lattice-matched GaInAsP SCH layers. The optical confinement factor in all the structures is kept almost constant and nearly equal to that of a previously described structure [9], that showed high quality characteristics at 85°C, hereafter referred as sample 1.

The active stacks are grown on 2" n-type InP wafers, and are cladded with a 0.5 μm thick n ([Si] = 2 × 10$^{18}$ cm$^{-3}$) type InP buffer and a 0.5 μm p ([Be] = 1 × 10$^{18}$ cm$^{-3}$) type InP layers. The growth is performed under high group V fluxes (PH$_3$ flow rate ≥ 10 scrm) at temperatures of 480°C to 500°C for the SL-GaInAsP/GaInAsP structures and of 470°C for the InAsP/tensile GaInAsP, in order to improve the tensile material quality [10]. Growth interruptions at each interfaces are necessary to allow for appropriate gas switching. As will be discussed later, such a process is of high importance regarding the control of the composition of low As concentration, high band gap quaternaries barriers.

The structures exhibit high room temperature photo-luminescence (PL) yield with low FWHM (25 to 30 meV) and sharp double crystal X-ray diffraction (DC-XRD) satellite peaks. The analysis of the positions and relative intensities of the satellite peaks, allows a precise measurement of the period of the MQW and the determination of the strain share between wells and barriers. In addition, due to the precise control of the III elements fluxes obtained in MBE, the well and barriers compositions are known with a good precision, as stated by the good agreement between experimental and calculated DC-XRD spectra. As mentioned above, the interface growth interruptions (IGI), for As-free barrier composition, are of major importance regarding the control of materials composition and crystalline quality, as clearly evidenced in Fig. 1. The DC-XRD spectra of two 8 periods InAsP/GaInP MQW are displayed together with their PL intensity maps. Growth interruption times have been increased to reduce arsenic residual incorporation in barriers. Sample B experiences an IGI four times longer than sample A and shows superior PL intensity and homogeneity. The comparison of the rocking curves indicates a lower tension in the barriers of sample A, cancelling the strain compensation effect. On the other hand,
Fig. 1. DC-XRD spectra and PL maps of 8 periods InAsP (8 nm)/Ga$_{0.1}$InP (8 nm) MQW structures grown with different interface growth interruption (IGI) times: (a) Sample A, (b) Sample B. IGI is 4 times than for sample A.
XRD rocking curves of 8 periods InP/Ga_{0.11}InP superlattices are easily fitted with the measured periods and the Ga concentrations calculated from the calibrated In and Ga flux ratio. We thus conclude to excess arsenic incorporation, decreasing with the duration of the growth interruption.

Using XRD, the arsenic concentration in InAsP can be calculated within ±1%, resulting in easy control of the PL wavelength of InAsP/GaInAsP MQWs as a function of hydrides flow rates, as shown in Fig. 2. Wavelengths calculated with the compositions determined by XRD are in good agreement with experiment.

Table 1 summarizes the parameters of the structures selected for lasers processing. $E_{th}$ values range from ≈70 to ≈170 meV for samples 1 to 6, respectively.
3. Device results

The structures are overgrown by GSMBE with p-type InP cladding (2.5 μm, \( p = 1 \times 10^{18} \text{ cm}^{-3} \)) and InGaAs contact (0.3 μm, \( p = 3 \times 10^{19} \text{ cm}^{-3} \)) layers, and are subsequently processed as 100 μm wide BA lasers. Their as-cleaved \( L(I) \) characteristics are measured from 20°C to 80°C, for cavity lengths ranging from 300 to 1200 μm. Fig. 3 shows a plot of threshold current densities (\( J_{th} \)) at 20°C as a function of inverse cavity lengths. Low \( J_{th} \) of 420 to 560 A cm\(^{-2} \) are measured on samples 1 to 5 for 1200 μm long lasers. External quantum efficiencies (\( \eta_{ext} \)) ranging from 0.31 to 0.36 W/A are measured for short cavity length (300 μm). Sample 6, with high barrier band gap of \( \approx 1.29 \text{ eV} \) (0.96 μm quaternary), exhibits threshold current densities in excess of 1 kA cm\(^{-2} \) for all lengths and poor external efficiencies for short cavity length. Calculation of current distribution in a MQW structure as a function of well number and barrier composition (lattice-matched GaInAsP) shows that inhomogeneous pumping may occur for barrier band gap in excess of 1.24 eV, because of the large valence bands offsets (\( \Delta E_v \approx 0.6 \Delta E_g \)) responsible for inhomogeneous hole injection. This effect has been reported by Kito et al. [11] to appear for barrier band gaps between 1.18 and 1.25 eV.

Internal losses (\( \xi_{int} \)) and quantum efficiencies (\( \eta_{int} \)) calculated from \( 1/\eta_{ext} \) (\( L \)) curves, are displayed in Table 2, as well as threshold current densities per well for infinite length (\( J_0 \)) at temperature of 20°C and 80°C. Low internal losses of 6 to 11 cm\(^{-1} \) are found up to 80°C, demonstrating the high quality of the active material. Such characteristics are among the best reported so far [12–15].

Average characteristic temperatures for 300 and 1200 μm long lasers range from 50 to 80 K and from 60 to 110 K, respectively, depending on the barrier band gap. A variation of \( T_0 \) with respect to

<table>
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<tr>
<th>Sample</th>
<th>20°C</th>
<th>80°C</th>
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<tr>
<td>( J_0 )</td>
<td>( \xi )</td>
<td>( \eta )</td>
</tr>
<tr>
<td>2</td>
<td>71</td>
<td>5.6</td>
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<tr>
<td>3</td>
<td>57</td>
<td>8.7</td>
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<td>5</td>
<td>64</td>
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calculated electron escape energy $E_{be}$ is seen in Fig. 4. Solid lines are exponential fits of the experimental points. A noteworthy improvement of $T_0$ is seen for $E_{be}$ higher than 140 meV. On the other hand, the comparison of the $T_0$'s measured for samples 1 to 4 indicates a minor influence of strain, in the 0.6–1.4% range, on characteristic temperatures. However, the lowest threshold current densities at 80°C are obtained for the InAsP sample 3, with $T_0 = 55$ K. This clearly demonstrates that $T_0$ is not the only parameter to be considered for high temperature laser operation.

4. Conclusion

In summary, 1.3 μm In(Ga)AsP SL-MQW broad area lasers, with QWs strains and InGaAsP barrier compositions ranging from 0.6% to 1.4% and from 1.10 to 0.96 μm, respectively, have been grown by GSMBE. Their high quality characteristics have been analyzed at temperatures up to 80°C, as a function of strain and electron escape energy. An improvement of the characteristic temperatures up to 110 K is shown for high band gap barriers, whereas strain does not influence $T_0$ within the studied range. However, inhomogeneous carrier injection limits the lasing characteristics of devices with band gap barriers higher than $\approx 1.24$ eV.

The lowest threshold current densities, at 80°C, of 2 kA cm$^{-2}$ and 0.96 kA cm$^{-2}$ for 300 μm and 1200 μm long cavities and as-cleaved facets has been obtained with an InAsP (1.4%/1.10 μm) GaInAsP (−0.2%) MQW structure, despite a $T_0$ of 55 K. These results show that the optimization of high temperature laser is a trade-off between low threshold current density at room temperature and $T_0$.

References


MBE grown mid-infrared type-II quantum-well lasers

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Abstract

We have employed molecular-beam epitaxy (MBE) for the growth of mid-infrared (MIR) type-II quantum-well laser structures. These lasers consist of strain-balanced InAs/InGaSb/InAs/AlSb type-II quantum wells lattice-matched to the AlSb cladding layers. We have demonstrated optically pumped lasers emitting from 3 to 4.3 \textmu m under pulsed operation. For the 3.2-\textmu m lasers, stimulated emission was observed at temperatures up to 350 K. The characteristic temperature $T_0$ at operation temperatures above ambient was 68 K. Here, we discuss the optimization of the MBE growth of MIR type-II quantum-well lasers, including substrate temperatures, V/III beam-equivalent pressure ratios, and shutter sequencing for better interface control and laser performance.

PACS: 42.55.P; 81.05.E

Keywords: Infrared; Semiconductor laser; MBE; Quantum well

1. Introduction

Efficient mid-infrared (MIR) diode lasers are highly desirable for a variety of applications including remote sensing, molecular spectroscopy, and environmental monitoring due to the strong characteristic absorption lines of many gas molecules in the MIR range. The detection sensitivity of trace gases is typically higher by two orders of magnitude at the fundamental absorption lines than at overtones in the near infrared (IR). Many commercial and military MIR systems are limited by a lack of adequate optical sources. Recently, significant progress has been achieved towards room-temperature operation for MIR diode lasers, promising compact and inexpensive sources more widely applicable than conventional technology such as optical parametric oscillators (OPOs).

MIR lasers have been demonstrated in a variety of material systems, including IV-VI and II-VI diodes, and III-V devices based on interband or intersubband transitions. Lead-salt IV-VI lasers emitting at wavelengths longer than 3 \textmu m have been commercially available for a number of years. These lasers with emission wavelengths of 4 \textmu m have exhibited pulsed operation at temperatures up to 290 K [1], and cw operation up to 223 K [2].
However, they are not expected to produce high output power (>1 mW) because of the poor thermal conductivity of lead-salt alloys, and the susceptibility to damage. For narrow band gap II–VI semiconductors such as HgCdTe alloys, relatively high powers have been reported [3], but for wavelengths beyond 3 μm the maximum operating temperatures have been ≤160 K. It is thus very desirable to develop semiconductor lasers based on III–V alloys, which offer good metallurgical and thermal properties with better-quality substrates readily available.

Room-temperature operation of diode lasers based on InGaAsSb/AlGaAsSb materials have been demonstrated at wavelengths shorter than 2.8 μm [4, 5]. However, the growth of InGaAsSb quaternary alloys with band gaps narrower than 0.37 eV (≥3.4 μm) turns out to be metastable and problematic because of a miscibility gap [6]. Until now, the highest reported operating temperature \(T_{\text{max}}\) of any III–V type-I interband semiconductor laser emitting at wavelengths ≥3.2 μm was 225 K [7]. The corresponding characteristic temperatures \(T_0\) were in the range of 17–30 K for operating temperatures above 120 K. Recently, type-I InAlAs/InGaAs quantum-cascade (QC) lasers based on intersubband transitions had lased up to 320 K at a wavelength of about 5 μm [8]. However, the demonstrated type-I QC lasers had low radiative efficiencies near threshold (<10⁻³) due to fast optical phonon scattering which led to high threshold current densities and substantial heating.

Recently, we reported an optically-pumped type-II quantum-well (QW) laser operating up to 285 K at 4.1 μm [9]. The characteristic temperature \(T_0\) was 35 K for 170 K ≤ \(T_{\text{op}}\) ≤ 270 K. The active region was composed of 35 periods of undoped InAs/In₀.₅Ga₀.₅Sb/InAs/AlSb (21 Å/31 Å/21 Å/43 Å) multiple quantum wells (MQWs) sandwiched by AlSb cladding layers. The maximum peak output power \(\lambda_{\text{pump}} = 1.06 \mu\text{m}\) was about 650 mW at 81 K, while about 250 mW was collected at 170 K. The type-II four-constituent QW structure combines the advantages of good carrier confinement, the potential of significant suppression of non-radiative Auger losses [10, 11], large optical matrix elements, and a two-dimensional density-of-states for both electrons and holes. Here we report results of 3.2-μm lasers and some preliminary results of 4.3-μm lasers. We also discuss key issues regarding the MBE growth of MIR type-II QW lasers, including substrate temperatures, V/III beam-equivalent pressure (BEP) ratios, and shutter sequencing for interface control and laser performance.

2. Experimental procedure

The type-II QW lasers were grown in a Riber 32 molecular beam epitaxy (MBE) system on p-type GaSb substrates using an EPI As valved cracker cell and an EPI Sb cracker cell. Growth rates for each layer composition were calibrated to within ±2% using reflection high-energy electron diffraction (RHEED) and were confirmed by double-crystal X-ray diffraction (DCXRD) measurements. The background doping was low \(10^{15} \text{ cm}^{-3}\) n-type for InAs and about \(2 \times 10^{16} \text{ cm}^{-3}\) p-type for GaSb.

![Fig. 1. PL spectrum of the GaSb/AlSb QWs at 10 K with GaSb well thicknesses of 40, 70, 100, 130 Å. The AlSb barrier thickness was 300 Å.](image-url)
From DCXRD spectra, the background Sb in InAs was about 1.2% and the background As in GaSb was about 3.2%, with the same growth rate and same V/III BEP ratio of 3 for InAs and GaSb. The BEP ON/OFF ratio with the shutter opened or closed was about 8 for As$_2$, and was about 160 for Sb$_4$. When the Sb/Ga V/III BEP ratio was increased, the background As in GaSb decreased. However, the background Sb in InAs was not sensitive to the As flux. This implies that the sticking coefficient of background Sb in InAs at 440°C is almost 100%. From photoluminescence (PL) spectra, the material quality of GaSb/AlSb QWs is not sensitive to the Sb/Ga V/III BEP ratio as long as the ratio is larger than 1.8. Fig. 1 shows the 10 K PL spectrum of GaSb/AlSb QWs with GaSb well thickness of 40, 70, 100, and 130 Å grown at 500°C. The full-width at half-maximum was 6.1 meV for the 100 Å GaSb well, 6.8 meV for the 70 Å GaSb well, and 9.0 meV for the 40 Å GaSb well.

For the laser samples, the substrate temperature was controlled at 530°C for the bottom AlSb cladding layer, 440°C for the MQW active region to optimize the material quality of InAs and InGaSb wells, and 510°C for the top AlSb cladding layer and GaSb cap to minimize degradation of the underlying layers. The MBE grown laser wafers exhibited excellent surface morphology. The substrate temperature was calibrated by optical pyrometry, the InSb melting temperature (525°C), and the GaSb oxide desorption temperature (575°C to 585°C). During the growth of AlSb cladding layers, we noticed that the RHEED patterns became more and more spotty. After growing 2-µm AlSb, the substrate temperature decreased by about 50 to 70°C as estimated from the optical pyrometry and InAs RHEED patterns, since InAs displays 2×1 RHEED patterns only when the substrate temperature is between 380°C and 470°C [12]. We believe that the decrease of the substrate temperature is due to the large difference in the refractive indices of the GaSb substrate and AlSb.

During the growth, the InGaSb layers in the active region and the AlSb cladding layers displayed good 1×3 RHEED patterns, while the InAs layers exhibited 2×1 patterns with a V/III BEP ratio of 10 using As$_4$ or 3 using As$_2$. However, the RHEED patterns for the AlSb barriers in the MQW active region were not as good, since the substrate temperature was about 440°C. We have also noticed that it took about 1.5 monolayers of InAs growth to change the RHEED patterns from 1×3 to 2×1. This suggests that the interface was InSb dominant and that there was some Sb floating on the InGaSb or AlSb surface and got incorporated into the InAs layers, which was subsequently confirmed by the TEM analysis. After the growth of InAs, the sample was exposed to As$_2$ only for two more seconds. Next, all of the shutters were closed for six seconds to reduce the background As. Then, the Sb shutter was open for two seconds before the growth of InGaSb or AlSb. When we grew InGaSb or AlSb on top of an InAs layer, the RHEED patterns changed from 2×1 to 1×3 right after we opened the Sb shutter. Therefore, the interface quality for InGaSb or AlSb grown on top of InAs is better than that for InAs grown on top of Sb-based materials. The buffer and cladding layers were grown at about 0.85 µm/h, and the InGaSb and AlSb layers in the MQW active region were grown at about 0.7 µm/h.

After characterized by DCXRD and PL, wafers were lapped down to a thickness of 125 µm, and then cleaved to laser bars with cavity lengths on the order of 500 µm. The lasers with uncoated facets were indium heat sunk to a copper cold finger, epitaxial-side-up, and mounted in an Air Products Heli-Tran dewar. The devices were optically pumped over a 90-µm wide strip by 70 ns pulses from a Q-switched 2.06 µm Ho–YAG laser. At this pump wavelength, it is estimated that only about 10% of the incident intensity is absorbed in the active region. The lasing output was collected and focused onto the slit of a ½ m monochromator and measured with an InSb detector.

For the 3.2-µm lasers, the active region is composed of 40 periods of undoped InAs/In$_{0.4}$Ga$_{0.6}$Sb/InAs/AlSb (17 Å/28Å/17 Å/43Å) MQWs sandwiched between two AlSb cladding layers. The PL line width was about 100 nm at 80 K, and about 180 nm at 300 K. The line width for edge emission above the lasing threshold narrowed to 45 nm at 300 K, which was limited by the instrument resolution. Fig. 2 shows the threshold pump intensity as a function of temperature. The slope of the data in
reported for a semiconductor laser emitting beyond 2.5 μm. Fig. 3 plots the peak output power as a function of pump intensity at 300 K. We find that more than 270 mW is generated at ambient temperature, which represents a significant step toward the realization of practical MIR sources. Correlation of the observed threshold intensities with estimated absorption coefficients and calculated threshold carrier concentrations leads to an estimated Auger coefficient of $5.0 \times 10^{-27}$ cm$^6$/s at 300 K [13]. Lasers from a second wafer using conventional As$_4$ and Sb$_4$ effusion cells lased only up to 310 K. It is possible that the use of the monomeric Sb source could have reduced the defect-related nonradiative recombination. Further investigations will be performed in the future.

Recently, we have also demonstrated 4.3-μm lasers emitting up to 270 K. The active region composed of 35 periods of undoped InAs/In$_{0.33}$Ga$_{0.67}$Sb/InAs/AlSb (24 Å/40 Å/24 Å/43 Å) MQWs. The DCXRD spectrum displayed clear satellite peaks which indicated good material quality, even though the MQW active region was lattice-matched to the AlSb cladding layers instead of the GaSb substrate. From the threshold pump intensity as a function of temperature, the characteristic temperature ($T_0$) was determined to be 36 K over the temperature range 160 K to 270 K. The detailed data will be presented elsewhere.

3. Conclusions

We have demonstrated that MBE is capable of growing high-quality strain-balanced type-II InAs/InGaSb/InAs/AlSb QWs which are well suited for the MIR lasers. Two optically pumped devices incorporating four-constituent MQWs lased at 3.2 μm with the maximum operating temperature of 350 K and a characteristic temperature of 68 K over the entire temperature range 90–350 K, and at 4.3 μm with the maximum operating temperature of 270 K and a characteristic temperature of 36 K. Higher operating temperatures and output powers are expected with further improvements in the material quality of the type-II QW.
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References

Waveguide modulator structures with soft optical confinement grown by the epitaxial shadow mask (ESM) MBE-technique


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Abstract

Excellent selective n- and p-contacts to highly doped and closely adjacent layers in n-i-p-i doping superlattices can be fabricated using the epitaxial shadow mask molecular beam epitaxy (ESM-MBE). With this method the growth within the window of a GaAs/AlGaAs mask obtained by selective wet chemical etching is controlled by the geometrical aspect of the effusion cells relative to the substrate. We demonstrate first results of using ESM-MBE for the in-situ fabrication of waveguide structures, including waveguide modulators. This double hetero p-i-n modulator exhibits a modulation contrast of more than 30 dB (for a 900 μm long device) for a voltage swing of 6.5 V. Propagation losses in the waveguide structure are less than 11 dB/cm and mainly due to free carrier absorption in the waveguide region.

1. Introduction

A key issue for the feasibility of opto-electronic integrated circuits (OEICs) is the possibility of smart integration of active components (laser diode, photodetector) and passive elements (intersection, crossing, coupler) with modulators [1, 2]. Epitaxial regrowth is widely used to fabricate buried ridge laser structures [3] or even vertical cavity surface emitting lasers [4]. Surface selective growth on masked substrates has been demonstrated for opto-electronic device applications [5–7]. The epitaxial shadow mask MBE technique [8] has been shown to yield excellent selective in situ ohmic contacts to the respective n- and p-layers of n-i-p-i doping superlattices. These structures are attractive for applications as (waveguide) modulators because high switching contrast can be achieved with low driving voltages [9]. Another important issue concerns the electro-optical effect used for intensity or phase modulation. Most work in the literature deals with the quantum confined stark effect (QCSE) [10–13] because of the very attractive and pronounced features of the field-dependent excitonic absorption in the quantum wells. Few papers report on the use of the Wannier–Stark localization in superlattices [14, 15] or on the bandfilling effect in electron transfer quantum well modulators [16]. The Franz Keldysh effect (FKE) has been barely used [17] despite of comparable absorption changes and rather simple double hetero p-i-n structures.

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polarization insensitive (in contrast to QCSE modulators) and exhibited a switching contrast of 40 dB (for a 760 μm long device) with propagation losses of less than 0.5 dB/cm without bias [18].

In this paper, we report the fabrication and the results of a double hetero p-i-n waveguide modulator structure grown by the epitaxial shadow mask MBE technique and based on the Franz-Keldysh effect.

2. Experimental procedure

The samples were grown in a Riber 32P machine with a CBr₄ gas line used for p-type doping. An EPI valved cracker cell was used in the As₂ mode ($T_{cracker} = 800^\circ$C) with a V/III-ratio of $\approx 10$ for the growth of samples containing AlGaAs layers. A quarter of a 2 in wafer with an epitaxial shadow mask (8 μm Al₀.₅Ga₀.₅As and 3 μm GaAs layer on top) was photolithographically patterned into stripes with different widths ($> 10 \mu m$). The GaAs layer was etched with an anisotropic etchant (NH₃·OH : H₂O₂ : H₂O = 3 : 1 : 15) and the Al-GaAs layer was then selectively etched with HF. The sample was extensively rinsed in clean water and blown dry before loading back into the MBE chamber. The standard growth position of the Riber 32P was used to have highest symmetry to all the cells. The sample was mounted on the In-free holder and grown under standard conditions ($T_{growth} = 600^\circ$C, growth rate = 1 μm/h, rotation speed = 12 rpm).

Starting with a 1 μm Al₀.₃Ga₀.₇As cladding layer, followed by a 1 μm GaAs layer and a 0.5 μm Al₀.₅Ga₀.₅As layer on top, a waveguide structure was fabricated in situ. Within the GaAs layer, a δ-n-doped bottom and a δ-p-doped top layer were grown while the rotation was stopped and the sample was properly aligned relative to the doping cells exploiting the shadowing effect for the doping beams [19]. The doping density of the δ-n- and δ-p-layer was $1 \times 10^{13}$ cm⁻² each. (The doping densities were calibrated by Hall measurements of homogeneous doped samples). In Fig. 1, a scanning electron picture of the as grown structure before the mask was removed and a cross section through the waveguide is shown. The GaAs waveguiding layer becomes gradually thinner at the edges of the structure resulting in soft optical confinement for the optical mode. The defect density on the surface of the waveguide is as low as for growth without a shadow mask. Some (sub-μm) roughness along the edges is still apparent which was transferred from the edges of the mask window.

After the growth, the shadow mask was selectively etched away and ohmic n-(Au/Ge) and p-contacts (Au/Zn) were evaporated to the respective edges. Finally, waveguide devices of about 1 mm length were cleaved.

Transmission experiments were carried out with spectrally dispersed light of a tungsten halogen lamp or with coherent light of a tunable Ti-Sapphire laser coupled into a fiber. The intensity was optionally tuned with a HP 8158 B optical fiber attenuator. The tapered fiber output was aligned to the front end of the waveguide structure while monitoring the output signal of the modulator at
the backside with a CCD camera. The camera was replaced by the Si detector head of a HP 8153 powermeter for measuring the transmitted power.

3. Results and discussion

The devices showed good rectifying behavior but relatively high contact resistance which is probably due to the fact that just one n- and p-δ-doped layer was used which are covered by the 0.5 μm thick undoped AlGaAs top layer. The normalized transmission changes for different wavelengths from 920 to 895 nm in 5 nm wavelength steps are shown in Fig. 2 for a 0.9 mm long and 10 μm wide waveguide structure. A modulation contrast of 30 dB could be achieved with a voltage swing between −6 and +0.5 V for the shortest wavelength used. The contrast decreases with increasing wavelength according to the diminishing absorption changes due to the FKE. We mention that in this 10 μm wide waveguide modulator at least one higher mode is propagating. The waveguide losses were determined by measuring the contrast of the Fabry-Perot oscillations of the modes while slowly heating the sample [20]. The relative transmission of a waveguide structure with the length \( L \) and the facet reflectivity \( R \) is given by

\[
T(\phi) = \frac{(1 - R)^2 \exp(-\alpha L)}{(1 - R \exp(-\alpha L))^2 + 4R \exp(-\alpha L) \sin^2(\phi)}
\]

(1)

with \( \phi = 2\pi n L / \lambda \) and the propagation loss \( \alpha \). The temperature dependence of the effective index \( n \) is about \( 4.5 \times 10^{-5} \) 1/K. The function \( T(\phi) \) is oscillating with \( \phi \) and from the contrast ratio \( k = (T_{\text{max}} - T_{\text{min}}) / (T_{\text{max}} + T_{\text{min}}) \) one can calculate an upper limit for the total losses. Fig. 3 shows the transmission at room temperature as a function of temperature change at the wavelength \( \lambda = 923 \) nm (solid line) together with calculated loss curves for different absorption values (dashed lines). The best correspondence to the experimental data was found for \( \alpha = 2.7 \) 1/cm which yields a loss of 11 dB/cm. This seems to be a still rather high value for waveguide modulator structures. But one has to bear in mind that the δ-doping layers lie in the active region and therefore absorption due to free carriers has to be considered. Also, leakage through the relatively thin (1 μm) lower cladding layer might still contribute significantly to the total loss.

4. Conclusion and perspectives

The fabrication of in-situ structured waveguides has been successfully demonstrated. Even with just one single δ-n and δ-p doping layer in the active layer satisfactory contrast ratios as high as 30 dB have been achieved. The waveguide losses for that structure are still relatively high compared to conventional waveguide structures which is mainly
due to free carrier absorption in the active layer. Further investigations are in progress to improve the results significantly. First, homogeneous doping layers have to be used in the cladding layers. This is made by the “flash-doping” technique, i.e. opening the respective doping shutters for a short time when the sample is properly oriented during continuous rotation [19].

The width of the waveguides has to be reduced also to achieve monomode propagation. This can be done by reducing the width of the shadow mask or by lateral variation of the Al-content. This can be done by the same “flash”-technique mentioned above by opening the Al-shutter for a short period in the proper position to give an increasing Al-content directed outside to the edges in the GaAs waveguiding layer.

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References

The growth and characterization of two new P-type compressively strained layer InGaAs/AlGaAs/GaAs quantum well infrared photodetectors for mid- and long-wavelength infrared detection

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Abstract

Investigation of two p-type compressively-strained layer (PCSL) InGaAs/AlGaAs/GaAs quantum well infrared photodetectors (QWIPs) grown on (100) semi-insulating (SI) GaAs substrate has been carried out. The first detector uses a step-bound-to-miniband (SBTM) transition scheme for long wavelength infrared (LWIR) detection and has a detection peak at 10.4 \textmu m with a full width at half-maximum bandwidth, $\Delta \lambda / \lambda_p = 20\%$. A responsivity of 28 mA/W was obtained for this detector at $T = 65$ K and $V = 3.0$ V, with a spectral detectivity $D^* = 1.4 \times 10^7$ cm Hz$^{1/2}$ W$^{-1}$ at $T = 65$ K and $V = 1.0$ V. The detector was under the background limited performance (BLIP) at $T = 40$ K and $V \leq 2.0$ V. The second detector is a two-color stacked QWIP composed of a PCSL In$_{0.2}$Ga$_{0.8}$As/Al$_{0.3}$Ga$_{0.7}$As QWIP for the MWIR detection at $\lambda_p = 4.8$ \textmu m and a PCSL In$_{0.15}$Ga$_{0.85}$As/Al$_{0.1}$Ga$_{0.9}$As for the LWIR detection at $\lambda_p = 10$ \textmu m. The peak responsivity for the LWIR QWIP was found to be 25 mA/W at $V_b = 2$ V, $T = 40$ K, $\lambda_p = 10$ \textmu m, a FWHM of $\Delta \lambda / \lambda_p = 40\%$, and the calculated detectivity was found to be $D^* = 1.1 \times 10^7$ cm Hz$^{1/2}$ W$^{-1}$. Two response peaks for the MWIR QWIP were found to be at 4.8 and 5.4 \textmu m with maximum responsivities of 12 and 19 mA/W (at $T = 77$ K and $V_b = 5$ V), and spectral bandwidths of 21 and 26\%, respectively. The detectivity ($D^*$) for the MWIR stack determined at $\lambda_p = 5.4$ \textmu m, $V_b = 1.0$ V and $T = 77$ K was found to be $5.5 \times 10^7$ cm Hz$^{1/2}$ W$^{-1}$.

1. Introduction

In recent years, a great deal of works have been published on the III–V semiconductor quantum well infrared photodetectors (QWIPs) for long wavelength infrared (LWIR) and mid-wavelength infrared (MWIR) focal plane arrays (FPAs) applications. Most of the studies have been centered on the n-type QWIPs [1, 2]. However, for n-QWIPs due to quantum mechanical selection rules, normal incidence absorption is forbidden without the use of metal or dielectric grating couplers. In contrast, the normal incidence absorption is allowed in p-type QWIPs due to the mixing between

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the off zone-center (k ≠ 0) heavy-hole and light-
hole states [3]. Because of the larger effective mass
(hence lower optical absorption coefficient) and the
lower hole mobilities, the performance of p-QWIPs
are in general lower than n-QWIPs [1, 4]. However,
if the biaxial compressive strain is introduced
into the quantum well layers of a p-QWIP, then the
effective mass of the heavy holes will be reduced,
which in turn can improve the overall device
performance [5]. In this work, we present a new p-type
compressively strained layer step-bound-to-miniband
(SBTM) transition InGaAs/AlGaAs/GaAs
QWIP for LWIR detection. The need to detect
simultaneously two different infrared wavelengths
in the MWIR and LWIR bands has led us to design
a stacked p-type compressively strained layer
(PCSL) QWIP structure with detection peaks in
the 3–5 μm MWIR band and 8–14 μm LWIR band.
The results of this multicolor stacked QWIP will also
be presented in this paper.

2. Design and growth of p-type strained layer
QWIPs

First, we report a new p-type SBTM CSL-QWIP
grown on a semi-insulating (SI) (1 0 0)GaAs by
molecular beam epitaxy (MBE) using InGaAs for
the quantum well and AlGaAs/GaAs material system
for the superlattice barrier layer. As illustrated
in Fig. 1a, the transition scheme for this p-QWIP is
from the localized ground bound heavy hole state
(HH1) in the wide Be-doped InGaAs quantum well
to the resonant coupled miniband of the undoped
GaAs/AlGaAs superlattice (SL) barrier. This structure
creates a potential difference between the SL
barrier region and the quantum well which blocks
part of the undesirable tunneling dark current from the
heavily doped heavy hole ground state, HH1
[2] in the quantum well. The physical parameters
were chosen so that the ground state in the wide
InGaAs quantum well is well above the top of the
GaAs/AlGaAs SL barrier, and the third excited
heavy hole state (HH4) is in resonance with the
ground level of the superlattice miniband (SL1) to
achieve a higher quantum efficiency. Since the super-
lattice consists of thin barriers, the photoexcited
holes can easily tunnel through the SL barrier layer
and transport along the aligned miniband to be
collected by the ohmic contacts.

The p-type SBTM CSL-QWIP consists of Be-
doped In0.12Ga0.88As quantum wells to populate
the ground heavy hole (HH1) state. The quantum
well layer is under compressive strain with a lattice
mismatch of nearly −0.8%. Surrounding the
quantum well layers are the SL barriers of undoped
Al0.35Ga0.65As alternating with undoped GaAs
quantum wells. The p-type ohmic contacts were
formed by using a cap layer and a buffer layer of
heavily Be-doped GaAs grown on top of the SBTM
QWIP and between the SBTM QWIP and the
GaAs substrate, respectively.

The second QWIP studied in this work is a
multicolor stacked P-QWIP for the MWIR and
LWIR two-band detection. Fig. 1b shows the energy band diagram of a stacked p-type compressively strained layer (PCSL-) InGaAs/AlGaAs QWIP for the MWIR and LWIR detection. This multicolor QWIP consists of two distinct multiquantum well stacks separated by a common ohmic contact layer and sandwiched between the two (top and bottom) ohmic contact layers. This stacked PCSL- QWIP was grown by the MBE technique on Si (1 0 0)GaAs substrate. The bottom contact consists of a heavily Be-doped GaAs contact layer. On top of the contact, the Al$_{0.5}$Ga$_{0.7}$As/In$_{0.2}$Ga$_{0.8}$As QWIP layer structure was grown for the MWIR stack. The LWIR QWIP stack was formed by using Be-doped In$_{0.13}$Ga$_{0.85}$As quantum wells surrounded with the undoped Al$_{0.1}$Ga$_{0.9}$As barrier layers. The whole stack was sandwiched between two thin Al$_{0.1}$Ga$_{0.9}$As blocking barriers. Finally, heavily Be-doped GaAs layers were grown for the top and middle ohmic contacts. Both the LWIR and MWIR quantum wells are in biaxial compression. This multicolor stacked QWIP used the bound-to-quasi-bound (BTQB) intersubband transition scheme for detection of MWIR and LWIR radiation.

\[ \Delta E_{hh} = b \frac{C_{11} + C_{12} \delta_0}{C_{11}}, \]

\[ \Delta E_{lh} = - \Delta E_{hh} + \frac{(\Delta E_{hh})^2}{2A_0}, \]

where \( c_1 \) is the combined hydrostatic deformation potential which characterizes the splitting of the \( \Gamma \) valence band under strain, \( b \) is the shear deformation potential, \( C_{ij} \) are the elastic constants, \( A_0 \) is the spin orbit split-off energy, and \( \delta_0 \) is the lattice mismatch or the in-plane strain, which is defined as \( \delta_0 = (a - a_0)/a \).

Another feature of the p-type QWIPs is the inherently larger quantum efficiency than that of n-QWIPs, which is given by \( \eta = P(1 - R)[1 - \exp(-2l)] \).

Given similar absorption coefficients, \( x \), and well thickness, \( l \), the prefactor \( P \) is equal to 0.5 for n-QWIPs and 1.0 for p-QWIPs, which gives us a doubling of the quantum efficiency for p-QWIPs. Due to the inherently low absorption coefficients of p-type materials, and the large hole effective mass, compressive strain must be used to reduce the hole effective mass in order to increase the intersubband absorption.

With the inclusion of compressive strain, the mobility of the heavy holes is enhanced by reducing the heavy hole effective mass [5]. Also associated with the presence of compressive strain is the reduction of the in-plane density of states in the InGaAs quantum well. Thus, with a significant lowering of the effective mass of the ground heavy holes, an increase in the absorption coefficient and the corresponding quantum efficiency is expected.

4. Results and discussion

In order to evaluate the performance of the QWIP devices, mesa structure with area of 216 × 216 \( \mu \)m$^2$ was formed on the MBE grown QWIP wafer by wet chemical etching for radiometric and electrical characterization. A narrow ring of Cr/Au film was deposited by E-beam evaporation to create the ohmic contacts. It is noted that in this type of mesa and ring contact structures, the
normal incidence IR radiation is allowed only one pass through the QWIP stack.

For the multicolor stacked QWIP, three mesa structures of different thicknesses were etched to allow separate characterization of the LWIR, MWIR, and the combined stacked QWIP devices. The LWIR QWIP mesa structure was formed using the top and middle ohmic contacts, while the MWIR QWIP mesa structure had the top LWIR stack etched away before the mesa formation. The combined stacked QWIP used the top and bottom ohmic layers for contacts.

The device characterization was performed in a closed cycle helium cryogenic dewar. An HP 4145B semiconductor parameter analyzer was used to measure the dark I-V characteristics and the 300 K background photocurrent. Under dark conditions, holes can be transferred out of the quantum wells and produce the observed dark current mainly due to two mechanisms: the thermionic emission out of the quantum wells and thermally generated carriers tunneling through the superlattice barriers. Given the high aluminum composition in the SL barrier layers, considerable indium content in the quantum well layers, and the effect of the compressive strain lowering the energy of the heavy hole states, the effective barrier seen by the ground heavy hole states was found to be 299 meV, which should suppress the thermionic emission of hole carriers out of the quantum well. However, due to the heavily doped ohmic contact regions, a large triangle potential might be formed to effectively lower the barrier with increasing thermionic emission, and hence results in a higher dark current than expected. Additional contributions to the higher dark current characteristics might also arise from the higher aluminum content used in the SL barrier layers, which has been attributed to the formation of DX centers in the AlGaAs barrier layers.

Fig. 2 shows the dark I-V curves measured over a wide range of temperatures from 40 to 77 K along with the 300 K background photocurrent (FOV = 180°) curve for the SBTM PCL-QWIP device. Relatively symmetrical I-V characteristics were obtained for this device under positive and negative biases, indicating that the dopant migration effect is negligible in this device. Fig. 3a shows the measured photoresponse of the same SBTM PCL-QWIP. A single peak was found at λ_p = 10.4 μm, which is in good agreement with the calculated value of 10 μm (see Fig. 1a). With a half-peak value at 12 μm, we derived a spectral bandwidth of Δλ/λ_p = 20%. This narrow responsivity bandwidth is consistent with that expected for a bound-to-miniband transition QWIP. A maximum responsivity of 28 mA/W was found at T = 65 K and V_b = +3.0 V. At an operating temperature of 65 K, the noise spectral density was
measured as $4.0 \times 10^{-26} \text{ A}^2/\text{Hz}$ at a bias of 1.0 V. Corresponding to this operating point, the measured responsivity at the 10.4 μm peak was found to be 13 mA/W. From the above data, the spectral detectivity of $D^* = 1.4 \times 10^9 \text{ cm Hz}^{1/2} \text{ W}^{-1}$ was obtained. Note that this is the detectivity achieved by a single pass of the normal incidence IR radiation through the SBTM PCL-QWIP. If the test structure is altered to include the backside thinning and a reflective top contact, then the responsivity will increase substantially. The corresponding detectivity will also increase, since the dark current and the noise spectral density remain constant. The variation of responsivity with applied bias is plotted in Fig. 3b. The results show that the responsivity and hence the photoconductive gain increase linearly with the applied bias at a fixed operating temperature.

The dark $I-V$ characteristic for the MWIR, LWIR, and the combined stacked QWIP measured at $T = 77 \text{ K}$ is shown in Fig. 4a. As expected the dark current of the LWIR QWIP stack is several orders of magnitude higher than the MWIR QWIP stack, due to the exponential dependence of the dark current on the barrier height. As clearly seen in Fig. 4a, most of the voltage drop is across the MWIR stack due to the much larger dynamic resistance of the MWIR stack. Fig. 4b shows the measured $I-V$ curves at $T = 40, 60,$ and $77 \text{ K}$ for the LWIR QWIP stack. The asymmetry in the dark $I-V$ characteristics observed in this device can be attributed to the dopant migration effect [8].

The spectral responsivity for the MWIR QWIP measured at $V = 5 \text{ V}$ and $T = 77 \text{ K}$ is shown in Fig. 5. The responsivity measurements were performed with the device mounted in a closed cycle helium dewar and illuminated by a blackbody source running through a grating monochromator and appropriate IR filters. The resulting photocurrent is amplified and detected by a lock-in amplifier. Results of the measurements revealed that two photoresponse peaks were observed in the MWIR band at $\lambda_{p,mw1} = 4.8 \text{ μm}$ and $\lambda_{p,mw2} = 5.4 \text{ μm}$. The 4.8 μm peak is in excellent agreement with the ground heavy hole (HH1) to second bound heavy hole (HH3) transition calculated by the TMM, which predicts a detection peak at 4.7 μm. The detectivity for this MWIR peak was determined to be $D^* = 3.3 \times 10^{11} \text{ cm Hz}^{1/2} \text{ W}^{-1}$ at $V_b = 1.0 \text{ V}$ and $T = 77 \text{ K}$. The measured spectral bandwidth for the first MWIR peak was found to be $\Delta \tilde{\lambda}/\tilde{\lambda}_{p,mw1} = 21\%$ and $\Delta \tilde{\lambda}/\tilde{\lambda}_{p,mw2} = 26\%$ for the second peak. The second, longer wavelength peak is attributed to the transition from the HH1 states to the second bound light hole (LH2) states within the quantum well. The calculated responsivity peak for this transition is 5.6 μm, which is also in good agreement with the measured value. The higher responsivity of the 5.4 μm peak is attributed to the...
higher absorption coefficient inherent with the HH to LH transition [4]. The spectral detectivity for this MWIR peak was found to be $5.5 \times 10^{11} \text{ cm Hz}^{1/2} \text{ W}^{-1}$ at $V_b = 1.0 \text{ V}$ and $T = 77 \text{ K}$.

The responsivity for the LWIR QWIP stack as a function of the wavelength is shown in Fig. 6. A peak detection wavelength at $\lambda_{p,16} = 10.0 \mu\text{m}$ was found for the LWIR QWIP device, which is in excellent agreement with the calculated value of 10 $\mu\text{m}$ from the TMM. The maximum responsivity measured at $T = 40 \text{ K}$ and $V = 2.0 \text{ V}$ was found to be 25 mA/W, with a detectivity of $1.1 \times 10^{10} \text{ cm Hz}^{1/2} \text{ W}^{-1}$ under the same conditions. It is interesting to note that a very broad response with full width at half-maximum bandwidth of 40% was achieved for this device. Fig. 7 shows the relative spectral response of the combined MWIR and LWIR QWIP stack, displaying one dominant response band at MWIR and two smaller response peaks at LWIR bands.

5. Conclusions

In conclusion, we have demonstrated two new p-type compressively strained layer QWIPs using InGaAs/AlGaAs/GaAs material systems for the MWIR and LWIR detection. It is shown that by using the compressive strain in the quantum well and the superlattice barriers, the step bound-to-miniband transition can be achieved, which could be useful in producing a new narrow bandwidth LWIR p-QWIP device. We have also demonstrated that simultaneous two-color two-band
detection can be achieved with strained layer p-type QWIPs by stacking the LWIR and MWIR QWIP devices in series and using a three terminal structure. Given the inherent benefit of normal incidence detection without the use of grating couplers, the simplicity of the p-QWIP design deserves further investigation. By further optimizing the quantum well doping density, biaxial strain strength, superlattice barrier parameters, and by using a triangle potential blocking layer in the ohmic contact regions for the SBTM p-QWIP, high performance LWIR and MWIR p-QWIPs can be developed for IR imagery applications.

Acknowledgements

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References

Large array of GaAs modulators and detectors flip-chip solder bonded to silicon CMOS using InGaP as the selective etch stop for GaAs substrate removal


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Abstract

We demonstrate the integration of a large (64 x 68) p-i(MQW)-n GaAs diode array to a silicon CMOS chip, using flip-chip solder bump bonding techniques together with concomitant GaAs substrate removal. The capability of removing a relatively large area of GaAs substrate cleanly and uniformly is attributed to the introduction of a new selective etch stop with a smooth isotropic etchant for the GaAs substrate removal. The presence of the lattice matched In_{0.5}Ga_{0.5}P selective etch stop layer has no detrimental effect on the performance of the GaAs/AlGaAs optical detector/modulator diodes, which gives the chip the large I/O count of 4352.

PACS: 42.79.Ta

Keywords: InGaP; Flip-chip solder bonding; Modulation/detector array

1. Introduction

In recent years, considerable effort has been devoted to the development of the integration of III-V optoelectronics with silicon electronic circuits. This material combination finds many applications in optical computing, optical information processing, and optical interconnect. The optical detector/modulator diodes based on the quantum-confined Stark effect (QCSE) with a GaAs/AlGaAs multiple-quantum-well (MQW) structure are good...
candidates for the photonic elements of this concept, since they can play the roles of surface normal optical input and/or output (I/O) devices for silicon integrated circuits (IC's) [1]. The integration of two-dimensional arrays of such diodes with Si CMOS circuits is desirable for massive parallelism and connectivity [2].

Possible technologies for achieving large arrays of GaAs/AlGaAs MQW devices on Si CMOS include heteroepitaxial growth, epitaxial lift-off (ELO) and flip-chip bonding. Heteroepitaxial growth of GaAs/AlGaAs modulators on silicon has been demonstrated, [3–5] however, it encounters the problems of the incompatibility of growth and processing technologies [6]. These problems can be alleviated by the technique of epitaxial lift-off [7, 8]. ELO is a process that can free the GaAs epilayers or devices from its growth substrate and reattach them to a new substrate. The high material quality of lattice-matched growth is therefore preserved. The key to the success of this technique is the selective etching of a sacrificial AlAs layer grown between the GaAs substrate and the bottom layer of the device by HF. ELO not only offers the advantage of placing a high-quality single crystalline GaAs film on Si without regard to lattice matching, but also allows GaAs devices and silicon IC's to be independently designed and fabricated for optimum performance. However, there are still some problems to be overcome, e.g. the presence of cracks or strain in the thin layers as well as the handling and adhesion of the thin films [9]. Another approach which is more manufacturable, is the so-called flip-chip solder bump bonding technique [1, 2]. In this approach, the hybridization process of GaAs/AlGaAs detector/modulator devices on silicon includes the chip fabrication, die separation, the alignment, and the solder bump bonding of the GaAs diode array chips onto silicon chips; etching off the optically opaque and thermally strained GaAs substrate; and finishing with the deposition of an anti-reflection coating. This approach offers the same advantages as ELO, but without the drawback of thin-film handling as is required in ELO.

Recently, flip-chip solder bump bonding of GaAs/AlGaAs MQW detector/modulator diodes on Si electronic IC's has been demonstrated by Goossen et al. [1]. In that process, a 1.5 μm thick Al0.3Ga0.7As was used as the selective etch stop for the GaAs substrate removal which was chemically removed by 1:100 NH4OH:H2O2 using a jet etcher. The low etching selectivity of this etchant limits the usefulness of the Al0.3Ga0.7As as an etch stop layer, because it cannot sufficiently compensate for any etching nonuniformities induced during the removal of a thick substrate. The GaAs chip size for successful substrate removal is therefore limited. Here we report a new selective etch stop approach which greatly increases the etching selectivity and gives uniform final etching results over a larger area compared to the Al0.3Ga0.7As etch stop. We use an In0.49Ga0.51P etch stop layer which is lattice matched to the GaAs substrate and which also acts as a buffer layer in the epitaxial growth. We have also demonstrated that the new In0.49Ga0.51P selective etch stop grown as the buffer layer has no detrimental effect on the performance of the GaAs/AlGaAs modulators. The entire substrate from a 64 x 68 diode array was removed cleanly with the new selective etch stop without using substrate thinning or jet etching techniques. Uniform photocurrent response across the array demonstrates that the In0.49Ga0.51P layer is indeed a good selective etch stop for GaAs substrate removal for the entire MQW diode array. The new selective etch stop layer thus enhances the manufacturability of the flip-chip solder bump bonding and substrate removal techniques.

2. Device fabrication

The epitaxial layers of the GaAs/AlGaAs p-i(MQW)-n diode structure as shown in Fig. 1 were grown using gas-source molecular beam epitaxy (GSMBE) on a 2” or 3” semi-insulating GaAs substrate. A 5000 Å In0.49Ga0.51P layer was first grown, followed by a 1 μm Al0.11Ga0.89As p+ layer and a 500 Å undoped Al0.11Ga0.89As spacer. Then the undoped MQW were grown, which consisted of 95 periods of 85 Å GaAs wells and 35 Å Al0.3Ga0.7As barriers. Finally, a 1000 Å n+-Al0.11Ga0.89As layer, followed by a 1000 Å n+ GaAs contact layer. The growth conditions and the switching procedure between AsH3 and PH3 gases
can be found elsewhere [10, 11]. Arrays of detector/modulator diodes are produced on the GaAs wafer via a coplanar ion-implantation process [12]. 3 μm thick Sn/Pb solder with dimension of 15 × 15 microns was deposited on both the n and p ohmic contacts of the modulators using 10 μm-thick photoresist and lift-off technique [13]. Each GaAs chip consists of 4352 GaAs/AlGaAs MQW diodes and 8704 solder bumps. The physical size of the GaAs modulator chip is 6.2 mm × 6.2 mm. The complete GaAs chips were then separated. The CMOS chips (7 mm × 7 mm) designed with 256 16 × 1 switching nodes, each with 16 optical inputs and 1 optical output, were made in the Lucent 1 μm technology. Mating aluminum pads from the modulators were designed on the chips, and Ti–Ni–Au metals were deposited on them to provide a solder-wettable surface. Then solder was deposited on those pads, by a process similar to the one for the GaAs chips. A precision bonder was employed to bond the chips together. A low viscosity epoxy (used to protect the circuitry from the substrate etchant) was flowed between the chips and was cured at 100°C. The bonded chips were then immersed in a H₃PO₄ based etching solution, which dissolves the GaAs substrate at a rate of approximately 3.5 μm/minute isotropically. The dissolution stopped at the In₀.₄₉Ga₀.₅₁P layer. Selectivity of ~290 was obtained. This etchant does not attack Si. Finally, a SiOₓ anti-reflection (AR) coating was deposited.

3. Results and discussion

With the new In₀.₄₉Ga₀.₅₁P selective etch stop, one question we have to answer first is whether

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**Fig. 2.** Nomarski contrast photomicrographs of (a) wavy-finished In₀.₄₉Ga₀.₅₁P surface caused by non-uniform etching during the GaAs substrate removal process by a H₃PO₄ based etching solution; and (b) smooth In₀.₄₉Ga₀.₅₁P surface with an isotropic H₃PO₄ etching solution for GaAs substrate removal.

---

<table>
<thead>
<tr>
<th>Layer</th>
<th>Thickness</th>
<th>Composition</th>
</tr>
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<tbody>
<tr>
<td>n+ GaAs</td>
<td>1000 Å</td>
<td></td>
</tr>
<tr>
<td>n+ Al₀.₁₁Ga₀.₈₈As</td>
<td>1000 Å</td>
<td></td>
</tr>
<tr>
<td>MQWs</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Al₀.₇Ga₀.₃As</td>
<td>35 Å</td>
<td></td>
</tr>
<tr>
<td>GaAs</td>
<td>85 Å</td>
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<td>Al₀.₇Ga₀.₃As</td>
<td>35 Å</td>
<td></td>
</tr>
<tr>
<td>undoped Al₀.₁₁Ga₀.₈₈As</td>
<td>500 Å</td>
<td></td>
</tr>
<tr>
<td>p+ Al₀.₁₁Ga₀.₈₈As</td>
<td>1 μm</td>
<td></td>
</tr>
<tr>
<td>In₀.₄₉Ga₀.₅₁P</td>
<td>5000 Å</td>
<td></td>
</tr>
</tbody>
</table>

**S.I. GaAs substrate**

Fig. 1. Structural schematic of the modulator/detector array used in this study.
a mirror-like smooth surface can be obtained after the removal of the GaAs substrate. Wavy-finished In$_{0.45}$Ga$_{0.51}$P surfaces, preferentially along the [0 1 1] crystallographic direction, are observed for some etching solutions we tried. The wavy surface is caused by non-uniform etching of the thick GaAs substrate and the attack of In$_{0.45}$Ga$_{0.51}$P by the etchant. Therefore, a selective etchant which can etch GaAs uniformly is the key to the success of this approach. Fig. 2 shows the difference of two Nomarski contrast photographs of the In$_{0.45}$Ga$_{0.51}$P surface after the removal of GaAs substrate by two different H$_3$PO$_4$ based etchants. Strong contrast of the finishing surfaces between the non-uniform and uniform etching solution is clearly revealed. Using the right etchant, we are able to achieve

Fig. 3. Photocurrent spectra of a GaAs/AlGaAs MQW modulator without AR coating at several wavelengths under different reverse bias conditions, revealing quantum-confined Stark effect in the modulator.

Fig. 4. A finished 64 x 68 GaAs/AlGaAs MQW diode array flip-chip solder bump bonded to silicon CMOS with switching nodes.
a mirror-like In$_{0.49}$Ga$_{0.51}$P surface with a 2 in diameter after the removal of the GaAs substrate.

Another important question to be answered is whether the In$_{0.49}$Ga$_{0.51}$P has a detrimental effect on the performance of the GaAs/AlGaAs modulators. Fig. 3 shows the photocurrent spectra of a modulator without AR coating at several wavelengths under different reverse bias conditions. As the wavelength increases, the peak of the photocurrent at different wavelength shifts to higher bias voltage. This is an expression of the quantum-confined Stark effect in the GaAs/AlGaAs MQW modulator. At the desired system operation wavelength of 850 nm, a photocurrent change from 12.5 μA to 42 μA for a 0–8.5 V bias swing has been measured. This result is comparable to the best performance of an otherwise identical GaAs/AlGaAs MQW modulator grown with an Al$_{0.3}$Ga$_{0.7}$As selective etch stop. A photo of a finished GaAs/AlGaAs MQW diode array bonded to silicon CMOS with switching nodes is shown in Fig. 4. The uniformity of the epitaxial growth and fabrication process is demonstrated through the virtually identical photocurrents measured from the center to the edge of a 2″ wafer as shown in Fig. 5. We also perform a visual test of the 64 × 68 SEED array by forward biasing the modulators in the light emitting diode (LED) mode. All but 2 out of the 4352 MQW diodes generate light, which is attributed to the improved yield (100%) of the new In$_{0.49}$Ga$_{0.51}$P selective etch stop. Missing contacts to the p-type metal cause the imperfect yield of the array. Switching nodes have been tested at rates above 400 Mb/s per channel. System operation from 155 to 208 Mb/s has been demonstrated using this chip. With their high optical signal I/O count, these chips will be the backbone of a 256 × 256 asynchronous transfer mode (ATM) switching fabric system demonstration performed at 155 Mbit/s data rate per channel.

4. Conclusions

We report a new In$_{0.49}$Ga$_{0.51}$P selective etch stop which can achieve smooth surface after the GaAs substrate removal. The In$_{0.49}$Ga$_{0.51}$P selective etch stop not only has no detrimental effect on the performance of the GaAs/AlGaAs modulators, but also gives more uniform etching over a larger area. This new method improves the yield of GaAs/AlGaAs MQW diodes integrated with silicon CMOS via a flip-chip solder-bonding technique. For a 64 × 68 diode array, all but 2 out of 4352 diodes generate photocurrent in response to light, which is attributed to the improved yield (100%) of the new selective etch. Switching nodes have been tested at rates above 400 Mb/s per channel. System operation from 155 to 208 Mb/s has been demonstrated using this chip.

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References


Molecular beam epitaxy of strain-compensated InGaAs/GaAsP quantum-well intersubband photodetectors

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Abstract

We report on the growth of quantum well intersubband photodetectors (QWIPs) with InGaAs wells grown on GaAs. Arsenic species and substrate temperature were varied to investigate their effects on the photoluminescence (PL) and lattice relaxation of 40 period InGaAs/GaAs QWIP samples. Both PL intensity and lattice relaxation decreased with decreasing substrate temperature – the PL dropping steeply below 500°C and the relaxation dropping gradually below 470°C. In all cases examined, however, the relaxation was at least 33% as measured by asymmetric (1 1 5) five crystal X-ray scans. Introducing a small percentage of phosphorus into the barriers to partially strain-compensate the structure was found to dramatically reduce the lattice relaxation while maintaining the highest PL intensity. Comparison of this strain-compensated InGaAs/GaAsP QWIP with the optimized InGaAs/GaAs QWIP revealed that the dark current was reduced by a factor of 4 while the photocurrent remained virtually unchanged resulting in an increase in the background limited operating temperature by approximately 3 K.

PACS: 62.40. + i; 85.60.Dw; 85.60.Gz

Keywords: Strain compensation; QWIP; MBE; GaAsP; GaAs; InGaAs

Quantum well intersubband photodetectors (QWIPs) are attractive devices for imaging at mid- and long-infrared wavelengths, especially for large staring arrays. The mature III–V material technology used to manufacture these devices results in much lower costs, larger array sizes, higher pixel operability, and better uniformity than are achievable in competing technologies, such as HgCdTe. The excellent uniformity is especially important for large arrays since the array uniformity, rather than the individual detector performance, ultimately limits the minimum resolvable temperature of state of the art arrays [1].

Typically, the active region of a QWIP detector consists of multiple 20–60 Å quantum wells each separated by several hundred angstroms of a wider bandgap barrier material. The absorption process takes place between states located entirely within either the conduction or valence band rather than
between bands, which allows the use of relatively wide bandgap semiconductors, such as GaAs, to detect light at wavelengths greater than 3 µm. The most widely studied QWIPs use GaAs for the quantum wells and AlGaAs for the barriers. Although the near lattice match of these materials at any AlGaAs composition reduces the difficulty of the growth, there are limitations to these devices which can be addressed by using other materials, such as InGaAs.

First, because the detection wavelength is limited by the conduction band offset, the detection wavelength of GaAs/AlGaAs QWIPs with direct AlGaAs barriers is limited to greater than 5.6 µm [2]. Although QWIPs can be made with indirect barriers, they generally experience unusually high dark current relative to the detection wavelength of the device [2, 3]. Using a smaller bandgap material in place of the GaAs well will give a larger band offset and allow detection in the 3–5 µm mid-IR without the need for indirect AlGaAs barriers. Second, the use of deeper wells at longer wavelengths can remove or lower the Al content of the barriers, potentially improving the material quality and carrier transport in the barriers. Finally, several groups have demonstrated normal incidence absorption — supposedly a forbidden transition in n-type QWIPs—using InGaAs quantum wells [4, 5]. The ability to absorb light at normal incidence with an intersubband transition would reduce the processing complexity and eliminate a likely source of non-uniformity by eliminating the need for diffractive gratings in QWIP arrays [6]. Although, strictly speaking, only the later two points are applicable to the present work near 15 µm, it is anticipated that the results will applicable to future work in the 3–5 µm band as well.

The use of InGaAs wells, however, is complicated by the associated lattice mismatch. Even with only 10% In in the wells, Lenchshyn et al., saw clear evidence of dislocations arising from the lattice mismatch in 32 period QWIPs [7]. To address this problem of growing a large number of strained quantum wells, we first varied the growth conditions attempting to optimize the material quality.

We grew the multiple quantum well (MQW) region of several QWIP samples, Fig. 1, at substrate temperatures ranging from 430°C to 530°C to assess the effects of substrate temperature on photoluminescence (PL) and lattice relaxation. The top and bottom GaAs layers were grown at 600°C. As_2 was used as the arsenic species since a sample grown at 530°C with As_4 resulted in almost an order of magnitude reduction in the photoluminescence intensity. The substrate thermocouple was calibrated by measuring the spectral position of the bandedge absorption. The calibrated thermocouple was then used to control the substrate temperature during the growth.

After growth, the samples were characterized by room temperature photoluminescence from the quantum wells and asymmetric (1 1 5) 5 crystal X-ray scans to assess the material quality. As can be seen in Table 1, the PL intensity, PL linewidth (FWHM) and lattice relaxation all increase with increasing substrate temperature. The PL peak wavelength was relatively constant near 986 nm indicating no systematic changes in quantum well composition or thickness. The change in PL intensity is most drastic, increasing by a factor of 5 as the growth temperature changed from 470°C to 500°C.

![Table 1](image.png)

**Table 1** Photoluminescence and X-ray measurements on QWIP samples with MQW region grown at different substrate temperatures

<table>
<thead>
<tr>
<th>Temperature (°C)</th>
<th>PL Intensity (au)</th>
<th>FWHM (meV)</th>
<th>Wavelength (nm)</th>
<th>Relaxation (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>430</td>
<td>0.256</td>
<td>26.6</td>
<td>986.5</td>
<td>33</td>
</tr>
<tr>
<td>450</td>
<td>0.295</td>
<td>29.5</td>
<td>983.5</td>
<td>35</td>
</tr>
<tr>
<td>470</td>
<td>0.335</td>
<td>27.1</td>
<td>982.5</td>
<td>39</td>
</tr>
<tr>
<td>500</td>
<td>1.05</td>
<td>28.8</td>
<td>988.5</td>
<td>39</td>
</tr>
<tr>
<td>530</td>
<td>1.25</td>
<td>30.0</td>
<td>987.5</td>
<td>39</td>
</tr>
</tbody>
</table>
The relaxation, on the other hand, increased slightly with temperature at lower temperatures, but was constant for temperatures above 450°C. Because the samples grown at 500°C and 530°C had similarly high PL intensities, we believed the smaller linewidth of the sample grown at 500°C indicated slightly better material quality. We acknowledge, however, that the small variation in linewidth and inconsistent point at 450°C might indicate that the FWHM of these samples is limited by variations among the 40 quantum wells in the structure rather than the FWHM of the individual quantum wells. A single quantum well would not have the same degree of relaxation, so it is difficult to verify this by a separate growth. It is interesting to note that although the measured relaxation of all these samples was greater than 30%, cross-hatching was only barely visible under a Nomarski microscope.

Because of the high degree of relaxation evident in these samples, we attempted to improve the material quality through the use of strain-compensation [8]. Adding phosphorous to the GaAs barriers will decrease their lattice constant to compensate for the larger lattice constant of the InGaAs wells. This results in a reduction of the accumulated stress from the large number of quantum wells in the structure.

As has been demonstrated [9, 10], the amount of phosphorous incorporated into GaAs$_{1-x}$P$_x$ layers is dependent not only on the As to P ratio, but also, because of the preferential incorporation of As, on the As to Ga ratio as well. To determine the As and P fluxes necessary to produce a strain-compensated structure, we grew three samples of the superlattice in Fig. 2 with different As and P fluxes. The structure was designed to have the same ratio of barrier thickness to well thickness as the previous growths, but with a superlattice period small enough so that the 0-order superlattice peak in (0 0 4) X-ray scans would be the first peak to the left of the substrate peak, even in a structure without any phosphorous in the barriers. This allows the X-ray scan to unambiguously determine the 0-order peak and thus amount of phosphorous incorporated into the barriers and degree of strain compensation.

Since we did not want to adjust the As flux during the growth, the minimum As flux examined was still sufficient to maintain As stabilized growth for the InGaAs quantum wells at a V : III BEP ratio of approximately 14 : 1. The phosphorous was obtained by heating bulk InP chunks to 524°C–527°C in an As effusion cell. In order to minimize the phosphorous background, we tried to keep the phosphorous flux as small as possible while still strain compensating the structure. Fig. 3 illustrates the effects of changing the As and P fluxes to vary the degree of strain compensation as evidenced by (0 0 4) X-ray diffraction scans. As the tensile strain in the barrier layer is increased, the average superlattice lattice constant approaches that of the substrate. In the X-ray scans, this shows up as a decrease in the separation between the 0-order superlattice peak and the substrate peak. The results are summarized in Table 23. The final case represents a reduction of the average strain by 60% compared to the non-strain-compensated case, indicating 2.3% phosphorous incorporation in the barriers. A more direct measurement of the reduction of the accumulated stress in the samples can be seen in the reduction of wafer warp as measured by a Tropel Autosort also shown in the table.

Although the final condition tested was not completely strain-compensated, we still expected to see a dramatic reduction of the lattice relaxation in a 40 period QWIP. An additional sample of the structure in Fig. 1 was grown except that the central 500 Å of the barriers was GaAs$_{0.977}$P$_{0.023}$ instead of GaAs. The MQW region was grown at 470°C. The resulting PL FWHM was 29.2 meV and the PL intensity was 1.23 – comparable to the best values in the non-compensated case. The lattice relaxation, however, was reduced from 39% to 5%. With the incorporation of slightly more phosphorous into the barriers, we feel that the relaxation can be eliminated entirely.
Fig. 3. Five crystal (004) X-ray scans of the superlattice structure in Fig. 2 grown with different As and P fluxes showing varying degrees of strain-compensation. These scans correspond to the data listed in Table 2.

Table 2
Degree of strain compensation measured by the difference in average lattice constant of the superlattice from that of the substrate (Δa/a); the reduction of accumulated stress as more strain is compensated is also evident in the reduction of wafer warp measured by a Tropel Autosort

<table>
<thead>
<tr>
<th>BEP (Torr)</th>
<th>SL Δa/a (ppm)</th>
<th>Wafer warp (μm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>As</td>
<td>P</td>
<td></td>
</tr>
<tr>
<td>1 × 10⁻⁵</td>
<td>3 × 10⁻⁷</td>
<td>1020</td>
</tr>
<tr>
<td>8 × 10⁻⁶</td>
<td>4 × 10⁻⁷</td>
<td>750</td>
</tr>
<tr>
<td>5 × 10⁻⁶</td>
<td>4 × 10⁻⁷</td>
<td>480</td>
</tr>
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</table>

This reduction of lattice relaxation can be seen qualitatively in (0 0 4) X-ray scans as well. Fig. 4 shows that more superlattice peaks are visible for the strain compensated case and the FWHM of the peaks is reduced from 92 to 28 arcsec. This is very close to the calculated FWHM of the superlattice peaks, assuming no relaxation, of 25 arc-seconds. The (0 0 4) scan of the strain-compensated QWIP also provides quantitative verification of the small relaxation of the strained quantum wells. The two peaks very close together in the center of the scan correspond to the substrate and the top GaAs contact layer. Because of the slightly larger in-plane lattice constant resulting from the very slight relaxation of the multiple quantum wells, the top GaAs layer is under tensile strain and produces the peak on the right. The peak separation is exactly that expected for the measured 25 ppm increase in the in-plane lattice constant of the MQW region. The small linewidth of this strained GaAs peak indicates that excellent crystalline quality is still maintained.

To assess the effects of the relaxation reduction on device performance, we processed QWIP devices from this strain-compensated wafer and the non-compensated wafer grown at 500°C. The dark current at different device temperatures along with the 300 K background photocurrent of typical devices from these wafers are shown in Fig. 5. The photocurrent for these devices is nearly identical, yet the
Fig. 4. Five crystal (004) X-ray scans of (a) a non-strain-compensated QWIP and (b) the strain compensated QWIP.

Fig. 5. Dark current and background photocurrent of (a) a non-strain-compensated QWIP and (b) the strain-compensated QWIP.

dark current is reduced by a factor of 4, leading to an increase of about 3 K in the background limited operating temperature of the strain-compensated device. Comparison of the photocurrent spectra of these devices shows only a small decrease in the cutoff wavelength, from 13.9 to 13.7 μm, indicating that the dark current reduction is not just due to an increase in the barrier height resulting from the addition of phosphorous to the barriers.

In conclusion, we have demonstrated the use of strain compensation to reduce the lattice relaxation resulting from the accumulated stress of a large number of strained quantum wells. In 40 period QWIP structures, the relaxation was reduced from greater than 30% to only 5%. Because only partial strain-compensation was used in this sample, it is likely that the relaxation can be eliminated entirely by a slight increase in the phosphorous content of the barrier layers. The improved material quality resulted in a factor of 4 decrease in the dark current and 3 K increase in the background limited operating temperature of QWIPs processed from the strain-compensated material compared to QWIPs processed from the non-compensated material grown.

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References

MBE growth and characterization of doped multiple quantum well avalanche photodiodes


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Abstract

In the present study, how certain growth, processing and surface treatment techniques can be used to dramatically lower the dark current behavior of APDs by several orders of magnitude and improve the depletion characteristics of the built-in structures is shown. Through careful dopant calibration, the devices could be grown such that they were essentially fully depleted as grown, even though doping levels in the avalanche region were in the low $10^{18}$ cm$^{-3}$. The full depletion characteristics of the APDs were verified using theoretical modeling conducted for devices with similar structure and doping profiles. After processing the devices into mesa diodes, various surface passivation treatments were investigated. These include both O$_2$ plasma and sodium sulfide treatments. Through the application of such treatments, a decrease in the reverse bias dark current by as much as a factor of 1000 was achieved. Dark currents as low as 1 pA were obtained near zero bias. In some APDs, the dark currents increased to only 12 pA at 20% of breakdown. In addition, these devices exhibited extremely high gains, which exceeded 35000 in some APDs. The effect of the dopant profile in the avalanche region was also investigated. It was found that by the proper choice of dopant density and separation the breakdown voltage of the devices could be reduced by as much as 25%.

Keywords: MBE; MQW; APD; Dark current; Surface passivation

1. Introduction

Extensive studies have been carried out on multiple quantum well (MQW) structures because of their potential applications in avalanche photodiodes (APDs) and tunneling devices [1]. The MQW structure was first proposed by Chin et al. [2] and Capasso et al. [3] as a method to enhance the electron–hole ionization ratio ($k = \alpha/\beta$) beyond that typically found in bulk semiconductor materials. Later, Blauvelt [4] proposed the doped MQW structure in an attempt to further increase $k$ through the introduction of localized electric field regions throughout the depletion layer. The desire for high performance optical detectors has resulted in several proposed MQW APD structures using different material systems and growth techniques in an attempt to minimize the high levels of inherent dark currents and optimize the APD gain, noise, and bandwidth characteristics. In addition, the

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recent interest in the use of APDs in imaging systems has necessitated the development of devices with low levels of dark currents and high gains for low light applications. Because of the high frame rate and high pixel density requirements in new imaging systems, it is desirable to provide some front-end gain in the imager to allow operation under low light level conditions. In order to maintain low dark currents and high gains for such applications, methods of minimizing dark current are needed. In addition, the APD must be capable of operating with low power, and low noise.

In previous studies of doped well APD's the results have been clouded by the effects of incomplete depletion of the avalanching region. This incomplete depletion led to premature avalanche breakdown and a voltage dependent gain due to a increased number of active stages as the reverse bias on the structure was increased [5].

Not only have previous studies of these structures been hampered by incomplete depletion, but inadequate surface passivation has lead to large dark currents. A number of techniques have been previously proposed for the passivation of GaAs/AlGaAs device structures. These include wet and plasma surface treatments [6], hydrogen ion-passivation [7], Se [8] and S surface treatments [9, 10].

In the following sections, the effects of careful dopant calibrations, and surface passivation on the properties of doped well MQW avalanche photodiodes will be presented.

2. Experimental technique

The structures that were analyzed were volume- and delta-doped MQW APDs that were grown using molecular beam epitaxy (MBE) in a Varian Gen II system and were fabricated using standard photolithographic techniques. All samples in this study were grown in a solid source Varian Gen II MBE system using solid As$_4$ as the arsenic species. Samples were grown on $\frac{1}{4}$ of a 2 in n-type GaAs wafer that were briefly etched in 5:1:1 H$_2$SO$_4$ : H$_2$O : H$_2$O$_2$ before being indium mounted on a standard molybdenum sample holder. In order to minimize dopant diffusion and segregation, while still maintaining the growth of high quality AlGaAs the temperature was kept near 580°C. A V/III beam equavalent pressure ratio of 15 was employed for all the growth runs.

All APD devices consisted of a 1 $\mu$m Be-doped (3 x 10$^{18}$ cm$^{-3}$) p$^+$ top layer, and a 1 $\mu$m Si-doped (3 x 10$^{18}$ cm$^{-3}$) n$^-$ back contact layer. In the volume-doped MQW devices, high electric fields were achieved in the GaAs wells of the avalanche region through the introduction of 50 Å of highly doped (3.0 x 10$^{18}$ cm$^{-3}$) adjacent p$^+$ and n$^-$ layers which resulted in the avalanche region consisting of a series of closely compensated p-n junctions. In the delta-doped APDs, this was accomplished using delta-doped p$^+$ and n$^-$ layers with a sheet charge density of (1-6) x 10$^{12}$ cm$^{-2}$, separated by undoped spacer layers ranging from 50 to 150 Å. The MQW region in all structures consisted of 10 sets of alternating layers of GaAs (500 Å) and Al$_{0.42}$Ga$_{0.58}$As (500 Å) with 1000 Å periods.

Before the actual APD device structures were grown extensive dopant calibrations were carried out. These included not only the growth and measurement of traditional Hall effect samples, but also the growth and testing of special dopant calibration structures each of which consisted of just an avalanche region, thereby eliminating the effect of flux transients on the measurements. These structures were also measured by Hall effect. In this way the n- and p-type doping could be precisely balanced in the actual device structures, such that even though the dopant concentration in the depletion region were in the 10$^{18}$ cm$^{-3}$ range, they were precisely balanced to insure complete depletion.

Dark current measurements (I-V) on the MQW APD devices were conducted using a fully automated computer-controlled Keithley Source-Measure Unit (SMU) which connects to the device under test using a probe station or an 8-pin test fixture. Both dark and light I-V measurements were completed and the data was then used to calculate the multiplication gain of the device as a function of applied bias. Either electron- or hole-injections could be achieved by focusing the HeNe laser beam inside the p-contact ring at the top p- or the bottom n-layer. Capacitance-voltage measurements (C-V) were made using a computer-controlled HP4277A LCZ meter which can be interfaced with the same
test fixtures used in the current–voltage experiment. The $C–V$ data was then used to calculate the doping concentrations and the depletion width profiles of the structures. $C–V$ modeling was conducted using Silvaco’s TCAD and device simulation software. Theoretical $C–V$ solutions were obtained using small signal analysis at a frequency of 1 MHz and with a signal magnitude of 0.03 V, similar to the testing conditions used in the experimental measurements.

3. Results and discussion

In Fig. 1 is shown the $C–V$ results for volume and delta-doped MQW APDs with a built-in 1 μm MQW region and device areas of $1.72 \times 10^{-4}$ cm$^2$ and $3.13 \times 10^{-4}$ cm$^2$, respectively. The large zero-bias capacitance of the volume-doped APD is an indication of a small initial depletion width due to dopant imbalance in the avalanche region. The dopant calibration for this sample was carried out using only traditional Hall samples. On the other hand, the capacitance for the delta-doped device is more than 50% lower at zero applied bias and remains relatively constant with applied voltage, which indicates that the structure is fully depleted. The theoretical $C–V$ data support this conclusion and shows a very good agreement (to within 0.2 pF) with the measured data. The slight discrepancy between the experimental and theoretical results is due to parasitic capacitance effects and uncertainties in the actual active area of the APD structure. In Fig. 2 is shown the calculated depletion width versus apparent carrier concentration profile obtained using the $C–V$ data presented in Fig. 1. The zero-bias depletion width for the delta-doped device is about 1 μm (full depletion) compared to only

![Graph showing C-V plots for a delta-doped MQW APD with 1 μm depletion region and a device area of 3.13 x 10^-4 cm^2.](image-url)
0.25 μm for the volume-doped APD. This implies that the p- and n-doping concentrations throughout the delta-doped device are very closely matched resulting in a symmetric electric field profile and a uniform depletion throughout all 10 stages of the MQW structure. This conclusion is supported by the corresponding theoretical data shown in Fig. 2 and by other previously conducted studies [11].

Dark current is one of the main parameters of concern in photodetectors. Dark current analysis and the reduction of dark current are very important for high sensitivity and low noise applications. Not only do large dark currents lead to saturation of read-out electronics, but they also have been directly correlated to large low frequency noise and device reliability [12]. Defect centers, heterojunction interface traps, as well as mesa surface leakage can generate high levels of excess dark current and reduce minority-carrier lifetime [13].

In a typical PN junction, the overall dark current is the sum of the bulk and surface components. The bulk component is usually made up of diffusion, generation-recombination, and tunneling currents. The surface component consists of generation-recombination, and leakage shunt currents usually formed at semiconductor and dielectric interfaces [14]. In a MQW device structure with top p and n contacts such as the APDs used in our experiments, there are additional sources of dark current components. These are due to defect centers at the GaAs/AlGaAs interface and most importantly to surface leakage currents along the mesa edge which can contribute significantly to the dark current.

After processing the APD devices into mesa diodes, various surface passivation techniques were investigated. These include both plasma ashing in an O₂ plasma and ammonium sulfide treatments. Through the application of such treatments,
a decrease in the reverse bias dark current by as much as a factor of 1000 was achieved in the low bias region. This can be seen in Fig. 3 where the dark current of an APD is plotted both before and after surface treatment by ammonium sulfide. The rapid increase in the dark current at low reverse bias and the large drop in its value achieved by surface treatment are indicative of surface leakage. The leakage in these heterojunction mesa diodes was dominated by generation/recombination currents near the intersection of the mesa surface with the GaAs/AlGaAs depletion region.

As a result of surface treatment, dark currents as low as 1 pA were obtained near zero applied bias. In some APDs, the dark currents increased to only 12 pA at 20% of breakdown. In addition, these devices exhibited extremely high gains which exceeded 10000 as shown in Fig. 4. In most traditional APDs, the presence of high dark currents usually presents a limiting factor preventing the further increase in a device's photocurrent gain beyond the onset of avalanche breakdown. By reducing the dark current in such device, it becomes possible to maintain the dark contribution at levels well below that of the photocurrent. This makes it possible to achieve and sustain high levels of gains well beyond the onset of breakdown.

The effect of the separation between the p- and n-delta doping layers has also been investigated. As shown in Fig. 5 the distance separation between the delta doped layers has a significant effect on the reverse bias at which the onset of breakdown occurs. By decreasing the distance between the delta-doped layers in the avalanche region from 150 to 50 Å the breakdown voltage of the devices could be reduced from more than 28 V to less than 22 V, a decrease of almost 25%.

The separation between the delta-doped layers in these devices could not only be used to reduce the reverse bias breakdown, but also had a significant
Fig. 4. Gain versus voltage characteristics of a delta-doped MQW APD.

Fig. 5. Dependence of the reverse bias breakdown voltage on the dopant separation in delta-doped MQW APDs.

effect on the dark current in the devices. The effect of the dopant separation on the dark current density at 25% of the reverse bias breakdown is shown in Fig. 6, where the dark current density is shown for a set of delta-doped APDs as a function of the dopant separation. In this figure, it can be seen that by proper choice of the dopant separation the dark current can be reduced by more than a factor of two.
4. Summary

In summary, it has been shown that by careful MBE dopant calibration and the use of delta-doping, doped well MQW APD's can be grown that are in excellent agreement with theoretical predictions. Through the application of various surface passivation treatments dark current in these devices could be reduced by three orders of magnitude. This reduction in dark current allows the devices to be operated past the onset of break down, and thereby achieve gains greater than 10000. These low dark currents also allowed for the first time the investigation of the effects of the exact dopant profile on the device properties. It was observed that significant decreases in reverse bias break down and dark current could be achieved by the proper choice of the dopant separation in these delta-doped MQW APDs.

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Low dark current AlGaInAs/InP waveguide photodiodes using hybrid MBE and MOCVD growth


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Abstract

Dark current of 7 pA with one cleaved facet and 20 pA with an anti-reflection coating at the reverse bias voltage of 3 V was obtained for AlGaInAs/InP pin waveguide photodiodes, in which an AlGaInAs absorption layer and optical confinement layers were grown by optimized molecular beam epitaxy and an upper InP cladding layer was grown by metal-organic chemical vapor deposition. This is the lowest known dark current reported for long-wavelength pin waveguide photodiodes to date.

1. Introduction

Hybrid integrated optical modules [1] consisting of a laser diode (LD), photodiodes (PDs) and a silica-based planar waveguide circuit (PLC), have been proposed as candidates for a low-cost optical module for an access network. As for the photodiodes used in the module, side-illuminated waveguide PDs (WGPDs) are promising since they can be integrated on the PLC in the same way as the LDs integration. The long-wavelength WGPDs with high sensitivity, large optical tolerance and low dark current are suitable for mass production of the optical integrated modules. High sensitivity and large optical tolerance have recently been achieved by using a large multimode waveguide [2] and a thick absorption layer up to 3 μm thickness [3]. This was because the thick absorption layer and thick optical confinement layers of which thickness summed up to 5.5 μm made a large multimode core and reduced the mode-field mismatch between the mode of the incident light and that of the waveguide. Dark current had been, however, very large because it is difficult to grow high-quality thick epitaxial layers and to decrease the leakage current on the sides of high-mesa for the waveguide. This large dark current had been one of problems in the development of the WGPDs.

In order to solve this problem, we optimized molecular beam epitaxy (MBE)-growth conditions and utilized hybrid-growth technique of MBE and metal-organic chemical vapor deposition
(MOCVD). The resulting dark current of the WGPDs was 7 pA with one cleaved facet at a reverse bias voltage of 3 V. This remarkably reduced dark current indicated generation-recombination which subsequently dominated the leakage and the surface-leakage component was eliminated.

2. MBE growth conditions optimization

It is necessary for the WGPDs to grow thick AlGaInAs layers of which thickness sums up to more than 5 μm. When we grow such thick epitaxial layers using MBE, the small defects increase. These defects are typically $4 \times 1 \mu m^2$ and 20 nm high. These small defects were not observed when we grew the epitaxial layer less than 2 μm thick. These increasing defects are a problem in growing thick AlGaInAs layer, but there are not any previous reports to optimize the growth conditions and solve this problem. Hence, we need to determine the best condition for the growth of thick AlGaInAs. We optimized three conditions namely, growth rate, arsenic pressure (P(As)) and growth temperature ($T_g$) to suppress the small defects. The examined epitaxial layers were 3 μm thick AlGaInAs with a band-gap wavelength of 1.25 μm. The defect type and density were measured with Nomarski differential interference microscope, scanning electron microscope, and atomic force microscope. The composition of the examined layers was controlled within 50 arcssec in high-resolution X-ray diffraction and the $T_g$ was monitored by a pyrometer.

Figs. 1–3 show the defect densities as a function of the growth rate, the P(As), the $T_g$, respectively, where the value of P(As) of $1 \times 10^{-5}$ Torr was accorded with the V/III ratio of 10. The defect density decreases as the growth rate is decreased. With increasing As pressure, the defect density decreases and reaches a minimum at the P(As) of $1 \times 10^{-5}$ Torr and increases with the P(As) above $1.5 \times 10^{-5}$ Torr. With increasing $T_g$, we can see a similar tendency that there is a minimum point of the defect density at the $T_g$ of 460°C. At high growth rate, high As pressure and low temperature, a defect density increase is caused by a decrease in the migration of atoms impinging on the substrate.
and a consequent increase in the inhomogeneity of atoms of the group III. The defect increase in the low As pressure and in the high temperature is caused by the increase in indium segregation due to the lack and the desorption of Arsenic.

As a result, with a growth rate of 1.0 μm/h, a V/III ratio of 10 and a T_G of 460°C, we have successfully obtained high-crystalline quality of 3 μm thick AlGaInAs layers with a defect density of 1.5 x 10^4 cm⁻².

3. Design and fabrication of waveguide photodiodes

The waveguide photodiode consists of a p-GaInAsP contact layer, a p-InP clad layer, a p-AlGaInAs optical confinement layer, a 0.4 μm thick undoped AlGaInAs absorption layer with a bandgap wavelength of 1.4 μm, and an n-AlGaInAs optical confinement layer. All layers were lattice matched to the (1 0 0) InP substrate. AlGaInAs epitaxial layers were grown by the optimized MBE and the upper InP cladding and the GaInAsP contact layer were grown by MOCVD. This was because the InP layer proud to be of stable material that had less leakage current than the AlInAs layer used ordinarily in MBE. This was also because we could etch selectively the InP cladding layer and the AlGaInAs layer and then we could obtain the high-mesa structure by a conventional wet etching. The surface made by the wet etching has less dark current than that by a dry etching.

We made a mesa stripe structure of which height was more than 5 μm and buried with polyimide and silicon nitride, as shown in Fig. 4. The junctions were 30, 50 and 100 μm long by 19 μm wide. The cleaved facet was coated with silicon nitride in order to prevent reflecting.

4. Dark current measurements

Fig. 5 shows the dark current in the resulting waveguide PDs at the reverse voltage of 3 V as a function of the absorption-stripe length. The dark currents for devices with one cleaved facet were 7, 10 and 16 pA with the stripe length of 30, 50 and 100 μm, respectively. The dark current with AR coating increased slightly to 20, 24, and 33 pA, respectively. These small values are the lowest reported to date for waveguide photodiodes and are smaller than that of well-designed long-wavelength surface-illuminated pin photodiodes.

Fig. 6 shows an illustration of the sources of dark current. The dark current consists of diffusion current, generation current [4], tunneling current,
surface leakage currents on the sides of the waveguide and on the illuminated surface. The diffusion current is estimated to be a few pA and the tunneling current is negligible at this low bias voltage.

The dark current value at the stripe length of 0 indicates the leakage on the illuminated surface. Dark current density that originated from the leakage on the illuminated surface can be evaluated to be 0.78 pA/µm. The increase with the stripe length consists of the leakage current originating from the generation-recombination and mesa surface leakage components. The increase rate can be evaluated to be 0.19 pA/µm. These results with the small values indicate that the generation-recombination component dominates and the surface leakage component is eliminated.

5. Conclusions

We optimized the MBE-growth conditions and utilized the hybrid-growth technique of MBE and MOCVD. As a result, we obtained waveguide photodiodes with dark current as low as 7 pA with one cleaved facet, and 20 pA with an anti-reflection coating at a reverse bias voltage of 3 V.

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References

Improved electroabsorption properties in 1.3 μm MQW waveguide modulators by a modified doping profile

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Abstract

In 1.3 μm electroabsorption (EA) waveguide modulators used for microwave photonic links, we have used a modified doping profile to provide a better confinement of the depletion region in the p-i-n structure under reverse bias, without causing serious free carrier absorption. Theoretical simulation of the electric field distribution and experimental result on surface-normal photocurrent measurement, as well as waveguide measurement, are shown in this paper. An improvement of 29% of the quantum confined Stark effect (QCSE) is obtained, and in the waveguide measurement, 15% improvement of the transmission–voltage slope efficiency is achieved.

1. Introduction

Electroabsorption (EA) modulators based on quantum confined Stark effect (QCSE) in multiple quantum wells (MQWs) are among the most promising candidates for components in microwave photonic links because of their high efficiency, high speed, and low drive voltage. We have demonstrated, in our previous work, high-efficiency 1.3 μm EA waveguide modulators utilizing 11 periods of strain-compensated InAs0.4P0.6/Ga0.13In0.87P MQWs as the active region [1]. More recently, we achieved improved RF efficiency by an optimized structure with 6 MQWs [2]. To achieve a high RF link efficiency, a low waveguide insertion loss and a large slope efficiency in the transmission–voltage (T–V) curves are two of the most desirable characteristics of the modulators. The insertion loss consists of two parts, coupling loss between the waveguide and the fibers and the propagation loss inside the waveguide. In this paper, we report a modified doping profile which increases the slope efficiency while having a low insertion loss.

2. Structure design

An EA modulator is in principle a reverse biased p-i-n structure with the active region in the undoped i-region. The i-region in our structures contains 6 periods of InAs0.4P0.6/Ga0.13In0.87P strain-compensated MQWs with the well and
barrier thicknesses of 95 and 105 Å, respectively. The EA properties of such MQWs have been reported in detail previously [3]. The active MQW region is sandwiched between a p-doped and an n-doped passive In_{0.87}Ga_{0.13}As_{0.29}P_{0.71} waveguiding layers with a band-gap wavelength of 1.15 μm. A schematic of the structure is shown in Fig. 1a. This structure provides a vertical optical mode size of approximately 2 μm, comparable with the mode size in a lensed fiber, so that the coupling loss is reduced. Notice that the doping level in the p- and n-type waveguiding layers is only 5 × 10^{17} cm^{-3}, low enough to reduce the propagation loss caused by free-carrier absorption. However, the problem with the above-mentioned structure is that the depletion region extends from the i-region into the guiding layers under reverse bias because of the relatively low doping level in these guiding layers and relatively thin i-region (thickness d_i = 0.13 μm). The wider depletion region causes a decrease in the electric field (E-field) in the MQWs, and thus the QCSE, at a given bias voltage V_m, resulting in a lower T–V slope efficiency. Our solution to this problem is to insert a highly doped (2 × 10^{18} cm^{-3}) and thin (200 Å) p^+ (n^+) In_{0.87}Ga_{0.13}As_{0.29}P_{0.71} layer (pulse-doped layer) in the p (n) guiding layer right on the side of the i-region as shown in Fig. 1b. Since the two pulse-doped layers are very thin compared to the waveguide thickness (~1.7 μm), they do not cause serious free carrier absorption.

Fig. 2 is a computer simulation using Simwin 3.2, a Poisson–Schrodinger solver, of the E-field distribution in the structures with and without pulse doping (PD) under a V_a of 3 V. The background doping in the i-region is assumed n-type, 8 × 10^{15} cm^{-3}, the same as that in our undoped InP epilayer. The built-in potential V_{bi} is 1 V from the simulation. It can be seen that the depletion region extends into the p and n regions in the conventional structure whereas PD provides a better confinement of depletion region within the i-region. Better confinement means that the depletion region is thinner under the same V_a, so a larger E-field (F) can be obtained in the MQWs. Fig. 3 shows three F–V curves from the simulation. Curve (a) is the case that the confinement is perfect, that is, there is no extension of depletion region into

![Diagram](image-url)

Fig. 1. Layer structures of 1.3 μm waveguide modulators with (a) conventional and (b) modified doping profiles.
3. Experimental procedure

The MQW samples were grown in a Varian GEN-II MBE system modified for gas-source MBE growth. Thermally decomposed AsH\textsubscript{3} and PH\textsubscript{3} were used as the group V sources, and elemental In and Ga as the group III sources. Elemental Si and Be were used as n- and p-type dopants, respectively. The growth temperature was 460°C, and the InP growth rate was typically 1 μm/h. The growth rate was determined by intensity oscillations of reflection high-energy electron diffraction (RHEED). The samples were grown on S-doped n\textsuperscript{+}-InP substrates.

For surface-normal photocurrent measurement, mesas with a 500 μm diameter were defined by wet chemical etching on the sample surface. A ring-shaped AuZn/Cr contact was evaporated onto the p\textsuperscript{+}-In\textsubscript{0.53}Ga\textsubscript{0.47}As cap layer. In was used as the n-type contact on the back side of the substrate. The breakdown voltage of the ring diodes is typically 6 V at 2 μA leakage current. The surface-normal photocurrent measurement was performed at room temperature. A beam of monochromatic light was focused onto the center window of the ring diode, and the photocurrent was amplified with a lock-in amplifier. The reverse electric bias was applied via the ring-shaped electrode and the back contact.

Waveguide modulators were fabricated using CCl\textsubscript{2}F\textsubscript{2} reactive ion beam etching. The 2 μm ridge height confines the waveguide mode laterally. The devices are 3 μm in width and 190–230 μm in length. Tapered single-mode fibers were used to couple TE polarized light into and out of the modulator. The measurement was carried out at 1.32 μm wavelength, approximately 27 meV detuned from the zero-bias exciton peak.

4. Results and discussion

For comparison, two waveguide modulator samples, #834 and #922 were grown. The structures are shown in Fig. 1a and Fig. 1b, respectively. #922 has the above-mentioned pulse-doped layers while #834 does not. In Fig. 4 we plot the QCSE energy shift of the n = 1 heavy hole exciton peak in
samples #834 and #922 against the reverse bias. The solid line is a theoretical calculation based on the effective well width model [4], assuming that the depletion region width equals the i-region width, $d_i$, and the E-field is uniform across the depletion region and is zero elsewhere. This situation is the ideal case where the confinement of the depletion region is perfect. From Fig. 4, one can see clearly that #922 has a much larger QCSE than #834 under a given bias. At 2 V reverse bias, the difference is as large as 29%. This result indicates that the pulse-doped layers provide a better confinement of the depletion region and therefore increase the efficiency of the bias voltage, as predicted by the simulation shown in Fig. 3. By comparing the experimental QCSE with the calculation value in Fig. 4, we obtained the E-field at each $V_r$. These E-field values are plotted in Fig. 3 as scattered symbols, together with the simulation result. The experimental data agree with the simulation very well. The E-field in #922 is approximately 20% larger than that in #834, and therefore it is expected to achieve a 20% improvement in the $T-V$ slope efficiency from waveguide measurement.

Fig. 5 shows the electrical-to-optical transfer curves of waveguide modulator devices fabricated on #834 and #922. The slope efficiency is obtained by taking the first derivative of these curves. The highest slope efficiency of #922 is 1.34 V$^{-1}$, 15% higher than 1.17 V$^{-1}$ of #834. Since the device of #922 is 190 μm in length, 35 μm shorter than #834 due to the fluctuation in cleaving, the contrast ratio of the former is a little smaller than that of the latter as seen in Fig. 4. If #922 had the same device length as #834, the slope efficiency difference would be even higher than 15%, which is in good agreement with the expected value, 20%. #922 exhibits not only a higher slope efficiency but also a lower bias voltage, 0.8 V, to achieve the peak slope efficiency, compared with 1.1 V in #834. From the result shown above, a higher RF link efficiency is expected in #922.

5. Conclusions

We have demonstrated an improved performance of waveguide modulators used for microwave photonic links by employing a modified doping profile. The new profile includes two pulse-doped layers on each side of the i-region, which provides a better confinement of the depletion region and increases the efficiency of the reverse bias without causing serious free carrier absorption. Twenty-nine percent improvement in QCSE and 15% improvement in $T-V$ slope efficiency are observed in the pulse-doped sample. Other device performance is still being investigated.
Acknowledgements

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Investigation of Si-doped p-type AlGaAs/GaAs, AlGaAs/InGaAs quantum well infrared photodetectors and multiquantum wells grown on (3 1 1)A GaAs

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Abstract

We have studied two Si-doped p-type quantum well infrared photodetectors (QWIPs) using AlGaAs/GaAs and AlGaAs/InGaAs grown on (3 1 1)A GaAs. The Si-doped AlGaAs/GaAs p-QWIP exhibits a symmetrical dark I–V characteristic at all the measured temperatures from 40 to 120 K. The strained p-type AlGaAs/InGaAs QWIP exhibits a slightly asymmetrical dark I–V characteristic, but is markedly less asymmetrical than that doped with beryllium. The slight asymmetry in dark I–V characteristic and the large blue-shift in responsivity spectra may be due to the thickness modulation observed from TEM and the red-shift of PL peak energy, where PL peak energies from (3 1 1)A AlGaAs/InGaAs multiple quantum wells are red-shifts of 7 and 22 meV to the side-by-side grown (1 0 0).

1. Introduction

P-type quantum well infrared photodetectors (QWIPs) have attracted much attention recently because of the inherent capability of normal incidence radiation and even enhanced intersubband transition [1, 2]. To date, most of the p-type QWIPs reported so far have used either beryllium (Be) or carbon (C) for the dopant species. However, both dopants have serious shortcomings. Be suffers from the problem of fast out-diffusion [3] and asymmetrical I–V characteristics measured in p-QWIP [2, 4] while C is not compatible to the normal solid source MBE. Because high p-type concentration has been reported by Si-doped GaAs grown in (3 1 1)A orientation [5], we have studied the Si-doped p-type QWIPs in this orientation. Other advantage of the growth of AlGaAs/GaAs or AlGaAs/InGaAs p-QWIP on (3 1 1)A GaAs is due to the reduced incorporation of deep recombination centers at the quantum well interfaces [6], which may in turn increase the quantum efficiency and improve the device performance. Further reduction of deep recombination centers can be expected in (3 1 1)A orientation because of the very-low diffusion coefficient of Si, and a higher growth
temperature can be used to grow the AlGaAs/GaAs multiple quantum wells (MQWs) in p-QWIP. It is well known that a high growth temperature can reduce the oxygen and growth-induced defect [7], while a higher efficiency of photoluminescence (PL) can be obtained.

In this paper, we report the first study of Si-doped p-QWIP grown on (3 1 1)A GaAs. Very symmetrical dark I–V characteristic was obtained in AlGaAs/GaAs p-QWIP, which demonstrates the low diffusion of Si impurity as compared to Be. The strained Si-doped AlGaAs/InGaAs p-QWIP shows a slightly asymmetrical dark I–V characteristic, but is markedly less than that of the same structure by Be-doping.

2. Experimental procedure

The layer structure of AlGaAs/GaAs p-QWIP consists of a 1.0 μm Si-doped GaAs bottom contact layer, 40 periods of 2.8 nm GaAs well with 30 nm Al0.25Ga0.75As barrier, and a 1.0 μm Si-doped GaAs top contact layer. The quantum wells and contact layers are doped with Si to a concentration of 3 x 10¹⁸ cm⁻³. The layer structure for strained AlGaAs/InGaAs p-QWIP is similar to the unstrained AlGaAs/GaAs, except that the quantum well region consists of 20 periods of 5 nm In0.12Ga0.88As well with 50 nm Al0.4Ga0.6As barrier. The growth temperatures were 600 and 550°C for AlGaAs/GaAs and strained AlGaAs/InGaAs p-QWIPs, respectively. A low V/III beam equivalent pressure (BEP) of ~ 10 is used to achieve high p-type doping. We have also grown AlGaAs/GaAs and AlGaAs/InGaAs multiple quantum wells (MQWs) in order to further characterize the quality of quantum well in (3 1 1)A orientation. The AlGaAs/GaAs MQWs consists of a 0.1 μm Al0.25Ga0.75As buffer, 5 periods of 5 nm GaAs well with 20 nm Al0.25Ga0.75As barrier, and a 0.1 μm Al0.25Ga0.75As top layer. The strained AlGaAs/InGaAs MQWs consists of a 0.1 μm Al0.4Ga0.6As buffer, 5 periods of 10 nm In0.12Ga0.88As well with 20 nm Al0.1Ga0.9As barrier, and a 0.1 μm Al0.1Ga0.9As top layer. Photoluminescence (PL) was used to characterize the optical properties of the MQWs. Hall measurement was used to measure the p-type doping for Si-doped (3 1 1)A GaAs. In order to measure the I–V characteristic and the responsivity, the p-QWIPs were processed into 216 x 216 μm² mesas.

3. Results and discussion

Fig. 1 shows the PL spectra for (3 1 1)A AlGaAs/GaAs MQWs grown at 600°C. A (1 0 0) AlGaAs/GaAs MQWs, grown at 550°C, is also shown for comparison. It is well known that significant Be out-diffusion occurs at a growth temperature of 600°C, and for that reason we have chosen the growth temperature of 550°C in (1 0 0) orientation for comparison. PL full-width-at-half-maximum (FWHM) line width of 9.2 and 11.2 meV are measured from an undoped AlGaAs/GaAs MQWs grown on (1 0 0) and (3 1 1)A GaAs, respectively, which indicates sharp interfaces can be obtained in both directions. However, one order of magnitude of PL intensity enhancement was measured from the 600°C grown (3 1 1)A MQWs than that grown on (1 0 0) at 550°C, which suggests better material quality can be achieved in (3 1 1)A direction. The increased PL intensity may be due to the

![Fig. 1. Low temperature (15 K) PL spectra of AlGaAs/GaAs MQWs grown on (3 1 1)A and (1 0 0) GaAs at 600°C and 550°C, respectively.](image-url)
improved material quality at a 50°C higher growth temperature or by the inherent lower interface recombination in (3 1 1)A orientation to (1 0 0) [6].

Fig. 2 shows the dark I–V characteristics of AlGaAs/GaAs p-QWIP grown at 600°C. Very symmetrical dark I–V characteristics are observed in both the forward and the reverse bias regions over the entire temperature range from 40 to 120 K, which is markedly improved when compared to Be-doped p-QWIP reported by Levine et al. [1]. The symmetrical I–V characteristic indicates the uniform doping within the quantum wells and the negligible impurity out-diffusion during growth. Similar low out-diffusion of Si impurity is also reported in a heterojunction bipolar transistor by Li et al. [8] even at a high growth temperature of 670°C. It is noticed that the dark current in the same order to similar structure by Levine et al. [1] of a 4 × 10¹⁸ cm⁻³ Be-doped AlGaAs/GaAs p-QWIP with 3 nm well separated by 30 nm Al₀.₅Ga₀.₅As, which indicates the excellent material quality of p-QWIP grown in (3 1 1)A direction. The measured hole mobility at a hole concentration of 3 × 10¹⁸ cm⁻³ is 75 cm²/(V s), which is comparable to the mobility value of ~ 100 cm²/(V s) doped by Be [5]. The slightly lower mobility is believed to be due to the compensation effect from the higher V/III BEP ratio. However, a higher As flux is required to grow the AlGaAs and InGaAs layers.

To further reduce the dark current, we have studied the strained AlGaAs/InGaAs p-QWIP with the same device size. Fig. 3 shows the dark I–V characteristic of the AlGaAs/InGaAs p-QWIP grown at 550°C. A low growth temperature of 550°C is chosen to reduce the In re-evaporation from the InGaAs layer. In spite of the expected reduction of dark current, a slight asymmetry was observed under forward and reverse biased conditions. However, the asymmetry is markedly less than that of the Be-doped QWIP. It is unlikely that the asymmetry is due to Si out-diffusion, because the growth temperature is lower than that of AlGaAs/GaAs p-QWIP. Furthermore, in sharp contrast to the Be-doped p-QWIP, the dark current asymmetry is opposite to that from doping migration effect [1].

Fig. 4 shows the measured photo-responsivity spectra of strained AlGaAs/InGaAs p-QWIP doped by Si. A strong absorption peak at 3.7 µm is observed, with a shoulder around 3.0 µm and a small absorption at 5.7 µm. The primary absorption peak at 3.7 µm is much shifted from the designed absorption peak at 10 µm, which may be due to either the mis-estimated parameters in
(311)A orientation or the formation quantum wires reported by Notzel et al. [9].

We have studied the AlGaAs/InGaAs MQWs to further understand the unusual $I-V$ asymmetry and the blue-shift of absorption wavelength. Fig. 5 shows the PL spectra of the strained AlGaAs/InGaAs MQWs grown on (311)A and (100) GaAs at 550°C. Although strong PL intensity was also observed for AlGaAs/InGaAs MQWs grown on (311)A GaAs, multiple PL transitions were measured with peak energy red-shifts of 7 and 22 meV to that of side-by-side grown (100) GaAs. We have used cross-sectional transmission electron microscopy (TEM) to study this effect. The PL energy red-shift is due to the growth-induced thickness modulation of quantum wells, which form quasi quantum wires in (311)A orientation [9]. More detailed study of TEM and the formation of quantum wires will be published elsewhere. Therefore the slight asymmetry in dark $I-V$ characteristic of AlGaAs/InGaAs p-QWIP may be attributed to the thickness modulation. The large blue-shift of peak absorption in responsivity spectra may be also due to the formation of wider quantum wells as observed from TEM and PL.

4. Conclusions

We have demonstrated that the Si can be used as an alternative dopant for p-type QWIP. The highly symmetrical dark $I-V$ characteristic from the AlGaAs/GaAs p-QWIP suggests the negligible dopant migration of Si impurity during growth. The strained p-type AlGaAs/InGaAs QWIP exhibits a lower dark current and a slightly asymmetrical dark current characteristic, but the asymmetry is markedly less than that doped by Be. The asymmetry in the strained p-QWIP is not due to the impurity out-diffusion, because the dark current asymmetry is opposite to that from doping migration. The slight asymmetry in dark $I-V$ characteristic and the large blue-shift in responsivity spectra may be due to the thickness modulation observed from the red-shift of PL peak energy and TEM.

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References


808 nm high-power laser grown by MBE through the control of Be diffusion and use of superlattice

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Abstract

808 nm high-power laser diodes are grown by MBE. In the laser structure, the combination of Si-doped GRIN (graded-index) region adjacent to n-AlGaAs cladding layer with reduced Be doping concentration near the active region has been used to diminish Be diffusion and oxygen incorporation. As compared with the laser structure which has undoped GRIN region and uniform doping concentration for Si and Be, respectively, in the cladding layers, the slope efficiency has increased by about 8%. Typical threshold current density of 300 A/cm² and the minimum threshold current density of 220 A/cm² for lasers with 500 µm cavity length are obtained. A high slope efficiency of 1.3 W/A for coated lasers with 1000 µm cavity length is also demonstrated. Recorded CW output power at room temperature has reached 2.3 W.

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Keywords: High-power; Semiconductor laser; MBE; Quantum well

1. Introduction

High-power laser diodes for the light source of solid-state laser pumping have been extensively investigated [1, 2]. To realize high-power operation, it is needed to achieve a lower operating current and higher quantum efficiency. The performance characteristics of the laser diodes are determined by the quality of the epitaxial layer, laser structure design and the control of growth condition. Beryllium (Be) diffusion plays an important role in the performance of GaAs/AlGaAs laser diodes grown by MBE. Both increased oxygen incorporation and p-n junction displacement are related to Be diffusion [3]. In addition, the Be is believed to be responsible for the failure of laser diode [4]. Although MBE technique can provide better reproducible control over composition, thickness and doping profile in the direction of growth, careful treatments are needed for Be doping. In this paper, we report the MBE growth of GRIN-SCH SQW
808 nm high-power laser diodes through the control of Be diffusion. The application of the superlattice buffer is also included.

2. Laser structure and epitaxial growth

The MBE machine used in this experiment is a Riber 32p system. Because the high-power laser diodes operate under crude conditions, it is required to improve the quality of the laser materials. It is believed that the growth of high-quality laser structure by MBE requires a more sophisticated method than that for MOCVD-grown lasers [5]. In our laser structure, superlattices in GaAs buffer and n-AlGaAs cladding layers were incorporated. The superlattice can bring some benefits such as the reduction in the propagation of defects from substrate, a relief of strain at the interface between high aluminum concentration cladding layer and the substrate [5]. The superlattice also offers additional defense against oxygen incorporation, makes the epitaxial layer flat and improves the quality of the lower interface of the QW [6]. The advantage of superlattice may improve the uniformity of the laser materials which is beneficial to the high-power or broad-area laser diodes. Low aluminum composition AlGaAs layer is introduced as a getter buffer to remove the transient oxygen because impurities including oxygen are released as the Mo holder is heated to higher temperatures.

Fig. 1 is the schematic diagram of GRIN-SCH AlGaAs SQW laser composition profiles. The laser was grown by MBE on an n+-GaAs substrate. The growth sequence is described as follows: a 1 µm thick n-GaAs buffer layer containing a five-period n-GaAs/Al_{0.6}Ga_{0.4}As (10 nm/10 nm) superlattice buffer layer, 0.5 µm thick n-Al_{0.1}Ga_{0.9}As layer, 1.3 µm thick n-Al_{0.6}Ga_{0.4}As cladding layer including a five-period GaAs/Al_{0.6}Ga_{0.4}As (10 nm/10 nm) superlattice; 0.2 µm thick slightly Si doped Al_{x}Ga_{1-x}As confining layer (linearly graded from x = 0.6 to 0.3), undoped 10 nm-thick Al_{x}Ga_{1-x}As active layer, undoped 0.2 µm thick Al_{0.6}Ga_{0.4}As as confining layer (linearly graded from x = 0.3 to 0.6), 1.3 µm thick p-Al_{0.6}Ga_{0.4}As cladding layer, and 0.3 µm thick p'-GaAs contact layer. Silicon was used as the n-type and beryllium as the p-type dopant. Growth rate was 1.2 µm/h for Al_{0.6}Ga_{0.4}As.

Beryllium has a high diffusivity in AlGaAs/GaAs structure [7]. The unstable behavior of this dopant during crystal growth, as well as during operation of laser diodes make the control of p-n junction difficult and affect the stability of lasers [8, 4]. On the other hand, Naresh Chandra et al. [3] reported that compared to undoped AlGaAs layer, the oxygen content is lower in the Si-doped AlGaAs layer and higher in the Be-doped one. Oxygen which forms nonradiative recombination centers degrades the performance and reliability of the laser diodes. The above research results indicate that beryllium doping and diffusion have important effects on the performance of laser diodes. Experimental analysis [3] also shows that the diffusion of Be can be retarded by increasing the doping of n-layer and doping the n-side of the GRIN region. Considering the Be, Si and undoping-related oxygen incorporation, it is reasonable to dope the GRIN region adjacent to the n-AlGaAs cladding layer with Si in order to reduce oxygen content in the active region and avoid the p-n junction displacement (a separation of p-n junction from the active layer heterojunction).

We adopted tailoring doping profiles in the epitaxial growth. In the laser structure (type I) as shown in Fig. 1, the Si doping concentration in the GaAs buffer and n-AlGaAs cladding layer (1) are $2 \times 10^{16}$ and $1 \times 10^{19}$ cm$^{-3}$, respectively. The GRIN region (2) adjacent to the n-AlGaAs cladding layer was doped with Si to $10^{16}$ cm$^{-3}$. GRIN region (3) was undoped. The low Be-doped region
structures, respectively. Enhancement in the slope efficiency is obtained by about 8% in type I as compared to type II. It was reported that doping modifications had no measurable effect on external quantum efficiency for a structure in which the optical field is tightly confined [9] because there were less changes in the free-carrier absorption. If this is true, for our laser structure (type I) in which the optical field is tightly confined [9], we consider the improvement of slope efficiency as the result of decreased concentration of nonradiative recombination center or the minimizing p–n junction displacement, or both, in type-I structure. The displacement allows majority carrier current flow out of the active layer. The current does not contribute to lasing [14].

Second, dielectric (SiO₂) defined metal-stripe lasers with 100, 200 μm wide stripes and 1000 μm cavity length were also fabricated for type-I structure. High- and low-reflectivity facet coatings were applied to each facet using SiO₂ and Si, which gives the reflectivity of 10% and 90% at the front and rear facets, respectively. Zn diffusion was performed to reduce the contact resistance. Ti/Pt/Au was used for the p-side contact, the n-side contact was formed with AuGe/Ni/Au. Laser chips were mounted on Cu heat sinks with In solder in the junction down configuration.

Typical slope efficiency of these lasers is 1 W/A. Output power versus current characteristics at room temperature under CW operation are shown in Fig. 2. These lasers are different in stripe width and same in the cavity length (1000 μm). Curve (a) represents one of coated lasers with 100 μm stripe width. Curve (b) represents one of coated-laser with 200 μm stripe width. In curve (a), laser can be operated up to 2 W. In curve (b), a high slope efficiency of 1.3 W/A for the coated-front facet alone is demonstrated, recorded CW output power reached 2.3 W. The typical emission wavelength of 100 μm wide stripe laser diodes under 1 W output power was measured to be 808 nm, as shown in Fig. 3.

The temperature dependence of CW light output versus current characteristics for 100 μm stripe laser is depicted in Fig. 4. The temperature dependence of Iₘ₉ is given by

\[ Iₘ₉ = I₀ \exp \left( \frac{T}{T₀} \right) \]
Fig. 2. Light output power vs. current characteristics of the lasers with 100 and 200 μm stripe width.

Fig. 3. Typical emission spectrum for the 100 μm x 1000 μm laser devices under output power. W at room temperature.

Fig. 4. Light output power vs. current characteristics of the high-power laser at different temperatures under CW conditions.

of the laser structure. Because shorter wavelength AlGaAs/GaAs lasers tend to have a small $T_0$ [12], this value is relatively high.

In summary, high-power 808 nm-AlGaAs quantum-well lasers are grown by MBE. Both reduced Be doping concentration near the active region and the Si doping in the n-side GRIN region are used to reduce the Be diffusion and oxygen incorporation near the active region. Compared with nontailoring-doping laser structure, the enhancement of the slope efficiency is observed. The minimum threshold-current density of 220 A/cm² is obtained. High slope efficiency of 1.3 W/A is demonstrated. Recorded CW output power at room temperature has reached 2.3 W.

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References

Continuously graded buffers for InGaAs/GaAs structures grown on GaAs

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Abstract

We report on the preparation under optimized conditions and on the study of InGaAs buffers grown on GaAs, intended for MQW structures for 1.3 and 1.5 μm optical operation at 300 K; the buffers have linear, square-root and parabolic composition profiles. They were designed so that MQWs grown atop the buffers are virtually unstrained, unlike those prepared following the conventional approach that are under compressive strain. The results obtained by the concomitant use of TEM, HRXRD, AFM and PL show that, by carefully designing the buffers: (i) the misfit dislocation (MD) profiles and thicknesses of the MD-free regions in the buffers can be predetermined, (ii) active structures atop the buffers are virtually unstrained and have efficient 300 K photoluminescence in the 1.3 and 1.5 μm windows of photonic interest, (iii) the structures have threading dislocation concentrations in the low 10⁴ cm⁻² range and show smooth and symmetric cross-hatchings.

1. Introduction

Lattice mismatched structures have interesting applications in spite of the potentially adverse effects related to strain relaxation, since they allow to extend the range of materials that can be considered and, then, of the properties that can be exploited. Moreover, the accurate control of the resulting strain may give significant advantages, since it can be used as a tool for band engineering. Furthermore, it is of interest to use substrates that, even if mismatched: (i) have higher crystalline perfection and definite electrical, optical and mechanical properties, or (ii) allow the preparation of structures suited for the fabrication of different devices with integrated functions. Structures consisting of InGaAs layers deposited on GaAs substrates are gaining increasing attention for optical modulators based on the quantum-confined Stark effect [1], quantum well infrared detectors [2] and metamorphic high electron mobility transistors [3], 1.3 μm quantum well lasers [4] and non-linear optical devices [5].
Buffer layers with either step-graded or continuously graded compositions have been proposed to (i) relax the strain resulting from lattice mismatch; (ii) bury the misfit dislocations (MD) away from the active part of the structure; (iii) prevent the onset of three-dimensional nucleation; and (iv) reduce the propagation of threading dislocations (TD) towards the active parts of devices [6]. Linearly graded buffers are generally used (with the exception of Refs. [7, 8]), however, we expect that continuously graded buffers with selected composition profiles may give additional benefits, such as the control of the MD profiles within the buffers.

The aim of this work was to evaluate buffers with compositions graded according to (a) linear, (b) square-root and (c) parabolic (with the vertex at the top of the buffer) dependences on the distance from substrates. The experimental results were obtained by the concomitant use of transmission electron microscopy (TEM), high-resolution X-ray diffraction (HRXRD), atomic force microscopy (AFM) and photoluminescence (PL). The results show that by proper buffer designing: (i) the thicknesses of the MD-free regions of buffers can be varied, (ii) the MDs can be concentrated far from the active structures grown atop the buffers, (iii) these structures are virtually unstrained, (iv) the TD concentrations can be reduced in the low $10^6$ cm$^{-2}$ range, (v) the structures grown under optimized conditions have smooth and symmetric cross-hatchings and (vi) the MQWs have efficient 300 K photoluminescence in the 1.3 and 1.5 µm spectral windows.

2. Experimental procedure

After careful studies on the optimization of growth conditions, the InGaAs buffers with different composition profiles were grown on semi-insulating (0 0 1) GaAs by solid-source MBE at 400°C using As$_2$ beams; instead, the InGaAs/GaAs MQWs were grown at 400°C by atomic layer MBE (ALMBE) that allows to select the compositions of the InGaAs wells by only adjusting the In and Ga supply times, without changing the temperature of the cation cells reached at the end of the growth of buffers [9]. Composition grading was obtained by varying the In and Ga cell temperatures in such a way to maintain a growth rate of 0.28 nm/s. Prior to each run, the In and the Ga fluxes were calibrated by RHEED oscillation measurements using InAs and GaAs substrates, respectively. The substrates were radiatively heated at growth temperatures $T_g$ measured by a suitable optical pyrometer for $T_g \geq 450°C$ and by a thermocouple (TC) for $T_g < 450°C$. For $T_g < 450°C$, the TC readings were corrected by the difference ($\sim 200°C$) between the TC and the optical pyrometer values measured at $T_g > 450°C$. Great care was paid to keep the growing surface temperature at the intended value during growth; this was done by lowering the electrical power to the substrate heater in a controlled way so as to counterbalance the increasing optical power absorbed during the growth of buffers and MQWs. The As$_2$ beam-equivalent pressure (BEP) was $5.5 \times 10^{-6}$ Torr, while the As$_2$/Ga BEP ratio was $\sim 9$.

TEM observations were carried out by an JEOL 2000FX instrument operating at 200 keV on (1 1 0) oriented cross sectional samples. The dislocation distribution at the buffer/substrate interfaces and across the buffer layers was studied under bright field, dark field and weak beam conditions using $g = 220$ type scattering vectors. Conventional $g \cdot b$ contrast analyses showed that the majority of the dislocations were 60° type in character. Dark field $g = 200$ type imaging conditions were used for studying the MQW structural quality and thicknesses. Finally, the MQW periods and interface-smoothness were studied in the lattice resolution mode.

High-resolution X-ray diffraction (HRXRD) measurements were performed using a diffractometer fitted with a four-crystal X-ray monochromator. (0 0 4) symmetrical and (3 3 5) asymmetrical CuK$_\alpha$ diffraction profiles were used to measure the strain and tilt angle of the MQWs; from these curves we also obtained the average lattice parameters of MQWs parallel and perpendicular to the interfaces, the compositions of the top of buffers and the average compositions and the period of MQWs. A good agreement between MQW periods determined by TEM and HRXRD was found, while the mean compositions of MQWs deduced by HRXRD are close to those determined
by Rutherford back-scattering experiments in selected samples.

Atomic force microscopy (AFM) was carried out under conventional conditions.

The photoluminescence (PL) spectra were taken at 10 and at 300 K with a spectral resolution of 0.5 meV, and were excited by the blue line of an Ar$^+$ laser, using power densities as small as 0.5 W/cm$^2$.

3. Design of the structures

According to the existing models on strain relaxation [6, 8, 10, 11] buffers with increasing compositions can be divided into two parts: (i) the one closest to the substrate (up to a distance $\delta_c$ from the substrate itself) is fully relaxed, due to the formation of misfit dislocations that have a concentration proportional to the composition gradient, and (ii) the topmost part that is void of misfit dislocations, (b) has a lattice parameter $a(x_c)$, where $x_r = x(\delta_c)$ ($x(\delta)$ giving the dependence of the In$_x$Ga$_{1-x}$As composition on the distance $\delta$ from the substrate and $a(x)$ being the lattice parameter as a function of $x$), and (c) is under strain in regions where $x(\delta) \neq x_r$. The values of $\delta_c$ can be found by means of the equation [8, 10, 11]

$$k_n = \left( \frac{\Delta a}{a} \right)^n \int_{\delta_c}^{\delta_r} (x(\delta) - x(\delta_c))^n \, d\delta$$

with $n = 1$ [10] or $n = 2$ [8, 11]. $\Delta a/a = 0.07$ is the lattice mismatch between InAs and GaAs, while $k_1 = 0.83$ nm [10], $k_2 = 3.68 \times 10^{-3}$ nm [8, 11] and $\delta_c$ is the total thickness of the buffer.

From Eq. (1) it follows that by choosing the composition profile $x(\delta)$ and the thickness $\delta_c$, it should be possible to predetermine (i) the thickness ($\delta_r - \delta_c$) of the MD-free region and (ii) the lattice parameter $a(x_c)$ of the upper part of the buffer; on the other hand, the concentration profile of MDs is determined by the composition profile in the buffers. It is worth noting that if additional structures deposited atop buffers can be considered as single layers with free-standing lattice parameters $a_i$ (and with a mean composition $x_i$), then, by choosing the composition profile of buffers, the active structures can be put under overall compressive, null or tensile strain, thus affecting their electronic, optical and transport properties. However, if the lattice parameter of the upper part of the buffer is significantly different from that of the active structure ($x_r \neq x_i$), the latter modifies the strain situation of the buffer that cannot be considered as an independent block any more; in that case, Eq. (1) should be used considering the whole structure consisting of buffer and active region.

We designed structures consisting of graded buffers and two types of MQW structures for 1.3 and 1.5 $\mu$m optical modulators, thereafter referred to as 1.3 and 1.5 $\mu$m MQWs. The 1.5 and 1.3 $\mu$m MQWs both have mean compositions $x_{m,\text{MQW}} = 0.34$ and consist of 30.5 periods of 30 and 35 $\AA$ GaAs barriers and of 65 and 75 $\AA$ thick In$_{0.5}$Ga$_{0.5}$As wells, respectively. The buffer structures were designed according to Ref. [8, 11] in order to avoid strain in the MQWs; therefore, the composition profiles with linear, square-root and parabolic profiles were graded in 3000 nm up to values $x_m$ of 0.41, 0.39 and 0.38, respectively, so that $x_m > x_r = x_{m,\text{MQW}}$. It should be noted that this approach is different from that generally used, where the buffer compositions are graded up to the mean compositions of the MQWs ($x_m = x_{m,\text{MQW}}$), thus resulting in MQWs under compressive strain.

4. Results and discussion

Fig. 1 shows a comparison among cross sections of linear, square-root and parabolic buffers; the most interesting features are: (i) it can be clearly seen that the topmost parts of buffers are void of MDs, as predicted by the existing models [6, 8, 10, 11], and as previously observed in Refs. [12, 13], (ii) the thicknesses of MD-free layers increase ranging from linear, square-root and parabolic buffers and are $\sim 600$ nm for the last ones; the values are in reasonable agreement with the results of Eq. (1) with $n = 2$; (iii) the misfit dislocations have different distributions in buffers with different composition profiles, being almost uniformly distributed (up to $x_r$) in the linear buffers and strongly accumulated towards the
substrate/buffer interface in square-root buffers; (iv) as predicted by the strain relaxation models [6], continuous grading of composition results in a relatively low concentration of TDs, that in our buffers is lower than $1 \times 10^6$ cm$^{-2}$.

When MQWs were grown on the buffers, only a few TDs and MDs at the MQW/buffer interface were found by TEM observations; their maximum density ranged between $1.5 \times 10^6$ and $4 \times 10^6$ cm$^{-2}$ for structures with linear and parabolic buffers, respectively. The thicknesses of the MD-free layers in structures with and without MQWs are approximately the same, thus implying that the MQWs do not contribute to the integral of Eq. (1), and, then, that the MQWs are closely lattice-matched to the topmost part of the buffers ($x_i \approx x_m$). Independent of the type of buffer, the MQWs have the same structural quality and interface smoothness (on atomic scale), as those shown in Fig. 2, relative to a MQW grown on a linear buffer.

As an example, in Fig. 3 we show the $(0 0 4)$ CuKα diffraction profile of a 1.5 µm MQW grown on a square-root buffer; the quality of the structures is assessed by the occurrence of peaks related to MQW periodicity. From the average lattice parameters of MQWs parallel and perpendicular to the interfaces we conclude that the MQWs are nearly lattice matched to the different types of buffers; the In compositions $x_m$ at the top of buffers evaluated by the diffraction profiles are in good agreement with those derived by using the experimental values of the mean composition of MQWs, the thicknesses of the barriers and wells (obtained by TEM observations) and the In and Ga supply times; it is interesting to note that the $x_i$ values obtained by Eq. (1) using the above $x_m$ values are consistent with the strain, whether slightly tensile or compressive, of the MQWs given by HRXRD experiments.

As for the surface smoothness, AFM observations show that both uncapped buffers and MWQs
Fig. 2. Dark field, $g = 002$ type (left panel) and bright field zone axis lattice resolution (right panel) micrographs of a 1.5 μm MQW (30 Å GaAs / 65 Å In$_{0.5}$Ga$_{0.5}$As) grown atop a linearly graded $(0.005 \leq x \leq 0.41)$ InGaAs buffer.

Fig. 3. (0 0 4) CuKα diffraction profile of a 1.5 μm MQW (30 Å GaAs / 65 Å In$_{0.5}$Ga$_{0.5}$As) grown atop a square-root $(0.005 \leq x \leq 0.39)$ InGaAs buffer.
have small and symmetric cross-hatchings along the two orthogonal \(<110>\)-type directions; the standard deviations of the surface heights (SDSH) depend only slightly on the type of buffer, being, for instance, 3.4 \(\pm\) 0.3 and 2.6 \(\pm\) 0.3 nm for linear and parabolic buffers, respectively. The MQWs tend to planarize the surfaces, as can be deduced by the SDSHs of 1.5 \(\mu\)m MQW surfaces that are 2.0 \(\pm\) 0.3 and 1.4 \(\pm\) 0.2 nm for linear and parabolic buffers, respectively.

In Fig. 4 we show the PL spectra at 10 and 300 K of 1.5 and 1.3 \(\mu\)m virtually unstrained MQWs grown on the same buffers as those shown in Fig. 1. It is worth noting that by carefully designing and growing the structures, efficient PL can be obtained at 300 K within or close to the 1.3 and 1.5 \(\mu\)m spectral windows of photonic interest. The PL full-width-at-half-maximums (FWHM) at 10 K relative to the 1.5 \(\mu\)m MQWs (9, 12 and 12 meV, for the parabolic, the square-root and the linear buffers, respectively) are significantly small; this result can be likely related to the design of buffers that virtually do not strain MQWs, unlike the approach that matches the compositions of the tops of buffers to the mean compositions of MQWs. This interpretation may also explain the reduction of the PL FWHMs for increasing \(x_m\) up to values larger than \(x_{MQW}\) reported in Ref. [14]; our 300 K FWHMs are smaller by a factor of \(~3-4\) than the 77 K FWHMs given in same paper [14].

5. Conclusions

We prepared under optimized conditions and studied InGaAs buffers grown on GaAs, intended for MQW structures for 1.3 and 1.5 \(\mu\)m optical operation at 300 K; the buffers have linear, square-root and parabolic composition profiles. The buffers were designed in such a way that the MQWs grown atop the buffers are virtually unstrained, unlike those prepared following the conventional approach, that are compressively strained. The results obtained by the concomitant use of TEM, HRXRD, AFM and PL show that (i) the MDs profiles and thicknesses of the MD-free regions in the buffers can be predetermined, (ii) the active structures atop buffers are virtually unstrained and have efficient 300 K photoluminescence in the 1.3 and 1.5 \(\mu\)m windows, and (iii) the structures have
TD concentrations in the low $10^6 \text{ cm}^{-2}$ range and smooth and symmetric cross-hatchings.

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References

Molecular beam epitaxy growth and characterization of In$_x$Ga$_{1-x}$As ($0.57 \leq x \leq 1$) on GaAs using InAlAs graded buffer

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Abstract

In$_x$Ga$_{1-x}$As ($x \geq 0.57$) thin layers were grown on GaAs by molecular beam epitaxy using an InAlAs graded buffer. Surface roughness of the graded buffer decreased with reduced buffer thickness and growth temperature. The InGaAs layers grown on a graded buffer revealed a larger tilting angle, a higher residual strain and electron mobility than those grown directly on GaAs. Increasing the graded buffer thickness above 1.2 μm did not improve the transport properties of InAs. The InAs mobility increased with layer thickness. This was explained by assuming two conducting channels, given $\mu_{300} = 1.8 \times 10^4$ cm$^2$/V s and $\mu_{T} = 5.1 \times 10^5$ cm$^2$/V s for bulk InAs.

PACS: 68.55. - a; 73.61.Ey; 81.15.Hi

Keywords: MBE; InGaAs; InAlAs graded buffer; GaAs

1. Introduction

The presence of threading and misfit dislocations is common in relaxed lattice mismatched semiconductor heterostructures and limits their use in device applications. In the last several years efforts have been made to reduce dislocation densities by first growing a graded alloy buffer [1–9]. In this case the nucleated dislocations glide to sample edges or interface more easily as compared with those in a uniform layer, because of a larger strain force and weaker pinning [10]. By using this technique, dislocation densities as low as $10^6$ cm$^{-2}$ and $10^5$ cm$^{-2}$ have been reported for Ge on Si [5] and for In$_{0.46}$Ga$_{0.54}$As on GaAs [7], respectively. Devices like InAlAs/In$_x$Ga$_{1-x}$As ($x \geq 0.52$) high electron mobility transistors on GaAs [11–13] and
In$_x$Ga$_{1-x}$P light emitting diodes on GaP [14] have been fabricated showing very promising performance. More important, this technique offers the possibility of growing metamorphic "substrates" with adjustable lattice constant. In this work, we report MBE growth and characterization of In$_x$Ga$_{1-x}$As ($x \gtrsim 0.57$) on GaAs using continuously graded InAlAs buffers. Incorporation of Al instead of Ga in the graded buffer is motivated by the fact that it reduces the background carrier density in subsequent layers [3]. The influence of growth temperature and graded buffer thickness on material quality is studied by atomic force microscopy (AFM), double crystal X-ray diffraction (XRD) and Hall effect measurements.

2. Material growth

We used quarter 2 in. semi-insulating (0 0 1) GaAs substrates and In-mounting Mo sample holders (MSHs). Samples were grown under As-rich growth conditions and rotated at 10 rpm in a solid source Varian Gen II MBE system. Each sample consisted of a GaAs buffer (0.1 to 0.5 μm thick) grown at 580°C, a graded In$_x$Al$_{1-x}$As buffer and in some samples an In$_x$Ga$_{1-x}$As bulk layer with the In content the same as the final value of the graded buffer, $x_{f}$. During growth of the graded buffer, both In and Al source temperatures were varied linearly. The initial $x$-value, $x_{i}$, was 0.005. For comparison, one sample ( #518) with a 1 μm thick In$_x$Ga$_{1-x}$As graded buffer was grown. The growth temperature, $T_g$, measured by a pyrometer, varied between 350°C and 480°C for the graded buffers and between 420°C and 500°C for the bulk layers. As the pyrometer measured temperature correctly only above 450°C and use of MSHs would change the pyrometer reading, a temperature calibration procedure was made. Fig. 1 shows the relation between the pyrometer and the thermocouple temperature, $T_{tc}$, for three MSHs and an In-free sample holder. The $T_g$-value was calibrated by the oxide desorption temperature of GaAs (583°C). As seen, the $T_g$-value is about 60–70°C lower for the MSHs than for the In-free holder, indicating heat dissipation of the MSHs. The pyrometer measures a higher $T_g$-value when the As$_2$ shutter is open than closed. This is due to reflection of heat radiation from the As$_2$ cracker (at 1000°C) by the MSH. The reflection is dominant at low $T_{tc}$-values which sets a minimum pyrometer reading of 520°C. Thus, the correct growth temperature is measured only when the As$_2$ shutter is closed. In the temperature range used in this work the difference of $T_g$-values between MSHs was within 10°C. The In content and the growth rate (around 1 μm/h) were calibrated by RHEED oscillations.

3. Characterization results and discussions

Atomic force microscopy was used to study the surface properties of the In$_x$Al$_{1-x}$As ($x_{f}=0.8$) graded buffer layers. We first checked the surface of the GaAs buffer and found that the root-mean-square (rms) roughness value decreased only slightly from 0.36 to 0.30 nm, when the GaAs thickness increased from 0.1 to 0.5 μm. Based on this fact, a 0.1 μm thick GaAs buffer was selected for the AFM samples, in which four samples had a graded buffer thickness, $L_{grad}$, of 1 μm grown at 300–480°C while two were grown at 350°C with thicknesses of 0.5 and 2 μm, respectively. The corresponding lattice mismatch slope, $k$, was between 2.9 and 11.5%/μm. All samples were mirror-like to the naked eye and showed random surface morphology under AFM. A typical AFM image is shown in Fig. 2 and the rms roughness values are collected in Fig. 3. The surface becomes more rough with
strained areas have a lower elastic energy and thus, a relatively higher growth rate. Since at high growth temperatures the adatoms have large kinetic energy and surface diffusion length, they can easily find such areas and form rough surface. For a 1 μm thick InAlAs graded buffer grown at 350°C, a rms roughness value of 2.5 ± 0.9 nm was obtained.

Double crystal XRD was performed in 2 μm thick In$_x$Ga$_{1-x}$As (0.57 ≤ x ≤ 1) layers, grown with and without the InAlAs graded buffer. Both symmetric (0 0 4) and asymmetric (1 1 5 ± ) diffraction geometry were used and the rocking curves were recorded at four different azimuthal angles. The XRD results are summarized in Table 1. For the InGaAs layers grown directly on GaAs, the alloy peak is narrow and the peak shift with the azimuthal angle is small. However, for the InGaAs layers grown on the graded buffer, the alloy peak is wide and the shift is strong. A typical rocking curve at the (0 0 4) diffraction geometry is shown in Fig. 4. The peak shift with the azimuthal angle indicates a tilting of the In$_x$Ga$_{1-x}$As layer relative to the GaAs substrate. This implies that the net value of the Burgers vector along the z-direction is not zero. The full-width at half-maximum (FWHM) is wider than that of layers without the graded buffer, indicating a wide distribution of the vertical lattice constant. This can be explained by the two factors: First, continuous alloy grading and non-uniform dislocation density along the z-direction in the InAlAs graded buffer cause a wide distribution of lattice constant. Second, strain could be further relieved in the 2 μm thick InGaAs layer by dislocation formation which broadens the InGaAs peak. In addition, the FWHM is about 2–3 times wider in one direction than in its perpendicular direction in all samples with the graded buffer. We believe that it is related to the asymmetric distribution of dislocations. The values of FWHM listed in Table 1 are for the narrow one.

Both vertical and lateral lattice constants are extracted from the asymmetric (1 1 5 ± ) XRD rocking curves. The in-plane residual strain, ε, and the In content are then obtained assuming Vegards law holds. The residual strain in the layers grown on a graded buffer is larger than in layers grown directly on GaAs, which is in line with the Tersoff
Table 1
Summary of XRD results

<table>
<thead>
<tr>
<th>Sample No.</th>
<th>Misfit slope ( k (%) )</th>
<th>In content ( x )</th>
<th>Tilting angle ( \theta, (^{\circ}) )</th>
<th>FWHM (mrad)</th>
<th>Residual strain ( \varepsilon (10^{-3}) )</th>
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<td>-</td>
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<td>6.0</td>
<td>1</td>
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<td>1.9</td>
</tr>
<tr>
<td>544</td>
<td>3.0</td>
<td>1</td>
<td>1.35</td>
<td>7.9</td>
<td>1.4</td>
</tr>
<tr>
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<td>2.0</td>
<td>1</td>
<td>1.19</td>
<td>5.7</td>
<td>1.2</td>
</tr>
<tr>
<td>552</td>
<td>-</td>
<td>0.741</td>
<td>&lt; 0.01</td>
<td>4.2</td>
<td>1.3</td>
</tr>
<tr>
<td>553</td>
<td>5.3</td>
<td>0.740</td>
<td>0.92</td>
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<td>3.4</td>
</tr>
<tr>
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<td>-</td>
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<td>&lt; 0.01</td>
<td>6.2</td>
<td>&lt; 0.1</td>
</tr>
<tr>
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<td>5.1</td>
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<td>1.01</td>
<td>11.1</td>
<td>0.7</td>
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<table>
<thead>
<tr>
<th></th>
<th>XRD</th>
<th>Tersoff model</th>
</tr>
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<tbody>
<tr>
<td></td>
<td></td>
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</tr>
</tbody>
</table>

Fig. 4. XRD rocking curve from a 2 \( \mu m \) thick InAs layer grown on a 3.6 \( \mu m \) thick InAlAs graded buffer.

The large strain force pushes nucleated dislocations towards sample edges and interface. Thus, the dislocation density is reduced. According to Tersoff [10] the residual strain near the surface is

\[
\varepsilon = (2 \lambda k/bc)^{1/2},
\]

where \( \lambda \) is the energy per unit length of dislocation, \( b \) is the Burgers vector and \( c \) is the appropriate elastic constant for biaxial strain. The calculated residual strain is also listed in Table 1. It increases with the misfit slope, in agreement with the observation. However, the calculated \( \varepsilon \)-values are larger than the measured data except for \# 553. This is due to further strain relaxation in the 2 \( \mu m \) thick InGaAs layer because of a small lattice mismatch between the graded buffer and the bulk layer. For the InAs samples with \( x_f = 0.995 \), this lattice mismatch is larger.

A comparative study of electron mobility in samples with and without a graded buffer was made. Since the mobility values depend highly on the cleanliness of the MBE system and the growth conditions, we only compare samples grown subsequently under similar growth conditions. The van der Pauw method was used in Hall-effect measurements and all data were measured at a magnetic field of 0.2 T and a current of 10 \( \mu A \). Table 2 lists electron mobility, \( \mu \), and carrier density, \( n \), of the InAs layers using a Hall coefficient of 1.3. Use of a graded buffer increases mobility by 10\%-15\% at 300 K for both As\(_4\) and As\(_2\) sources. Compared with the InGaAs graded buffer, use of an InAlAs graded buffer reduces background carrier density without significant decrease of mobility. When the \( L_{grad} \) is increased from 1.2 to 3.6 \( \mu m \), the mobility is nearly unchanged at 300 K and decreases slightly at 77 K. Hence, a thick graded buffer does not improve transport properties.

We have also grown InAs layers with different thicknesses on a 1.2 \( \mu m \) thick InAlAs graded buffer. The mobility at 300 K increases from \( 1.08 \times 10^4 \text{ cm}^2/(V \text{ s}) \) for the 2 \( \mu m \) layer to \( 1.39 \times 10^4 \text{ cm}^2/(V \text{ s}) \) for the 10 \( \mu m \) layer. The measured conductivity, \( \sigma \), and Hall factor, \( R \), are shown in Fig. 5. It is well known that for InAs the Fermi level is pinned 0.2 eV above the conduction band minimum [15] and a surface accumulation layer exists with a high sheet electron concentration [16]. Hence, the measured mobility
Table 2
Summary of transport properties from InAs layers with and without InAlAs graded buffer

<table>
<thead>
<tr>
<th>Sample No.</th>
<th>Description</th>
<th>300 K</th>
<th>77 K</th>
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</thead>
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<tr>
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<td></td>
<td>$\mu$ (10^4 cm^2/(V s))</td>
<td>$n$ (10^{16} cm^{-2})</td>
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</tr>
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<td>501</td>
<td>reference</td>
<td>1.04</td>
<td>2.09</td>
</tr>
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<td>on 1 $\mu$m InGaAs buffer</td>
<td>1.21</td>
<td>4.00</td>
</tr>
<tr>
<td>519</td>
<td>on 1 $\mu$m InAlAs buffer</td>
<td>1.11</td>
<td>1.77</td>
</tr>
<tr>
<td>520</td>
<td>on 3 $\mu$m InAlAs buffer</td>
<td>1.14</td>
<td>1.79</td>
</tr>
<tr>
<td>As$_x$ source, $T_s = 460-470^\circ$C, d = 2 $\mu$m</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>541</td>
<td>reference, $L_{\text{grad}} = 0$</td>
<td>0.87</td>
<td>2.91</td>
</tr>
<tr>
<td>542</td>
<td>$L_{\text{grad}} = 1.2 \mu$m</td>
<td>1.08</td>
<td>2.24</td>
</tr>
<tr>
<td>544</td>
<td>$L_{\text{grad}} = 2.4 \mu$m</td>
<td>1.13</td>
<td>2.43</td>
</tr>
<tr>
<td>545</td>
<td>$L_{\text{grad}} = 3.6 \mu$m</td>
<td>1.03</td>
<td>2.42</td>
</tr>
</tbody>
</table>

is an averaged value due to this parallel conduction. For two conducting channels the $\sigma$ and $R$ can be expressed by [17]

$$R = \frac{(d_b + d_s)(R_b\sigma_b^2 + R_s\sigma_s^2)}{(\sigma_b d_b + \sigma_s d_s)^2},$$  \hspace{1cm} (2)

$$\sigma = \frac{(\sigma_b d_b + \sigma_s d_s)}{d_b + d_s},$$  \hspace{1cm} (3)

where subscripts $b$ and $s$ denote bulk and surface layer, respectively, and $d$ is the layer thickness. This is a good assumption here, since no abrupt band offset exists at the interface between the graded buffer and the InAs layer. The best theoretical fits using Eqs. (2) and (3) are shown in Fig. 5 by lines using $\mu_b = 1.8 \times 10^4$ cm$^2$/V s, $n_b = 1.4 \times 10^{13}$ cm$^{-3}$, $\mu_s = 8.9 \times 10^3$ cm$^2$/V s, $n_s = 4.4 \times 10^{12}$ cm$^{-2}$ at 300 K and $\mu_b = 5.1 \times 10^4$ cm$^2$/V s, $n_b = 1.1 \times 10^{15}$ cm$^{-3}$, $\mu_s = 2.7 \times 10^4$ cm$^2$/V s, $n_s = 1.8 \times 10^{12}$ cm$^{-2}$ at 77 K. From this we can estimate the maximum dislocation density. Theoretical calculations have revealed that the lattice scattering limits the 300 K mobility of InAs to be $3.5 \times 10^4$ cm$^2$/V s [18]. The mobility due to the ionized impurity scattering in our case is over $1 \times 10^6$ cm$^2$/V s and thus negligible. If we assume that among the other scattering mechanisms only dislocation scattering is significant, this gives us $3.7 \times 10^4$ cm$^2$/V s. The maximum dislocation density can then be calculated [19] to be $4 \times 10^7$ cm$^{-2}$.

Fig. 5. Measured conductivity and Hall factor as a function of inverse InAs thickness. Lines are theoretical fits.

Fig. 6 shows the transport properties of In$_x$Ga$_{1-x}$As layer as a function of In content measured at (a) 300 K and (b) 77 K. Again, a higher electron mobility is found in the layers grown on an InAlAs graded buffer. The mobility decreases rapidly when the x-value is reduced from 1. This can be partly explained by the fact that the growth conditions are not optimized for the In$_x$Ga$_{1-x}$As layers and partly by the increased alloy scattering. For x = 0.57, the mobility at 77 K is smaller than at 300 K. To understand the scattering mechanisms behind this abnormal phenomenon, a temperature dependent mobility measurement is necessary.
4. Conclusions

We have studied surface, structural and transport properties of MBE grown In₀.₅₋ₓGaₓ₁₋ₓAs (x = 0.8) graded buffers on GaAs and bulk In₀.₅₋ₓGaₓ₁₋ₓAs (0.57 ≤ x ≤ 1) layers with and without such a graded buffer. The surface of the graded buffers revealed random and rough morphology. The rms roughness value increased with growth temperature and graded buffer thickness. A small roughness value of 2–3 nm was obtained for $L_{\text{grad}} \leq 1 \mu\text{m}$ and $T_g \leq 350^\circ\text{C}$. The In₀.₅₋ₓGaₓ₁₋ₓAs layers grown on an InAlAs graded buffer showed a larger tilting angle, a broader FWHM and a higher residual strain as compared with the layers grown directly on GaAs. Transport properties were improved in layers grown on a graded buffer. Increasing the graded buffer thickness did not increase the electron mobility of InAs for $L_{\text{grad}} \geq 1.2 \mu\text{m}$. The thickness dependent mobility of InAs can be well explained by assuming two conducting channels. The extracted electron mobility of the bulk InAs is $1.8 \times 10^{4} \text{cm}^2/(\text{V} \cdot \text{s})$ at 300 K and $5.1 \times 10^{4} \text{cm}^2/(\text{V} \cdot \text{s})$ at 77 K.

Acknowledgements

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References

Improved nucleation and spiral growth of PbTe on BaF$_2$ (1 1 1)

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Abstract

Molecular beam epitaxy of PbTe on BaF$_2$(1 1 1) is studied by in situ reflection high-energy electron diffraction and atomic force microscopy. It is shown that a 4 monolayer thick EuTe nucleation layer predeposited on the BaF$_2$ substrate drastically improves the nucleation and coalescence of the PbTe layers. After complete coalescence of the layers, PbTe growth is found to be completely dominated by growth spirals formed around threading dislocations that originate from the growth on the 4.2% lattice-mismatched BaF$_2$ substrates. Due to annihilation processes, the threading dislocation density rapidly decreases with layer thickness, which results in drastic increase of the 10 K electron mobilities in the layers. Thus, thick PbTe buffer layers are well suited as “virtual” substrates for IV–VI compound multilayers and heterostructures.

_PACS:_ 68.55.Bd; 61.16.Ch; 61.14.Hg; 68.55.Jk

1. Introduction

Advances in epitaxial growth of lead-salt (IV–VI) semiconductors [1] have recently lead to room temperature operation of mid-infrared lead-salt diode lasers in pulsed mode [2] and to cw operation at 223 K [3]. While these devices are usually grown on lead-salt substrates, (1 1 1) oriented BaF$_2$ has proven to be an excellent alternative as substrate material for lead-salt heterostructures [4]. It is readily available in high crystalline quality and because of the well-matched thermal expansion coefficients no significant degradation of epitaxial layers occurs during thermal cycling. In addition, while lead-salt substrates are always highly conducting, BaF$_2$ is insulating and optically transparent, which is an essential prerequisite for the study of the electronic properties of heterostructures [5, 6].

In the present work, we carried out a systematic study of molecular beam epitaxy (MBE) of PbTe on BaF$_2$(1 1 1) using in situ reflection high-energy electron diffraction (RHEED) and ex situ atomic force microscopy (AFM). It is demonstrated that by using a thin EuTe nucleation layer predeposited on the BaF$_2$ substrate, the 3D nucleation of PbTe can be drastically improved and extremely smooth surfaces can be obtained already after 300 Å layer
thickness. Due to the 4.2% lattice-mismatch of PbTe relative to the BaF$_2$ substrate we observe a high density of threading dislocations on these surfaces. These give rise to a spiral-type growth mode during further deposition. While such types of growth mode are well known and understood for bulk crystal growth at low supersaturation [7], it is surprising that this growth mode is also present under MBE conditions far from thermodynamic equilibrium. From AFM and scanning tunneling microscopy we find a very rapid decrease of the threading dislocation density with increasing layer thickness. This decrease is correlated with an increase of the low-temperature electron mobilities in the epitaxial layers by two orders of magnitude, reaching more than $2 \times 10^5$ cm$^2$/V·s for layers larger than 3 µm in thickness.

2. Experimental procedure

Epitaxial growth was carried out in a Riber 1000 MBE system with standard effusion cells for PbTe, Eu, Te$_2$, and Bi$_2$Te$_3$ [5], where the latter was used for n-type doping. Cleaved BaF$_2$(1 1 1) substrates were baked at 500°C for 15 min prior to epitaxial growth. For PbTe a substrate temperature of 340°C and growth rates of about 1.4 µm/h were used. Additional Te$_2$ flux was used in order to improve PbTe growth [8]. In situ RHEED was carried out using a 35 keV electron gun and the surface structure of the samples were studied using a Park Scientific Autoprobe CP AFM operated in contact mode. With this, the monolayer step structure could be clearly resolved in air. In situ UHV scanning tunneling microscopy (STM) studies were also performed [9].

3. Nucleation of PbTe on BaF$_2$ (1 1 1)

Due to the much larger (1 1 1) free surface energies of the lead salts as compared to BaF$_2$, epitaxial growth always starts in a 3D Volmer–Weber growth mode, where large triangular-shaped pyramidal islands with low-energy (1 0 0) side facets are formed on the substrate surface [10]. In situ RHEED studies show that coalescence of these islands begins only for layer thicknesses larger than 1000 Å, and a completely streaked RHEED pattern characteristic of a 2D surface is usually restored not before 0.5 µm layer deposition [11, 12]. Therefore, for high-quality lead-salt heterostructures always thick buffer layers have to be deposited on BaF$_2$. In principle, the same nucleation behavior is observed also for other lead-salt compounds, although it is noted that the coalescence behavior can be quite different for BaF$_2$ surface areas preirradiated by the RHEED high-energy electron beam.

For MBE growth of the ternary compound Pb$_{1-x}$Eu$_x$Te on BaF$_2$ (1 1 1), however, we found that a complete layer coalescence takes place already at 500 Å layer thickness, even when the Eu content is below 1%. This is a factor of 10 faster as for the pure binary compound. Since such a small Eu content does not change the overall growth properties, the drastic improvement can be only due to a different interaction between Eu and the BaF$_2$ substrate. Indeed, from separate experiments we find that a few monolayers (ML) of pure EuTe predeposited on BaF$_2$ just before PbTe growth results in a similar improvement of the coalescence behavior. The pure EuTe compound has the same rock salt crystal structure as the lead-salt compounds and the lattice-mismatch with respect to PbTe is only 2.1%. EuTe is grown using elemental beam flux sources with excess Te$_2$ flux in order to retain the right stoichiometry of the compound [13].

Fig. 1 shows the RHEED patterns recorded for such a growth sequence. In this case, after the usual substrate preheating procedure, first 4 ML EuTe is deposited on the BaF$_2$ surface. Instantaneously, the RHEED patterns changes to a 3D transmission diffraction pattern (see Fig. 1), indicating the presence of a 3D growth mode also for the EuTe case. When PbTe is deposited on top of this EuTe nucleation layer, initially, the surface roughens even more, as is indicated by a slight faceting of the diffraction spots (see RHEED patterns for 10 Å PbTe in Fig. 1). Then, however, a rapid smoothening of the surface takes place and already at 50 Å layer thickness the onset of streaking is observed in the RHEED patterns, and very rapidly a completely streaked RHEED pattern is restored (Fig. 1). Thus, already at 200–300 Å PbTe layer...
thickness a smooth 2D epitaxial surface is obtained by this modified growth sequence.

4. Spiral growth and structural properties

AFM images taken after complete coalescence of the PbTe layers show that the epitaxial surface at 500 Å layer thickness is very smooth even on a large length scale (see Fig. 2a). These surfaces consist only of monolayer steps due to epitaxial growth. In addition, detailed STM studies reveal that most of the surface step segments are terminated by threading dislocations (TD), as indicated in Fig. 2b and Fig. 2c by the arrows. These threading dislocations originate from the growth on the 4.2% lattice-mismatched substrate and connect the misfit dislocation segments at the PbTe/EuTe/ BaF$_2$ interface with the free surface, as shown schematically in Fig. 3.

If one draws a closed loop around a threading dislocation on the surface, there remains a net surface normal displacement of 3.7 Å, which indicates that the threading dislocations are of screw-type character. In the lead-salt compounds, the dislocation Burgers vector is usually of $\frac{1}{2}[110]$ type, which is inclined by 54.7° with respect to the (111) interface. Therefore, it has 3.72 Å surface normal component, which corresponds exactly to the height of the resulting step on the surface. As evident from Fig. 2b and Fig. 2c, most of the surface steps produced by the TDs are terminated at the other end by a second dislocation with opposite Burgers vector.

Since in our AFM images, we usually do not observe 2D islands nucleated on the epitaxial surface, PbTe growth proceeds in a pure step flow growth mode where deposition of atoms results in a lateral growth at the step edges. However, since the end points of these surface steps are pinned by the threading dislocations, this lateral step growth eventually leads to the formation of growth spirals on the surface (see Fig. 2), similar as in dislocation-mediated growth of bulk single crystal. As more material is deposited on the surface, the TD density rapidly decreases. Since the diameter of the growth spirals is equal to the spacing of the TDs on the surface, larger and larger
Fig. 2. AFM images of PbTe epitaxial layers with different layer thicknesses deposited on BaF$_2$(1 1 1) using a 4 ML EuTe nucleation layer. From top left to bottom right: 500 Å PbTe, 1000 Å PbTe, 2000 Å PbTe, 1 µm PbTe, 3 µm PbTe and 8 µm PbTe. Note the different scales of the images which is $4 \times 4$ µm$^2$ for the top row and $8 \times 8$ µm$^2$ for the bottom row. The arrows in the images indicate threading dislocations originating from the growth on the 4.2% lattice-mismatched BaF$_2$ substrate.

Fig. 3. Schematic illustration of the dislocation configuration.

growth spirals are produced on the surface (see Fig. 2d–Fig. 2f).

At 3 µm layer thickness, the surface is completely covered by large growth spirals, which are typically 2–3 µm apart (Fig. 2e), and at 8 µm thickness spirals with diameter as large as 8 µm are observed (note the different length scales in the AFM images shown in Fig. 2). Since throughout the whole growth process the average step spacing within the spirals remains essentially constant (≈2000 Å, see Fig. 2), the large growth spirals consist of many turns and as a result, the centers of these spirals are elevated by more than 50 Å above the rest of the surface. This gives rise to a gentle overall hillock structure of the PbTe surfaces that is usually observed by Nomarski microscopy.

As is evident from Fig. 2, both right- and left-handed growth spirals are observed on the surface. Thus, threading dislocations with different orientations of the Burgers vector are present in the layers. Due to the elastic strain fields of dislocations, dislocations with anti parallel Burgers vector are strongly attracted to each other. As a result, they move
closer and closer together and when they meet annihilate each other, i.e., the threading arms of the misfit dislocations are removed. This is the mechanism responsible for the reduction of the TD density as growth proceeds.

From STM and AFM images taken at various layer thicknesses, we have determined the TD density as a function of layer thickness and the result is shown in Fig. 4. We find a surprisingly rapid decrease of the TD density with increasing layer thickness from above $1.5 \times 10^6$ cm$^{-2}$ for 1000 Å layer thickness, to a few times $10^6$ cm$^{-2}$ for layers of several μm in thickness, following roughly a power law dependence with a negative exponent of 1.2 (dashed line in Fig. 4). This rapid decrease implies a quite high dislocation mobility at the elevated growth temperatures, which is consistent with recent thermal cycling studies of lead-salt epitaxial layers on Si(1 1 1) substrates [14].

Fig. 5 shows the carrier mobilities of PbTe epitaxial layer as a function of layer thickness obtained from temperature-dependent Hall effect measurements. In this sample series, Bi-doping was used to adjust the n-type carrier concentration to be in the range of the mid-$10^{17}$ cm$^{-3}$, whereas not intentionally doped layers of less than 0.5 μm in thickness are usually p-type of the order of $10^{18}$ cm$^{-3}$. While the phonon-limited room temperature mobility of the layers is essential constant for all layers, we observe a drastic increase of the 10 K mobilities by two orders of magnitude from $2 \times 10^4$ for layers of 1000 Å in thickness to above $2 \times 10^6$ cm$^2$/V s for layers exceeding 3 μm. This rapid increase clearly indicates a strong correlation between carrier mobility and TD density in PbTe epitaxial layers.

5. Conclusions

In conclusion, it was shown that MBE growth of PbTe on BaF$_2$(1 1 1) substrates can be greatly improved by the use of a thin EuTe nucleation layer predeposited on the BaF$_2$ substrate. From AFM studies, the surface structure of these layers is found to be determined mainly by threading dislocations originating from the PbTe/BaF$_2$ interface. This, in combination with persisting step flow growth results in a unique spiral type of growth mode that is usually not observed for strained-layer heteroepitaxy of other semiconductors. Continued growth results in a drastic decrease of the threading dislocation density, leading to significant improvements in the electronic properties of the layers.
Acknowledgements

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References

Molecular beam epitaxy of Al\(_{0.48}\)In\(_{0.52}\)As/Ga\(_{0.47}\)In\(_{0.53}\)As heterostructures on metamorphic Al\(_x\)Ga\(_y\)In\(_{1-x-y}\)As buffer layers

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Abstract

Ternary Al\(_{1-x}\)In\(_x\)As/Ga\(_{1-y}\)In\(_y\)As heterostructures with a lattice mismatch up to 4% are grown on GaAs by molecular beam epitaxy. Two buffer layer concepts to compensate the lattice misfit between the Al\(_{1-x}\)In\(_x\)As/Ga\(_{1-y}\)In\(_y\)As layers and the GaAs substrate using the quaternary Al\(_x\)Ga\(_y\)In\(_{1-x-y}\)As in a linear graded and two-step graded fashion, respectively, are presented. The Al and Ga content of the ternary layers were chosen to be x = 0.48 and y = 0.47, respectively, in order to obtain the same heterostructures identical to those grown lattice matched on InP as a reference. The surface morphology and the transport properties of Al\(_{0.48}\)In\(_{0.52}\)As/Ga\(_{0.47}\)In\(_{0.53}\)As high-electron mobility transistor structures were studied by atomic force microscopy and Hall measurements, respectively. Optical properties were investigated by low-temperature photoluminescence on quantum well structures. The use of the two step graded buffer layers resulted in three-dimensional layer growth and inferior layer quality. In contrast, the linear graded buffer approach was found to result in superior heterostructure properties due to the two-dimensional growth mode during the whole growth process resulting in the typical cross-hatched surface morphology.

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Keywords: Molecular beam epitaxy; Metamorphic growth; AlInAs/GaInAs heterostructures

1. Introduction

Ternary Al\(_{0.48}\)In\(_{0.52}\)As/Ga\(_{0.47}\)In\(_{0.53}\)As heterostructures lattice matched to InP offer a wide range of applications for electronic and optoelectronic devices. This material system is especially suitable for applications in long-wavelength optical-communication range at wavelengths between 1.3 and 1.55 µm. Interesting as well are the high-electron mobility and concentration achieved in modulation-doped Al\(_{0.48}\)In\(_{0.52}\)As/Ga\(_{0.47}\)In\(_{0.53}\)As heterostructures for the design of high-speed field effect transistors [1]. InP substrates are generally more expensive than high-quality GaAs substrates, it is therefore desirable to combine the advantages of Al\(_{0.48}\)In\(_{0.52}\)As/Ga\(_{0.47}\)In\(_{0.53}\)As heterostructures

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with the low cost and availability of large GaAs substrates. To achieve successful growth of possible $\text{Al}_{0.48}\text{In}_{0.52}\text{As}/\text{Ga}_{0.47}\text{In}_{0.53}\text{As}$ device structures, the large mismatch between the GaAs substrate and the $\text{Al}_{0.48}\text{In}_{0.52}\text{As}/\text{Ga}_{0.47}\text{In}_{0.53}\text{As}$ high-electron mobility transistor (HEMT) and quantum well (QW) structures has to be accommodated by a suitable epitaxial buffer layer sequence. This resorts to a more complicated growth process. A successful epitaxy of relaxed $\text{Al}_{x}\text{In}_{1-x}\text{As}/\text{Ga}_{y}\text{In}_{1-y}\text{As}$ on GaAs with good material quality can be achieved by a stepwise or continuous change of the $\text{In}$ content (and thus the lattice constant) during growth of a ternary $\text{Al}_{x}\text{In}_{1-x}\text{As}$ or $\text{Ga}_{y}\text{In}_{1-y}\text{As}$ buffer layer and with different growth temperatures [3–7]. The growth of a relaxed buffer layer sequence provides a 'new substrate', and if the lattice constants of both $\text{Al}_{x}\text{In}_{1-x}\text{As}$ and $\text{Ga}_{y}\text{In}_{1-y}\text{As}$ were the same, one could grow on this new substrate lattice-matched heterostructures with an *arbitrarily chosen In content*. This work presents two buffer layer concepts to compensate the lattice misfit using the quaternary $\text{Al}_{x}\text{Ga}_{y}\text{In}_{1-x-y}\text{As}$ in a step and linear graded fashion, respectively. To preserve the advantage of GaAs substrate in terms of epitaxial cost, the buffer thickness was kept at 1 $\mu$m. The $\text{In}$ content of the layers was chosen to be $1-x=0.52$ and $1-y=0.53$ in order to have the same heterostructures grown lattice matched on InP as a reference.

2. Experimental procedure

The heterostructures were grown by molecular beam epitaxy (MBE) in a Varian Modular Gen II on InP substrates. The calibration of the growth rates was done by measuring RHEED intensity oscillations and performing beam equivalent pressure measurements before and after growth of the layers. With additional X-ray measurements (HRXRD) one is able to determine the composition of the lattice matched layers. Control of the substrate temperature was achieved with an Ircon Mod 2 infrared pyrometer. The substrates, GaAs (Freiberger Inc.) and InP (Sumitomo Inc.) were etched with $\text{H}_2\text{O} : \text{NH}_3\text{OH} : \text{H}_2\text{O}_2$ and $\text{H}_2\text{O} : \text{HCl} : \text{H}_2\text{O}_2$, respectively, before loading into the MBE system. On top of the buffer structure, which is kept at a thickness of 1 $\mu$m, the appropriate structures described below were grown. The growth rate of the buffer layers was 2 $\mu$m/h. The HEMT structure has a $\delta$-doping concentration of $5 \times 10^{12}$ cm$^{-2}$ separated by a 5 nm spacer layer from the 40 nm $\text{Ga}_{0.47}\text{In}_{0.53}\text{As}$ electron channel (Fig. 1). To obtain good morphological and electrical properties, the growth of a superlattice ($10 \times 4$ nm $\text{Al}_{0.44}\text{In}_{0.56}\text{As}/4$ nm $\text{Ga}_{0.47}\text{In}_{0.53}\text{As}$) prior to growth of the HEMT structure is necessary. The optimum growth temperature of the HEMT structure was at a thermocouple readout of 600°C (pyrometer readout: 550°C) for both, lattice matched growth on InP as well as lattice relaxed growth on GaAs. The quantum well (QW) structure for the optical studies consisted of five quantum wells with well thicknesses of 20, 10, 4, 3 and 1 nm grown at the same optimized growth temperature. Before growing the actual buffer layer for the linear grading beginning with $\text{Al}_{0.51}\text{Ga}_{0.49}\text{As}$, a 100 nm GaAs layer is grown. The grading was performed using computer controlled temperature ramping for Ga and In, finally conforming to the lattice constant of $\text{Al}_{0.48}\text{In}_{0.52}\text{As}$ (Fig. 1a). The step graded buffer consists of two layers beginning with 500 nm of $\text{Al}_{0.33}\text{Ga}_{0.31}\text{In}_{0.36}\text{As}$ and continuing with 500 nm of $\text{Al}_{0.48}\text{In}_{0.52}\text{As}$ (Fig. 1b).
3. Results and discussion

Focusing on the surface morphology, Fig. 2a shows an atomic force microscope (AFM) surface plot obtained from a linearly graded HEMT structure. The growth temperature $T_B$ of the buffer for this sample was 400°C (thermocouple readout). One observes a 'cross-hatched' morphology of the surface, which indicates the existence of misfit dislocations with a two-dimensional growth (2D) mode or layer by layer growth still present [8]. The 3D–AFM surface plot shows that the RMS...
roughness of the 'cross-hatched' structure in the (0 1 1) direction is significantly higher (RMS value: 2.6 nm) than that of perpendicular to this direction (RMS value: 3.1 nm). For the step-graded buffer the AFM surface plot is shown in Fig. 2b. The morphology of the surface is the result of a three-dimensional (3D) island growth (RMS value: 4.1 nm). An In content of 36% results in a major change of the lattice constant after which no 2D growth of the layers occurs. The 3D growth is caused by the quaternary Al$_{0.53}$Ga$_{0.31}$In$_{0.36}$As having a lattice mismatch beyond 2% with respect to GaAs [4]. As shown by Schweizer et al. [2] and Yao et al. [6], it is necessary for 2D growth of the layers to keep the In content clearly below 35%. This requirement cannot be fulfilled with all effusion cell temperatures fixed and with only one intermediate step in the lattice constant. The range between 300°C and 400°C for the buffer growth temperature $T_B$ has been found to yield nearly identical RMS roughness values for both step and linear grading with a slightly better surface morphology of the heterostructure utilizing the linear graded buffer. The lowest RMS roughness value of 2.6 nm was found at a substrate temperature $T_B$ of 350°C for the linear graded buffer [9]. The same HEMT structure grown lattice matched on InP yielded a RMS roughness of 0.2 nm.

Hall mobilities of the HEMT structures with both buffer types for 300 and 77 K are given in Fig. 3. The Hall mobility $\mu_{77 \text{ K}}$, plotted as a function of the temperature $T_B$, apparently shows higher values for the linear-graded structures with a maximum of 38 000 cm$^2$/V s (9800 cm$^2$/V s at 300 K) at 450°C. For the step-graded structures a maximum of 26 000 cm$^2$/V s (9200 cm$^2$/V s at 300 K) at a $T_B$ of 400°C (Fig. 3) is measured. An electron concentration of $3.0 \times 10^{12}$ cm$^{-2}$ was measured for all samples. A decrease of the mobilities for $T_B$ below 350°C and above 450°C is observed for both buffer types. The same HEMT structure grown lattice matched on InP yielded an electron mobility of 50 000 cm$^2$/V s at 77 K (11 000 cm$^2$/V s at 300 K), slightly better than the values achieved on the linear-graded buffer. This difference may be attributed to the large increase in surface roughness in the latter case, and therefore to an increase in the interface roughness of the electron channel.

The line width data of a series of Al$_{0.48}$In$_{0.52}$As/Ga$_{0.47}$In$_{0.53}$As QWs with well widths of 20, 10, 4, 3, and 1 nm are shown in Fig. 4. The superior interface quality of the QWs grown on the linear-graded buffer, compared to the growth on the step-graded buffer, is obvious from the reduced PL line width. Line widths obtained of the structures grown on linear-graded buffer have values between
17 and 33 meV (Fig. 4). The same structure grown on step-graded buffer yielded line widths above 35 meV. The maximum intensities of the PL spectra for the QWs grown on linear- and step-graded buffer layers each differ by a factor of ten compared to the QWs grown lattice matched on InP [9]. Together with the Hall mobility data the linear-graded buffer concept yielded results close to those achieved on InP. In addition, Al$_{0.48}$In$_{0.52}$As/Ga$_{0.47}$In$_{0.53}$As quantum wells grown lattice matched on InP show the lowest line widths in the range of 7–15 meV (Fig. 4) which are comparable to published data [10].

However, after the successful growth of Al$_{0.48}$In$_{0.52}$As/Ga$_{0.47}$In$_{0.53}$As heterostructures on GaAs which yielded the best results grown on a linear-graded lattice-relaxed buffer an experiment in terms of varying the thickness of the lattice relaxed buffer was performed. Intuitively, one would suggest an increase in electron mobility and a decrease in RMS roughness, respectively, by the use of very thick (2–4 μm) lattice-relaxed buffer layers due to fewer misfit dislocations. As it is clearly observed from Fig. 5, the growth of thicker buffer material only yields a small increase of the electron mobility and a small decrease in RMS roughness, respectively, but to a large expense in epitaxial material.

4. Conclusions

Two buffer layer concepts for the lattice relaxed growth of Al$_{0.48}$In$_{0.52}$As/Ga$_{0.47}$In$_{0.53}$As heterostructures on GaAs have been compared and related to the growth on InP. Electrical data yield the optimum growth temperature of the buffer at 400°C. Based on electrical and optical data, the two buffer concepts in their extreme linear- (Ga and In furnace temperatures are varied) and step-graded form (all furnace temperatures are constant) clearly show that it is favorable to vary the lattice constant during buffer growth as smoothly as possible. Increasing the total relaxed buffer thickness up to 4 μm only results into marginal improvements of electrical and morphological properties of the Al$_{0.48}$In$_{0.52}$As/Ga$_{0.47}$In$_{0.53}$As heterostructures.

References

Incorporation of As$_2$ in InAs$_x$P$_{1-x}$ and its application to InAs$_x$P$_{1-x}$/InP quantum well structures

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Abstract

We report studies on the incorporation of arsenic in InAs$_x$P$_{1-x}$ layers grown on InP substrates by solid-source MBE (SS-MBE) using group V valved cracker sources. Under phosphorus stabilised growth conditions we observe that As$_3$ is incorporated into InAs$_x$P$_{1-x}$ with near-unity efficiency up to As fractions ($x$) of about 0.4. For higher As$_2$ fluxes the incorporation is increasingly non-linear and the composition saturates at a value of $x \sim 0.8$. Similar behavior has been observed for the growth of InAs$_x$P$_{1-x}$ on InAs. The results are applied to the growth of InAs$_x$P$_{1-x}$/In(GaAs)P quantum wells and to graded composition InAs$_x$P$_{1-x}$/InP buffer layers. The results indicate excellent potential for optical device applications over the wavelength range 0.9–2.0 μm.

1. Introduction

InAs$_x$P$_{1-x}$/InP quantum well (MQW) structures have emerged as an important material system for long wavelength optoelectronics, at wavelengths such as 1.06, 1.3 and 1.55 μm [1–4]. A favourable strain-induced valence band splitting and large conduction band discontinuity ($\Delta E_C \sim 0.6$–0.7$\Delta E_G$) offers considerable advantages for lasers and optical modulators. Devices reported include 1.3 μm separate-confinement heterostructure (SCH) lasers with low threshold current density and high characteristic temperature ($T_0$) [5] and 1.3 μm electroabsorption waveguide modulators [6]. Tensile-strained InGaP can be used to replace InP as the barrier layer, offering increased electron confinement and suppression of strain relaxation in strain-balanced structures. In addition the relative ease which group V compositions can be varied, together with the relatively low strain difference between InAs and InP (+3.2%) makes InAs$_x$P$_{1-x}$ an attractive system for graded composition buffer layers. The growth of detectors or emitters on strain-relaxed buffer layers offers access to wavelengths beyond that of conventional InP-based heteroepitaxy.

From a more fundamental point of view it is interesting to contrast the properties of InAs$_x$P$_{1-x}$/InP, where the interface is formed due to substitution on the group V sub-lattice, with group III alloy QW systems and in particular with strained In$_x$Ga$_{1-x}$As/GaAs. Although high quality In$_x$Ga$_{1-x}$As/GaAs QW's have been widely reported there are problems, particularly for high
indium compositions, as a consequence of indium desorption and surface segregation. The effects can alter both indium content and the compositional profile of In₃Ga₁₋ₓAs quantum wells. In contrast the interfacial quality of the group V alloy QW system InAsₙP₁₋ₙ/InP may be influenced by intermixing or exchange between As and P and the complex incorporation behaviour of As₂ and P₂ makes compositional calibration at first sight seem very difficult.

2. Experimental procedure

Growth was carried out using a VG V80H MBE system equipped with group III effusion sources and group-V valved cracker cells. Elemental arsenic and red-phosphorus are evaporated at temperatures of 385 and 340°C, respectively, in the evaporation zone of the valved cracker and subsequently cracked by tantalum baffles in a high temperature zone (950°C). Growth was carried out under P-stabilised (2 × 4) conditions at a temperature of 480–500°C using a P₂ beam equivalent pressure ≈5 × 10⁻⁶ mbar. For the growth of InAsₙP₁₋ₙ, As₂ was co-evaporated during the growth of InP with the flux controlled by the valved cracker position. The performance characteristics of the EPI VCIII-As source are well documented. We find that the cracker cell provides a stable reproducible As₂ flux at a specific valve position. However steps must be taken to prevent the build-up of arsenic around the cracker zone by regular high temperature outgassing. The addition of a computer controlled servo-motor valve drive enables the flux to be changed from zero to a set value within a timescale ≈1 s.

For a fixed evaporation temperature (385°C) we have calibrated the arsenic flux as a function of cracker valve position by performing As-induced Reflection high-energy electron diffraction (RHEED) measurements on gallium terminated GaAs substrates at a temperature of 520°C. The results give a value for the incorporated arsenic flux in terms of monolayers of GaAs/ů. The indium flux is measured using ‘conventional’ group V stabilised RHEED oscillation measurements on InAs. Both the As and In growth rates are modified to take into account the differences between the host lattice constant and InP.

3. Arsenic incorporation in InAsₙP₁₋ₙ

It has long been known that arsenic incorporates much more readily than phosphorus into III–V compounds. The effect was first quantified by Foxon et al. using modulated beam techniques [7] who showed that probability of incorporation of As₄ into GaAs or InAs is up to 50 times higher than that of P₄. The behaviour is attributed to a significantly greater surface lifetime for As₄. Their results suggested a growth scheme for mixed group V alloys, such as InAsₙP₁₋ₙ, in which the supply of the most easily incorporated group V (in this case arsenic) is restricted with respect to the group III flux. The remaining group III sites are then available for incorporation with the weakly incorporated group V (phosphorus), which is provided in excess. The growth scheme has been exploited by Hou and Tu [8] who first suggested the use of As-induced RHEED measurement technique for in situ InAsₙP₁₋ₙ composition determination.

Fig. 1 shows the relationship between the incident (As/In) ratio and the incorporated arsenic fraction (x) in a series of InAsₙP₁₋ₙ/InP multi-quantum well (MQW) structures, with (x) determined by post-growth X-ray diffraction analysis of single thin bulk layers and multi-quantum well structures. Two regions are observed; for x ≤ 0.4, As₂ is incorporated with near-unity efficiency, whilst for higher x the dependency is increasingly nonlinear and the InAsₙP₁₋ₙ composition tends to an asymptotic value of around 0.8. Our results are very similar to those of Hou and Tu [8]. We find the As composition (x) of our layers is relatively invariant of variations in P₂ flux. This is fortunate since the stability of the phosphorus cracker source is not as good as that of the arsenic source. We have also investigated the growth of thin InAsₙP₁₋ₙ multilayers on InAs substrates, for which we used (As/In) values of up to, and exceeding, unity.

Despite the presence of sufficient As₂ to account for all of the available group III sites, we find that arsenic compositions of between 0.7 and 0.8 are obtained, or conversely that P₂ incorporates into
InAs at fractions of between 0.2 and 0.3. The arsenic fractions are slightly higher than those achieved on InP for similar (As/In) values. The results indicate a rapid reduction in the As\textsubscript{2} sticking coefficient at high surface coverage.

4. Optical properties of multi-quantum well p-i-n diodes

To investigate the optical and electrical properties of InAs\textsubscript{1-x}P\textsubscript{x}/InP QW's a number of p-i(MQW)-n structures have been grown with values of arsenic composition from \( x = 0.15 \)–0.66. Table 1 lists the details of several of these structures, grouped together as structures with room temperature exciton peak positions close to the wavelengths 1.06, 1.3 and 1.55 \( \mu \)m. These are the wavelengths of technological interest for optoelectronic devices. All the structures had 80 Å QW thicknesses. The samples differ however in terms of barrier material, barrier thickness and total number of periods. In the absence of strong relaxation, InAs\textsubscript{1-x}P\textsubscript{x}/InP quantum well structures exhibit intense, narrow, room temperature photoluminescence (PL) emission. Fig. 2 shows relative PL data from several samples and emphasizes the wide wavelength range that can be covered by InAs\textsubscript{1-x}P\textsubscript{x} QW's. The data includes for example, samples \#1108 and \#1125 in Table 1 which provide a comparison between 1.3 \( \mu \)m strained InP and strain-balanced InGaP barriers. The strain balanced 1.55 \( \mu \)m (\#1217) is also shown in the figure. PL linewidths as low as 12 meV at room temperature and 3.5 meV at low temperature have been observed in the 1.3 \( \mu \)m structures.

Electrical measurements have been carried out on p-i(MQW)-n+ samples fabricated into 200 \( \mu \)m diameter mesa diodes. Typical reverse leakage values of \( I_R \) < 10 nA at -30 V are observed in the low strain \( \lambda \approx 1.06 \mu m \) samples. For the \( \lambda \approx 1.3 \mu m \) samples with InP barriers, barrier thickness \( \geq 400 \) Å and \( \leq 20 \) periods were required to obtain reasonable

![Graph showing As fraction (x) in InAs\textsubscript{1-x}P\textsubscript{x} as a function of incident flux ratio.](image)

**Table 1**

InAs\textsubscript{1-x}P\textsubscript{x}/In(Ga)AsP multiquantum well structures

<table>
<thead>
<tr>
<th>#</th>
<th>( d_{bar} ) (Å)</th>
<th>Barrier</th>
<th>Periods</th>
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<th>300 K Photoluminescence</th>
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<td>11</td>
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</table>
dark currents. Values of $I_R$ of 1 mA at $-10$ V are fairly typical of these structures. Strain-balanced InAs$_{0.41}$P$_{0.69}$/In$_{0.96}$Ga$_{0.04}$P MQWs show $I_R$ values one or two orders of magnitude lower than those with InP barriers.

Although the wavelength of InAs$_x$P$_{1-x}$ MQW's can be extended to 1.55 $\mu$m ($x \approx 0.36$) and the optical properties of these structures are excellent (300 K linewidths of 18–20 meV) the reverse leakage of the devices is rather high. The values are $\sim 1$ mA at $-20$ V for the InP barrier structure (#1216) and $\sim 50$ $\mu$A at $-20$ V for the InGaP barrier sample (#1217). These high reverse leakage values may preclude the use of such structures in applications such as optical modulators.

5. Structural characterisation

High resolution X-ray diffraction has been used to assess well composition and strain relaxation in these samples, using (0 0 4) and high/low incidence asymmetric (1 1 5) reflections. (0 0 4) reflection data from several InAs$_x$P$_{1-x}$/InP MQW structures is shown in Fig. 3. The data also shows the InAs-based structure, which has thin InAs$_x$P$_{1-x}$ tensile barriers, and which was described in relation to the As incorporation measurements of Fig. 1. The data shows the excellent structural quality of InAs$_x$P$_{1-x}$/InP which is at least comparable to In$_{0.4}$Ga$_{0.6}$As/GaAs. Values of relaxation obtained from X-ray measurements are given in Table 1. It can be seen that the level of relaxation increases with increased $x$, with reduced barrier width or with increasing number of periods. Plan view TEM of wedge shaped specimens has been used to image the buffer-MQW, MQW and MQW-cap interface planes of these structures. Even in the most highly relaxed samples no dislocations were observed within the MQW region. Misfit dislocation arrays were present at the buffer-MQW and at the MQW-cap interfaces, with the latter at a much lower density. The behaviour is normal for low strain MQW's, with partial relaxation occurring at the buffer-MQW interface whilst the MQW itself remains coherent [9]. The MQW acts as a single strained layer of average well-barrier composition. The relaxation data can be well explained using the geometrical theory of strain relaxation developed for single strained layers [10]. The model predicts accurately, for example, a critical layer thickness (CLT) of 420 nm for the InAs$_{0.23}$P$_{0.77}$ As/InP MQW's which have an average strain ($\epsilon$) = $3.1 \times 10^{-3}$. This value of CLT is almost 10 times higher.
than that predicted by the Matthews and Blakeslee single-kink model for the same strain value [11].

6. Compositional graded buffer layers

Graded InAs$_x$P$_{1-x}$ compositions can be easily grown by slowly ramping the position of the arsenic cracker valve, the variation of which is approximately linear with composition. We have grown strain-relaxed p-i-n diode structures on linear graded InAs$_x$P$_{1-x}$ buffer layers. For the buffer a grading rate of $x = 0.2/\mu$m was used. The thickness of the buffer layers varies from 2.5 to 4.8 $\mu$m. After the buffer layer a step back in composition of $x \approx 0.07$ is performed to cancel the residual strain in the buffer layer. The design scheme for the linear graded buffer follows that of reference [12]. On top of the buffer layer a p-i-n diode is grown at the relaxed InAs$_x$P$_{1-x}$ composition. The structure is 1.5 $\mu$m Si-doped (n$^-$), 1.5 $\mu$m intrinsic region and 0.5 $\mu$m Be-doped (p$^+$) InAs$_x$P$_{1-x}$. A series of structures has been grown with compositions ($x$) of 0.32–0.85. The final compositions, as measured by post-growth X-ray analysis, are within $x \approx 0.05$ of the target composition from calibration of the valved cracker source.

Fig. 4 shows room temperature PL from a number of graded p$^+$-i-n$^+$ diode structures. The PL peak wavelength reaches 2.32 $\mu$m for the highest composition, $x \approx 0.85$. The PL wavelength is in good agreement with the expression $E_g(300 \text{K}) = 1.351 - 1.315x + 0.32x^2$ eV for relaxed InAs$_x$P$_{1-x}$ layers in reference [13]. Room temperature PL linewidths are typically 35–40 meV. We can further increase the wavelength by growing In$_x$Ga$_{1-x}$As lattice-matched to the relaxed buffer layer. We have grown In$_{0.15}$Ga$_{0.85}$As structures on relaxed InAs$_{0.7}$P$_{0.3}$ buffer layers which have PL emission at wavelengths $\geq 2.5 \mu$m.

Since the primary interest in such structures is for mid-IR detectors the reverse dark currents need to be fairly low. Analysis of 200 $\mu$m mesa diode structures shows a general increase in dark current and degradation in breakdown as the composition in increased. Diodes with $x \approx 0.32$ and 0.42 show dark currents in the range 1–100 nA at $-10 \text{ V}$, whilst for example for the $x \approx 0.7$ sample the value is in excess of 100 $\mu$A for the same voltage. The increase suggests that threading dislocations are not solely confined to the mismatched buffer layer in the higher strain samples and that some further improvement may be required to use these structures in applications such as avalanche photodetectors. For $x < 0.5$ however the performance is quite satisfactory and compares well with previous reports of planar pn diodes grown on generally much thicker buffer layers by MOVPE or VPE techniques [14, 15].

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References

Growth of abrupt GaAs/Ge heterointerfaces by atomic hydrogen-assisted molecular beam epitaxy

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Abstract

High-quality epitaxy of GaAs-on-Ge system has been developed at low growth temperatures around 400°C by atomic hydrogen-assisted molecular beam epitaxy (H-MBE). The segregation of Ge is significantly reduced and abrupt GaAs/Ge heterointerface is successfully formed. Using reflection high-energy electron diffraction (RHEED), secondary ion mass spectroscopy (SIMS) and atomic force microscopy (AFM) as characterization tools, it is demonstrated that enhanced layer-by-layer two-dimensional growth mode is realized from the earliest stages of growth and evolution of single-domain (2 × 4) GaAs(0 0 1) is achieved on vicinal Ge substrates at 400°C by H-MBE technique. The modification of growth mode and consequent improvement in material quality are explained at the simplest level by the surfactant properties of atomic H in MBE growth of GaAs.

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Keywords: MBE; Surfactant-mediated epitaxy; Heteroepitaxy; Atomic hydrogen; Segregation; Growth mode; Surface reconstruction; GaAs-on-Ge; RHEED; AFM; SIMS

1. Introduction

GaAs-on-Ge material is gaining particular interest for application to low-cost, high-efficiency solar cells [1]. Not only is the lattice-mismatch between GaAs and Ge favorably small ( < 0.1%), the difference in their thermal expansion coefficients $\Delta \alpha$ is also small. This is in contrast with a more popularly studied GaAs-on-Si system (lattice-mismatch of $\sim 4.0\%$ and $\Delta \alpha \approx 60\%$). However, in GaAs-on-Ge epitaxy, a serious problem of segregation and interdiffusion of Ge and As at the GaAs/Ge heterointerface arises when GaAs is grown on Ge substrates at high temperatures ($> 450^\circ$C) [2]. Consequently, the initial stages of growth generally proceed by a three-dimensional (3D) islanded growth mode, thereby resulting in the degradation of surface flatness and interface abruptness [3, 4].

In this work, high-quality epitaxy of GaAs-on-Ge is achieved at a lower growth temperature of
~400°C by atomic hydrogen-assisted molecular beam epitaxy (H-MBE), in which atomic H is generated by cracking of molecular hydrogen at a hot tungsten filament heated to > 1500°C inside the MBE growth chamber. It is shown that Ge segregation is minimized and layer-by-layer two-dimensional (2D) growth mode is realized from the early stages of growth. Previously, we showed that atomic H acts as a surfactant in MBE growth of GaAs, where both the kinetics and energetics of growth can be modified in an accurately controlled manner [5–8]. The modification and control of elementary processes of growth at an atomic scale has historically been difficult to accomplish, until when the concept of surfactant epitaxy was first reported in SiGe/Si heteroepitaxy using As as a surfactant [9]. The use of atomic H is regarded as favorable since self-doping, or auto-doping could be a problem with other surfactant elements, and from a technological viewpoint, it is advantageous that sources of atomic H can be easily installed into MBE systems. Furthermore, irradiation of atomic H during MBE growth brings a number of additionally favorable effects, such as in-situ surface cleaning and inactivation of deep-level defect centers [7, 8]. As a result, layer-by-layer 2D growth and high material quality can be achieved even for the MBE growth of highly lattice-mismatched systems, and are recently reported for GaAs/Si [7, 8], SiGe/Si [9] and other important materials.

Using reflection high-energy electron diffraction (RHEED), atomic force microscopy (AFM) and secondary ion mass spectroscopy (SIMS) as characterization tools, it is demonstrated that H-MBE growth results in significant reductions of Ge segregation and Ge/As interdiffusion in GaAs-on-Ge heteroepitaxy. Hence, atomically smooth surfaces and abrupt GaAs/Ge heterointerfaces are formed as required for a variety of applications. Also the mechanisms responsible for the observed reductions of Ge segregation in H-MBE technique are discussed in terms of surfactant properties of atomic H.

2. Experimental procedure

Continuous generation of atomic H inside our MBE growth chamber was achieved with a self-assembled cracker that fits into one of the effusion cell ports of the chamber. The hydrogen cracker consists of spirally wound tungsten filament, which when heated ≥1500°C, dissociates molecular hydrogen into atomic H through catalytic process with cracking ratios of ≥1% [10]. The surface structure was monitored by RHEED during the substrate cleaning and growth. The surface morphologies and roughness were characterized by AFM in ambient atmosphere and depth profiles of compositional elements were measured by SIMS with Cs+ ions with an acceleration voltage of 1 kV.

Vicinal Ge(0 0 1) substrates tilted 2° off toward [1 1 0] azimuth were arbitrarily chosen for this work. Before the deposition of GaAs, each Ge substrate was first cleaned in the organic solvents followed by a wet-etch in H2SO4 : H2O2 : H2O = 5 : 1 : 1 in volume. Once placed in the chamber, the native oxide was removed either by atomic H irradiation at ~350°C for 15 min, or by conventional thermal cleaning at 550°C. We found reproducibly that atomic H cleaning results in a more reliable surface with improved flatness than by thermal cleaning [11]. The growth was then carried out under various growth parameters. The growth temperature was varied from 250°C to 600°C, though the temperature of greatest interest was around 400°C in order to minimize the Ge segregation and interdiffusion of Ge and As while achieving as high a material quality as possible. The growth rate was between 0.1 and 1.0 µm/h, and H2 backpressure was kept constant at 6 × 10^-6 Torr by manual control of a variable leak-valve, both during substrate cleaning and MBE growth.

3. Results

Fig. 1 summarizes the changes of RHEED pattern measured for Ge(0 0 1) and GaAs during MBE growth for both with atomic H (H-MBE), and without atomic H, i.e., conventional MBE. The growth temperature and growth rate were 400°C and 0.5 µm/h, respectively. First, the RHEED pattern observed after the substrate cleaning was typically a (2 × 2) streak or two orthogonal (2 × 1) reconstructions indicative of double-domain structure regardless of the cleaning method, atomic H or
Fig. 1. Changes of RHEED pattern of GaAs(0 0 1) with growth measured for H-MBE and conventional MBE. The growth temperature and growth rate are 400°C and 0.5 μm/h, respectively.

thermal cleaning. But, the streak was sharper and brighter for the atomic H-cleaned samples showing improved surface flatness compared to thermal cleaning. The improvement in surface flatness of Ge substrates after atomic H cleaning was also confirmed by AFM measurements. Second, the RHEED pattern changed from (2 × 2) streak to a spotty (1 × 1) pattern immediately after the onset of MBE growth without atomic H. It gradually recovered to a streaky (2 × 2) pattern after continued growth of ~1200 ML. Third, the RHEED pattern changed from clear (2 × 2) streak to a (2 × 2) spotty streak during ~5 ML deposition for H-MBE case. After the growth of ~150 ML, it recovered to a clear streak pattern, but this time, evolution of single-domain (2 × 4) reconstruction was obtained. As the RHEED results show, streak diffraction pattern is maintained and layer-by-layer 2D growth is enhanced by H-MBE. However, it is noted that RHEED patterns were strongly dependent on the growth rate of GaAs. The recovery to a streak after becoming spotty during the initial stage of deposition was faster for higher growth rates.

Fig. 2 plots the results of average surface roughness measured by AFM for GaAs grown on Ge substrates by H-MBE and MBE. The measurements were performed in air immediately after cooling the samples rapidly down to room temperature. The growth temperature and growth rate were 400°C and 0.5 μm/h, respectively, in both cases. It can be observed that for H-MBE case as marked by open circles, the surface roughness does not increase appreciably with growth and is ~0.6 nm after 1800 ML of deposition, which corresponds to about 2 ML thickness roughness. On the other hand, surface roughing worsens with growth for the MBE sample as marked by closed triangles. The average roughness is as much as ~3.0 nm after 1800 ML deposition, which therefore consist of 3D multilayers of ~10 ML thickness. Thus, it is shown that the ideal layer-by-layer growth achieved by H-MBE technique as from the RHEED analysis in Fig. 1, also results in significantly improved surface smoothness.

Figs. 3 and 4 show the SIMS depth profiles of compositional atoms, Ge, Ga and As around the GaAs/Ge heterointerface measured for samples grown by H-MBE and MBE, respectively. The growth temperature, growth rate, and film thickness in both cases were 400°C and 0.5 μm/h, and 1.0 μm, respectively. It is clearly observed that Ge segregation into GaAs layer and diffusion of As into Ge substrate are more severe in MBE samples even for a relatively low growth temperature of 400°C. On the other hand, the interface abruptness is significantly improved in H-MBE samples. For example, it is seen that after a fast initial decrease of Ge concentration at the GaAs/Ge interface with a slope of ~7.6 nm/dec, which gives
the resolution limit of our SIMS measurement, the slope levels off to $\sim 17$ nm/dec at a concentration of $\sim 1.4 \times 10^{20}$ cm$^{-3}$ at the bending point in the MBE sample. In comparison, the slope is unchanged for the H-MBE sample and Ge concentration drops abruptly down to a measurement limit of $\lesssim 2.2 \times 10^{18}$ cm$^{-3}$. Furthermore, both the Ge concentration at the bending point and level-off slope in the SIMS profiles were found to be smaller for H-MBE samples than for MBE in a temperature range of 330–560°C, as plotted in Fig. 5. It is also noted that the Ge segregation decay length, instead of reducing exponentially with temperature, which one would expect because of kinetic/thermal limitation imposed to segregation process, showed a minimum around 400°C. Lastly, it was observed, as shown in Fig. 6, that irradiation of atomic H during MBE growth reduces oxygen concentration in GaAs layer to below measurement limit of $9.0 \times 10^{16}$ cm$^{-3}$ for H-MBE sample grown at 400°C. Hence, reduction of oxygen-related defect
centers is expected, though a detailed analysis needs to be performed to be more quantitative.

4. Discussion

As presented, Ge segregation and Ge/As interdiffusion are significant even for a low growth temperature of 400°C by MBE. Though the segregation decay length depends strongly on growth conditions such as growth rate, it is shown that Ge segregation can be reduced significantly for the given conditions, by the use of atomic H continuously irradiated on the surface during MBE growth. It is known that segregation process is generally governed by the strength of bonding energy [3], the surface, interface as well as strain energies associated with heteroepitaxial system as a whole [9], and the densities of dislocations and types of defects present in the system. Kroemer also pointed out the importance of charge neutrality fulfillment in polar-on-nonpolar heteroepitaxy [12]. Our aim for introducing atomic H into MBE growth is to try to modify and control the growth mode by modifying the surface energetics and kinetics in such a manner that layer-by-layer 2D nucleation mode would be promoted and hence segregation and interdiffusion be minimized. Such a growth concept known as surfactant epitaxy, is attracting increasing attention in recent years and widely reported for GaAs/Si [7, 8], and SiGe/Si [9] systems.

Recently, we reported on the basic analysis of atomic-scale growth mechanisms for homoepitaxial MBE of GaAs using atomic H as a surfactant [5]. In conventional MBE, the surface free energy of Ge(0 0 1) being lower than the barely As-stabilized GaAs(0 0 1) surface favors 3D islanded growth and Ge segregation and Ge/As interdiffusion are accelerated even for low substrate temperatures. In H-MBE growth of GaAs-on-Ge, it is thought that the effect of reduction of surface energy of H-adsorbed GaAs(0 0 1) is predominant and, consequently, a preference for 2D layer-by-layer growth mode is provided, whereby successfully resulting in atomically smooth and abrupt heterointerfaces as required for a variety of device fabrication. It is also noted that the growth model discussed here should be applicable only in a relatively limited range of growth temperature. This is because the desorption rate of atomic H from GaAs(0 0 1) surface is found to increase rapidly above ~450°C and the growth kinetics becomes thermally limited and crystal quality becomes poor below ~300°C, even though the effectiveness of atomic H is still observable at these temperatures. And this is probably the reason for Ge segregation decay length, as plotted in Fig. 4, to have a minimum point at ~400°C, though additional work is necessary to clarify completely the role of atomic H. Finally, in situ removal of foreign contaminants such as carbon and oxygen during H-MBE also serve to suppress heterogeneous nucleation and formation of impurity-related defect centers.

5. Summary

We have developed high-quality epitaxy of GaAs-on-Ge system at low growth temperatures by H-MBE technique. It is demonstrated that enhanced layer-by-layer 2D growth mode is realized from the earliest stages of MBE growth and single-domain (2 x 4) GaAs(0 0 1) surface is evolved at 400°C by H-MBE. The Ge segregation and Ge/As interdiffusion are reduced and an abrupt GaAs/Ge heterointerface is successfully formed. In H-MBE growth of GaAs-on-Ge, the reduction of surface energy of H-adsorbed GaAs(0 0 1) seems to be the predominant effect that as a result, provides preference for 2D layer-by-layer growth mode. Modification of growth mode and consequent improvements in material quality presented here are explained at the simplest level by the surfactant properties of atomic H in MBE growth of GaAs.

References


Growth and characterization of In$_2$Se$_3$ epitaxial films by molecular beam epitaxy

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Abstract

Structural control of In$_2$Se$_3$ films was attempted by molecular beam epitaxy (MBE). In$_2$Se$_3$ epitaxial films with layered structure based on zincblende structure were successfully grown on (0 0 1)GaAs substrates at low temperatures ($T_s \leq 550{\text{C}}$ at VI/III ratio of 10) and at low VI/III ratios (VI/III $<$ 30 at 500{C}), and polycrystalline In$_2$Se$_3$ films with defect wurtzite structure were grown at high temperatures ($T_s \geq 540{\text{C}}$ at VI/III ratio of 10) and high VI/III ratios (VI/III $\geq 90$ at 500{C}). In the epitaxial In$_2$Se$_3$ films with layered structure, the direction of the $c$-axis corresponded to [1 1 0] and [1 1 1] directions in the (0 0 1)GaAs substrates. Furthermore, exciton emission located at around 579 nm was observed in the photoluminescence spectra of the In$_2$Se$_3$ films at 4.2 K.

1. Introduction

III$_2$–VI$_3$ compound semiconductors have several crystal structures. For example, Ga$_2$Se$_3$ has a defect zincblende structure, but $\frac{1}{3}$ of cation sites are vacant. Up until now, we investigated the molecular beam epitaxial (MBE) growth of Ga$_2$Se$_3$ films on (0 0 1)GaAs and (0 0 1)GaP substrates, and it was found that a spontaneous superlattice was formed and the unique properties such as large optical anisotropy were shown by the ordering of the native Ga vacancies in Ga$_2$Se$_3$ films [1–6]. On the other hand, In$_2$Se$_3$ has two types of crystal structure different from that of Ga$_2$Se$_3$, i.e., layered structure ($\alpha$-phase) and defect wurtzite structure ($\gamma$-phase) [7–9]. The layered structure consists of rather loose stacks of covalently bonded layers, including five atomic layers of Se–In–Se–In–Se, by van der Waals force. The basic structure of the layered In$_2$Se$_3$ is wurtzite or zincblende structure. Therefore, it is considered that the native In vacancies form planes in every three In planes in the layered In$_2$Se$_3$ and the layered In$_2$Se$_3$ is a kind of vacancy-ordered structure. Meanwhile, the defect wurtzite structure is basically a wurtzite structure, but $\frac{1}{3}$ of the cation sites are vacant as well as the defect zincblende structure. In this structure, the In vacancies take screw arrangement. Therefore, $\gamma$-In$_2$Se$_3$ shows large optical rotary power [9]. However, there are no reports on the epitaxial
growth of In$_2$Se$_3$ films. In this paper, we attempted the structural control and characterization of In$_2$Se$_3$ films on (0 0 1)GaAs substrates, and In$_2$Se$_3$ epitaxial films with layered structure based on zincblende structure were successfully grown on (0 0 1)GaAs substrates. This is the first report on the successful growth of In$_2$Se$_3$ epitaxial films. Furthermore, we observed an exciton emission located at around 579 nm in the low-temperature photoluminescence spectra of the In$_2$Se$_3$ films for the first time.

2. Experimental procedure

In$_2$Se$_3$ films were grown on SI-(0 0 1)GaAs substrates by MBE technique using EIKO Engineering Model EV-100. The growth temperature was varied in a range of 450–540 °C. VI/III ratio was varied from 5 to 90 by increasing Se beam equivalent pressure (BEP), keeping In BEP constant. In these growth conditions, the growth rate was limited by the supply of In atoms. The growth rate of In$_2$Se$_3$ films was around 800 nm/h, and the film thickness was around 800 nm. Before being inserted into the MBE system, the GaAs substrates were degreased, chemically etched in a 3 : 1 : 1 H$_2$SO$_4$ : H$_2$O$_2$ : H$_2$O solution at 60 °C for 1.5 min. After acid etching, the GaAs substrates were heated at 550 °C for 15 min in the growth chamber to remove surface oxides. The [1 1 0] and [1 1 0] direction of the GaAs substrates were determined by anisotropic etching. The crystallinity of In$_2$Se$_3$ films was characterized by reflection high-energy electron diffraction (RHEED) and Raman spectroscopy. An Ar$^+$ laser (514.5 nm, 50 mW) was used for Raman spectroscopy. Surface morphology and cross-sectional image were observed by scanning electron microscopy (SEM). Photoluminescence (PL) measurement was carried out at 4.2 K. The 488.0 nm emission line of an Ar$^+$ laser was used as an excitation light.

3. Results and discussion

First of all, we investigated the effects of VI/III ratio on crystal structure of In$_2$Se$_3$. Fig. 1 shows RHEED patterns of In$_2$Se$_3$ films for various VI/III ratios. The growth temperature was kept at 500°C. At the VI/III ratios below 30, zincblende-type spots are observed for [1 1 0] incidence. This result indicates that epitaxial In$_2$Se$_3$ films with zincblende structure were successfully grown on GaAs substrates. For [1 1 0] incidence, spot pattern is not observed. This result is considered to be attributed to the surface morphology, which will be discussed later. On the other hand, at VI/III ratio of 90, spot pattern is not observed. We confirmed by X-ray diffraction that the film prepared at VI/III ratio of 90 was polycrystalline.

In order to clarify the crystal structure of the In$_2$Se$_3$ films, we characterized the In$_2$Se$_3$ films by Raman spectroscopy. Fig. 2 shows Raman spectra of In$_2$Se$_3$ films for various VI/III ratios. At VI/III ratios below 30, a peak located at 108 cm$^{-1}$ is observed. This peak appears at the same wavelength as in the reported Raman spectrum of the bulk α-In$_2$Se$_3$ with layered structure [10]. Therefore, In$_2$Se$_3$ epitaxial films are found to have layered structure based on zincblende structure. A peak located at around 205 cm$^{-1}$ in the films with VI/III ratios below 30 was also observed in the Raman spectrum of the bulk α-In$_2$Se$_3$ [10]. On the other hand, at VI/III ratio of 90, a peak is located at 151 cm$^{-1}$, which is the same wavelength as observed in the spectrum of the bulk γ-In$_2$Se$_3$ with defect wurtzite structure [10]. This result indicates that polycrystalline In$_2$Se$_3$ with defect wurtzite structure was obtained under high VI/III ratio conditions. Besides, we investigated the growth of In$_2$Se$_3$ films with VI/III ratio below 5. In the growth conditions, the growth rate of In$_2$Se$_3$ decreased with decreasing VI/III ratio, and no deposition of In$_2$Se$_3$ was observed with VI/III ratio below 1. This result can be considered to be attributed to the reevaporation of In and Se compounds with high vapor pressure such as In$_2$Se, and is similar to the MBE growth of Ga$_2$Se$_3$ films [1].

Furthermore, we observed surface morphology and cross-sectional image of the epitaxial In$_2$Se$_3$ film by scanning electron microscopy (SEM). Fig. 3 shows the surface morphology and the (1 1 0) cross-sectional SEM image of the epitaxial In$_2$Se$_3$ film grown at 500°C with VI/III ratio of 10. A schematic drawing of (1 1 0) cross-sectional view of the
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Fig. 1. RHEED patterns of In$_2$Se$_3$ films for various VI/III ratios.

epitaxial In$_2$Se$_3$ with layered structure on (0 0 1)GaAs substrate is also shown in Fig. 3. In the layered In$_2$Se$_3$, c-axis corresponds to [1 1 1] direction in zincblende structure. In the cross-sectional image, it is found that the layered structure is formed and the direction of c-axis corresponds to [1 1 1] and [1 1 1] direction in the (0 0 1)GaAs substrate. This structure was not observed in (1 0 0) cross-sectional image, which indicates that c-axis of the layered In$_2$Se$_3$ is parallel to [1 1 0] direction. In the surface morphology of the epitaxial In$_2$Se$_3$ film, a lot of lines parallel to the [1 1 0] direction are observed. This morphology is considered to be attributed to the a-axis direction. These features are similar to the epitaxial GaSe films on (0 0 1)GaAs substrates [11, 12]. We consider that triangular ridges run along [1 1 0] direction and that the anisotropy in the RHEED patterns is attributed to the anisotropic features in the surface structure. The details are still under study.
Next, we investigated growth temperature dependence. Fig. 4 shows the dependence of Raman spectrum on growth temperature. VI/III ratio was kept at 10. At the temperatures below 520°C, the peak located at 108 cm⁻¹, which originates in In₂Se₃ with layered structure, is observed. In these films, spot pattern based on zincblende structure was observed in RHEED patterns. These results suggest that epitaxial In₂Se₃ films with layered structure based on zincblende structure are formed at the temperature below 520°C. On the other hand, the peak located at 151 cm⁻¹, which originates in In₂Se₃ with defect wurtzite structure is observed at the temperature of 540°C. A peak located at around 180 cm⁻¹ was also observed in the Raman spectrum of the bulk γ-In₂Se₃ [10]. In this film, ring pattern was observed in RHEED pattern, which indicates that the polycrystalline In₂Se₃ with defect wurtzite structure is obtained at high temperature. From these results, it is found that the epitaxial In₂Se₃ films with layered structure are obtained at low temperatures and low VI/III ratios and that the polycrystalline In₂Se₃
films with defect wurtzite structure are obtained at high temperatures and high VI/III ratios.

Furthermore, we investigated the optical properties of the epitaxial In$_2$Se$_3$ films by PL measurement at 4.2 K. Fig. 5 shows PL spectrum of the In$_2$Se$_3$ epitaxial film grown at 520°C with VI/III ratio of 10. Detailed spectrum in the band edge region is shown in the inset. A sharp peak located at around 579 nm (2.14 eV) is observed in the epitaxial In$_2$Se$_3$ films. This peak is also observed in the polycrystalline γ-In$_2$Se$_3$ films. In order to clarify the origin of the PL emission, we measured wavelength dependence of photoconductivity of the polycrystalline γ-In$_2$Se$_3$ film on glass substrate at 4.2 K. It was found that a peak due to an exciton absorption was observed near absorption edge in the wavelength dependence profile and that the wavelength of the exciton absorption peak was almost the same as that of the PL emission peak. Therefore, the PL emission peak located at around 579 nm is found to be exciton emission. Furthermore, intensity of deep level emission peak observed at around 900 nm is very weak. A peak located at around 830 nm is an emission from GaAs substrate. These results suggest that the quality of the In$_2$Se$_3$ films is quite good in terms of optical properties. This is the first report on the exciton emission in the In$_2$Se$_3$. Moreover, the band gap of In$_2$Se$_3$ estimated from Fig. 5 is about 2.15 eV at 4.2 K. This value is larger than the reported band gap of the bulk α-In$_2$Se$_3$ with layered structure ($E_g \approx 1.4$ eV) [13, 14], but almost the same as the reported γ-In$_2$Se$_3$ films with defect wurtzite structure ($E_g \approx 2.0$ eV) [15]. Therefore, we can consider that the band gap of In$_2$Se$_3$ with layered structure is larger than the reported value.

4. Conclusions

We attempted the structural control of In$_2$Se$_3$ films by molecular beam epitaxy (MBE). In$_2$Se$_3$ epitaxial films with layered structure based on zincblende structure were successfully grown on (0 0 1)GaAs substrates at low temperatures ($T_g \leq 520$°C at VI/III ratio of 10) and at low VI/III ratios (VI/III = 30 at 500°C), and polycrystalline In$_2$Se$_3$ films with defect wurtzite structure were grown at high temperatures ($T_g \geq 540$°C at VI/III ratio of 10) and high VI/III ratios (VI/III = 90 at 500°C). In the epitaxial In$_2$Se$_3$ films with layered structure, the direction of c-axis corresponded to [1 2 1] and [1 1 1] direction in the (0 0 1) GaAs substrates. Furthermore, exciton emission located at around 579 nm was observed in the photoluminescence spectra of the In$_2$Se$_3$ films at 4.2 K. This is the first report on the exciton emission in the In$_2$Se$_3$.

References

Effects of strain on the growth and properties of CuInSe₂ epitaxial films

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Abstract

CuInSe₂ films have been grown on both GaAs(0 0 1) and In₀.₂₅Ga₀.₇₅As pseudo-lattice-matched substrates by solid-source molecular beam epitaxy and the effects of strain on the properties of epitaxial films have been investigated. For CuInSe₂ epitaxial films grown on GaAs(0 0 1), the intensities of excitonic emissions become stronger and the line widths of the predominant defect-related emissions become smaller on increasing the thickness of the films from 0.4 to 2 μm, suggesting that the defect density decreases with the film thickness. Drastic improvements in optical and transport properties have been achieved by using In₀.₂₅Ga₀.₇₅As pseudo-lattice-matched substrates; free-exciton emissions were predominantly observed, indicating that high-quality CuInSe₂ epitaxial films have been grown by controlling the misfit strain between the substrates and the epitaxial films.

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Keywords: Molecular beam epitaxy; Photoluminescence; X-ray diffraction; Transmission electron microscopy; CuInSe₂

1. Introduction

CuInSe₂ (CIS)-based chalcopyrite semiconductor solar cells have already demonstrated high conversion efficiencies of up to η = 17.7% [1]. Thin films with thickness of only 2–3 μm are required to absorb sunlight effectively and no degradation of solar cell performance for more than 4 years has been reported [2], suggesting that CIS-based solar cells are promising for practical low-cost thin film solar cells. CIS-based poly-crystalline solar cells typically consist of a ZnO window layer, a CdS buffer layer, and a CIS or CuInGaSe₂ (CIGS) absorber layer grown, in turn, on Mo-coated soda-lime glass substrates [1]. Significant mismatch in lattice constants, thermal expansion coefficients as well as crystalline structure still exist among the layers as shown in Table 1 despite a remarkable improvement reported in solar cell performance. The effects of such a mismatch on the properties of CIS and consequently on the characteristics of the

*Corresponding author.
Table 1
Crystalline structure, lattice constants, and thermal expansion coefficients of the materials used for fabricating CuInSe$_2$-based solar cells

<table>
<thead>
<tr>
<th>Material</th>
<th>Crystalline structure</th>
<th>Lattice constants (Å)</th>
<th>Thermal expansion coefficients ($\times 10^{-6}$ K$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CuInSe$_2$</td>
<td>Chalcopyrite</td>
<td>$a = 5.7821$, $c = 11.6191$</td>
<td>$\alpha_a = 11.4$, $\alpha_c = 8.6$</td>
</tr>
<tr>
<td>CdS</td>
<td>ZnSblende</td>
<td>$a = 5.825$</td>
<td>$\alpha_a = 6.5$, $\alpha_c = 4.0$</td>
</tr>
<tr>
<td>ZnO</td>
<td>Wurtzite</td>
<td>$a = 4.136$, $c = 6.714$</td>
<td>$\alpha_a = 4.75$, $\alpha_c = 2.92$</td>
</tr>
</tbody>
</table>

Solar cells cannot be ignored for further extending the solar cell performance and for manufacturing large-size reproducible modules.

In our earlier work, we have reported the growth and properties of CIS films grown by molecular beam epitaxy (MBE) [3]. Sharp photoluminescence (PL) spectra including near-band-edge emissions have been observed from the CIS epitaxial films grown on GaAs(0 0 1) though defect-related emissions were still dominant [4]. CIS has a chalcopyrite structure with the lattice constants of $a$-axis ($a$) and $c$-axis ($c$) being $a = 5.7821$ Å and $c = 11.6191$ Å based upon the JCPDS standards. A significant lattice mismatch of $\Delta a/a = 2.2\%$ at room temperature (RT) is present between GaAs and CIS ($a$-axis); therefore, a large number of defects were observed both at the CIS/GaAs interface and in the active CIS layers [3]. Strain-induced novel phenomena such as the formation of interfacial layer caused by strain-induced interdiffusion between GaAs and CIS have been observed [5].

In addition, a significant mismatch in thermal expansion coefficients exists between $11.4 \times 10^{-6}$ K$^{-1}$ for CIS ($a$-axis) and $6.86 \times 10^{-6}$ K$^{-1}$ for GaAs, therefore the lattice mismatch at growth temperature becomes more significant than $\Delta a/a = 2.2\%$ at RT.

In this work, effects of strain on the properties of CIS epitaxial films have been investigated by changing the film thickness and by using two kinds of substrates with different lattice mismatch. As shown in Fig. 1, lattice constants of the ternary alloy, In$_x$Ga$_{1-x}$As, can be controlled between those of two binary alloys, GaAs and InAs, as a function of InAs mole fraction ($x$); lattice constant of In$_x$Ga$_{1-x}$As is described as $a_{InGaAs}(x) = 5.6533 + 0.4051x$ Å based upon the Vegard’s law [6]. Use of a linearly or step-graded buffer layer in order to provide nearly 100% strain relaxation and a reduction in threading dislocation densities below $10^6$ cm$^{-2}$ has produced promising results and has been applied for large lattice-mismatched systems such as SiGe/Si [7], In$_x$Ga$_{1-x}$As/GaAs [8, 9], etc. A similar technique was applied to investigate the strain effects on film properties and to grow high-quality CIS films.

2. Experimental procedure

CIS epitaxial films with a range of growth parameters have been grown by solid-source molecular
beam epitaxy on both GaAs(0 0 1) (CIS/GaAs) and pseudo-lattice-matched In$_{0.29}$Ga$_{0.71}$As substrates (CIS/InGaAs) at the growth temperatures $T_s = 350$–600°C. More detailed growth procedures can be found elsewhere [3]. For CIS/GaAs, the S-passivated (2 × 1) GaAs surface has been used, and epitaxial growth with the c-axis normal to the substrate has been demonstrated. Thickness of the films was varied from 0.4 to 2.0 μm which is significantly thicker than the calculated critical layer thickness limit of ~80 Å for CIS on GaAs [5].

Pseudo-lattice-matched substrates (Pseudo-substrates) were prepared by gas-source MBE using elemental In, Ga, and gas-source As (AsH$_3$) [10]. A 200 Å GaAs buffer was first grown on a semi-insulating GaAs(0 0 1) substrate, followed by a linearly composition-graded In$_x$Ga$_{1-x}$As buffer layer, where the In composition was varied linearly with the depth from $x = 0$ to 0.29 at a grading rate of ~1% In/400 Å. A final 1 μm-thick In$_{0.29}$Ga$_{0.71}$As layer was grown as a pseudo-substrate layer. The surface of the pseudo-substrate was passivated by As$_2$ before removal from the vacuum chamber for transport to the CIS growth system.

The results using various characterization methods such as low-temperature PL, cross-sectional transmission electron microscopy (TEM) and high-resolution X-ray diffraction (HRXRD) will be discussed in comparison with those of the CIS films grown directly on GaAs(0 0 1) (CIS/GaAs). X-ray rocking curves have been obtained using a three-axis configuration and a high-resolution four-crystal Ge(2 2 0) X-ray diffractometer (Philips: Materials Research Diffractometer). A two bounce Ge(2 2 0) channel cut analyzer crystal was also placed between the CIS layers and the detector in order to improve the resolution of the measurements. Typical resolution in the $\omega$ direction was ~0.003° and ~0.006° in the 2θ direction. PL experiments were carried out at 2 K with a 1 m single-pass monochromator, Ar$^+$-laser ($\lambda = 514.5$ nm) excitation, and a liquid nitrogen-cooled Ge detector (North Coast: Model E0-817L). Cross-sectional TEM images were taken using a JEOL 4000 F/X microscope operating at 350 kV.

3. Results and discussion

Fig. 2 shows the PL spectra of CIS/GaAs grown at $T_s = 450$°C as a function of film thickness. The appearance of more distinct near-band-edge emissions with increasing film thickness indicates a reduction in defect density. Fig. 3 shows the dependence of the measured c/a ratio of CIS films as measured by X-ray diffraction and the line width of the conduction band to acceptor transition (peak 'c') by PL measurements on film thickness. The line width of peak 'c' decreases from 12.0 meV at 0.4 μm to 6.5 meV at 2.0 μm consistent with the above results. On the other hand, the c/a ratio deviates further from the JCPDS standard value of c/a = 2.0095; tensile strain along the direction parallel to the surface becomes larger with increasing film thickness. This tensile strain is attributed to the difference in thermal expansion coefficients between the CIS layer and the substrate of about a factor of two. This leads to a large, but imperfect lattice relaxation near the CIS/GaAs interface and is thought to appear during the cooling process [11].

![Fig. 2. Photoluminescence spectra of CIS/GaAs grown at $T_s = 450$°C as a function of film thickness.](image-url)
The c/a ratio changes substantially from c/a = 2.009 at 0.4 μm to 2.000 at 2.0 μm, and the expected change in crystal field, ΔE, with respect to the change in the c/a ratio is calculated to be about 4 meV, using the equation, ΔE = \frac{1}{2}b(2 - c/a) where b is the deformation potential [12]. However, peak ‘c’ showed an energy shift of only less than 1.1 meV for varying film thicknesses from 0.4 to 2.0 μm. The band gap of CIS (E_g) can be defined from the energy of peak ‘c’ (E_x) using the relation E_g = E_x + ΔE, provided that the ionization energy (E_i) is independent of the c/a ratio. The c/a ratios in the CIS films were measured at RT, thus the data may not be directly compared with the PL data obtained at 2 K. Assuming that a similar difference in the c/a ratio is sustained at 2 K, the band gap of CIS is likely to be insensitive to a change in the crystal field in this range of c/a ratios.

Fig. 4 shows the X-ray scan in the [001] direction of the CIS(0 0 8) and the GaAs(0 0 4) reflections for both (a) GaAs and (b) InGaAs pseudo-substrates. The results of XRD are also listed in Table 2. It is clear from the figure that the CIS layer was grown epitaxially on the pseudo-substrate. Somewhat unexpectedly, no noticeable improvement in the line width of CIS/InGaAs has been observed in comparison with that of CIS/GaAs; the line width of the CIS(0 0 8) peak of CIS/InGaAs is apparently limited by the mosaic of the pseudo-substrate layer. The measured lattice constants for CIS/InGaAs were in good agreement with the JCPDS standard values, while substantial tensile strain along the surface was present in CIS/GaAs consistent with the results discussed in the earlier paragraph.

Table 2
List of lattice parameters by means of high resolution X-ray diffraction

<table>
<thead>
<tr>
<th>Substrate</th>
<th>a-axis (Å)</th>
<th>c-axis (Å)</th>
<th>FWHM (degrees)</th>
</tr>
</thead>
<tbody>
<tr>
<td>GaAs</td>
<td>5.7920</td>
<td>11.6149</td>
<td>0.117</td>
</tr>
<tr>
<td>InGaAs</td>
<td>5.7835</td>
<td>11.6175</td>
<td>0.106</td>
</tr>
<tr>
<td>Standard (JCPDS)</td>
<td>5.7821</td>
<td>11.6191</td>
<td>–</td>
</tr>
</tbody>
</table>
A more significant difference in the in-plane lattice constants along two $\langle 100 \rangle$ orthogonal directions of $\Delta a_//a_{AVE} \sim 0.2\%$ was observed in CIS/InGaAs with respect to $\Delta a_//a_{AVE} \sim 0.1\%$ for CIS/GaAs. This difference may be due to the existence of a tilt in the pseudo-substrate; however, it is not considered to be sufficient to dominate the film properties.

Fig. 5 shows a cross-sectional TEM image of CIS/InGaAs grown at $T_s = 550^\circ$C. Most of the plastic deformation takes place in the linearly composition-graded buffer layer. Such a large relaxation is substantial enough to isolate the upper layers from the GaAs substrate reducing the residual strain in the upper layers. It also shows that the number of misfit dislocations is significantly reduced in the In$_{0.29}$Ga$_{0.71}$As and CIS layers.

PL spectra obtained from CIS/GaAs and CIS/InGaAs grown at 550°C are shown in Fig. 6a and Fig. 6b, respectively. Peak 'c' (0.970 eV) is dominant in Fig. 6a, which is typical for near stoichiometric and Cu-rich CIS epitaxial films grown on GaAs. Distinct but weak free-exciton emissions, $E_{X1}$ (1.039 eV) and $E_{X2}$ (1.045 eV), are present, and an emission due to an exciton–donor complex, $I_{X0}$ (1.031 eV), can also be observed [4, 13]. Strong defect-related emission lines (peaks ‘c’ and ‘d’) indicate the presence of a large number of point defects in CIS/GaAs.

On the other hand, emissions due to free excitons ($E_{X1}$ and $E_{X2}$) were predominantly observed in Fig. 6b; the density of optically active residual defects has been substantially reduced. The decrease in the intensity of the defect-related peak ‘c’ made possible the observation of new emission lines such as peaks ‘b’, $I_{X2}$ and $I_{X3}$. Temperature-dependent PL analysis on $E_{X1}$ and $E_{X2}$ confirmed our

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Fig. 5. Cross-sectional image of CuInSe$_2$ grown on In$_{0.36}$Ga$_{0.71}$As pseudo-lattice-matched substrates.

Fig. 6. Photoluminescence spectra at 2 K obtained from CuInSe$_2$ grown on (a) GaAs and (b) In$_{0.36}$Ga$_{0.71}$As pseudo-lattice-matched substrates.
assignment of these peaks to the free exciton emissions. $I_{X_2}$ and $I_{X_3}$ are attributed tentatively to emissions associated with an exciton-defect complex from their temperature dependence. Electrical characterization of the CIS/InGaAs (CIS/GaAs) film showed $\mu \sim 110$ cm$^2$/Vs ($\mu \sim 38.6$ cm$^2$/Vs) with a hole concentration of $p \sim 1.5 \times 10^{17}$ cm$^{-3}$ ($p \sim 3.4 \times 10^{17}$ cm$^{-3}$) at RT, also suggesting improvement in crystalline quality consistent with the PL results. Further improvement in the film quality can make possible the control of transport and optical properties of the material extrinsically by intentionally incorporating impurities.

4. Summary

Defects in CIS were found to be strongly sensitive to the misfit strain with respect to the substrate, suggesting that the residual defects created during the process of solar cell fabrication can be further controlled by minimizing the strains among the substrate, absorber and window layers, and metal contacts as well as the surface strain at the grain boundaries. In addition, such high-quality CIS epitaxial films can make possible the investigation of the properties of intrinsic defects as well as the behavior of impurities which are critical parameters to be controlled for device applications.

Acknowledgements

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References

MBE growth of PbSe/CaF$_2$/Si(1 1 1) heterostructures

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Abstract

Successful epitaxial growth of PbSe/CaF$_2$/Si(1 1 1) heterostructures using just one growth chamber of a molecular beam epitaxy system is described. Sharp (1 x 1) streaks in the RHEED pattern confirm the high structural quality of PbSe, and SEM and Nomarski microscopy reveal smooth PbSe surfaces. The PbSe epilayers show mobilities as high as 22,000 cm$^2$ V$^{-1}$ s$^{-1}$ and electron concentrations of 1 - 2 x 10$^{17}$ cm$^{-3}$ at 77 K. In addition, observation of RHEED intensity oscillations confirm layer-by-layer growth modes for both CaF$_2$ and PbSe. The PbSe/CaF$_2$ interface has also been characterized by using X-ray photoelectron spectroscopy. All the Ca 2p, F 1s, Pb 4f and Se 3d peaks show chemical shifts as PbSe coverage on CaF$_2$ increases. Results show that the PbSe/CaF$_2$ interface consists of a thin transition layer dominated by Pb-F and Ca-Se bonds.

1. Introduction

Despite the large thermal expansion mismatch between PbSe and silicon, high-quality epitaxial layers of PbSe can be grown on (1 1 1)-oriented silicon substrates when fluoride buffer layers are used [1, 2]. Dislocation glide along {1 0 0} slip planes, which can preserve the crystalline quality of the PbSe epilayer, appears to be responsible for this thermal strain relief [3]. Such heteroepitaxial growth offers a number of benefits that can lead to fabrication of improved infrared optoelectronic devices (PbSe has a band gap of 175 meV at 77 K) such as photovoltaic infrared sensor arrays [4]. Initial work in this area focused on using a stacked BaF$_2$/CaF$_2$ buffer layer [5], but recent work has shown that just a thin CaF$_2$ layer produces good PbSe epitaxy [6]. This fluoride buffer layer, even when as thin as 50 Å, does appear to be necessary for good PbSe epitaxy since attempts to grow PbSe directly on Si(1 1 1) have resulted in poor layer quality [7]. In this work, we show that high quality PbSe/CaF$_2$/Si(1 1 1) heterostructures can be grown using just one growth chamber of a molecular beam epitaxy system. In addition, we present reflection high energy electron diffraction (RHEED) intensity oscillation data that confirm layer-by-layer growth.
modes for both CaF$_2$ and PbSe. We so present X-ray photoelectron spectroscopy (XPS) data which show that the PbSe/CaF$_2$ interface consists of a thin transition layer dominated by Pb–F and Ca–Se bonds.

2. MBE growth procedures and results

Epitaxial growth of CaF$_2$ on Si commonly employs a high-temperature ( > 850°C) oxide desorption step. High background pressure, as is the case when both CaF$_2$ and PbSe are grown in the same chamber, are a cause for concern since unwanted reactions between selenium and silicon or calcium may occur. Reactions between silicon and Group-VI elements are possible [8], and prior work [9] has shown that selenium vapor does react with BaF$_2$ forming a BaSe compound at moderate temperatures and a selenium-rich Ba$_2$Se$_3$ compound at temperatures above 690°C. Formation of similar reaction products involving selenium and calcium, and/or silicon, would certainly interfere with the silicon-calcium bonding that is required for good CaF$_2$ epitaxy on Si(111) [10]. It is therefore desirable to reduce processing temperatures to as low a value as possible such that unwanted reactions are inhibited. Accordingly, we have reduced the silicon preparation temperature more than 300°C by adding an HF : H$_2$O (1 : 10 in volume) dipping step to the standard Shiraki cleaning procedure [11]. This produces a hydrogen-passivated silicon surface [12], which can be thermally cleaned at a much lower temperature than an oxide-passivated surface. Successful preparation of three inch diameter Si(111) wafers was confirmed by observing sharp (7 × 7) streaks in the RHEED patterns after they were annealed for 20 min at 520°C.

CaF$_2$ growth was accomplished by heating bulk CaF$_2$ in a dual zone effusion cell held at 1280°C. This cell temperature produces a beam equivalent pressure for CaF$_2$ of 6.4 × 10$^{-8}$ Torr and a growth rate of 0.07 ML/s. A two-stage procedure [13] was used to grow approximately 100 Å of CaF$_2$ for subsequent PbSe epitaxy. Two monolayers of CaF$_2$ were first grown on the thermally-cleaned Si(111) at 700°C, and then the substrate temperature was lowered to < 250°C where additional CaF$_2$ growth occurred. To minimize the extent of any unwanted reactions, the total time for the initial growth step, which includes: (1) heating up, (2) about 30 s of growth, and (3) cooling down, is kept to less than 15 min. Fig. 1 shows RHEED intensity oscillation data for growth at the second stage temperatures. As the substrate temperature is lowered, the number of intensity oscillations increases showing that layer-by-layer growth continues longer at lower temperatures. As many as ten oscillations are observed for a substrate temperature of 70°C, an indication that the surface migration length for CaF$_2$ molecules is large, as expected for an ionic compound. Observation of RHEED intensity oscillations is a useful technique for determining epilayer quality during CaF$_2$ growth since

![Figure 1](image-url)

**Fig. 1.** RHEED intensity oscillations for MBE growth of CaF$_2$ on Si(111). Enhanced layer-by-layer growth at low temperatures is the evidence of large surface migration length of CaF$_2$ molecules.
surface reconstruction does not occur with this material and is thus not available as a technique for determining epitaxial quality. Moreover, observing RHEED oscillations allows accurate growth rate calibration.

PbSe growth was accomplished by heating bulk PbSe in a low-temperature effusion cell held at 605°C. This cell temperature produces a beam equivalent pressure for PbSe of $6.1 \times 10^{-7}$ Torr and a growth rate of 0.26 ML/s. For this work, PbSe layers of approximately 2.3 μm thickness were grown at 310°C on 100 Å of CaF₂ on Si(1 1 1). A small amount of additional PbSe was then grown at various substrate temperatures ranging from 310°C to 210°C for RHEED intensity oscillation experiments, see Fig. 2. Our results are in general agreement with those of Fuchs et al. [14] who observed optimal PbSe growth temperatures above 220°C, where surface diffusion is sufficient for layer-by-layer growth, but below 260°C, where step propagation is not enhanced. However, our results show that layer-by-layer growth below 200°C may be possible since no degradation in RHEED oscillations is seen down to 210°C. This would be consistent with recent results for growth of PbTe where as many as 155 RHEED oscillations have been observed at temperatures as low as 95°C [15]. RHEED oscillation data therefore suggest that both CaF₂ and PbSe may grow via a layer-by-layer mechanism at substrate temperatures below 200°C. Overlap in optimal growth temperatures for CaF₂ and PbSe, as well as accurate growth rates provided by the RHEED intensity oscillation frequencies, will facilitate the fabrication of heterostructure devices composed of thin layers of CaF₂ and PbSe.

Fig. 3 is a Nomarski micrograph showing the surface morphology of a typical PbSe layer. The smooth, featureless surface indicates that unwanted reactions such as those observed by Mathet et al. [2] did not occur during the growth procedure. Hall-effect measurements show that our PbSe layers are n-type and have mobilities as high as 23 200 cm² V⁻¹ s⁻¹ at 77 K, which are comparable to those obtained by Zogg et al. [16]. Even though the layers are undoped, electron concentrations are in the range of $1-2 \times 10^{17}$ cm⁻³ due to native defects in the form of selenium vacancies, which each donate two electrons to the conduction band [17]. These vacancies are believed to be created by selenium evaporation after PbSe molecules condense on the substrate surface. Although the concentrations reported here may seem large, they are more than an order of magnitude smaller than electron concentrations for undoped IV–VI semiconductor layers grown by liquid phase epitaxy (LPE) on BaF₂ substrates [18]. It is possible to control selenium vacancy concentration during MBE growth by providing an additional source of selenium as shown by Mathet et al. [2] who demonstrated that even p-type PbSe layers, corresponding to a condition of lead vacancies, could be obtained by increasing the selenium flux. We have just begun experiments in controlling selenium vacancy concentrations using a selenium flux provided by
an EPI Inc. valved-cracker source. Initial results suggest that carrier type and concentration can be controlled by varying the PbSe to Se flux ratio.

3. XPS characterization of the PbSe/CaF₂ interface

Earlier work involving LPE growth of PbSe on BaF₂ substrates revealed that selenium vapor reacts with BaF₂ to form a BaSe reaction layer [9]. Evidence for interfacial reactions between IV–VI semiconductors and group-II-A fluorides has also been seen in MBE growth of PbSe on BaF₂/CaF₂ buffer layers on silicon; blisters are observed to occur when the CaF₂ layer is greater than the critical thickness of 20 Å [2]. It was suspected that reaction with BaF₂ allowed migration of selenium to the CaF₂ layer where it penetrated threading dislocations and reduced the layer’s plasticity. Possible reaction product formation at the PbSe/CaF₂ interface is worth investigating since the chemistry is similar to the PbSe/BaF₂ interface. In addition, it is important to know whether the PbSe/CaF₂ interface is structurally abrupt. If so, these materials could be used to fabricate quantum effect heterostructure devices having band edge discontinuities greater than 3 eV.

The PbSe/CaF₂ interface was studied using an XPS analysis chamber attached via a buffer chamber to our MBE growth chamber. XPS spectra were taken using an AlK₃₁,₂ source (hv = 1486.6 eV) at a potential of 15 kV and with an emission current of 10 mA. Spectra of sample surfaces were taken following growth at 220°C of small amounts of PbSe onto 100 Å of CaF₂ on Si(1 1 1). Fig. 3 shows several XPS spectra for various amounts of PbSe coverage ranging from 0.27 to 26 ML. Monolayer coverage is calculated from the known PbSe growth rate as determined from the RHEED intensity oscillation frequency, see Fig. 4. Shifts of 1.65 eV and 1.70 eV, for the Pb 4f₁/₂ and Se 3d binding energies are seen, respectively, as PbSe coverage increases. Note that most of the chemical shift occurs between 0.81 and 6.5 ML and that photoelectron peak energies for Pb and Se shift in tandem. This shows that both Pb and Se interact with the CaF₂ surface and that this interaction extends for only a short distance into the PbSe layer. Binding energies for Ca 2p₃/₂ and F 1s are also observed to shift in tandem as PbSe coverage increases, although the amount of shift is less, 1.25 eV for both elements.

Our XPS results do not provide any evidence for the existence of a reaction product at the PbSe/CaF₂ interface. The fact that all photoelectron peak energies shift in tandem implies that there is an orderly compositional transition through the interface. The thickness of this transition layer, about 6.5 ML, can be partially accounted for by the open three-dimensional structure of the CaF₂(1 1 1) surface. Three layers of atoms are available for bonding with deposited material: (1) a layer of fluorine at the surface, (2) a layer of calcium 0.79 Å below the surface, and (3) another layer of fluorine 1.58 Å below the surface. This structure allows interpenetration of the PbSe and CaF₂ layers, which necessarily results in a thin compositional transition layer.

Based on electrostatic arguments, it is reasonable to assume that only Pb–F and Ca–Se bondings occur at the PbSe/CaF₂ interface. This leaves two possible bonding arrangements for the interface. The first one consists of lead atoms residing directly over and bonding to the top layer of fluorine atoms and of selenium atoms residing directly over the
lower layer of fluorine atoms but bonding to the three surrounding calcium atoms. The second consists of lead atoms residing directly over and bonding to the lower layer of fluorine atoms and of selenium atoms residing directly over and bonding to the layer of calcium atoms. Coexistence of these two bonding arrangements results in the growth of PbSe epilayers with twinned domains azimuthally rotated by (2n-1)π/3 with respect to each other. Diffraction from such a layer would show sixfold symmetry. This was not observed in our RHEED patterns showing that only one bonding arrangement exists for our structures. It is not clear from the XPS data which of the two bonding arrangements exists; both are supported by the data. It has, however, been suggested that equilibrium interactions favor the second bonding arrangement [19].

buffer layers, where both materials were grown in the same MBE chamber. SEM and Nomarski microscopy show that the layers are smooth with no evidence of unwanted reactions associated with a selenium background pressure. RHEED intensity oscillations were observed during growth of both the CaF$_2$ and the PbSe showing that both materials have layer-by-layer growth modes. Hall mobilities for the PbSe layer were as high as 23,200 cm$^2$ V$^{-1}$ s$^{-1}$. XPS data showed that the PbSe/CaF$_2$ interface consists of a thin transition layer and lacks any distinct interfacial compound. Our results show that this materials system is a promising candidate for fabrication of infrared optoelectronic devices on silicon substrates.

4. Summary

In summary, we have grown high-quality PbSe layers on silicon substrates using 100 Å thick CaF$_2$

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References


(GaMn)As:
GaAs-based III–V diluted magnetic semiconductors
grown by molecular beam epitaxy

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Abstract

We have grown novel III–V diluted magnetic semiconductors, (Ga\textsubscript{1-x}Mn\textsubscript{x})As, on GaAs substrates by low-temperature molecular beam epitaxy using strong nonequilibrium growth conditions. When the Mn concentration x is relatively low (\(\leq 0.08\)), homogeneous alloy semiconductors, GaMnAs, are grown with zincblende structure and slightly larger lattice constants than that of GaAs, whereas inhomogeneous structures with zincblende GaMnAs (or GaAs) plus hexagonal MnAs are formed when x is relatively high. Magnetization measurements indicate that the homogeneous GaMnAs films have ferromagnetic ordering at low temperature.

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\textit{Keywords:} GaMnAs; Diluted magnetic semiconductor; Low temperature MBE; Ferromagnetic order

1. Introduction

Molecular beam epitaxy (MBE) has made it possible to grow a variety of new heterostructures and alloy semiconductors with novel properties. Among them, diluted magnetic semiconductors (DMS) have attracted much interest, because they are expected to combine the properties both of magnetic materials and of semiconductors. DMSs are alloy semiconductors in which a fraction of cations are replaced by magnetic ions (transition metals or rare earths). So far, the studies of DMSs were mainly concentrated on II–VI-based materials [1], because some magnetic ions, particularly Mn\textsuperscript{2+}, can be easily incorporated into II–VI compounds by substituting group-II cations. In II–VI DMSs such as CdMnTe and related heterostructures, magneto-optical properties were extensively studied, and recently optical isolators were fabricated utilizing their large Faraday effect [2].
On the other hand, the study of III–V based DMSs has been limited to InMnAs [3, 4], in which some interesting properties associated with carrier induced magnetism were found in p-type samples. Since these InAs-based DMSs have very narrow band gap (≈ 0.36 eV) and much larger lattice constant (≈ 0.605 nm) than that of GaAs, the applications to III–V electronics and optoelectronics will be limited. For a wide range of application in III–V systems, a new type of DMS based on GaAs, which is the most widely used and well understood III–V semiconductor, is needed to be prepared. Very recently, De Boeck et al. [5, 6] reported that ferromagnetic MnAs particles were controllably embedded in GaAs by annealing homogeneous GaMnAs layers. However, little is known so far on the epitaxial growth, structure, and magnetic properties of GaMnAs [7, 8]. This paper presents the study of MBE growth and properties of this new class of GaAs based DMS.

2. Molecular beam epitaxial growth

MBE growth of (Ga1–xMnx)As thin films on semi-insulating (0 0 1) GaAs substrates were performed with our ULVAC MBC-508 system using solid sources of Ga, As, and Mn. After growing a GaAs buffer layer at 600°C under normal growth conditions, the substrate temperature was cooled to 200–300°C while the As2 flux was kept on. The reflection high-energy electron diffraction (RHEED) pattern changed from (2 × 4) to (4 × 4). Then Ga and Mn shutters were successively opened to grow (Ga1–xMnx)As at 200–300°C with various Mn content x ranging from 0 to 0.17. The growth rate and the film thickness of GaMnAs were 0.35–0.5 μm/h and 0.5–1.4 μm, respectively. The Mn flux was determined from the growth rate of epitaxial MnAs thin films grown on GaAs and/or on Si, which was already established by our group [9]. All the homogeneous GaMnAs samples were grown under the As-rich conditions at low temperature, as will be described later. Although equilibrium solubility of transition metals in GaAs is generally limited to the order of doping levels (∼ 10¹⁹ cm⁻³), this low-temperature MBE realizes strong nonequilibrium growth conditions, allowing a large amount of Mn atoms to be incorporated into the host GaAs lattice far above its solubility.

We found that homogeneous (Ga1–xMnx)As ternary alloys were grown when the Mn content x and growth temperature were relatively low. Fig. 1a shows a typical RHEED pattern of the homogeneous samples, which was taken during the growth of GaMnAs (x = 0.007) at 250°C with the azimuth of [1 1 0]. At the growth temperatures of 250°C and of 300°C, homogeneous GaMnAs alloys were obtained with x up to 0.078 and 0.005, respectively. In this case the RHEED patterns were very streaky like Fig. 1a, with (1 × 2) reconstruction (twofold streaks were weak). RHEED observations with various azimuths,

Fig. 1. RHEED patterns during the growth of (Ga, Mn)As with the azimuth of [1 1 0]: (a) the Mn content x = 0.007 grown at 250°C, (b) x = 0.10 grown at 300°C.
3. Structural characterizations

To characterize the structure of the (Ga, Mn)As films, X-ray diffraction measurements were done on all the samples. Fig. 3a–Fig. 3c shows θ–2θ spectra of the various homogeneous Ga$_1$-$x$Mn$_x$As films grown at 250°C: (x, thickness) = (0.037, 1.4 μm), (0.064, 0.44 μm), (0.074, 1.4 μm). The peaks of GaMnAs were observed very close to the peak of GaAs (4 0 0) reflection. The splittings are due to Cu-Kα1 and Cu-Kα2. One can see that the lattice constants of the GaMnAs films are slightly larger than that (a$_0$ = 0.6533 nm) of GaAs. No indication of MnAs formation was seen in these samples. From double-crystal X-ray diffraction measurements, the values of lattice mismatch Δa/a$_0$ to the GaAs substrate were estimated to be 0.59% at x = 0.074. The values of lattice mismatch measured by X-ray at various x are plotted by the solid circles in Fig. 3d. Here, we plotted the data of some samples for which we measured x by EMPA (the data of other samples that we did not measured by EMPA are not plotted, and the values of x of those samples were estimated by the experimental formula derived from Fig. 3d). The line-widths of the GaMnAs peaks were ~ 150 s, only a little wider than that of GaAs substrate. The narrow line width of the GaMnAs peaks indicates high epitaxial quality. The GaMnAs films on GaAs substrates are under the compressive strain, therefore, the lattice spacing of these sample is extended in the vertical direction. We calculated intrinsic lattice constants of cubic GaMnAs assuming that the Poisson’s ratio of GaMnAs is equal to that of GaAs, as plotted by the crosses in Fig. 3d. The intrinsic lattice constants are on a nearly straight line, i.e. following Vegard’s law.

In contrast, for the inhomogeneous samples whose RHEED patterns showed MnAs clustering like Fig. 1b, MnAs formation was also observed in X-ray spectra, as shown in Fig. 3e which was taken from the sample with nominal x of 0.08 grown at 300°C. One can see an intense peak of MnAs(1 1 0) as well as that of GaAs (2 0 0). A small peak of MnAs(1 0 0) was also seen. This indicates that the crystal orientation of the MnAs clusters in GaAs (or possibly GaMnAs) is mainly the (1 1 0) of the
hexagonal structure, in agreement with the results of RHEED.

4. Magnetic properties

First, magnetotransport properties were studied for the homogeneous GaMnAs samples. In Hall measurements at room temperature and at 77 K, all samples showed p-type conduction with the hole concentration of $3.8 \times 10^{17} \text{ cm}^{-3}$ (Mn composition $x = 0.004$, $T_e = 300^\circ\text{C}$) $- 2.6 \times 10^{20} \text{ cm}^{-3}$ (Mn composition $x = 0.074$, $T_e = 250^\circ\text{C}$) with no ferromagnetic behavior. Within our experiments, there was no special correlation between the hole concentration and their growth parameters. At low temperature range (4.2 K or below), however, clear hysteretic behaviors have appeared in magnetoresistance ($R_{xx}$-$H$) measurements, and Hall resistance ($R_{xy}$-$H$) measurements showed extraordinary Hall effect (EHE), suggesting the presence of ferromagnetic ordering.
Magnetization measurements were performed on the homogeneous GaMnAs films using a superconducting quantum interference device (SQUID). Fig. 4a shows a magnetization ($M$–$H$) curve of a 1.4 μm-thick (Ga$_{1-x}$Mn$_x$)As film with $x = 0.074$ measured at 1.9 K, when the magnetic field $H$ was applied in plane along the $\langle 110 \rangle$ axis. The hole concentration and mobility of this sample at room temperature were $2.6 \times 10^{20}$ cm$^{-3}$ and 7.24 cm$^2$/V·s, respectively. A square-like hysteresis in the $M$–$H$ characteristics directly indicates the presence of ferromagnetic ordering. The values of coercive field $H_c$ and saturation magnetization $M_s$ were estimated to be 56 Oe and 33 emu/cm$^3$, respectively. The clear hysteresis behavior remained when the temperature was increased up to 55 K, above which the hysteresis behavior disappeared. Fig. 4b shows the temperature dependence of residual magnetization $M_r$, indicating that the Curie temperature of this sample is 55–60 K, much higher than that (7.5 K) of p-type InMnAs [4].

To characterize the magnetic anisotropy, we have done magnetization measurements on the same sample with the magnetic field $H$ applied perpendicular to the film plane. Fig. 4c shows the result of $M$–$H$ characteristics measured at 1.9 K, exhibiting much higher saturation field (1500–2000 Oe). Though a little hysteresis behavior was observed, its shape is far from square. The results of Fig. 4a and Fig. 4c indicate that the easy magnetization axis of the present GaMnAs film is not perpendicular but aligned in the film plane.

5. Summary

We have grown new GaAs-based diluted magnetic semiconductors (Ga, Mn)As by low-temperature MBE. Homogeneous ternary alloys were obtained when the Mn content $x$ and growth temperature were relatively low. The crystal structure was zincblende with slightly larger lattice constants than that of GaAs. All the GaMnAs samples showed p-type conduction with the hole concentration of $3.8 \times 10^{17}$–$2.6 \times 10^{20}$ cm$^{-3}$. The homogeneous GaMnAs film with $x = 0.074$ was shown to have ferromagnetic ordering at low temperature.
with the easy magnetization axis in plane, and its Curie temperature was estimated to be 55–60 K.

References


Epitaxy of (Ga, Mn)As, a new diluted magnetic semiconductor based on GaAs

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Abstract

GaAs-based diluted magnetic semiconductor, (Ga, Mn)As, with Mn composition $x$ up to 0.07 was prepared by molecular-beam epitaxy on GaAs substrate at temperatures ranging from 160 to 320$^\circ$C. Clear reflection high-energy electron diffraction oscillations were observed at the initial growth stage, indicating that the growth mode is two-dimensional. The lattice constant of (Ga, Mn)As films determined by X-ray diffraction showed a linear increase with the increase of Mn composition. Well-aligned in-plane ferromagnetic order was observed by magnetization measurements. Magnetotransport measurements also revealed the presence of ferromagnetic order in the (Ga, Mn)As layer. The easy axis of magnetization can be reversed by changing the strain direction in (Ga, Mn)As. GaAs/(Ga, Mn)As superlattice structures with high crystal perfection and good interface quality were also prepared.

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\textit{Keywords}: Molecular-beam epitaxy; III–V compounds; Diluted magnetic semiconductors; Ferromagnetic order; Magnetic anisotropy; Superlattice

1. Introduction

Due to the limited low solubility of magnetic ions in III–V semiconductors [1–3] the study of diluted magnetic semiconductors (DMSs) has long been concentrated mostly on the II–VI-based materials [4]. By using non-equilibrium growth conditions of low-temperature molecular-beam epitaxy (LT-MBE), a III–V-based DMS, (In, Mn)As, was successfully prepared in 1989 [5] and substitution of Mn atoms for the In sites was found in homogenous (In, Mn)As films by X-ray-absorption fine-structure measurements [6]. Detailed magnetotransport study of p-type homogeneous...
(In, Mn)As films revealed the presence of carrier-induced ferromagnetic ordering at low temperatures [7]. The study of III–V DMSs, however, has so far been limited mostly to (In, Mn)As [5–11], except for the works on (Ga, Mn)Sb [12] and on the formation of MnAs clusters in GaAs host [13]. Recently, we have successfully prepared a new III–V-based DMS, (Ga, Mn)As [14], which is based on the most well understood and extensively used III–V compound, GaAs. Here we report in detail the MBE growth and properties of the (Ga, Mn)As films.

2. MBE growth of (Ga, Mn)As films

The (Ga, Mn)As films were grown by solid-source MBE on semi-insulating (0 0 1) GaAs substrates using constituent elements of Ga, Mn, and As. All the growth was carried out under As stabilized condition with the growth rate of 0.6–0.8 μm/h. The Mn content x in the (Ga$_{1-x}$Mn$_x$)As was calibrated by electron probe microanalysis measurements. Reflection high-energy electron diffraction (RHEED) was used for in situ monitoring of the growth procedures.

At a substrate temperature $T_s = 250^\circ$C, (Ga, Mn)As films were obtained with x varying from 0.015 to 0.071. At a fixed x = 0.035 epitaxial (Ga, Mn)As layers can be grown at $T_s$, ranging from 160°C to 320°C. Hereafter we will concentrate mostly on the samples grown at $T_s = 250^\circ$C. A 100 nm GaAs layer was first grown at $T_s = 700^\circ$C. After lowering $T_s$ to 250°C, another 100 nm GaAs was grown prior to the growth of 150 nm thick (Ga, Mn)As. The first GaAs layer grown at 700°C showed the well-known (2 × 4) surface reconstruction, which changed to c(4 × 4) when $T_s$ was lowered to 480–520°C and remained c(4 × 4) below. The surface reconstruction of LT GaAs and that of (Ga, Mn)As were (1 × 1) and (1 × 2), respectively, as shown in Fig. 1.

Clear RHEED oscillations were observed at the initial growth stage, indicating that the growth mode is two-dimensional. Fig. 2 shows the RHEED specular beam intensity oscillations for GaAs grown at 600°C and 250°C and for (Ga, Mn)As. One striking feature is that GaAs grown at 250°C can still keep the layer-by-layer growth. At first, we thought the oscillation of GaAs at LT might be caused by Mn, which acted as a surfactant. But the results are reproducible even
in the growth chamber without Mn cell. We have done works on the RHEED oscillations of GaAs systematically and found that GaAs can keep the layer-by-layer growth throughout the temperature range from 150°C to 730°C. Detailed results on RHEED oscillation of GaAs will be published elsewhere.

3. Metallurgical evaluations

θ–2θ scans of X-ray diffraction were performed at room temperature using Cu Kα radiation. Clear diffraction peaks from (Ga, Mn)As were observed for all the samples along with diffraction peaks from the underneath GaAs. Fig. 3 shows the diffraction curves of three samples with \( x = 0.015, 0.035 \) and 0.071. As can be seen from the figure, the peak position of (Ga, Mn)As shifts to lower angle as \( x \) increases, indicating that the lattice constant \( a \) increases with the increase of \( x \). Dependence of \( a \) on the Mn composition \( x \) of (Ga, Mn)As is shown in the inset (a) of Fig. 3. \( a \) was calculated from the peak position of (Ga, Mn)As (0 0 4) diffraction assuming that (Ga, Mn)As is fully strained and that (Ga, Mn)As has the same elastic constant as GaAs. The assumption that (Ga, Mn)As is fully strained on GaAs was further confirmed by asymmetric X-ray diffraction by using a double crystal diffractometer with (1 1 5) reflection. As can be seen from inset (a), \( a \) increases linearly with \( x \), showing that it follows Vegard’s law. The extrapolated lattice constants for GaAs (0.566 nm) and MnAs (0.598 nm) are in good agreement with the known GaAs lattice constant (0.56533 nm) and with the MnAs lattice constant extrapolated from the (In, Mn)As work (0.601 nm) [5], suggesting that all the Mn atoms were incorporated in the zincblende alloy. There is no indication of the formation of the second-phase MnAs in the film from the X-ray diffraction measurements.

Inset (b) of Fig. 3 shows the diffraction curves from two samples with the same nominal Mn composition \( x = 0.035 \). One sample was annealed at
700°C for 2 min, the other one is the as-grown sample. The annealing was done in situ under As flux. The complete disappearing of (Ga, Mn)As peaks in the annealed sample suggests the formation of MnAs clusters in GaAs host [13].

4. Magnetic and magnetotransport properties

Magnetization measurements done by SQUID (superconducting quantum interference device) magnetometer showed the presence of a ferromagnetic order in the (Ga, Mn)As films. Fig. 4 shows the magnetic field dependence of the magnetization of a sample with $x = 0.035$ at 5 K, with magnetic field $B$ applied both parallel and perpendicular to the sample surface. A clear square hysteresis was observed when $B$ was applied parallel to the sample surface, indicating the existence of a well-ordered ferromagnetic structure in the film. On the other hand, when $B$ was applied perpendicular to the surface the hysteresis is less pronounced, suggesting that the magnetic anisotropy favors in-plane direction. From the vanishing of the remanent magnetization, which was obtained by temperature-dependent measurements, the Curie temperature of the sample was determined to be 60 K. Absence of remanence above 75 K also confirms that the concentration of MnAs, whose Curie temperature is 310 K, is extremely low.

Magnetotransport measurements were performed at temperatures ranging from 1.5 to 300 K under magnetic field up to 9 T. All samples showed p-type conduction. At low temperatures, the Hall resistivity $\rho_{\text{Hall}}$ is dominated by the anomalous Hall effect and the $B$ dependence of $\rho_{\text{Hall}}$ showed a hysteresis loop very similar to what was observed in magnetic measurements when $B$ was applied perpendicular to the sample surface. The carrier concentration in the sample with $x = 0.035$ was determined to be $3 \times 10^{20} \text{ cm}^{-3}$ from the high magnetic field region ($6 \text{T} < B < 9 \text{T}$) at 1.5 K, which is of the same order as the Mn concentration ($7.7 \times 10^{20} \text{ cm}^{-3}$). The inset of Fig. 4 shows the magnetization $M$ determined by SQUID measurement and that deduced from magnetotransport measurement by taking $\rho_{\text{Hall}} = c \rho M$ (where $\rho$ is the resistivity, $c$ a constant). Very good agreement was obtained with $c = 1.75 \text{ T}^{-1}$. The Curie temperature $T_c$ was also obtained from the temperature dependence of susceptibility determined from the transport measurements in the paramagnetic region.

Fig. 4. Magnetic field dependence of magnetization $M$ at 5 K for a (Ga, Mn)As film with $x = 0.035$, with the field applied parallel (solid circle) and perpendicular (open square) to the sample surface. Inset shows a comparison of $B$ dependence of $M$ between SQUID measurement (open square, at 5 K) and magnetotransport measurement (solid line, at 4.2 K).

Fig. 5. Hall resistance, $R_{\text{Hall}}$, of (Ga, Mn)As/(In, Ga)As heterostructure as a function of $B$ with an angle between $B$ and the sample surface normal as a parameter. A square hysteresis loop was observed. The coercive field changes with the angle while the height of the hysteresis loop keeps unchanged, suggesting the easy axis of magnetization is perpendicular to the sample surface.
(T > 120 K), where the susceptibility follows the Curie-Weiss law. Tc determined from transport measurements for the samples with x = 0.035, 0.043 and 0.053 are 83, 99 and 97 K, respectively.

The in-plane easy axis of magnetization in the (Ga, Mn)As grown on GaAs was thought to be induced by a compressive strain present in the film, although in general the easy axis favors in-plane direction due to the demagnetization field. By growing (Ga, Mn)As on a relaxed (In, Ga)As buffer layer with larger lattice constant than that of (Ga, Mn)As, which makes the (Ga, Mn)As under tensile strain, the easy axis was changed to be perpendicular to the plane, as demonstrated in Fig. 5.

5. GaAs/(Ga, Mn)As superlattice structures

GaAs/(Ga, Mn)As superlattice structures were also prepared by MBE. The introduction of superlattice structures is expected to provide additional parameters for controlling the spatial confinement of both carriers and magnetic ions, possibly involving dimensional effects. The nonmagnetic/ferromagnetic multilayer system is also an interesting system for studying the exchange coupling between the ferromagnetic layers by adjusting the thickness of the nonmagnetic layers.

Double crystal X-ray diffraction measurements were performed for the superlattice samples. Clear satellite peaks were observed, indicating the existence of a periodic structure in the samples. The high perfection of the structure was confirmed by the observation of thickness fringes due to perfect crystal diffraction from thin films. Magnetotransport measurements revealed the existence of ferromagnetic order in the current superlattice structures. Detailed results concerning the superlattice structures will be published elsewhere [15].

6. Summary

In summary, we have successfully prepared a new GaAs-based III–V DMS, (Ga1-x,Mn)xAs, by MBE, with x varying from 0.015 to 0.07. Clear RHEED oscillations were observed at the initial growth stage, indicating that the growth mode is two-dimensional. X-ray diffraction measurements showed that the lattice constant of the (Ga, Mn)As films increases linearly with Mn composition, suggesting homogenous incorporation of Mn atoms in the zinc-blende alloy. Well-aligned in-plane ferromagnetic order was observed by SQUID measurements. Magnetotransport measurements revealed the occurrence of the anomalous Hall effect in the (Ga, Mn)As layer and the hole concentration to be of the order of Mn concentration. The easy axis of magnetization was found to be reversible by changing the strain direction in the (Ga, Mn)As films. GaAs/(Ga, Mn)As superlattice structures were also prepared.

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References

Growth parameter dependence of step patterns in AlGaAs molecular beam epitaxy on vicinal GaAs(110) inclined toward (111)A

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Abstract

We have studied growth parameter dependence of macrostep patterns in AlGaAs molecular beam epitaxial (MBE) growth using As$_2$ molecular beams with the use of atomic force microscopy. It is found that MBE growth by using As$_2$ molecular beams leads to the formation of coherently aligned macrosteps. From the analysis of the macrostep patterns, we deduce the characteristic that increasing growth temperatures leads to the formation of wider macrosteps. In contrast, MBE growth using solid As source (As$_4$) gives rise to highly connected zigzag networks of macrosteps. We will discuss the difference in macrostep patterns between MBE growth using As$_2$ and As$_4$ in terms of the presence of As$_4$ physisorption states and the difference in the diffusion length of As species.

I. Introduction

We have demonstrated that molecular beam epitaxial (MBE) growth on vicinal GaAs(110) induces macrosteps several tens of nm high [1], and that such macrostep system is one of fairly good candidates as substrates for fabricating nanostructures such as quantum wires, since macrosteps formed by

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1 The misorientation direction of (111)B in our previous papers [1,10] is incorrect. It should be corrected to (111)A.

MBE using AsH$_3$ gas source are coherently aligned along [110] direction perpendicular to the misorientation direction of the vicinal (110) substrates [2]. However, MBE growth by using solid As source (As$_4$) produces rather fluctuated macrostep patterns. In order to fabricate excellent nanostructures, it is necessary to control macrostep patterns, i.e., to know what growth parameters make straight and uniform macrosteps stable.

Mainly, there are two differences between MBE growth by using AsH$_3$ and As$_4$: one is the presence of atomic hydrogen and/or hydrogen molecules, the other is the difference in the constituent of arsenic molecules (i.e., As$_2$ and As$_4$). Recently, Takeuchi [3] using conventional MBE plus hydrogen
source showed that MBE growth under atomic hydrogen and/or hydrogen molecules leads to little improvement of the macrostep patterns. He also reported that cracking of AsH₃ produces As₂ and hydrogen, while the generation of AsHₓ is negligible.

In this paper, we present growth parameter dependence of macrostep patterns in AlGaAs MBE growth using As₂ molecular beams with the use of atomic force microscopy (AFM). In particular, our interest is concentrated both on how the macrostep patterns are changed by varying growth temperature and V/III ratio and on what leads to the difference in the resulting step patterns when compared with the results using As₄ molecular beams.

2. Experimental procedure

Al₀.₅Ga₀.₅As layers of 400 nm thickness were grown by MBE on vicinal GaAs(1 1 0) substrates inclined by 3° toward (1 1 1)A using the following arsenic sources: one, a tetrameretic arsenic (As₄) beam sublimated from solid arsenic source, the other a dimeretic arsenic (As₂) beam produced by dissociating GaAs polycrystalline at temperatures ranging from 860 to 890°C using Knudsen cell. Dissociation of GaAs at these temperatures produces arsenic beams almost consisting only of As₂ (As₂/As₄ is usually larger than 1 0 0) [4–7]. Simultaneously, Ga beams are apparently produced by the dissociation of GaAs but negligible order of below one 10th of its As₂ flux. Growth temperature T_G was varied from 430 to 580°C. V/III pressure ratio R_V/I for AlGaAs growth using As₂ and As₄ beams was also varied in the range of 20–80 and 50–260, respectively. Growth rate of AlGaAs layers was 200 nm/h. Before AlGaAs growth, chemically etched substrates were thermally cleaned at 630°C for 20 min under As₄ flux, then buffer layers of 200 nm thick GaAs were grown at 580°C by using As₂ beams. This growth condition for GaAs buffer layers was found to provide atomically smooth growth surfaces consisting of diatomic steps. All AFM measurements were carried out in air in contact mode.

3. Results and discussion

To demonstrate how the difference between As₂ and As₄ beams affects patterns of macrosteps formed by MBE growth, shown in Fig. 1a and Fig. 1b are typical 5 µm × 5 µm AFM images of AlGaAs surfaces grown by using As₂ and As₄ molecular beams, respectively. They were taken for the AlGaAs films grown at 580°C by using As₂ beams at R_V/I = 30 and As₄ beams at R_V/I = 152. From the AFM image shown in Fig. 1a, it can be seen that macrosteps formed by using As₂ beam are coherently aligned along [1 1 0] direction, and that their terrace width and step height are 300 nm.

![AFM images (5 µm × 5 µm) of AlGaAs films grown by MBE at 580°C on vicinal GaAs(1 1 0) substrates when using (a) As₂ molecular beams and (b) As₄ molecular beams.](image-url)
wide and 15 nm high on an average, respectively. This macrostep pattern is very similar to those formed by gas source (AsH$_3$) MBE [1, 2]. In contrast, MBE growth by using As$_4$ produces highly connected zigzag networks of macrosteps, as shown in Fig. 1b. We, therefore, consider that MBE growth by using As$_4$ is essential for the formation of fairly straight and uniformly spaced macrosteps on vicinal GaAs (1 1 0) substrates.

Next, to make clear the growth parameter dependence of the macrostep patterns when using As$_2$, we have carried out AFM measurements for the AlGaAs films grown at various growth condition ($T_G = 430–580$°C, $R_{III} = 20–80$).

The AFM images for the films grown at 480–580°C showed that AlGaAs growth leads to the formation of macrosteps similar to the one shown in Fig. 1a, while such macrosteps were not observed for the films grown at 430°C. To deduce the characteristics of the macrosteps formed at 480–580°C, we plotted the growth temperature dependence of the average terrace widths $<W>$ and the terrace width distribution $<W>P(W/<W>)$ scaled to the average terrace width shown in Fig. 2. On the micrographs, we measured the local distance between adjacent steps along [0 0 1] direction. The present analysis allows us to deduce the following two characteristics of the macrosteps. Firstly, it is clear from Fig. 2a that with increasing growth temperature, the average terrace width increases showing little dependence on the V/III ratio. Secondly, it is obvious from Fig. 2b that all the terrace width distributions plotted for the AlGaAs film surfaces fall on the same curve regardless of the growth temperature and the V/III ratio. In an equilibrium picture, the terrace width distribution for a given vicinal surface is determined by energetic interactions between steps. Bartelt et al. [8] and Joós et al. [9] have theoretically derived the relationship between the terrace width distributions and the strength of step-step interactions. The solid curve shown in Fig. 2b is the scaled terrace width distribution calculated for many non-interacting wandering steps except for the non-crossing condition for the steps, i.e. the free-fermion model. Surprisingly, the observed distributions $<W>P(W/<W>)$ almost fit the solid curve, although some data are scattered around the curve. In other words, the present results for the macrosteps formed by using As$_2$ seem to be well described by the model that the steps do not interact with each other except that they cannot cross. Of course, macrosteps formed by MBE growth do not represent equilibrium structures. However, the above analysis is still useful for deducing the characteristics of step patterns, as will be discussed later.

As for AlGaAs growth by using As$_4$, the surface morphologies including the step patterns show rather complicated dependence both on the growth
temperature $T_G$ and on the V/III ratio $R_{V/III}$. From the AFM and reflection high-energy electron diffraction (RHEED) observations, the $T_G$-$R_{V/III}$ growth parameter plane ($T_G = 430-630^\circ C$, $R_{V/III} = 50-260$) seems to be divided into the 5 regions (A-E) in terms of the surface morphologies. The characteristic surface morphologies corresponding to the regions (A-E) are as follows. AlGaAs growth under the parameter of the regions (A: $T_G \leq 530^\circ C$, $R_{V/III} \geq 200$ or $R_{V/III} \sim 50$ and $T_G \sim 580^\circ C$, $R_{V/III} \geq 150$) and (B: $T_G = 450-550^\circ C$, $R_{V/III} \sim 150$) leads to the macrosteps shown in Fig. 1b and Fig. 3b, respectively. For the growth under the region (C: $T_G \geq 630^\circ C$, $R_{V/III} \geq 150$) of high-growth temperatures and sufficient V/III ratio, rather smooth surface morphologies were observed by AFM. In addition, RHEED observations suggested that the growing surfaces consist of diatomic steps, as previously reported [10], although such diatomic steps were not observed in the AFM images. In the region (D: $T_G \geq 580^\circ C$, $R_{V/III} \leq 50$) of high-growth temperatures but low V/III ratio, however, the surface morphology became macroscopically rough. For the growth under the region (E: $T_G \leq 430^\circ C$) of low growth temperatures, flat surface morphologies were observed by AFM, while the growing surface displayed the non-split (1×1) RHEED patterns. Judging from the AFM and RHEED observations, it is thought that the growth under the region (E) causes multi-atomic steps of several tens of nm wide. Among these five regions, contrasted with the growth by using As$_2$, are the results for the regions (A) and (B). As mentioned above, the growth under the parameter of the region (A) causes highly connected zigzag macrosteps. Obviously, the terrace width distributions for such macrostep patterns are different from the distribution shown in Fig. 2b. It is peaked near the half of the average terrace width, then almost linearly decreases with increasing the normalized terrace width $W/\langle W \rangle$. On the other hand, the macrosteps formed under the parameter of the region (B) yields the terrace width distribution very similar to the one for the macrosteps formed by using As$_2$. If we now look over the AFM images shown in Fig. 3a and Fig. 3b, however, we note that there is the difference between the macrosteps formed by using As$_2$ and As$_4$. Almost all the As$_4$ steps run along [\bar{1}10] direction perpendicular to the miscut direction over 5 µm without touching each other, while the macrosteps formed by using As$_4$ just extend to at most 1 or 2 µm along [\bar{1}10] direction. Analyzing the distance between step-step collisions would extract a characteristic of those macrostep patterns. However, it is hard to obtain such data because of the complicated connected patterns. Instead, we measured the terrace length distributions along [\bar{1}10] direction. As a result, it turns out that the terrace length distributions for all macrosteps formed by using As$_4$ fall on the same
exponentially decay curves with increasing terrace length. This indicates that the macrostep patterns formed by using As$_4$ are characterized by the terrace length distribution. In other words, the macrosteps formed by using As$_4$ have the same fluctuation in terrace length. We, therefore, consider that complex and highly connected zigzag networks of steps result from the fluctuation along the step edges originating in the use of As$_4$ beams.

In order to deduce the origin of the fluctuation along the step edges leading to complex and highly connected networks of steps, we summarize the difference between As$_2$ and As$_4$ molecular beam. Firstly, the sticking coefficient of As$_2$ to Ga-rich GaAs(0 0 1) surface tends to unity [11, 12] while that of As$_4$ never exceeds the value of 0.5 [13, 14]. Secondly, there exists the physisorption state for As$_4$ molecules even at high temperatures [15]. Finally, As$_4$ and As$_2$ differ in the mass and the temperature of the molecular source. It is expected that this affects thermalization process of As species, i.e., energy transfer between As molecules and substrate atoms, and energy conversion process of adsorbing molecules such as translational to vibrational energy transfer.

Based on the present results and the above-mentioned difference between As$_2$ and As$_4$ molecules, one possible explanation is that the formation of zigzag steps originates in the presence of the physisorption state of As$_4$ and fluctuates along the step edges. It is conceivable that step motion is influenced by the presence of such physisorbates. In other words, the physisorbates act as impurities and impede the step motions. Recently, Kandel and Weeks [16] have studied the impurity-induced step bunching suggested by Frank [17] using two-dimensional models that allow fluctuations along the step edges. Their Monte Carlo simulation shows that impurity-induced step bunching leads to the formation of very complex and highly connected step patterns which resemble the present AFM image shown in Fig. 1b very much.

Another explanation is that the observed morphological difference comes from the difference in the diffusion length between the As species, which is expected from the third difference between As$_2$ and As$_4$ mentioned above. Consider the initial stages of AlGaAs growth proceeding in step-flow mode on the diatomic steps of GaAs(1 1 0) surfaces. Taking account of the direction of Ga–As bonding and step structures of the surface, AlGaAs growth is considered to proceed along step edges directed toward [T 1 0]. Initially, Al(Ga)As nuclei stick to GaAs diatomic step edges. An average distance between the adjacent nuclei usually depends on the diffusion length of As and/or Ga atoms. This means that when the diffusion length is short, diatomic steps largely meander on the surface because of the short average distance between nuclei. In fact, the RHEED observation of the surfaces grown at 630°C on the vicinal GaAs(1 1 0) inclined toward (1 1 1)A by 6° revealed that diatomic steps formed by using As$_4$ wander largely [10]. We, therefore, expect that such fluctuation along the step edges affects the macrostep pattern.

4. Conclusions

We have studied growth parameter dependence of macrostep patterns in AlGaAs MBE growth using As$_2$ molecular beams with the use of AFM. It is found that MBE growth by using As$_2$ leads to the formation of coherently aligned macrosteps, as is in the case of MBE growth by using AsH$_3$. From the growth temperature dependence of the average terrace width and the terrace width distribution, we deduce the characteristic that increasing growth temperatures leads to the formation of wider macrosteps regardless of V/III ratio. Besides, the terrace width distribution for these macrosteps is well described by the free-fermion model. In contrast, MBE growth by using As$_4$ leads to highly connected zigzag networks of macrosteps. By analyzing the macrostep patterns it is found that its patterns are characterized by the terrace length distribution showing the exponentially decay with increasing terrace length. These characteristics of the macrostep patterns indicate that complex and highly connected zigzag networks of steps result from the fluctuation along the step edges originating in the use of As$_4$ beams.
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References

Effects of tensile strain and substrate off-orientation on the growth of GaInAs/InP multiple quantum well structures by CBE


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Abstract

GaInAs/InP multiple quantum well (MQW) layers with high tensile strain in the GaInAs layers have been grown by chemical beam epitaxy (CBE). The samples were analysed by high-resolution X-ray diffraction (XRD), photoluminescence (PL) spectroscopy and transmission electron microscopy (TEM). The influence on layer properties of the tensile strain and the substrate off-orientation of the (100) InP substrates is investigated. XRD studies reveal that the growth rate of GaInAs as well as that of InP is larger for off-oriented substrates than for exactly oriented substrates. This behaviour is independent of the strain in the GaInAs layers and of the growth temperature in the chosen range between 485°C and 545°C. We attribute the observed changes of the growth rate to the different growth modes on exact (100) and off-oriented substrates. MQW structures grown on off-oriented substrates show a linear increase of the PL line width of the GaInAs MQW peak with increasing tensile strain in the GaInAs layers. In contrast, it stays constant for exactly oriented substrates. This increase in the PL line width for off-oriented substrates is explained by the lattice tilting occurring when strained layers are deposited on stepped surfaces. The tilt angle is proportional to the amount of incorporated elastic strain as shown by the azimuthal dependence of the position of the zeroth-order satellite reflection in XRD.

1. Introduction

Ga_{x}In_{1−x}As/InP multiple quantum well (MQW) layers are becoming increasingly important for applications in optical telecommunications in the near infrared region (1.3–1.5 μm). For many applications it is highly desirable to realize polarization-independent optical components. This can possibly be achieved by growing GaInAs superlattice structures with high tensile strain, i.e. with high Ga content in the GaInAs well layers [1, 2].

For strained MQW structures, however, the critical thickness for misfit dislocation generation has to be taken into account. Additionally, the thicknesses of wells and barriers as well as the Ga
content of the GaInAs layers have to be controlled precisely. Also a smooth surface morphology over large areas is necessary.

Therefore, it is of importance for device applications to investigate the influence of the epitaxial growth parameters on the structural and optical properties of strained structures and optimize the epitaxial growth with respect to these results.

We present here results obtained on a series of 10-period MQW structures grown by chemical beam epitaxy (CBE) in the growth temperature range between 485°C and 545°C. We discuss the effect of tensile strain and substrate off-orientation on the growth rate and on the structural and optical properties. Such test structures provide valuable insight in the relevant properties of highly strained MQW structures, which have to be taken into account for designing MQW-based device structures.

2. Experimental procedure

The samples were grown in a Riber CBE 32P system using triethylgallium (TEGa), trimethylindium (TMIn), arsine (ASH₃) and phosphine (PH₃). The TEGa and TMIn flows were combined without carrier gas in a low temperature (LT) injector to form a single emerging beam. Arsine and phosphine were introduced through a high-temperature (HT) cracker. The growth temperature varied between 485°C and 545°C and was measured with an optical pyrometer. The V/III ratio was 3.8 for the InP barriers and 1.75 for the GaInAs wells. The (100) InP substrates were either exactly oriented (±0.2°) or 2° off-oriented towards the [110]-direction. As test structure a MQW structure consisting of 10 periods nominally 5 nm GaInAs/5 nm InP/1 monolayer (ML) InAs/5 nm InP with various Ga concentrations (xGa) was chosen. The compressively strained InAs ML was obtained by exposing the InP surface to a beam of AsH₃ during 1 s [3] to provide some strain compensation for the tensile-strained GaInAs layers. The growth interruption sequence was chosen in a way that the upper and the lower interface of the GaInAs layer were symmetrically strained in compression [3]. Therefore, the interfacial layers also contribute to the compensation of the tensile strain in the GaInAs layers. X-ray diffraction studies show that at the bottom and at the top of the GaInAs layers 1 ML InAs₀.₅P₀.₅ is formed. Details about the interface structure have been published elsewhere [3–5].

The epitaxial layers were investigated by high-resolution X-ray diffraction (XRD), low-temperature photoluminescence (PL) and cross-sectional transmission electron microscopy (TEM). The XRD studies were performed by using a Bede model 200 double-crystal diffractometer (Bede Scientific, UK). The PL spectra were recorded at 5 K at a power density of 20 mW/cm² of the 514 nm Ar-ion laser line. A CCD camera was used as detector. The cross-sectional TEM images were taken in a Philips CM30T electron microscope operating at 300 kV.

3. Results and discussion

3.1. XRD characterization

The thicknesses of the individual MQW layers as well as the Ga concentration (xGa) in the GaInAs layers could directly be calculated from the XRD rocking curves by using the previously mentioned interface structure and the relation \(t_{\text{InAs}} = t_{\text{InP}}\), which is valid for the same TMIn flow. \(t_{\text{InAs}}\) and \(t_{\text{InP}}\) are the binary growth rates of InAs and InP, respectively. To verify these calculations the measured rocking curves of the samples were compared to dynamical simulations using the simulation package RADS (Bede Scientific, UK). Both measurements and simulations show good correspondence up to \(x_{\text{Ga}} = 0.77\). This corresponds to a lattice mismatch perpendicular to the surface \((\Delta d/d)^2 = -4 \times 10^{-2}\) in the individual GaInAs layers and an average mismatch of the whole MQW structure \((\Delta d/d)^2 = -1.1 \times 10^{-2}\). The values of the layer thickness and strain for the individual MQW layers as well as for the entire MQW structure are approximately equal to the critical thickness for misfit dislocation generation [6,7]. However, the good agreement of the linewidth of the MQW satellite reflections of approximately 110 arcsec between measurement and
simulation indicates that no relaxation of the layers by formation of misfit dislocations has occurred. This becomes clear from Fig. 1 giving an example of a measured and a simulated XRD rocking curve for $x_{Ga} = 0.74$.

To investigate a possible tilting of the MQW layers the dependence of the XRD peak positions on the azimuthal angle $\Phi$ was analysed. The tilt was determined by the dependence of the distance between the substrate reflection and the zeroth-order satellite reflection on the azimuthal angle $\Phi$ following [8, 9]. Fig. 2 shows the dependence of the tilt of the structures grown on off-oriented substrates on the average lattice mismatch $(\Delta d/d)^2$ of the respective MQW structures. For an increase of $(\Delta d/d)^2$ the tilt increases linearly. Additionally, the relationship Nagai [8] derived for an elastically strained layer, namely $\tan \Delta \alpha = (\Delta d/d)^2 \tan \alpha$, where $\Delta \alpha$ is the angle of tilt and $\alpha$ the substrate off-orientation angle, is included into Fig. 2 as solid line. The measured values show good correspondence to the theory again confirming that the samples are not relaxed by dislocations. Because the tilt is a direct consequence of surface steps due to substrate off-orientation [8, 9] the samples grown on exactly oriented substrates show no lattice tilting.

Fig. 2. Dependence of $\tan \Delta \alpha$ on the average lattice mismatch $(\Delta d/d)^2$ using the relationship $\tan \Delta \alpha = (\Delta d/d)^2 \tan \alpha$ [8]. $\Delta \alpha$: tilt angle, $\alpha$: substrate off-orientation angle (here: 2°). The measured data and the theoretical relation show good correspondence.

3.2. Dependence of the growth rate of InP and GaInAs on growth temperature and substrate off-orientation

Fig. 3 shows the InP (Fig. 3a) and GaInAs (Fig. 3b) growth rates $v_{InP}$ and $v_{GaInAs}$ and the Ga content ($x_{Ga}$) in the GaInAs well layers (Fig. 3c) in dependence of the growth temperature for 2° off-oriented substrates (solid circles) and for exactly oriented substrates (open circles). The growth rates $v_{InP}$ and $v_{GaInAs}$ were obtained by dividing the layer thicknesses determined by XRD by the respective growth times in CBE. For all temperatures in the range between 485°C and 545°C the growth rate for InP $v_{InP}$ as well as for GaInAs $v_{GaInAs}$ is larger for the off-oriented substrates than for the exactly oriented substrates, although the individual values are dependent on the growth temperature. The Ga content in the GaInAs well layers also changes with temperature, but is almost independent of the substrate off-orientation in agreement with Ref. [10]. Simultaneously, as noted in Ref. [11] the use of off-oriented substrates improves the surface morphology. Details about the influence of the substrate off-orientation on the surface morphology will be published elsewhere.
The InP growth rate \( v_{\text{InP}} \) is almost constant for growth temperatures from 500°C to 530°C and shows a slight decrease for temperatures beyond this temperature range. For high temperatures this decrease is due to the desorption of In species from the surface [12]. For low growth temperatures \( v_{\text{InP}} \) is reduced because of an incomplete dissociation of TMIn [13].

The variations of \( v_{\text{GaInAs}} \) with the growth temperature are mainly due to changes in the GaAs growth rate for temperatures below 545°C. This is indicated by the temperature dependence of \( x_{\text{Ga}} \) (Fig. 2c) which shows a similar dependence as the GaInAs growth rate. For temperatures above 500°C \( v_{\text{GaInAs}} \) is reduced by increased DEGa desorption from the surface whereas the decrease in \( v_{\text{GaInAs}} \) for temperatures below 500°C is caused by the blocking of free surface sites by adsorbed ethyl groups [14]. For \( T = 545°C \) the desorption of DEGa is lower than the evaporation of In causing the slight increase of \( x_{\text{Ga}} \) although \( v_{\text{GaInAs}} \) is further reduced.

The differences in the growth rates between off-oriented substrates and exactly oriented substrates can be explained by the different growth modes taking place on either type of substrate. For off-oriented substrates mainly step flow growth can be assumed while for exactly oriented substrates layer by layer growth takes place. The then required nucleation of 2D islands of a critical size leads to a higher probability of evaporation of group-III atoms and therefore to a reduced growth rate on exactly oriented substrates.

Additionally, Fig. 3a shows that for off-oriented substrates \( v_{\text{InP}} \) decreases only for \( T > 530°C \) and stays almost constant between 500°C and 530°C. For exactly oriented substrates, however, \( v_{\text{InP}} \) already starts to decrease within this temperature range. This indicates a continuous In evaporation with increasing temperature for exactly oriented substrates, while for off-oriented substrates the surface steps act as a barrier retarding the In evaporation up to a critical temperature \( T > 530°C \) in our case.

From Fig. 3c it becomes clear that for our growth conditions the evaporation rate of In and Ga atoms is approximately the same for off-oriented and exactly oriented substrates. Only at \( T = 545°C \) the Ga content is higher on off-oriented than on exactly oriented substrates suggesting a delayed Ga evaporation from off-oriented substrates. Our results additionally indicate that \( v_{\text{GaInAs}} \) is independent of the incorporated strain and simply remains proportional to the total group-III flux at a given temperature.

### 3.3. Effects of tensile strain on the optical properties of the structures

Fig. 4 shows the dependence of the PL line width of the MQW peak on the tensile strain in the...
GaInAs well layers for exactly and off-oriented substrates. The energy of the respective transitions is approximately 1 eV and depends on the thickness and Ga content of the individual MQW structures.

For exactly oriented substrates the line width does not change up to the maximum Ga content \( x_{Ga} = 0.77 \). For off-oriented substrates the PL line width is the same as for exactly oriented substrates for lattice matched structures (\( \Delta d/d_{GaInAs} = 0 \)), but increases linearly with the strain. Note that the PL line width shows no dependence of the growth temperature in the range from 500°C to 530°C.

The increase in the PL line width can be caused by a number of effects such as changes in alloy distribution, roughening of the interfaces or lateral well thickness fluctuations. However, the very clear trend shown in Fig. 4 leads us to the conclusion that the strain induced lattice tilting is the dominant contribution on substrates with a relatively high step density. Due to the different step heights for materials with different lattice constants a pure tetragonal distortion at the step edges is not possible. This leads to the formation of local strain fields being highest at the step edges with increasing strain in the GaInAs layers. Thus, the layers show lateral and periodic strain fluctuations. The non-tetragonal distortion at the step edges eventually leads to the lattice tilt [8].

From literature it is known that GaInAs layers with high tensile strain can have a wavy surface [15]. Such surface undulations are known to decrease the strain by elastic deformation. However, cross-sectional TEM reveals that this applies for both exactly and off-oriented substrates in our case. Therefore, these quantum well thickness variations cannot explain the trend observed in Fig. 4, although they might also lead to a broadening of the PL peaks. Further investigations are necessary, how the PL line width is affected by the undulations. Details about the layer morphology of the structures in dependence of strain and substrate off-orientation will be published elsewhere.

4. Summary

We have investigated the influence of the process parameters on the CBE growth of tensile-strained GaInAs/InP MQW structures. The samples were characterized by HR-XRD, PL and cross-sectional TEM. The exact layer structure was obtained by comparing XRD measurements to simulations. The GaInAs as well as InP growth rate on off-oriented substrates is found to be higher than on exactly oriented substrates. This is attributed to the fact that for off-oriented substrates mainly step flow growth can be assumed while for exactly oriented substrates layer by layer growth takes place. The then required nucleation of 2D islands of a critical size leads to a higher degree of evaporation of group-III atoms and, therefore, to a lower growth rate for exactly oriented substrates over the whole temperature range under investigation.

PL measurements on samples grown on exactly oriented substrates show that the line width of the GaInAs MQW peak stays constant with increasing tensile strain. For off-oriented substrates the line width increases linearly with the tensile strain in the GaInAs layers. This strain-induced broadening of the PL peaks is explained by the increase of the lattice tilt with increasing strain for epitaxial growth on off-oriented, i.e. stepped substrates.

Fig. 4. Photoluminescence line width of the GaInAs MQW peak as a function of the lattice mismatch \( (\Delta d/d)^{GaInAs} \) of the GaInAs layers for off-oriented (solid markers) and exactly oriented (open markers) (100) InP substrates. The included lines are guides to the eye.
leading to an increase of the local strain fields at the step edges. These strain fields eventually causes the observed broadening of the PL peaks.

Our results indicate that for high-quality devices based on highly strained MQW structures the substrate off-orientation has to be reduced as far as possible.

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References


Organized growth of GaAs/AlAs lateral superlattices on vicinal surfaces: where are the limits?


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Abstract

We report on three issues concerning the growth of GaAs/AlAs lateral superlattices on vicinal surfaces to obtain quantum wires with negligible inter-wire tunnel coupling. We show that the ledge roughness increases with increasing terrace length and decreases for slower growth rates. The tilt angle of lateral superlattices is not the relevant parameter as long as electronic properties are concerned: one has to consider the product of the coverage error by the lateral superlattice thickness. The Ga flux stability is a critical parameter to be controlled. The use of dual-filament cells improves it by a factor two compared to single-filament ones.

1. Introduction

Despite an efficient Al/Ga vertical atomic exchange [1], ordered GaAs/AlAs growth on vicinal surfaces is now a well-recognized approach leading to one-dimensional electron systems (1D ES) with new and fascinating electronic properties [2]. The strength of the lateral potential modulation is of the order of the Fermi energy (20 meV) whereas the lateral period is of the order of the Fermi wavelength (30 nm). Electronic properties are described in a lateral superlattice (LSL) picture, i.e., multiple quantum wires with significant inter-wire tunnel coupling. In order to investigate such 1D ES without inter-wire tunnel coupling, one has to increase the potential energy over confinement energy ratio either by increasing the LSL Al content or by increasing the lateral period using vicinal surfaces with smaller misorientation angle.

Increasing the terrace length of the vicinal surface up to 80 nm raises several problems concerning the LSL growth. The first issue is related to the morphology of the step array, the periodicity of which has to be optimized. The second issue concerns the LSL tilt. Indeed, if the LSL coverage (GaAs and AlAs) differs from one by $\epsilon$ then the LSL tilt angle $\beta$ is given by $\beta = \arctan \frac{\epsilon}{\alpha}$ where $\alpha$ is the misorientation angle. Since $1/\alpha$ is typically 200–300, one would think that vertical LSL cannot be made. We will show here that the relevant parameter is not $\beta$ but the product $N\epsilon$ as far as electronic properties are concerned (here $N$ is the number of LSL

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MonoLayers (ML)). Since $N$ is typically 20–30, this makes the flux control ten times easier! A third issue concerning LSL growth lies around the flux stability so that the Ga and Al fluxes are constant throughout the LSL. This is to prevent the formation of LSL with random tilt variations. With respect to this, we will show that the proper use of a dual-filament Ga cell improves the flux stability compared to a single-filament one.

2. Steps periodicity at low misorientation angles

Our purpose is not to discuss the reasons of the build-up of a periodic step array (presented elsewhere [3]) but to investigate the limits for such a build-up, the adjustable parameter being the misorientation angle. Fig. 1 shows AFM images of GaAs surfaces misoriented toward [1 1 0] by 0.2° and 0.3°. These images are obtained in a N$_2$ dry box in less than 1 h after being taken out of the molecular beam epitaxy (MBE) system. The epitaxy consists of a 1 μm GaAs buffer grown at 2 Å/s with a substrate temperature $T_{sub} = 620$°C followed by 200 nm of GaAs grown at 0.5 Å/s with $T_{sub} = 580$°C. For these two temperatures, the As$_4$ flux is minimized in order to keep an As-rich surface during MBE. To enhance surface diffusion, migration enhanced epitaxy (MEE) (alternate supply of element III and V) [4] has been used to grow the last 10 nm of GaAs in one sample at a growth rate of 0.1 Å/s while keeping $T_{sub}$ at 580°C.

For the two MBE grown samples, step edges along [1 1 0] are not straight because steps move by incorporating GaAs islands that develop on large terraces under these growth conditions. The ledge roughness is more pronounced for lower misorientation (0.2°) because nucleated islands are larger on wider terraces. The improvement of the surface of the MEE grown sample is readily apparent in Fig. 1, where the ledge roughness is considerably reduced, whereas the step periodicity is more well-defined at 50 nm. On this MEE sample, regions with periodicities close to 60 nm are observed with rough ledges so that 50 nm is considered as the largest terrace width for which we obtain smooth ledges and periodic steps at 580°C by MEE.

3. Scaling laws in tilted LSL

As LSL are designed to be active regions so as to confine 1D ES, it is of primary importance to understand the effect of an imperfect coverage of the surface by LSL layers in order to specify flux control. Fig. 2 depicts a tilted LSL in which each LSL ML covers a fraction $1 + \varepsilon$ of the surface. As

![Fig. 1. Atomic force microscopy of GaAs vicinal surfaces. The misorientation and the final growth mode (MBE or MEE) used prior to sample cooling are indicated.](image-url)
mentioned above, the tilt angle is given by $\beta = \arctan c/a$. If we consider that in each LSL ML the Al composition along $[1 0 1]$ (or $x$) is given by a cosine law, the stacking of $N$ such MLs results in the following lateral Al content:

$$X_{Av}(x, z) = X_{Av}(1 + \lambda \cos \left( \frac{2\pi}{L_x} - \frac{2\pi x}{a} \right)),$$  \hspace{1cm} (1)

where $X_{Av}$ is the LSL mean Al content and $\lambda$ describes the amplitude of lateral ordering ($\lambda = 0$ in the case of a random growth, $\lambda = (2/\pi)X_{Av}$ in the case of a perfectly ordered growth). In Eq. (1), the tilt or imperfect coverage acts through the phase term $2\pi x/a$. If the product $Ne$ is much smaller than 1, all LSL layers are considered in-phase so that the LSL can be treated as a vertical one at least as far as electronic properties are concerned. If $Ne$ is much larger than one, the electronic envelope wave function will average the tilted LSL composition profile to $X_{Av}$ so that the modulated part goes to zero.

To estimate this averaging effect quantitatively, we insert this LSL potential in an infinite square quantum well (QW) and compute matrix elements between envelope functions associated to the subband defined by the infinite QW. Envelope functions $\Psi_n(z)$ are of the form:

$$\Psi_n(z) = \sqrt{\frac{2}{Na}} \sin \left( \frac{n \pi z}{Na} \right).$$  \hspace{1cm} (2)

Matrix elements $V_{z,n}^m(x)$ can be considered as effective potentials scattering an electron of the subband $n$ into the $m$ one. We get

$$V_{z,n}^m(x) = -\frac{V_0^2}{\pi} A_{z,n}^m$$

$$\times \cos \left( \frac{2\pi x}{L_x} + \pi Ne + (m + n) \frac{\pi}{2} \right).$$  \hspace{1cm} (3)

$$A_{z,n}^m = \frac{16\pi m N_e}{[(m - n)^2 - 4N_e^2\varepsilon^2][(m + n)^2 - 4N_e^2\varepsilon^2]}$$

$$\times \sin \left( \pi Ne + (m + n) \frac{\pi}{2} \right).$$  \hspace{1cm} (4)

where $V_0$ is a term related to the band offset and proportional to $X_{Av}$. Matrix elements preserve the $2\pi/L_x$ periodicity defined by the step array as it appears in Eq. (3). The phase term $\pi Ne + (m + n)\pi/2$ is of no importance in Eq. (3), whereas in Eq. (4), it governs the dependence of the effective lateral potential amplitude $A_{z,n}^m$ in Eq. (4). This amplitude in Eq. (4) depends only on $\varepsilon$ and not on $x$ or $L_x$, unlike the tilt angle $\beta$. This demonstrates that the relevant scaling parameter for the electronic properties is $\pi Ne$ not $\beta$.

Alternatively, if a large and constant tilt angle is chosen, let us say $\beta = 80^\circ$, a large lateral potential is obtained only if $N$ is smaller than 1. This means that the LSL has to be thin in order to exhibit 1D electronic properties, with a thickness smaller than $N_e = \cot \beta/\pi$ with $N_e = 10$ for $x = 1^\circ$ and $\beta = 80^\circ$.

Fig. 3 shows the variations of $A_{z,n}^m$ with $Ne$ for $\{m, n\} = \{1, 2\}$. It is clear that the intra-subband potential ($m = n$) decreases for large $Ne$, whereas the first inter-subband potential exhibit a resonance for $Ne = 1$. This inter-subband coupling resonance has been recently evidenced in optical experiments performed on doped tilted LSL [5]. This demonstrates that a tilted LSL cannot be considered as a vertical one of smaller amplitude: a vertical LSL provides a separable potential.

![Fig. 2. Schematic view of a tilted lateral superlattice.](image)

![Fig. 3. Universal amplitude of the effective potential for the first two subbands.](image)
(i.e. depending only on x), whereas a tilted LSL does not.

4. Ga flux stability

From the previous considerations it is clear that the coverage should be constant throughout the LSL growth. This requires highly stable Ga and Al fluxes. It is widely accepted that the main emphasis should be put on the former. Dual filament effusion cells allow to operate a Ga cell with the tip of the crucible heated independently and hotter than the base of the crucible to prevent accumulation of Ga droplets at the tip. This avoids cold droplets to change the thermal equilibrium of the Ga bath when they flow from the tip to the base of the crucible. This mechanism is the main source of flux instability in Ga effusion cells. We show here that the use of a dual filament cell requires an adequate regulation strategy of both tip and base filaments to get highly stable fluxes. These fluxes are measured by reflection high-energy electron diffraction (RHEED) oscillations on a 2×2 mm GaAs substrate at $T_{\text{sub}} = 560^\circ\text{C}$ to ensure a unity Ga sticking coefficient. The RHEED oscillation trace is then analyzed by mean of Fourier transform in order to get the frequency with a high accuracy. Then the dispersion of the data is analyzed over almost an hour.

Fig. 4a shows flux measurements starting 1 h after outgassing Ga cells 100°C above their operating temperature. The single-filament cell displays a noisy flux (statistical standard deviation $\sigma = 0.25\%$), whereas the dual-filament one is less noisy but shows a systematic drift ($\sigma = 0.2\text{–}0.6\%$). Fig. 4b displays similar measurements recorded 4 h after outgassing. The systematic drift of the dual-filament cell has disappeared and the stability is excellent ($\sigma = 0.1\%$ or less), whereas the stability of

![Fig. 4 (a) Ga flux is plotted versus time after shutter opening for two different cells (circles and triangles) and two different runs (open and filled symbols). Each set of data is normalized to 1 and shifted vertically for clarity. (b) Ga flux is plotted versus time for two different cells (filled circles and triangles). Open circles show the Ga flux when the tip power is regulated with the tip thermocouple.](image-url)
the single-filament cell has only slightly improved ($\sigma = 0.18\%$). This confirms that Ga droplets at the crucible tip are the main source of Ga flux instability since visual inspection of both crucibles reveals a large Ga accumulation at the tip of the single-filament cell and the absence of any Ga droplet at the tip of the dual-filament one.

Concerning the regulation strategy we omitted to say that the dual-filament cell was operated with a base filament regulated with the base thermocouple and a constant power injected in the tip filament to ensure identical temperatures on both base and tip thermocouples when the cell shutter is closed. This strategy yields the most stable fluxes. As a matter of comparison we display on Fig. 4b the measured fluxes when both filaments are regulated on their respective thermocouple with identical set-points. We observe then a very long transient: the Ga flux never stabilizes at a given value over 45 min! This is certainly due to the change in the Ga bath thermal equilibrium when the power increases in the tip filament after the shutter is opened. When operating this cell with a constant power in the tip filament (i.e. with a tip 30\(^\circ\)C colder than the base) we never detected any Ga droplets accumulation provided the cell is outgassed every day with a tip at the same temperature than the base.

5. Conclusion

We have considered three important issues related to LSL growth on vicinal surfaces. To obtain periodic step arrays with large periodicities requires to grow GaAs and AlAs in the MEE mode. Even if large periodicities (i.e. small misorientation angles) would lead to strongly tilted LSLs we have shown that the flux control is determined only by the LSL thickness probed by the electronic wave function as long as electronic properties are concerned. This makes the flux control ten times easier. Finally, the use of dual-filament Ga cell improves the flux stability compared to single-filament ones provided a constant power and not a regulated one is injected in the filament heating the tip of the crucible.

References

Formation of multi-atomic steps and novel n-AlGaAs/GaAs heterojunctions on vicinal (1 1 1)B substrates by MBE and anisotropic transport of 2D electrons

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Abstract

We have investigated the MBE growth of GaAs on vicinal (1 1 1)B substrates and examined in detail the multi-atomic step structures appearing at the growth front as well as at the n-AlGaAs/GaAs heterojunction. It has been found that the uniformity and periodicity of such steps depend strongly on the growth condition, especially As\textsubscript{4} flux, and that their morphology drastically affects transport of electrons flowing along heterojunctions. The use of high As\textsubscript{4} flux is effective in improving the step uniformity, and in drastically reducing the channel resistance perpendicular to the steps. Key factors for the fabrication of high-quality devices such as quantum wires on (1 1 1)B planes have been discussed.

To fabricate several semiconductor devices, it is important to prepare GaAs/AlGaAs structures with very smooth (1 1 1)B surfaces and interfaces. For example, quantum wells (QWs) on (1 1 1)B are useful to reduce the threshold current of lasers due to its unique valence band structure [1]. As for piezo-electric effect, it becomes maximum on the (1 1 1) plane among low-index planes [2]. To fabricate very narrow quantum wires (QWRs), this plane is attractive, since ridge QWRs and edge QWRs can be fabricated on GaAs/AlGaAs-facet structures having two (1 1 1)B side planes and (0 0 1) plane [3–5].

Especially for the edge QWRs, the smoothness on (1 1 1)B facets is essential to make high-quality wires. Earlier, we investigated the morphology of slightly misoriented (1 1 1)B facet by atomic force microscopy (AFM) and detected multi-atomic steps with \sim 30 nm periodicity [6]. In making the edge QWR, one must clarify the role of such steps and the way to control them, since electrons flow across these steps.

In this paper, we investigate these issues to show that very regular steps can be prepared on vicinal (1 1 1)B planes and that a highly conductive channel can be formed for electrons flowing across these steps. In particular, we study by AFM how multi-atomic steps on vicinal (1 1 1)B substrates depend on As\textsubscript{4} flux. We also investigate how the morphology of these steps affects the transport of...
two-dimensional (2D) electrons in selectively doped n-AlGaAs/GaAs heterojunctions [7].

In this work, we use GaAs substrates having vicinal (1 1 1)B plane with misorientation angle $\Delta \theta$ of 2° toward the nearest (1 0 1) plane. Fig. 1a is an illustration of the top view of (1 1 1) or (1 1 1)B orientation with respect to other orientations. Fig. 1b shows the cross-sectional view cut along line $\alpha-\beta$ in (a) and also the misorientation $\Delta \theta$. As will be described later, multi-atomic steps are parallel to the intersection of (1 1 1) and the (1 0 1) plane [6, 7]. If another misorientation direction is used, (1 1 1)B surfaces will be rough due to faceting.

We have studied first how the morphology of 0.6 $\mu$m-thick GaAs layer grown at 580°C depends on As$_4$ fluxes. The As$_4$/Ga flux ratios of beam-equivalent pressure were set 50 or 130, respectively. GaAs growth rate is 0.26 $\mu$m/h. Fig. 2a and Fig. 2b show AFM images of a GaAs surface grown with low As$_4$ flux. In the height image (a), the height of the structure is indicated by the brightness. In the derivative image (b), dark lines and bright regions correspond to multi-atomic steps and flat terraces, respectively. Note that the step periodicity is not uniform. For the sample grown with high As$_4$ flux, however, steps are quite straight and uniformly spaced with the main period of $\sim$ 20 nm as shown in Fig. 2d and Fig. 2e.

We plot the cross-sectional height profiles of these two samples in Fig. 2c and Fig. 2f, where two distinctive differences are noteworthy. Large amplitude steps having the height of 3 nm and the long spacing of several hundred nanometers are seen only in Fig. 2c, whereas regular steps with the average height of 0.4–0.7 nm are seen in Fig. 2f. Thus, steps of Fig. 2f are about two monolayers in height. The other point to note is the presence of very steep steps in Fig. 2c, whose slope is as large as 20°. In contrast, no such steps are seen in Fig. 2f with the slopes being several degrees at most. As will be discussed later, a large anisotropy in electron transport appears only in the presence of high-amplitude steps.

We discuss the relation between As$_4$ flux and the step uniformity. It has been pointed out that step bunching will appear when adatoms are incorporated preferentially at down-steps. The spacing of steps will be equalized when adatoms are incorporated selectively at up-steps. In case of vicinal (1 1 1)B plane, As atoms stick with single bond at step edges as shown in Fig. 1c, since the surface orientation at the step is close to (1 1 0) or (1 1 1)A. The probability of an As atom to occupy this site at the step edges may not be high as the site is unstable but it can be enhanced by increasing the As$_4$ flux. Hence, the existence of these As atoms at the step edges may enhance the possibility of Ga-atom incorporation at up-steps as indicated by open arrow in Fig. 1c and may contribute to an equalization of the step spacing.

To investigate how the morphology of steps affects the transport of electrons, we deposited an n-type AlGaAs layer on two kinds of GaAs layers.

Fig. 1. (a) Top view of crystal orientations near (1 1 1), or (1 1 1)B plane. The cross-sectional view seen at line $\alpha-\beta$ of (a) is shown in (b) together with the misorientation angle $\Delta \theta$. (c) Atomic-scale structure of the step on the vicinal (1 1 1)B plane.
having multi-atomic steps, as shown in Fig. 3a. After growing a 0.6 μm GaAs, we grew a 20 nm-thick undoped Al$_{0.33}$Ga$_{0.67}$As spacer layer, a 90 nm-thick Si-doped Al$_{0.33}$Ga$_{0.67}$As with the dopant density of $6 \times 10^{17}$ cm$^{-3}$, and a 20 nm-thick GaAs cap layer. The growth condition was similar to that for the earlier study, where As$_4$/Ga flux ratios of beam-equivalent pressure were 55 and 170 for the two samples. By using each of these wafers, two kinds of FETs with the channel of 50 μm in width and 600 μm in length were formed as shown in Fig. 4a. Note that 2D electrons at these
corrugated heterojunctions will flow either perpendicular (\(\perp\)) to or parallel (\(|\rangle\)) to the multi-atomic steps. The Hall measurement done at 14 K on a sample grown with high As\(_4\) flux has shown that the mobility \(\mu\) and concentration \(N_e\) of 2D electrons are \(80,000 \text{ cm}^2/\text{V s}\) and \(1.9 \times 10^{11} \text{ cm}^{-2}\), respectively. From the Shubnikov de Haas measurement on this sample, \(N_e\) was found to be \(3.6 \times 10^{11} \text{ cm}^{-2}\), though the origin for this difference is not clear yet. Hall measurement on the other sample will be described elsewhere, since their behavior is complicated as shown below.

Fig. 4b shows current \(I_d\) – voltage \(V_d\) characteristics of these devices measured at 4 K for the two current directions. In the sample grown with low As\(_4\) flux, the current \(I_{d//}\) flowing parallel to the steps (blank squares) is high but current \(I_{d\perp}\) perpendicular to the steps is extremely small and nonlinear for \(V_d\) as shown by blank circles, indicating an extremely large anisotropy. Only the current \(I_{d\perp}\) perpendicular to the steps is blocked, probably by large potential barriers. Here, we confirmed that source and drain electrodes are good and ohmic by measuring the channel conductance parallel to the steps. When high As\(_4\) flux is used, currents for both directions are linear and almost isotropic, and are as large as the parallel current \(I_{d//}\) of the sample grown with low As\(_4\). These results indicate that very smooth (1 1 1)B planes can be prepared and that highly conductive channels can be formed even for the direction perpendicular to the steps. Hence, high-quality transport devices and QWRs can be prepared on (1 1 1)B planes.

In the sample grown with low As\(_4\), potential barriers appear for electron motion perpendicular to the multi-atomic steps as mentioned above. Mechanisms of the barrier formation are not clear yet, but it should be related to the presence of large steps or steep steps as shown in Fig. 2c. One possible reason is the roughness itself at the heterojunction. As illustrated in Fig. 3a, 2D-electrons flowing across the steps will be reflected at the steps, since the potential \(V(x, z)\) at points A and B are displaced as shown by solid and dashed lines in Fig. 3b. Although the details of potential profile

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![Fig. 3. Cross-sectional diagram (a) of modulation doped n-AlGaAs/GaAs heterojunction with a corrugated interface. Potential profiles (b) at points A and B, indicating the appearance of a potential barrier at the steps.](image)

![Fig. 4. FET patterns (a) to measure the current \(I_d\) perpendicular to and that \(I_{d\perp}\) parallel to the steps. The drain current \(I_{d\perp}\) and \(I_{d//}\) were measured at 4 K as a function of \(V_d\). The data for the sample prepared under high As\(_4\) flux are shown by solid circles \((I_{d\perp})\) and squares \((I_{d//})\). The data of the sample grown under the low As\(_4\) flux are shown by open circles \((I_{d\perp})\) and squares \((I_{d//})\).](image)
is not known, earlier models on interface roughness scattering have shown that the effective potential \( \Delta V \) caused by the interface displacement is of the order of the interface electric field \( F \) multiplied by the roughness height \( \Delta z \). Hence, by setting \( \Delta z = 3 \text{ nm} \) and \( F = 50 \text{ kV/cm} \) at sheet electron density \( N_s \) of \( 3.6 \times 10^{11} \text{ cm}^{-2} \) [8], we estimate that the effective potential \( \Delta V \) introduced by roughness may be of the order of 15 meV. Since this value overestimates the effect of roughness, a more rigorous analysis is certainly required not only by refining the mathematical model but also by considering the possible variations of spacer layer thickness and the donor concentrations.

Another possible reason for the appearance of potential barriers is the anomalous incorporation of Si atoms in the region of multi-atomic steps. Since (1 1 0), (1 1 1)A, and crystallographic planes other than (1 1 1)B show up at the steps of our sample as shown in Fig. 1c, some fraction of Si atoms, which fall directly on or migrate to these planes, can be either inactivated or incorporated into acceptor sites. If such processes take place, the local potential near the steps is raised and the electron concentration is locally lowered, resulting in a kind of potential barriers. This anomalous incorporation of Si dopants is quite probable, since Si atoms on (1 1 1)A and related planes of GaAs are known to work as acceptors [9], unless the layer is grown with very high As\(_4\) flux at low temperatures.

In summary, it is shown that the growth of GaAs with low As\(_4\) flux on vicinal (1 1 1)B planes results in aperiodic multi-atomic step structures. When the layer is grown with high As\(_4\) flux, multi-atomic step become evenly spaced with typical period of 20 nm. The transport of electrons is found to be scarcely affected by evenly spaced step structures but strongly blocked by aperiodic steps with high amplitude, leading to a huge in-plane anisotropy of conductance in excess of 100. The key factor for high-quality QWRs using (1 1 1)B facets is clarified.

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References

MBE growth of two-dimensional electron gases on (110) GaAs

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Abstract

We have studied the influence of growth temperature on the electrical properties of two-dimensional electron gases, grown by molecular beam epitaxy on (1 1 0) oriented GaAs. The mobility and carrier density of the grown layers depend strongly on the chosen substrate temperature, having a narrow maximum around 470°C. Growth at 10–20°C below the optimum temperature results in insulating layers. We show that this steep drop in mobility and carrier density is strongly correlated with the carbon incorporation observed in photoluminescence. In the temperature range 480–520°C the mobility is only about 60% of the peak value and both mobility and carrier density change only slightly. Above 520°C the mobility drops again whereas the carrier density stays virtually unchanged. We believe this to be caused by interface roughness.

1. Introduction

Motivated by the current strong interest in low-dimensional systems, grown by cleaved edge overgrowth (CEO) \[^{1}\], we have investigated how the properties of two-dimensional electron gases (2DEG’s) grown by molecular beam epitaxy (MBE) on (1 1 0) GaAs are influenced by the substrate temperature.

For the realization of high-quality one-dimensional (1D), or modulated 2DEG samples, by CEO \[^{2}\], one has to grow a good-quality 2DEG on the cleaved (1 1 0) edge of an already grown GaAs/AlGaAs structure.

The growth of high-quality material on the (1 1 0) surface has already been reported \[^{3–8}\]. The optimal growth conditions for the (1 1 0) surface use a high V/III flux ratio and a low substrate temperature compared to growth on a conventional (1 0 0) surface. These conditions compensate for the low incorporation coefficient of the As to the non-polar (1 1 0) surface. An additional improvement in the quality of the grown layers can be achieved by using As\textsubscript{5} instead of As\textsubscript{4} \[^{6–8}\]. In this paper we investigate a total of 16 2DEG samples grown in the temperature range 440–595°C. We find that the low-temperature transport properties are very sensitive to small variations (± 10°C) in the growth temperature. The grown layers were also characterized by low-temperature photoluminescence (PL), and in the temperature range 440–480°C we find a strong correlation with the transport properties.
2. Experimental procedures

The layers were grown in a Varian Modular Gen II, 3-inch MBE machine using a Varian cracker source for the arsenic (As$_2$).

The (110) oriented substrates used were semi-insulating GaAs 'epi-ready' 2 in. diameter wafers, cut with an orientational accuracy of $\pm 0.25^\circ$, from American Xtal Technology.

The growth rates used were about 0.2 $\mu$m/h for AlAs and 0.5 $\mu$m/h for GaAs, as measured by RHEED oscillations on a (100) calibration wafer prior to growth. The V/III beam equivalent pressure (BEP) ratio was 20-30. A more detailed description of the system and growth procedure can be found in Ref. [6].

All substrate temperatures in this work are the temperatures indicated by the thermocouple ($T_{TC}$), located behind the wafer. Using an IRCON series V infrared pyrometer we obtain the following relation between the temperatures read by the pyrometer ($T_{pyr}$) and the thermocouple ($T_{TC}$): $T_{pyr} = 0.78T_{TC} + 56.4$ ( emissivity set to 0.7).

All 16 samples were grown with an identical structure schematically shown in Fig. 1.

The structure used was not designed for obtaining maximum mobility [9], but instead designed as a test structure, insuring a high carrier density in the 2DEG under the growth conditions used in this work. We have employed the technique of $\delta$-doping in these structures to reduce the effect of auto-compensation [6].

After growth the samples were cut in 5 x 5 mm pieces, and in order to make electrical measurements, ohmic contacts were formed by alloying small Sn beads at 300°C for 1 min, in an atmosphere of dry N$_2$.

The sample configuration used was a standard van der Pauw square sample, with a contact in each corner.

3. Electrical characterization

Of the 16 samples we succeeded in contacting 13, grown in the temperature range 440–560°C. Two samples grown at 582°C and 595°C were insulating. The samples grown at 440°C and 560°C were difficult to contact, giving non-linear $I/V$ characteristics, and the exact values of electron mobility and

![Fig. 1. Schematic view of $\delta$-doped 2DEG structure. $\delta$-1 and $\delta$-2 are Si delta-doped layers with densities of $3.5 \times 10^{12}$ and $1.5 \times 10^{12}$ cm$^{-2}$.](image)

![Fig. 2. Electron mobility (a) and carrier concentration (b) as a function of substrate growth temperature. Measurement temperature is 10 K.](image)
carrier density are therefore uncertain. Fig. 2 shows mobility and carrier density as measured at 10 K in the dark. The two shaded regions indicate where the samples gradually become insulating. The curves are intended to guide the eye only.

As seen in the plot of mobility versus growth temperature (Fig. 2a), the mobility has a maximum around a temperature of 470 ± 5°C. At lower temperatures the mobility drops drastically, and at 440°C the samples become insulating. The drastic drop in mobility is followed by an equally steep drop in the carrier concentration. This drop in mobility is correlated with the PL measurements which are discussed in detail in Section 4 (Fig. 3).

In the temperature range 480–520°C the mobility and carrier density change only little. The mobility is, however, only about 60% of the peak mobility.

At temperatures higher than 520°C there is again a steep drop in mobility. However, in this range the carrier concentration stays virtually unchanged, and we see no significant changes in the PL spectra.

The surface morphology is excellent and mirror like for the samples grown at temperatures below 470°C, while the surfaces of the samples grown in the range 480–520°C show a gradual increase in the number and size of facets, as also observed by others [8–10]. At temperatures higher than 520°C the surface displays a clear faceting, indicating a Ga-rich growth caused by a poor As sticking coefficient. The limiting factor of the 2DEG mobility at the high growth temperatures thus seems to be the roughness of the GaAs/AlGaAs interface.

For reference, we have also grown the same 2DEG structure on a (1 1 0) oriented substrate, under optimum (1 0 0) conditions (TTC = 730°C and minimally As rich). This sample had a carrier density of 6.0 × 10^{11} cm^{-2} and a mobility of 300 000 cm^{2}/(V s). The carrier densities of the best (1 1 0) samples are thus only about 10% lower than the (1 0 0) reference. The mobility, however, is only one third of the (1 0 0) mobility.

4. Optical characterization

The optical quality of the (1 1 0) layers was investigated by low-temperature photoluminescence. The samples were mounted in a continuous-flow cryostat using liquid He as coolant. During the measurements the sample temperature was kept at 4 K, and the luminescence was excited by the 632.8 nm line of a HeNe laser (intensity 4 W/cm^{2}). A 0.66 m focal length spectrometer was used to disperse the luminescence. The signal was detected by a GaAs photomultiplier and recorded using lock-in techniques.

From the total of 16 samples, we chose nine samples grown in the temperature range 440–560°C for elaborate PL characterization. All
nine samples were characterized using three different excitation intensities: 4, 0.4 and 0.04 W/cm², thus obtaining a total of 27 spectra.

In Fig. 3, we display two representative spectra, one taken at the high-temperature side of the mobility peak and one taken on the low-temperature side.

The spectrum in Fig. 3a is taken from the sample grown at 480°C and shows a single dominant line at 1.5152 eV with a width of 1.0 meV. This is assigned to the emission from the n = 1 state of the free exciton, and is labeled by Xₙ₋₁. The peak at 1.5183 eV, on the high-energy side of the free exciton, indicates the n = 2 state of the free exciton (Xₙ₋₂). At the low-energy side we observe a peak, which we attribute to excitons bound to neutral donors (D¹X).

The signal at 1.495 eV has an intensity 15 times lower than the dominant free-exciton line. It is assigned to transitions involving carbon acceptor states, as commonly observed in PL of MBE-grown GaAs material [11] and labeled C₅ in the figure.

The spectrum taken from the sample grown at 460°C is quite different as seen in Fig. 3b. In this sample the peak related to carbon acceptor states is the dominant one. The intensity ratio of the free-exciton line to the carbon-related luminescence (Xₙ₋₁/C₅) is in this case 0.7.

As can be seen, there is a substantial difference in the two spectra, the spectrum from the sample grown at 480°C indicates a high-quality material, whereas the spectrum from the 460°C sample is dominated by the carbon-related impurities. This is, as mentioned earlier, strongly correlated with the drop in carrier concentration and mobility.

We have also investigated the samples grown at higher temperatures and in this case the spectra remain virtually unchanged, even past the point where the mobility displays a steep drop. However, for the samples grown at 480°C and higher we start to see a small signal at 1.48 eV (Fig. 3a), which we attribute to Si acceptors. At higher temperatures the sticking coefficient of As starts to drop, and the chance of Si incorporating as an acceptor on an As site increases. This is also consistent with the observed small drop in carrier concentration as seen in Fig. 2b.

Fig. 4 shows the ratio of PL intensity between the free-exciton emission (Xₙ₋₁) and the carbon-related transitions (C₅) versus growth temperature. Data from all the 27 spectra are shown.

The three curves, taken at different excitation intensities (4, 0.4 and 0.04 W/cm²), all show the same general tendency, namely that the intensity ratio Xₙ₋₁/C₅ changes drastically when the growth temperature is reduced from 480°C to 460°C. All spectra taken from samples grown at temperatures > 480°C are dominated by the free-exciton emission, whereas all spectra taken from samples grown at temperatures ≤ 460°C are dominated by the emission from the carbon-related transitions.

5. Conclusion

We have shown that using the optimum growth temperature is extremely important for the quality of 2DEG's on (1 1 0) GaAs. A growth temperature which is 10-20°C below the optimum temperature, yields an almost insulating layer. In CEO, it is common practice to mount the samples to be cleaved, on a bracket, mounted on a modified
substrate holder [12, 13]. This bracket sticks out 3–5 mm from the holder at an angle of 90° towards the sources. This mounting procedure will inherently lead to a lower temperature of the cleaved surface, as compared to a reference sample mounted in the normal position [13].

As we have shown, this difference in temperature will affect the low-temperature transport properties of the CEO samples seriously.

The cause of the drop in mobility and carrier density at temperatures lower than 470°C has been shown to be related to the enhanced incorporation of carbon, at the low growth temperature used, whereas the drop in mobility at temperatures higher than 520°C is probably related to the roughness of the grown layers.

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References

The building up of terrace periodicity by MBE growth on (0 0 1) GaAs vicinal surfaces


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Abstract

The morphology of MBE grown GaAs vicinal surfaces is studied using ex situ AFM. Analysing the statistical distribution of terrace width, we succeed in explaining our measurements either by a thermodynamical equilibrium model taking into account the step by step interaction or by a 2D Monte Carlo simulation of the growth. In the latter case, an anisotropic Schwoebel barrier at the step edge has to be introduced in order to get a good agreement. We argue that growth is the efficient process to get a low disordered step array at the usual time scale and temperature range of MBE. In addition the effect of a GaAs/AlAs buffer superlattice, or of pure AlAs layer, or of Si doping in GaAs is also discussed.

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Keywords: Molecular beam epitaxy; Vicinal surfaces; Monte Carlo simulation; Atomic force microscopy

1. Introduction

Organized growth on vicinal surfaces appears as a simple and attractive way to realize, in a single technological step, high quality 1D GaAs/AlAs quantum structures [1] with novel and interesting optical [2] and electrical [3] properties. A necessary requirement is obviously to get and to keep a surface with a nearly periodic arrangement of the step array. In this work we use vicinal surfaces with monomolecular Ga terminated steps (type A) for a large range of misorientation angle (the corresponding mean terrace length \( L \) varies from 8 to 75 nm). We used solid source MBE (Riber 2300) and a commercially available Atomic Force Microscope (Nanoscope III). The AFM measurements are performed outside the MBE system with the entire scanning unit placed in a plexiglass glove box purged with pure and dry nitrogen gas. We worked with a silicon cantilever in the contact mode and the applied force is less than 10 nN [4].

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2. Building up of terrace periodicity

Before any growth, but after oxide sublimation and a 5 min thermal annealing at 650°C under As flux in the growth chamber, although the RHEED pattern is 2D-like and reconstructed, the AFM scan reveals that the substrate surface is in fact very rough and exhibits large anisotropic structures elongated along [1 1 0] (they are typically 100 nm wide, 500 nm long, 3–6 nm deep). During subsequent GaAs buffer growth, the splitting of the RHEED specular spot (observed for misorientation angle \( \alpha > 0.5^\circ \)) appears rapidly (20–60 monolayers (ML)) indicating that some nice terrace periodicity is recovered. An AFM scan made after 180 s growth at 620°C and 2 Å/s gives a confirmation to this, although some deeper holes still remain. Note that the post-growth oxydation during the transfer of the sample limits the AFM lateral resolution (4 nm here instead of better than 1 nm on a Au surface) and kinks on step edges are not resolved. After a thicker buffer grown with weak As/Ga flux ratio (the surface is reconstructed 3×1) and a reduced growth rate (0.5 Å/s at 620°C), a peaked terrace distribution is obtained for terrace length \( L \) below 50 nm \( (L = a/\tan \alpha, a = 2.83 \text{ Å in GaAs}) \). On low misoriented substrates \( (\alpha = 0.2^\circ) \), the growth is indeed not in the step flow mode and the partial island nucleation on terrace disrupt the periodicity. No step bunching occurs in our growth conditions (see \( \alpha = 1^\circ \) data on Fig. 1), at variance with previous reports [5]. The terrace width

![AFM scan of a vicinal surface misoriented by an angle of 1° towards [1 1 0] (Ga step) after the growth of a GaAs buffer.](image1)

![Distribution of terrace width for five misorientation angles (nominal values: 2°, 1°, 0.5°, 0.5°, 0.3°) obtained from the analysis of an area comprising 4000–10 000 terraces. The two sets of data for 0.5° are within the accuracy of the misorientation. In the insert, variation of the standard deviation with mean terrace length for 8 samples.](image2)
distributions obey to a Gaussian law with a mean value close to the nominal terrace periodicity (misorientation accuracy ± 0.1°). A nearly linear relationship between the standard deviation and the mean value of the Gaussian (σ = 0.24 ± 0.04 ⟨L⟩) is obtained as shown in Fig. 2. There is no reduction in σ by including a superlattice GaAs/AlAs in the buffer. Our results are very similar to vicinal Si (1 1 1) where it was found σ = 0.25 ⟨L⟩ [6]. This close identity with Si is really surprising if we consider the conditions used in that case (a few hours at T = 900°C) which are far away from ours (T = 580°C and 2000 Å grown at 0.18 ML/s).

3. Equilibrium models

In Si, the results have been explained by thermodynamical equilibrium between interacting steps. In order to understand how this works, we have considered four equilibrium models based on: (i) straight noninteracting steps, (ii) single step meandering between fixed walls, (iii) terrace-step-kink (TSK) model taking into account only entropic repulsion, (iv) entropic and energetic step-step repulsion. Fig. 3 displays the terrace width distributions observed on Ga (and also on As terminated steps misoriented by 0.5°) and the distributions predicted by the different equilibrium models. In these models, the step configuration is only determined by the state of lower energy without any consideration of the kinetics of the incorporation at the step edges. For a sequence of straight non-interacting steps, an exponential distribution is expected. The distribution, obtained for a single step confined between two rigid walls separated by 2 ⟨L⟩ [7], already captures most of the physics of the entropic repulsion which drastically reduces the probability of finding small terraces. For purely entropic interactions, the distribution obeys the “universal law” calculated by Joos et al. in the one-dimensionnal free fermion model [8]. If entropic and energetic repulsion coexist, the distribution is narrow and obeys to a Gaussian law when energetic repulsion outweighs the entropic one. An energetic repulsion with a L−2 decay, as in the last model, is needed to explain the linear relationship between the standard deviation and the mean value of the Gaussian observed on Ga surfaces. For As terminated steps, note that the observed distribution is skewed (with a peak value slightly below the mean terrace width) and that the step edges are more disordered. Some terraces are as large as twice the nominal periodicity. But the probability of finding a terrace narrower than 10 nm remains small.

![Fig. 3. Fit of the data obtained on a 0.5° A (Ga step) sample using four equilibrium models discussed in the text: short dashes (model i), long dashes (model ii), dots (model iii), plain line (model iv). The insert shows the data for a 0.5° B (As step) sample.](image)
4. Role of the kinetics

In GaAs around 600°C, thermal annealing is not efficient to get a periodic step array and growth is unavoidable. Although this is partly due to the fact that the starting surface is very far from a smooth surface, we believe that the atom kinetics on the surface will be more important to determine the terrace distribution rather than the minimization of a free energy accounting only for the interactions discussed above. In our growth conditions, weakly bound mobile atoms are mainly provided by impinging atoms rather than by atoms thermally excited out of the surface.

In order to determine the critical parameters of the step array ordering, we have grown on 0.5° A misoriented samples a GaAs buffer in the conditions optimized for the terrace periodicity and then we have studied the change of the surface morphology as a function of specific growth conditions. Thus, a 60 nm thick GaAs layer growth at a much lower temperature (540°C) leads to the formation of mounds elongated in the [1 T 0] direction (4 ML’s high, 60 nm wide and 300 nm long). These results are in good agreement with those observed by Orme et al. [9] on nominal surfaces. In the [1 1 0] direction, the terrace width defined by these structures (~ 6 nm) is approximatively equal to the diffusion length at 540°C if we refer to the transition temperature, between island nucleation and step flow, measured by RHEED on 2° A (⟨L⟩ = 8 nm).

Growing a Ga1-xAlxAs (x = 0.33) barrier, the step periodicity remains quite good despite the lower lateral resolution of the AFM measurement (~ 6 nm) due to the stronger surface oxidation. On AlAs layer (900 Å at 0.25 Å/s), buried by 20 Å of GaAs to avoid the AlAs oxidation, the step array is more disordered in spite of a sharper specular spot splitting in RHEED. The surface morphology is explained by a partial incorporation at the step edge for pure AlAs growth, in agreement with the fact that RHEED oscillations can always be resolved during AlAs homoepitaxy. A surprising point is that we did not observe any mounds after the epitaxy of AlAs grown at low temperature (100 nm at 500°C). We are led to the conclusion that the Schwobel barrier evidenced on GaAs is much weaker on an AlAs surface. This agrees with the well-known fact that AlAs layers grown at low temperature are mirror like, but not with the impossibility to smooth out a rough surface by AlAs growth at higher temperature. We believe that then impurities (O, C) floating on the AlAs surface hamper the adatom diffusion.

Fig. 4. AFM scan of a GaAs:Si surface showing step bunching. The substrate is a 0.5° A surface.
To evidence the role of impurities in the building up of the terrace periodicity, we have grown Si doped GaAs layers at 580°C. After the growth of a 200 nm GaAs layer at low doping \( (2 \times 10^{17} \text{ cm}^{-3}) \) which is typically used in order to provide STM measurements on GaAs, we observe a more irregular step array \( (\sigma = 0.31 \langle L \rangle, \text{i.e.} \text{25–30\% larger}) \) and some elongated islands on the terraces. The poor terrace periodicity observed in ultra-vacuum STM measurements [5] can be explained by impurity step pinning. However no difference with pure GaAs growth can be evidenced on the RHEED oscillations (the transition temperature to step flow is just the same) but this can be fortuitous and misleading because of these islands. For very high Si concentration \( (2 \times 10^{19} \text{ cm}^{-3}) \), irregular macrosteps (up to 30 ML’s high for 60 nm thick layers) which might be unresolved step bunching are observed (Fig. 4). The surface morphology is quite different from the mounds. We have not yet a clear understanding of how to explain this. At this high doping level the surface As coverage is lower [10] and a change of the surface reconstruction from 3 \( \times \) 1 to a 3 \( \times \) 2 is observed.

5. 2D Monte Carlo simulation

We developed a Monte Carlo simulation to mimic the MBE growth and to understand the results observed on pure GaAs. Our model is very close to that developed by Vvedensky et al. [11]. Only 5 parameters are introduced: the binding energy to the substrate \( (E_b = 1.58 \text{ eV}) \), the binding energy to in-plane nearest neighbour \( (E_n^{(A)} \sim 0.16 \text{ eV} \text{ along } [1 \bar{1} 0]) \) and \( E_n^{(B)} \sim 0.32 \text{ eV} \text{ along } [1 \bar{1} 0] \) and a Schwoebel barrier \( (E_b^{(A)} \text{ and } E_b^{(B)}) \). \( E_b \) is determined by the temperature required for step flow growth mode whereas the anisotropy ratio \( E_n^{(A)}/E_n^{(B)} \) is determined by the anisotropy of islands nucleating on the terraces. The values of the parameters are those taken in Ref. [12] but we have then to increase the temperature of the calculation by 50°C. We did not use additional rules such as the instantaneous search by impinging atoms of the most favourable site in a given area. This rule destroys at once any roughness at low temperature which does not correspond to experimental observation. Our calculations are done on a 100 \( \times \) 100 array of “GaAs site” \((400 \text{ Å} \times 400 \text{ Å}) \) with periodic boundary conditions. We investigated the smoothing of initial rough surfaces and the step array ordering on 2° misoriented substrates.

We are able to account for all the observations using an anisotropic Schwoebel barrier: (i) the formation of the mounds observed at low growth temperature requires this barrier [9], (ii) it is impossible to smooth out a rough surface if this barrier is strong in both directions [13], (accepting this, the barrier anisotropy is consistent with the mound anisotropy) and (iii) as will be discussed next, the terrace periodicity is not good enough without a Schwoebel barrier. We do not claim that we are able to distinguish between an anisotropic Schwoebel barrier and an anisotropic diffusion: either the barrier is weaker or the hopping rate is larger in the [1 \( \bar{1} \) 0] direction. The published results on the hopping rate anisotropy are somehow contradictory [14, 12]. So far we have decided to ignore it. In order to minimize the number of parameters we took \( E_b^{(B)} = 0 \). With \( E_b^{(A)} = 0.175 \text{ eV} \), it is possible to smooth a 10 ML’s deep hole in a 20 ML’s at 500°C and 0.1 ML/s and to develop mounds at 500°C. Simulation at low temperature (500–550°C) without any Schwoebel barrier followed by 60 s relaxation showed a surface morphology similar to our AFM study of AlAs growth (to simplify the analysis the other parameters remained unchanged).

To study the terrace ordering, our array comprised 5 steps which are bunched together (4 nm apart each other) and separated by a large terrace (24 nm wide) as initial conditions. The error function \( f(t) = 1/N \sqrt{\sum (L_i - \langle L \rangle)^2} \), where \( N \) is the number of steps and \( L_i \) the mean distance between two neighbouring steps, gives us a quick quantitative estimation of the ordering \( (f(t) = 0 \) for a perfectly ordered surface). \( f(0) \) is close to 1 in our initial conditions and \( f(t) \) should become smaller than \( \sigma/\langle L \rangle = 0.24 \) to reproduce AFM measurements. Without any Schwoebel barrier this is not possible. With an isotropic barrier, if some island nucleation occurs on the large terrace then the roughness develops further and \( f(t) \) increases. But with an
Fig. 5. Time evolution of the error function $f(t)$ (used to have a quick estimation of the terrace ordering) as a function of the number of monolayers for two cases: no Schwoebel barrier and anisotropic Schwoebel barrier. The initial and a typical final surfaces are also shown.

Anisotropic barrier, $f(t)$ decreases below the required value in less than 60 ML's as shown in Fig. 5. The time scale is then also correct. The oscillations of $f(t)$ could be reduced if the step interactions were taken in account.

6. Conclusions

Studying by AFM the terrace distribution of misoriented substrates after MBE growth of undoped or doped GaAs and undoped AlAs, we report several new results. Step bunching on GaAs vicinal surfaces can be avoided in the usual growth conditions but can be induced at high Si doping. In optimized conditions the terrace distribution of pure GaAs is sharply peaked around the mean value for a large range of misorientation angle. The results can be well explained quantitatively by equilibrium and dynamical models (Monte Carlo simulation). Lower migration length (AlAs or moderate Si doping) increases the terrace disorder. We give also some evidence that the Schwoebel barrier at step edges is strongly anisotropic in GaAs and negligible in AlAs.

References

Photoluminescence investigation of p-type Si-doped AlGaAs
grown by molecular beam epitaxy on (1 1 1)A, (2 1 1)A and
(3 1 1)A GaAs surfaces

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Abstract

A systematic study of Si-doped AlGaAs grown on (1 1 1)A, (2 1 1)A, and (3 1 1)A GaAs surfaces by molecular beam epitaxy is reported. Both electrical and photoluminescence measurements have been carried out to investigate the physics of the Si incorporation. The optical identification of the incorporated impurities together with the determination of free carrier concentration and mobility allowed us to suggest a simple kinetic model for the Si incorporation in AlGaAs grown on substrates with different orientations.

1. Introduction

The possibility of exploiting in device manufacturing the amphoteric character of the Si impurity in (Ga,Al)As by simply using different planes has recently attracted great interest. It has been demonstrated that under identical growth conditions Si incorporates preferentially as a donor for growth on \((N \, 1 \, 1)B\) and as an acceptor on \((N \, 1 \, 1)A\) surfaces \((N = 1, 2, 3)\) [1, 2]. However, Si is mainly a donor when the growth is on the conventional \((1 \, 0 \, 0)\) plane.

This has opened the possibility to realise lateral p-n junction and field-effect transistors [3] on patterned substrates with the use of a single dopant species. Moreover, the study of the molecular beam epitaxy (MBE) growth on substrates with different orientations could provide useful information on the epitaxial growth mechanism and on the physics of dopant incorporation.

Here, we will report on a systematic study of the optical properties of AlGaAs samples grown on \((1 \, 1 \, 1)A\), \((2 \, 1 \, 1)A\) and \((3 \, 1 \, 1)A\) surfaces with different Si doping levels.

2. Experimental details and samples

The epitaxial layers have been grown on GaAs substrates oriented to within \(\pm 0.5^\circ\) for the \((2 \, 1 \, 1)A\) and \((3 \, 1 \, 1)A\) planes and \(3^\circ\) off towards \((1 \, 0 \, 0)\) for the \((1 \, 1 \, 1)A\) surface using a Varian Gen-II MBE system. The growth temperature and rate were 630°C and 1.5 monolayer/s, respectively. The As/(Ga,Al)
beam-equivalent pressure ratio determined by an ionisation gauge was ~12. After growth the epitaxial surfaces were examined using a Nomarski phase-contrast optical microscope and were found to be mirror smooth and nearly defect free, except for the (1 1 1)A surface which had a milky finish. This bad morphology is expected for the growth on the (1 1 1)A orientation under the above growth conditions. In this study the samples were grown simultaneously adopting the growth parameters which gave optimal electrical properties of (3 1 1)A and (1 0 0) heterostructures [4]. For each substrate orientation a set of samples with four different silicon dopings has been grown: the silicon flux was set in order to give intended free carrier concentrations, \( N_i \). of \( 1.0 \times 10^{15} \), \( 2.0 \times 10^{16} \), \( 1.3 \times 10^{17} \), and \( 1.3 \times 10^{18} \) cm\(^{-3} \) in (1 0 0) oriented AlGaAs. These were characterised at 77 K and 300 K using Hall measurements (see Table 1 for details).

The photoluminescence (PL) was excited by the 488 nm line of an Ar\(^+\) laser with an excitation power density \( (J_{\text{exc}}) \) in the \( 2 \times 10^{-5} \) W/cm\(^2\) to 20 W/cm\(^2\) range. The temperature \( (T) \) of the samples has been varied between 12 K and 300 K using a cryocooler. The luminescence was analyzed by a single grating monochromator and detected by a photomultiplier tube with a cooled GaAs photocathode followed by a lock-in amplifier.

The sample characteristics and the electrical measurements results are summarized in Table 1. All the samples show a p-type conductivity and the room temperature Hall mobility \( \mu_H \) does not show a significant dependence on the substrate orientation [5]. In Ref. [5] we reported that carbon was present in high concentration in (2 1 1)A and (3 1 1)A samples and not in (1 1 1)A surfaces.

3. Photoluminescence spectra

The energy gap \( (E_g) \) and hence the Al mole fraction \( x \) (see Table 1) were determined by PL measurements at different values of \( T \) and \( J_{\text{exc}} \). By comparing the spectral position and the evolution with \( J_{\text{exc}} \) and \( T \) of the various PL bands with the literature [6], we identified the responsible impurities. More data and details are reported in Ref. [7].

In Fig. 1 the spectra at \( T = 12 \) K of the (3 1 1)A samples are shown. The spectrum of sample e1 \( (N_i = 1.0 \times 10^{15} \) cm\(^{-3} \) is characterized by several overlapping transitions broadened by the alloy

<table>
<thead>
<tr>
<th>Sample</th>
<th>Surface orientation</th>
<th>( d ) (μm)</th>
<th>( N_i ) (cm(^{-3} ))</th>
<th>( p_H ) (cm(^{-3} ))</th>
<th>( \mu_H ) (cm(^2)/V·s)</th>
<th>( x )</th>
<th>( E_g ) (eV)</th>
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</thead>
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<td>insulating</td>
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<td>1136</td>
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disorder: the bound exciton (BE) recombination at 1.919 eV, the free electron to bound hole recombination \( N_{\text{e}} \) at 1.907 eV and the donor-acceptor pair transition \( N_{\text{d}} \) at 1.894 eV involving the C acceptor. The same transitions involving Si, namely \( N_{\text{d}} \) (e, Si) and \( N_{\text{d}} \) (D, Si), are peaked at 1.876 and 1.854 eV, respectively. The spectrum of sample e2 \( (N_{\text{e}} = 2.0 \times 10^{16} \text{ cm}^{-3}) \) consists of a wide band from which at least two structures can be resolved by varying \( T \) or \( J_{\text{exc}} \): the (e,C) and (e, Si) transitions. In the spectrum of sample e3 \( (N_{\text{e}} = 1.3 \times 10^{17} \text{ cm}^{-3}) \) we only observe the (e, Si) recombination. Finally, the band in the spectrum of sample e4 \( (N_{\text{e}} = 1.3 \times 10^{18} \text{ cm}^{-3}) \) is peaked at the (D, Si) transition. In all the observed donor to acceptor transitions the donor is likely to be Si on the Ga (or Al) site.

The low temperature spectra of the (2 1 1)A samples are shown in Fig. 2. The spectra of d1 \( (N_{\text{e}} = 1.0 \times 10^{15} \text{ cm}^{-3}) \) and d2 \( (N_{\text{e}} = 2.0 \times 10^{16} \text{ cm}^{-3}) \) consist of a single narrow band, peaked at an energy lower than that of the BE, which shifts to higher energies by increasing \( J_{\text{exc}} \). Thus the narrow peak in the spectra of samples d1 and d2 could be attributed to the overlap of the BE and the defect bound excitons (d, X) recombinations. The variation of the relative intensity of the two transitions with \( J_{\text{exc}} \) may be responsible for the energy shift. The (d, X) transitions have been tentatively associated with complexes consisting of impurities and intrinsic defects, such as vacancies and interstitials [8]. In the spectrum of sample d3 \( (N_{\text{e}} = 1.3 \times 10^{17} \text{ cm}^{-3}) \) the main peak has the same attribution as that of the samples d1 and d2 and the presence of Si acceptors is revealed by a weak (D, Si) emission band at 1.866 eV. Decreasing \( J_{\text{exc}} \) the main peak shifts to 1.899 eV corresponding to the (e, C) recombination and the (D, Si) band intensity rises. The spectrum of sample d4 \( (N_{\text{e}} = 1.3 \times 10^{18} \text{ cm}^{-3}) \) is characterized by an asymmetric band with a long low energy tail. Increasing \( T \) the peak energy follows the thermal energy gap shift as given by the Varshni equation [9]. This suggests that free hole with free electron recombinations (BB) are responsible for this emission. Moreover, some weak low energy structures which could be attributed to (d, X) and C acceptor related transitions, can be resolved with increasing \( T \).

The PL spectra of the (1 1 1)A samples shown in Fig. 3 consist either of a single peak at an energy much lower than that of the C or Si related emissions (samples c1 and c3) or of a wide band (samples c2 and c4). We argue that these PL spectra are attributed to the low sample quality due to the growth conditions used, which are not the optimal for the (1 1 1)A oriented substrates [2].

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**Fig. 1.** Photoluminescence spectra of the (3 1 1)A samples at different doping levels measured at 12 K with an excitation power density of 20 W/cm². For each sample the intended doping \( N_{\text{e}} \) (in cm⁻³) is given. The relative sensitivity factor is given on the left and the energy gap \( E_g \) values are indicated by the hollow arrows. The numbers on the spectra stand for the following processes: (1) bound exciton recombination; (2) free electron to carbon acceptor transition; (4) donor to carbon acceptor pair transition; (5) free electron to silicon acceptor transition; (6) donor to silicon acceptor pair transition.
Fig. 2. Photoluminescence spectra of the (211)A samples at different doping levels measured at 12 K with an excitation power density of 20 W/cm². For each sample the intended doping \( N_i \) (in cm⁻³) is given. The relative sensitivity factor is given on the left and the energy gap \( E_g \) values are indicated by the hollow arrows. The numbers on the spectra stand for the following processes: (2) superposition of bound exciton and defect bound exciton recombinations; (6) donor to silicon acceptor pair transition.

Fig. 3. Photoluminescence spectra of the (111)A samples at different doping levels measured at 12 K with an excitation power density of 20 W/cm². For each sample the intended doping \( N_i \) (in cm⁻³) is given. The relative sensitivity factor is given on the left and the energy gap \( E_g \) values are indicated by the hollow arrows.

4. Discussion

Since the samples were grown simultaneously, it is not possible to make a direct comparison between our PL data and those published in the literature. The latter in fact generally refer to samples grown using conditions optimized for a given substrate orientation.

We found that the morphological and optical quality continuously improve by changing the substrate orientation from (1 1 1)A, to (2 1 1)A and to (3 1 1)A. However, in spite of the differences in the PL lineshapes at \( T = 12 \) K, the luminescence efficiency, taken as the integral of the luminescence spectrum at \( T = 300 \) K, is comparable for all the samples and increases with increasing \( p_{\text{H}} \). The nearly equal luminescence intensities are reflected in the mobility values of all the samples which compare reasonably well with each other. On the other hand, the low temperature PL spectra are rather different. This is not surprising. In fact, small
concentrations of efficient radiative channels can dominate the optical properties without affecting the electrical ones.

A clear dependence of the acceptor incorporation in the samples on the substrate orientation has been found. The (1 1 1)A samples show a relevant Si incorporation on As sites without evidence of other unwanted acceptors. In fact, the free carrier concentration increases markedly with increasing \( N_r \). These conclusions cannot be supported by the PL data because the spectra are dominated by unresolved bands and deep transitions.

The acceptor incorporation in (3 1 1)A samples is different. Firstly, the electrical measurements of the samples with low \( N_r \) reveal the presence of a residual acceptor impurity. From PL this impurity can be identified as carbon. Secondly, the PL spectra show that the (D, Si) transitions become more and more evident by increasing \( N_r \) revealing the presence of Si self-compensation.

The electrical data of the (2 1 1)A samples have a similar trend as those of the (3 1 1)A ones but the PL spectra show a more relevant defectiveness of the former as revealed by the presence of the (d, X) recombination.

The phenomenology of the incorporation of silicon on III or V sites in AlGaAs has been described in a previous work [2]. However the physics of such incorporation is still a matter of debate. To our knowledge, no detailed theoretical work is present in the literature. Only a phenomenological model, which has been applied in different cases [3, 10], has been developed to explain the amphoteric behaviour of group IV impurities in GaAs grown on high index surfaces [11, 12]. This model does not take into account the surface reconstruction. However, it works quite well and we use it for the discussion of our data. The main point of the model is that, among the different \( (N 1 1)A \) surfaces, the efficiency of the Si incorporation on the As site varies due to the specific bonding geometry of each of the surfaces, in particular to the different single to double dangling bonds ratio. For surfaces with a higher density of double dangling bonds, As predominates in the competition with the Si for the group V site occupation. For instance, the (1 1 1)A plane presents on the Al/Ga rich surface only single dangling bonds for the As incorporation. Hence a good incorporation efficiency of Si in As site is expected. Also a lower quality of the grown epilayers follows [13].

In terms of single to double dangling bonds ratio, the (1 0 0) and the (1 1 1)A surfaces present the two extreme situations (Fig. 4). Going from the (1 1 1)A to the (1 0 0) surface one passes through the (2 1 1)A and the (3 1 1)A ones, which have a periodic step structure with As step edge atoms and Al/Ga terrace atoms. This structure can be viewed as a sequence of (1 0 0)-like surfaces with double dangling bonds and (1 1 1)A-like surfaces with single dangling bonds. Let us consider, for example, the growth on the (2 1 1)A plane (Fig. 4). A newly chemisorbed Al/Ga atom in position (2) together with the pre-existing Al/Ga atom in position (1) makes two dangling bonds available for the incorporation of an As atom in position (3). In proximity of these sites the As\(_2\) molecule easily decomposes [11] and the As incorporation is favoured with respect to the Si one.

The observed differences in the Si incorporation between the (2 1 1)A and the (3 1 1)A grown samples are explained by the extent of the (1 1 1)A-like terraces which increases from the (3 1 1)A

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**Fig. 4.** Atomic arrangement projected on the (1 1 0) plane of the ideal (1 0 0), (1 1 1)A, (2 1 1)A and (3 1 1)A surfaces for the zinc blende crystal structure showing the single and double dangling bonds sites. Full dots and open dots are the group V atoms and the group III atoms, respectively. During the growth, a newly chemisorbed Al/Ga atom in position (2) together with the pre-existing Al/Ga atom in position (1) makes two dangling bonds available for the incorporation of an As atom in position (3).
surface to the (2 1 1)A one. Hence, passing from (1 1 1)A to (2 1 1)A and finally to (3 1 1)A surfaces the Si incorporation in As sites is less and less favoured by the increasing As competition and the Al/Ga sites occupation by Si is more and more probable.

It is tempting to apply this simple model also to the incorporation of C. It is well known that C incorporates quite easily in the MBE growth of (1 0 0) layers while we have found that in the (1 1 1)A layers it does not. Thus it is expected that the C incorporation is favoured in presence of (1 0 0)-like steps. This is confirmed by the PL data giving a clear evidence of C in the (3 1 1)A samples while no strong evidence of C presence is found in the spectra of the (2 1 1)A samples.

5. Conclusions

We have observed that the (1 1 1)A surfaces incorporate less carbon than the (2 1 1)A and the (3 1 1)A ones, which need a higher silicon flux before the presence of Si clearly manifests itself in the electrical and optical measurements. We have found a lower optical and morphological quality in the (1 1 1)A samples; however this does not affect markedly the values of the Hall mobility, free carrier concentration and PL efficiency.

Our results can be explained using a simple model in which the single to double dangling bond ratio characterizing the different planes mainly determines the impurities incorporation.

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Patterned growth on GaAs (3 1 1)A substrates

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Abstract

We report on a new type in the selectivity of growth found during MBE of (AlGa)As on patterned GaAs (3 1 1)A substrates. For mesa stripes oriented along the [0 1 − 1] direction a smooth convex curved surface profile develops on one side enclosing thicker regions of GaAs along the sidewall between thinner regions on the mesa top and bottom. Cross-sectional TEM directly images the evolution of this unique profile to rely on the preferential migration of Ga atoms from both sides toward the sidewall (opposite to the case of patterned GaAs (1 0 0) or (1 1 1) substrates), via size reduction of a fast growing side facet close to the next (1 0 0) plane. On the opposite side, a rough slow growing (1 1 1)A side facet is established. Mesa stripes oriented along [−2 3 3] exhibit two slow growing side facets without roughening of the growth front. The surface morphology of the different sidewalls is related to the anisotropy of the surface diffusion length of Ga adatoms along [−2 3 3] and [0 1 − 1] determined by AFM line-scans from the variation of the GaAs layer thickness close to the edges of shallow square shaped mesas. As confirmed by TEM and CL spectroscopy, the uniformity and the selectivity of the growth mode across the fast growing sidewall is not reduced for step heights in the nanometer range allowing the formation of quasi-planar sidewall quantum-wire arrays.

1. Introduction

We study the selectivity of growth during molecular beam epitaxy (MBE) of (AlGa)As on patterned GaAs (3 1 1)A substrates. As expected from the unique growth modes in MBE and metal-organic vapor phase epitaxy (MOVPE) on planar GaAs(3 1 1)A & B substrates to naturally produce ordered arrays of nanometer-scale quantum structures [1, 2], the selectivity of growth on patterned GaAs (3 1 1)A substrates is qualitatively different from that on patterned low-index GaAs (1 0 0) and (1 1 1) substrates [3, 4]. This offers a new flexibility for the design of devices and quantum structures. During MBE of (AlGa)As on patterned GaAs (3 1 1)A substrates we have found the formation of a fast growing sidewall on one side of mesa stripes oriented along the [0 1 − 1] direction [5]. Preferential migration of Ga adatoms from the mesa top and bottom toward the sidewall develops a smooth convex curved surface profile without faceting. This unique growth mode that does not occur for the perpendicular stripe orientation nor on other
patterned GaAs (1 1 1)A&B substrates [6] remains stable for step heights down to the quantum-size regime to produce lateral quasi-planar quantum wires on patterned GaAs (3 1 1)A substrates. The wires exhibit high structural perfection and uniformity, narrow emission lines in photoluminescence (PL) spectroscopy and high PL efficiency up to room temperature [7]. Quantum confinement of excitons in the wires has been demonstrated by the changeover from one-dimensional confinement to magnetic confinement with increasing magnetic field [6]. The PL energy position of the wires critically depends on the initial step height providing an additional parameter to control the size of the wires that has no correspondence in V-groove- or ridge-type structures. Finally, it has been shown that, for a given step height, the wires can be stacked in growth direction without any increase in interface roughness or wire size fluctuations indicating a self-limiting lateral growth mechanism [6].

In this paper we present an atomic force microscopy (AFM) study of the surface profiles across the edges of mesa stripes along [0 1 −1] and [−2 3 3] on patterned GaAs (3 1 1)A substrates after overgrowth. The surface morphology of the convex curved fast growing sidewall is as smooth as that of the flat parts of the mesa. On the other hand, the slow growing (1 1 1) side facet on the opposite side shows pronounced roughening of the growth front. The evolution of the fast growing sidewall is investigated by cross-sectional transmission electron microscopy (TEM). The convex curved surface profile is found to develop via size reduction of a fast growing side facet close to the next (1 0 0) plane. For the perpendicular mesa stripes along [−2 3 3] two symmetrically arranged slow growing side facets close to {3 3 1} planes develop on both sides. Besides macroscopic defects on one side no roughening of the growth front occurs. The surface diffusion lengths of Ga adatoms along the two perpendicular [−2 3 3] and [0 1 −1] azimuths are deduced from the variation of the GaAs layer thickness close to the edges of shallow square shaped mesas. The anisotropy of the diffusion

Fig. 1. AFM images of the edges of the 400 nm high mesa stripes after overgrowth. (a) Fast growing sidewall and (b) slow growing sidewall along [0 1 −1]. (c), (d) AFM images of both sidewalls along [−2 3 3].
length is brought in connection with the surface morphology of the slow and fast growing sidewalls of the mesa stripes. For step heights in the quantum-size regime an array of sidewall quantum wires with 4 μm pitch is fabricated and characterized by cathodoluminescence (CL) spectroscopy.

2. Sample preparation and experimental setup

The GaAs (311)A substrates were patterned by optical lithography and wet chemical etching using the H₂SO₄ : H₂O₂ : H₂O (1 : 8 : 40) preferential chemical etching solution. The mesa stripes were oriented along the two perpendicular [0 1 −1] and [−2 3 3] directions. After cleaning in concentrated H₂SO₄ and rinsing in deionized water the samples were introduced into the MBE growth chamber. The GaAs/Al₀.₅Ga₀.₅As multilayer structures (see below) were grown at 620°C while the substrates were rotated with 6 rpm. The growth rates for GaAs and Al₀.₅Ga₀.₅As were 0.5 and 1 μm/h, and the group-V-to-III-flux ratio was about 5. The structural and optical properties of the samples were investigated by AFM, TEM, and CL spectroscopy.

3. Results and discussion

Fig. 1a–Fig. 1d show three-dimensional AFM images of the sidewalls of 80 μm wide and 400 nm high mesa stripes along [0 1 −1] Fig. 1a, Fig. 1b, and along [−2 3 3] Fig. 1c, Fig. 1d after overgrowth. The layer sequence comprises 5 periods 100 nm Al₀.₅Ga₀.₅As/100 nm GaAs. In the center of the fourth (AlGa)As layer a 8 nm thick GaAs layer is inserted. The AFM image in Fig. 1a shows the surface profile of the convex curved fast growing sidewall of the mesa stripe along [0 1 −1] that develops in the sector toward the next (1 0 0) plane. As confirmed by scanning electron microscopy (SEM) [5] the surface is not roughened compared to the flat parts of the mesa. The surface profile in

![Figure 2](image)

Fig. 2. (a) Cross-sectional TEM image viewed along [0 1 −1] of the fast growing sidewall of the mesa stripe along [0 1 −1]. In (b) the scale is enhanced.
Fig. 1b of the opposite side in the sector toward the next (111) plane shows the evolution of a rough slow growing (111) side facet with the characteristic concave curved bottom corner. Smooth slow growing side facets are formed at the sidewalls of the mesa stripes along [−233] (Fig. 1c and Fig. 1d).

The evolution of the fast growing sidewall is investigated by cross-sectional TEM shown in Fig. 2a and Fig. 2b. At the sidewall, first a fast growing side facet close to the next (100) plane is developed which reduces in size during growth of GaAs to end up with the convex curved surface profile with a stable bottom step corner. The adjacent areas in the top and bottom areas of the mesa exhibit a reduced layer thickness due to migration of Ga adatoms toward the sidewall as has been confirmed by spatially resolved CL from the quantum-well layer [5]. For (AlGa)As almost no variation of the layer thickness and contrast across the step is observed due to the small surface diffusion length of Al suggesting an almost uniform Al composition.

Lateral growth can be obtained also in MOVPE and metalorganic MBE, in particular, on patterned GaAs (111)A&B substrates [8, 9] due to the dependence of the chemical reactivity of the precursors on different crystallographic planes and growth conditions. The shape of the side facets, however, that is determined by the migration of adatoms, has a concave curved surface profile at the bottom corner of the mesas similar to that of slow growing side facets. Hence, also in these cases the surface diffusion of adatoms is away from the side facets highlighting the unique growth mode on patterned GaAs (311)A substrates.

Shallow mesas, 30 nm high, have been prepared to determine the surface diffusion lengths $L_{[−233]}$ and $L_{[−133]}$ of Ga adatoms along the perpendicular [−233] and [01−1] directions. As the mesa height is more than one order of magnitude smaller than usual surface diffusion

![AFM LINESCAN](image)

Fig. 3. AFM linescans along [−233] and [01−1] across shallow square shaped mesas after overgrowth. The arrows mark the diffusion lengths determined from the exponential variation of the layer thickness close to the edges of the mesa. The inset shows the AFM top view of the mesa.
lengths, the shape of the sidewalls is essentially determined by the diffusion length that can then be measured from the exponential variation of the layer thickness, i.e., the growth rate close to the edges of the mesa in analogy to the experiments using $\mu$-RHEED [10, 11]. Fig. 3 shows the AFM linescans along $[-2\bar{3} 3]$ and $[01\bar{1}]$ across the mesa depicted in the inset. The thickness of the GaAs layer grown over the mesa is 70 nm. From the exponential variation of the layer thickness in the linescan along $[-2\bar{3} 3]$ close to the bottom corner of the convex curved surface profile acting as a sink for Ga atoms, $L_{[-2\bar{3} 3]}$ amounts to 1–1.5 $\mu$m. The linescan along $[01\bar{1}]$ close to the top of the mesa acting as a source of Ga atoms gives $L_{[01\bar{1}]}$ to be 0.3–0.5 $\mu$m. The anisotropy of the diffusion length suggests a relationship with the surface morphology of the sidewalls. Smooth surface profiles are obtained for either a slow growing side facet together with the larger diffusion length parallel to the step or a fast growing sidewall with the larger diffusion length perpendicular to the step. A slow growing sidewall with the larger diffusion length perpendicular to the step produces a rough surface morphology. This finding is essential for choosing the optimum pattern alignment on arbitrarily oriented substrates. The anisotropy of the diffusion length, however, is comparable to that on (100) surfaces [10, 11]. Hence, on patterned GaAs (311)A substrates the highly anisotropic microscopic surface corrugation along $[-2\bar{3} 3]$ [1] is assumed to stabilize the respective growth fronts additionally.

Fig. 4. (a) Schematic of the growth mode on shallow mesa stripes along $[01\bar{1}]$ to form lateral quantum wires at the fast growing sidewall. (b) Filtered CL intensity mapping detected at the wire peak position at 1.540 eV of the array of sidewall quantum wires with 4 $\mu$m pitch.

wire around 60 meV has been estimated from the red shift of the PL from the wire compared to that of the adjacent wells. Fig. 4a presents a schematic of the quantum wire formation deduced from TEM. The high uniformity of the wire array over a large area is demonstrated by the filtered CL intensity mapping detected at the wire peak position at 1.540 eV shown in Fig. 4b.

4. Conclusion

We have investigated the MBE growth of (AlGa)As on patterned GaAs (311)A substrates. On one side of mesa stripes along $[01\bar{1}]$ a fast growing sidewall with a smooth convex curved
surface profile is developed. On the opposite side a slow growing (1 1 1) side facet with pronounced roughness is formed. Smooth slow growing surface profiles are formed at the sidewalls of the mesa stripes along [−2 3 3]. Cross-sectional TEM has shown the convex curved surface profile to evolve during size reduction of a fast growing side facet close to the next (1 0 0) plane due to preferential migration of Ga adatoms from both sides toward the sidewall. The surface diffusion lengths of Ga adatoms along [−2 3 3] and [0 1 −1] have been determined from AFM linescans and related to the surface morphology of the sidewalls. For step heights in the quantum-size regime an array with 4 μm pitch of lateral quantum wires along the fast growing sidewalls has been fabricated that exhibits high uniformity over a large surface area.

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References

Strain-balanced AlGaInAs/InP heterostructures with up to 50 QWs by MBE

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Abstract

Using solid-source MBE, strain-balanced quaternary AlGaInAs/AlGaInAs multiple quantum well (MQW) structures have been grown and studied by photoluminescence (PL) and X-ray diffraction (XRD) measurements, showing high epitaxial quality and excellent homogeneity in growth direction. The PL linewidths are low and almost independent of the number of QWs \( (1 \leq N_{QW} \leq 50) \). With respect to photonic device applications at 1.55 \( \mu \)m, structures with quaternary QWs seem to be superior compared to structures with ternary QWs: structures with quaternary QWs enable more degrees of freedom in device design and reveal a five times larger lateral wafer area in which the PL wavelength is constant.

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From the epitaxial point of view and for applications in advanced optoelectronic and electronic devices, strain-balanced QWs are very attractive. For the Bragg reflectors of surface emitting lasers, e.g., high-quality multiple heterostructures with a large refractive index contrast and a large number of periods are required. Strain-balanced structures allow numerous periods, although highly strained layers are involved [1–5]. Furthermore, for the active layers of lasers, multiple QW structures are desirable, allowing as many degrees of freedom as possible in selecting compositions, well widths and number of wells. E.g., in order to obtain ultra-high modulation bandwidths, a sufficiently high number of wells is required in order to guarantee high differential gain. In recent years, QW structures including quaternary material in both wells and barriers have been grown and studied several times using the rather well known GaInAsP/GaInAsP/InP material system (see, for example, Refs. [4–9]). In contrast, AlGaInAs/AlGaInAs/InP heterostructures have been grown, investigated and applied only occasionally [10–14].

Using a Modular GEN II solid source MBE, we have grown AlGaInAs/AlGaInAs MQW structures on semi-insulating, n- and p-doped (1 0 0)-oriented InP- substrates. We used two In-furnaces and one furnace, respectively, for the Al, Ga and As source.

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materials at 510°C substrate temperature. In the structures studied here, no growth interruptions were applied at the QW heterostructure interfaces. During the growth of the MQW structure the Al-flux and the Ga-flux are kept constant. The variation of the well and barrier compositions is realized by changing the In-fluxes (switching between the In-I and In-II furnace) at the interfaces, respectively. Combining XRD studies, low temperature PL measurements and corresponding simulations, respectively, we determined the well widths, barrier widths and compositions of all layers involved, using also results from wafers with isolated bulk layers grown under nominally identical conditions. To obtain strain-balanced structures, this procedure was used to adjust the geometric and compositional parameters such as the In-content in the wells and barriers (lattice mismatch) as well as the Al-content in the wells and barriers. The number of QWs is defined by the number of reproductions of a unit cell in growth direction. The unit cell consists of an Al0.093Ga0.197In0.71As well (width \( L_w \)) embedded by two Al0.208Ga0.445In0.347As barriers of thickness \( L_b/2 \). This enables a strain-balancing of the compressively strained well (\( \Delta a/a \sim +1.2\% \)) by two symmetrically positioned tensile strained barrier parts (\( \Delta a/a \sim -1.4\% \)) of widths \( L_b/2 \). A strain-compensation can also be obtained where the product of barrier lattice mismatch and the barrier width \( L_b \) is about the well lattice mismatch times \( L_w \). Finally, there are lattice-matched Al0.227Ga0.243In0.53As layers (\( L_{pl} \sim 1.21 \mu m \)) cladding the two bordering barrier layers asymmetrically on both sides. The inset of Fig. 3 schematically displays the variation of the conduction band edge of our samples. In Ref. [12] we have studied several single QW samples having the same compositions and identical ratio of \( L_w/L_b \), but different well widths. These samples show a variation of the PL linewidths as a function of well width which can be attributed to the changing influence of interface roughness and alloy disorder broadening mechanisms.

To investigate the stability of the strain-balanced MQW structures, samples containing different numbers of wells (1 \( \leq N_{QW} \leq 50 \)) were grown. Fig. 1 depicts a scanning electron microscope (SEM) image of a cleaved strain-balanced MQW structure, containing 50 compressively strained quaternary wells (light stripes), 49 tensile strained barriers of width \( L_b \) (dark stripes) and 2 barriers of the same composition but half the width \( L_b/2 \) (thin dark stripes, first and last of the stack). The geometrical parameters (\( L_w, L_b \)) and compositional parameters (In-content, Al-content, ...) are intended for a laser structure emitting at 1.55 \( \mu m \).

The corresponding PL spectrum is displayed in Fig. 2. We measured a PL linewidth of 9.3 meV at 10 K which can quantitatively be attributed to broadening mechanisms involving interface.

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**Fig. 1.** SEM micrograph of a cleaved strain-balanced heterostructure including 50 QWs.
roughness and alloy disorder. This low value indicates that the variations in both compositional and geometrical parameters from well to well are very low. The emission at room temperature is thermally broadened and shifted to longer wavelengths according to the temperature dependence of the fundamental band gaps of the individual layers.

Fig. 3 depicts the experimental PL linewidth at 10 K as a function of \(N_{\text{QW}}\). The layer widths of these structures are \(L_x = 7.2 \text{ nm}\) and \(L_B = 5.8 \text{ nm}\) (for \(N_{\text{QW}} = 1, 10, 15, 20\)) as well as \(L_x = 8.2 \text{ nm}\) and \(L_B = 4.8 \text{ nm}\) (for \(N_{\text{QW}} = 30\) and 50). We observe no dependence of the PL linewidth on the number of QWs for \(N_{\text{QW}} \leq 30\), except for statistical fluctuations. It is interesting to observe that the lowest linewidth is measured for \(N_{\text{QW}} = 50\), which might originate from a smaller influence of interface roughnesses, since the well width is slightly higher than in the structures with \(1 \leq N_{\text{QW}} \leq 20\). However, we suppose another physical reason for this low PL linewidth as explained below. For the barrier widths considered here tunneling processes through the potential barriers play a minor role. However, reducing the barrier width, the two-dimensional character of the carriers slightly weakens due to increasing coulomb interaction towards a growing three-dimensional character [15]. Therefore, the sensitivity of the wave functions to interface roughnesses is decreased, causing a slight reduction of the interface roughness induced PL linewidth broadening.

Fig. 4 depicts a measured XRD rocking curve of a structure containing 50 QWs. The corresponding simulated spectrum, which is obtained from dynamical linewidth theory, is shown below and is in good agreement. This indicates that the geometrical and compositional parameters as well as the material properties give a good description of the system. Fig. 5a displays a blow-up of Fig. 4 indicating the \(50 - 2 = 48\) side-maxima corresponding to 50 wells. Note that the total strained stack has a very large width of 0.65 \(\mu\text{m}\). A blow-up of an XRD spectrum measured for a sample including 30 wells is shown in Fig. 5b. Also in this case the number of wells (30) can be identified by the number of side-maxima. Between the 1st and the 2nd satellite reflection we observe \(30 - 2 = 28\) side-maxima. In Fig. 5a the resolution of all structures is affected by the final resolution of the quadrupole-monochromator and the noise at low count rates. All our samples containing different numbers of wells \((1 \leq N_{\text{QW}} \leq 50)\) show a very good crystalline quality and excellent homogeneity in growth direction with respect to layer thicknesses and compositions (Figs. 2–5).

Finally, the homogeneity of our structures is studied in lateral direction on the wafer, which is essential for device yield and the reproducibility of specific device properties. PL scans at 300 K were performed on several uncleaved 2"-wafers. Fig. 6 displays the wavelength of the PL maximum in [0 1 1] direction on (100) oriented surfaces.
MQW structures with 10 ternary and 10 quaternary wells are compared, having an absolute emission wavelength of $\lambda_0 = 1.55 \, \mu m$ at 300 K. Note that the profiles are vertically shifted and the vertical scale is blown up (one tick corresponds to a wavelength variation of 0.06%). The two lower profiles correspond to compressively strained Ga$_{0.29}$In$_{0.71}$As wells ($L_z = 3$ nm) and lattice matched Al$_{0.225}$Ga$_{0.25}$In$_{0.525}$As barriers ($L_B = 8$ nm). The two upper profiles correspond to compressively strained Al$_{0.093}$Ga$_{0.197}$In$_{0.71}$As wells ($L_z = 7.2$ nm) and tensile strained Al$_{0.205}$Ga$_{0.444}$In$_{0.347}$As barriers ($L_B = 5.8$ nm). The lateral wavelength variations of all samples with ternary wells are similar. Two typical examples are shown in Fig. 6. All structures with quaternary wells also show a similar behaviour among each other, which is reverse, however, to the ternary wells. With respect to device fabrication it is important to note that the structures with quaternary wells show a much larger part in the center of the 2'' wafer where the wavelength is almost constant. For structures with quaternary wells this area exceeds 5 cm$^2$ and, thus, is five times larger than for ternary wells. That difference seems to be partly influenced by the ratio of the well widths of 3 nm/7.2 nm. Assuming in both cases an identical absolute fluctuation in well width, the emission wavelength of the wider wells is less sensitive to these changes. For the second profile from the top the standard deviation is only 0.065 nm in the central region.

In the following, possible explanations will be given for the different radial variations of the wavelength profiles of the ternary and quaternary wells. Since variations in the layer thickness and composition of the well materials cause much larger changes in the quantization energies, we do not consider the respective changes in the barriers. Now, we list possible mechanisms which in combination may explain the observed lateral wavelength variations in Fig. 6.

1. Compositional changes due to slightly varying desorption rates may be caused by a possible small temperature gradient in radial direction. Since the desorption of Al is negligible, no gradient in the Al-content will occur. In contrast, the desorption
Fig. 5. Blow-ups of experimental and simulated XRD-rocking curves of strain-balanced heterostructures including 50 QWs (a) and 30 QWs (b).

Fig. 6. Variation of the wavelength of the PL maximum at 300 K in lateral direction for (1 0 0) oriented wafers. The lasers structures include 10 ternary wells (open symbols) or quaternary wells (closed symbols) and have an emission wavelength of $\lambda \approx 1.55 \mu m$. For a better distinction the profiles are vertically shifted.

rates of In and Ga are considerably higher. For a given substrate temperature AlGaInAs shows a weaker In desorption than GaInAs, due to the presence of Al. However, we believe that this effect is still very small for 510°C.

(ii) The layer thicknesses are mainly determined by the fluxes of the molecular beams, and possible radial temperature gradients do not have a significant influence. We believe that the differences in the profiles of structures with ternary and quaternary wells most likely result from spatial flux variations (the centers of the beam characteristics probably do not coincide with the center of the wafer, i.e. the rotation axis).

The use of quaternary QWs in photonic devices allows strain and well width to be varied independently of each other while maintaining e.g. 1.55 $\mu m$ emission wavelength. The use of strain compensation overcomes the total critical layer thickness restrictions and enables a free choice of the number of QWs in photonic device design. Recently, we obtained a record modulation bandwith of 26 GHz for 1.55 $\mu m$ AlGaInAs lasers [16–18] with either ternary or quaternary QWs.

In conclusion, strain-balanced high-quality AlGaInAs/AlGaInAs MQW structures were grown by solid source MBE. The PL linewidths are low and do not vary with the number of QWs (1 \lesssim N_{QW} \lesssim 50). With respect to photonic device applications, structures with quaternary QWs seem to be superior compared to structures with ternary QWs, e.g. due to more efficient carrier capture from the cladding layers [19] and due to a five times larger lateral wafer area in which the PL wavelength is constant.

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References


New route to reduce ionized impurity scattering in modulation-doped GaAs quantum wells

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Abstract

We propose a new concept for the reduction of impurity scattering in remotely doped GaAs single quantum wells by using heavy-mass X-electrons in the short-period AlAs/GaAs superlattice barriers which smooth the potential fluctuations of the ionized Si dopants. Electron mobilities as high as 120 m²/V·s and electron densities of 1.6 × 10¹⁶ m⁻² are obtained in 10 nm GaAs single quantum wells in the one-subband occupation mode without any parallel conductance. To demonstrate the applicability of our concept magnetotransport and photoluminescence measurements together with self-consistent calculations are presented. Limitations of the effectiveness of the reduction of impurity scattering by impurity segregation and intermixing in the superlattice are discussed.

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Keywords: Electronic materials; Heterojunctions; Quantum wells; Superlattices; Molecular beam epitaxy

1. Introduction

For the application in low-noise and high-frequency devices a high conductivity of the two-dimensional electron gas (2DEG) in a heterojunction is required. Material combinations providing high barriers allow for high 2DEG concentrations \( n \) and make the devices work at higher temperatures [1–4]. However, in a given material system the scope to increase the conductivity either by simply increasing the dopant density or decreasing the dopant sheet-conductive channel separation is limited. The occurrence of parallel conductivity in the doping channel as well as the population of excited subbands with additional intersubband scattering [5] and, most importantly, the increased remote impurity scattering [6, 7] result in a lowering of the electron mobility \( \mu \) and hence, the conductivity. In addition, in high mobility samples other limitations of the mobility such as interface roughness scattering in (Al,Ga)As systems [8] and alloy disorder scattering in ternary systems [9] become important. Recently, low temperature experiments with a high electron mobility of about 400 m²/V·s in the (Al,Ga)As system with densities*

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of $0.64 \times 10^{15}$ m$^{-2}$ per quantum well [2] or up to $3 \times 10^{15}$ m$^{-2}$ have been reported [3]. In the pseudomorphic (AlGa)As/(InGa)As material system conductivities approaching $n \times \mu = 9 \times 10^{16}$/V·s up to temperatures as high as $T = 77$ K have been obtained [4] by optimizing the quantum well thickness and the dopant separation from the quantum well.

In this work we propose a new concept for both the enhancement of the carrier concentration and the reduction of remote impurity scattering (RIS) in GaAs single quantum wells (SQW), thereby significantly increasing the $n \times \mu$ product, i.e., the conductivity. The concept is based on the occurrence of heavy-mass X-electrons in short-period superlattices (SPSL) barriers, which are located near to the dopant sheet and are very effective in smoothing potential fluctuations caused by randomly distributed dopant atoms. The barriers for the GaAs SQW are formed by AlAs/GaAs SPSL. We present magnetotransport experiments in order to confirm the applicability of this concept. Results of photoluminescence (PL) measurements show the presence of X-electrons in our structures.

2. Experimental procedure

The structures under investigation are grown by solid-source molecular beam epitaxy on seminsulating GaAs (0 0 1) substrates. The free carriers in the 10 nm GaAs SQW are provided by remote $\delta$-doping with Si at a distance of 14 or 10 nm. The barriers consist of 60 periods of a 4 monolayers (ML) AlAs/8ML GaAs SPSL. Single Si $\delta$-doping sheets with a dopant concentration of $N^{2D} = 2.5 \times 10^{16}$ m$^{-2}$ are placed on both sides of the SQW into a GaAs layer of the SPSL. The growth temperature of the lower barrier and the SQW is 580°C, whereas the temperature for the $\delta$-doping sheet and its vicinity is 510°C to suppress Si segregation into the SQW. The growth rate and the beam-equivalent As$_4$-to-Ga pressure ratio are 0.66 μm/h and 8, respectively. SPSL and SQW are grown in the two-dimensional island nucleation growth mode as was proven by atomic force microscopy and by reflection high-energy electron diffraction in additional experiments. For comparision structures doped on one side as well as on both sides of the SQW are grown at a constant temperature of 580°C. For more flexibility in placing the doping sheet at certain distances to the SQW in these structures the SPSL period is decreased from 12 ML to 6 ML. We study the low-temperature magnetotransport properties of samples with a Hall-bar geometry. A Ti/Au gate electrode is used to change the electron density. In all measurements the magnetic field was perpendicular to the SQW interfaces.

3. Results and discussion

The dependence of the parallel $\rho_{xx}$ and transverse $\rho_{xy}$ components of the resistivity tensor on the magnetic field of a remotely doped SQW with SPSL barriers is shown in Fig. 1. A very high mobility at surprisingly high electron densities is observed. The one-subband occupation is manifested by the one frequency Shubnikov–de-Haas (SdH) oscillation of the corresponding $\sigma_{xx}$ component of the conductivity tensor up to the highest gate voltages. We use the procedure described in [10] to derive the electron densities $n_{SdH}$ from the SdH frequency. These values are equal to the carrier

![Fig. 1. Magnetic field dependence of $\rho_{xx}$ and $\rho_{xy}$ at $T = 0.33$ K for a remotely doped SQW with a spacer thickness of 14 nm.](image-url)
densities \( n_H \) calculated from the low field Hall-effect \( \rho_{xy} = \gamma/(enH) \) using a Hall-factor \( \gamma = 1 \). Together with the exact value \( \rho_{xx} = \hbar/(e^2v) \) and \( \rho_{xx} = 0 \) in the quantum Hall regime for the integer filling factor \( v \) these results demonstrate the absence of any parallel conductance in the doping region. The conductance is determined exclusively by electrons in the lowest 2DEG subband of the SQW. Therefore, we can determine the Hall mobilities of the 2DEG as \( \mu_H = \rho_{xy}/\rho_{xx} \) at low magnetic fields. Fig. 2 reveals an unusual dependence of \( \mu_H \) on the electron density \( n_H \). While we observe a power-like dependence \( \mu_H \sim (n_H)^k \) with \( 1.5 < k < 2 \) at lower densities (for example \( n_H < 0.95 \times 10^{16} \text{ m}^{-2} \) in sample S1 in Fig. 2) the mobility increases drastically up to \( 120 \text{ m}^2/\text{V} \cdot \text{s} \) for \( n_H > 0.95 \times 10^{16} \text{ m}^{-2} \) within a very narrow electron density region. The power-like behaviour of \( \mu_H \) at lower densities is usually related to the remote impurity scattering (RIS) by randomly distributed impurities in the doping plane [9]. At higher densities the power \( k \) exceeds 8, which is difficult to model by scattering theories based on random potentials. We find that the maximum product \( n \times \mu = 1.35 \times 10^{18}/\text{V} \cdot \text{s} \) and \( 4.20 \times 10^{17}/\text{V} \cdot \text{s} \) at temperatures \( T = 0.3 \) and 77 K, respectively. These values are comparable to the highest conductivities in experiments with SQW's at low temperatures, \( T < 1 \text{ K} \) [2, 3]. The \( n \times \mu \) product at \( T = 77 \text{ K} \) in this work is even several times larger than the highest reported value for 2DEG [4]. To explain the increased mobility in our system we adopt a RIS model with reduced fluctuating scattering potential at higher densities and take into account additional carriers with very low mobilities in the SPBSL very close to the doping layers. By voltage dependent capacitance measurements we found that additional low mobility carriers are present in the SPBSL nearby the doping plane [11]. To clarify the nature of the additional carriers we show in Fig. 3 the results of model calculations of the potential and charge distribution in our structures [12]. These calculations account for two different kinds of electrons. First, we
consider the usual \( \Gamma \)-electrons with the isotropic effective mass \( m_\parallel/m_e = 0.07 \) which form the 2DEG ground subband. These electrons with high mobility are located in the SQW. The edge of the higher \( \Gamma \)-electron subbands in the SQW as well as in the SPSL is more than 170 meV above the Fermi-energy and, therefore, these subbands cannot be occupied at low temperatures. This is the reason for the possible enhancement of the electron density in the SQW up to \( 1.6 \times 10^{16} \text{ m}^{-2} \) without causing any parallel conductance. The maximum density for single subband occupation is not yet known. Our model calculations show this value to be much higher than \( 2 \times 10^{16} \text{ m}^{-2} \). Next we have to include the higher conduction band minima of the AlAs/GaAs SPSL system into the calculation. It is known from experiments with AlGaAs/AlAs multiple quantum wells [13] that at AlAs-thicknesses \( d_{\text{AlAs}} < 3-4 \text{ nm} \) the \( X \)-electrons govern the transport properties. These are electrons with the lower in-plane effective mass of about \( m_{\text{eff}}/m_e = 0.25 \) while the heavy mass component \( m_{\text{eff}}/m_e = 1.1 \) in AlAs accounts for the subband quantization. The period in our SPSL is well below the \( \Gamma \)-\( X \) crossover and our calculations show that states with the lower in-plane mass component are occupied first. The subband edge of states with the heavy in-plane mass is too high to be occupied by electrons at low temperatures. Our simple calculations do not account for any spatial lateral relief in the SPSL, but show quantitatively that in this system the \( X \)-electrons appear in the AlAs ‘well’ closest to the \( \delta \)-doped GaAs sheet. Their density and spatial distribution, however, depend sensitively on the interface structure and the lateral potential fluctuations. Our results show conclusively that with \( X \)-electrons in the SPSL the mobility of the electrons in the GaAs SQW is considerably increased. The explanation follows directly from our calculations. Fig. 3 shows that the \( X \)-electrons are located in the AlAs ‘well’ on one side of the doped GaAs sheet in the SPSL. Their expected Bohr radius \( a_B \approx 2-3 \text{ nm} \) as well as their nominal distance from the doping layer \( d \approx 1.7 \text{ nm} \) is considerably smaller than the average distance between the Si-dopant atoms \( d_0 = 1/\sqrt{N_D} \approx 8-9 \text{ nm} \). Therefore, the \( X \)-electrons can be very easily localized at the minima of the fluctuating potential caused by the randomly distributed dopant atoms. This localization explains the absence of any parallel conductance. The high effective mass of the \( X \)-electrons results in a screening parameter which is several times larger than that of the \( \Gamma \)-electrons. Therefore, the \( X \)-electrons screen the fluctuating potential of the ionized Si impurities more effectively and selectively, leading to an increase of the mobility of the \( \Gamma \)-electrons in the GaAs SQW.

Mobility enhancement is not observed in similar samples which are \( \delta \)-doped only on one side. Obviously, deep trap centers are present at the SPSL interfaces which now decrease the mobility. Their potential fluctuations cannot be smoothed because of the missing \( X \)-electrons on this side.

To study the energy spectrum of the \( X \)-electrons we have carried out photoluminescence (PL) measurements. In all our samples the PL spectra for the type-II recombination region show two peaks (see inset in Fig. 3). This structure follows directly from the subband-splitting of the \( X \)-electrons in the SPSL, which is caused by the electric field of the separated \( X \)- and \( \Gamma \)-electrons. The indirect recombination is possible from AlAs-‘wells’ on both sides of the GaAs-‘well’ with a small energy difference. According to our calculations this subband-splitting with an energy separation of nearly 20 meV depends weakly on the position and the doping density (see Fig. 3). This splitting is close to the observed peak separation in the PL. The higher density of \( X \)-electrons at the lower energy sublevel explains the higher intensity of the low-energy side peak in the PL-spectra.

This new route to reduce RIS in modulation-doped GaAs QW is effective unless dopant segregation and intermixing of the SPSL barrier take place. In order to be more flexible in placing the dopant sheet at discrete separations to the well, the SPSL period is reduced from 12 ML to 6 ML. The carrier concentration is not affected, but the mobility decreases by a factor of about 8. This mobility reduction is probably caused by intermixing and dopant segregation. It is known that there is an unidirectional segregation of Ga from GaAs into AlAs at the GaAs/AlAs interface [14]. This segregation leads to an intermixing of the SPSL components. The intermixing process develops inhomogeneously depending on the actual defect structure of the
surface (terrace-step edge structure). The process will be more pronounced the smaller the individual SPSL layers are. As a consequence the dopant segregation towards the SQW interface in the growth direction is enhanced leading to an increase of RIS. Additionally, strong intermixing can lift the type II SPSL character. Growing the whole structure at a constant substrate temperature (580°C) also leads to a loss of the extreme high electron mobility because of similar reasons. Therefore, the achievement of smooth SQW interfaces and the suppression of dopant segregation in combination with intermixing effects should be balanced by an adequate choice of growth temperatures.

4. Conclusions

In conclusion, we have shown that the impurity scattering in remotely doped GaAs SQWs can be reduced effectively by the presence of heavy-mass X-electrons in the direct vicinity of the dopant atoms in the SPSL barriers. These X-electrons exhibit an extremely high screening capability and are able to smoothen the potential fluctuations caused by the random distribution of the dopant atoms. They are localized and do not contribute to the conductivity. PL studies as well as self-consistent calculations confirm the existence of X-electrons in the AlAs/GaAs SPSL barriers. The high electron conductivity is lost by dopant segregation in combination with intermixing effects in ultra-short period superlattices. With an adequate choice of the growth procedure these detrimental effects can be minimized.

Acknowledgements

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References

[12] K.-F. Friedland and R. Zimmermann, unpublished. The calculations are performed by the one-dimensional self-consistent solution of the Schrödinger and Poisson equations in the effective mass approximation, including exchange and correlation interaction energy.
The growth of AlGaAs–InGaAs quantum-well structures by molecular beam epitaxy: observation of critical interdependent effects utilizing the design of experiments approach

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Abstract

The traditional approach to determine relationships between growth conditions and material properties has rested on the standard experimental approach – varying one parameter while holding all others constant. This technique does not effectively allow the observation of important interactions in complicated multilayer structures. Critical interdependent effects have been observed in the growth of AlGaAs/InGaAs quantum-well structures by molecular beam epitaxy (MBE). It is shown that statistical experimental design is an effective method for optimizing complex multilayer structures quickly. This technique is very useful for the optimization of processes with a large number of interdependent parameters and allows for the clear visualization and separation of complex interwoven effects. In the present work, we show for the first time the importance of the oxide desorption process for the optimal growth of AlGaAs-containing structures. The choice of an optimized oxide desorption process can lead to a decrease in the interfacial oxygen by almost two orders of magnitude.

1. Introduction

Molecular beam epitaxy (MBE) is a well-developed and versatile technique for the growth of compound semiconductor heterostructures and devices. Although this process has the capability for stringent control of layer composition, purity, and thickness [1], the process optimization for a given device structure is complicated by a large number of coupled control parameters, such as the substrate temperature, V/III flux ratio, growth rates of alloys, etc. For this reason the optimization of a new device structure can often take tens to hundreds of growth runs.

The traditional approach to determine relationships between growth conditions and material properties has rested on the standard experimental approach – varying one factor while holding the
others constant. The basic assumption of this approach is that the coupling of the various growth parameters is small. This technique does not effectively allow the observation of important interactions in device structures. For example, growing high-quality AlGaAs requires high substrate temperature, unless other parameters are important, such as the minimization of dopant diffusion, which requires lower substrate temperature. The results of such approaches have been reported in the literature, where the effect of growth temperature on the quality of AlGaAs epitaxial layers has been correlated to growth temperature [2, 3] and possibly interfacial oxygen [4], which leads to a dramatic decrease in the recombination lifetime in these materials [5, 6].

Response surface modeling is a powerful technique for process optimization [7] and has been used to optimize the growth of InGaAs by chemical beam epitaxy [8] and is now standard practice in the silicon processing industry. In the present work, we have utilized a Resolution IV, 2⁴-2 fractional factorial experimental design (with 16 trials) to ascertain important interactions in a generic AlGaAs/InGaAs single-quantum-well structure. We have observed interactions which have never been quantified before. In addition, we show, for the first time, the importance of effective oxide removal techniques for the growth of high-quality structures. These results clearly show that the optimum growth conditions are strongly coupled, and for this reason the traditional approach of varying one parameter while holding all others constant is ineffective.

2. Experimental procedure

The structure investigated in the present work consisted of a 0.1 μm GaAs interface layer, a 0.5 μm AlGaAs buffer layer followed by a 75 Å InGaAs quantum well, which was capped by 0.25 μm AlGaAs cap layer. All samples in this study were grown in a solid source Varian Gen II MBE system using solid As₄ as the arsenic species. Samples were grown on 1/4 of a 2" semi-insulating GaAs wafer that was briefly etched in 5 : 1 : 1 H₂SO₄ : H₂O : H₂O₂ before being indium-mounted on a standard molybdenum sample holder. In the 16 trial runs the following factors were varied: oxide removal temperature from 580 to 650°C, oxide removal time from 30 to 300 s; AlGaAs growth temperature from 580 to 630°C, InGaAs growth temperature from 480 to 520°C, the interface

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interrupt time from 30 to 90 s, and the arsenic cell temperature from 325 to 340°C (1, 2 × 10^{-5} Torr). The 16 sample runs and the order in which they were performed are summarized in Table 1. It is important to note that the RHEED pattern indicated complete oxide removal for all the 16 samples in the experiments.

All 16 samples were characterized by surface defect density, AlGaAs structural and optical quality, and InGaAs optical quality by 77 K photoluminescence (PL) and double-crystal X-ray diffraction (DCRC).

3. Results and discussion

The chosen factors were all found to be statistically significant and large variations in material properties were observed. For example, the defect density varied by one order of magnitude, the AlGaAs PL full-width at half-maximum (FWHM) varied by a factor of six, the AlGaAs X-ray FWHM varied by a factor of two, and the InGaAs PL FWHM varied by a factor of six. The experimental data were analyzed using the RS/Discover commercial software package [9]. Table 2 shows the main effects and two-factor interactions found to be significant at a 95% confidence level using the Student’s t-test statistic [7]. This table shows one of the significant advantages over the statistically designed experiment approach: it not only allows the visualization of the results, but also provides a numerical estimate to the reliability of the results.

The results for the InGaAs photoluminescence (PL) full-width at half-maximum (FWHM) as a function of the InGaAs growth temperature and arsenic cell temperature is shown in Fig. 1. The narrowest photoluminescence linewidths are obtained at the lowest substrate temperature and the highest arsenic fluxes. This observation is in agreement with results published in the literature [10], and gives confidence that our approach agreement with the known phenomena.

We also observe a strong relationship between the growth temperature of the AlGaAs buffer layer and the resultant quality of the InGaAs quantum wells. The InGaAs PL FWHM as a function of the AlGaAs buffer layer growth temperature and the

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arsenic cell temperature is shown in Fig. 2. The best InGaAs photoluminescence is obtained when the AlGaAs buffer layer is grown at the higher end of the range of the AlGaAs growth temperatures investigated. This is consistent with the observation that a higher AlGaAs growth temperature improves surface morphology, which, in turn, should improve quantum-well interface smoothness.

Fig. 1. InGaAs photoluminescence FWHM as a function of substrate temperature during the InGaAs growth and arsenic flux.

Fig. 2. AlGaAs photoluminescence FWHM as a function of substrate temperature during the growth of the AlGaAs and arsenic flux.
The effect of the oxide desorption process prior to the initiation of growth on the quality of the AlGaAs buffer layer is shown in Fig. 3, wherein the AlGaAs PL FWHM is plotted as a function of the desorption temperature and time. As can be seen, the quality of the AlGaAs epitaxial layer is dramatically affected by the exact desorption process. To our knowledge, this is the first quantitative observation of the critical importance of the oxide removal conditions on AlGaAs quality.

Not only are the temperature and time of the desorption process important, but as can be seen in Fig. 4, there is a coupling between the desorption temperature and the arsenic flux. The photoluminescence FWHM of the AlGaAs buffer is shown as a function of the desorption temperature and arsenic flux. This relationship may be due to growth kinetic processes during the AlGaAs growth and/or changes in surface roughness during the oxide desorption process which are arsenic-flux-dependent.

Secondary ion mass spectrometry measurements (SIMS) have confirmed that the higher desorption temperatures lead to films with lower oxygen content. The oxygen concentration at the substrate/GaAs buffer layer interface as a function of oxide desorption temperature and time is shown in Fig. 5. The critical nature of the oxide desorption is readily evident. At low oxide desorption temperatures (580°C) a decrease in the interfacial oxygen concentration by also two orders of magnitude can be obtained by increasing the hold time before the initiation of growth from 30 to 300 s. Increasing the temperature at which the oxide desorption takes place also reduces the amount of interfacial oxygen to a lesser extent.

The exact oxide desorption process employed not only affects the amount of oxygen at the film/substrate interface, but also has a dramatic effect on the oxygen incorporation in the AlGaAs buffer layer. The oxygen concentration in AlGaAs buffer layer as determined by SIMS is shown in Fig. 6 as a function of the desorption temperature and desorption time. Similar to the results for the interfacial oxygen, a strong decrease in the concentration of oxygen in AlGaAs buffer layer is seen upon increasing the hold time at the oxide desorption temperature. It is also observed that the concentration of oxygen in AlGaAs buffer layer actually increases slightly as the oxide desorption temperature is raised from 580 to 650°C. This may be due to the oxygen segregation at the growth front and/or increased outgassing of the surrounding growth environment.

![Fig. 3. AlGaAs photoluminescence FWHM as a function of oxide removal temperature and time for a growth temperature of 630°C.](image-url)
Fig. 4. AlGaAs photoluminescence FWHM as a function of oxide removal temperature and arsenic flux.

Fig. 5. Substrate/GaAs interfacial oxygen concentration as a function of the oxide desorption temperature and desorption time as determined by SIMS.
Fig. 6. Concentration of oxygen in the AlGaAs buffer layer as a function of the oxide desorption temperature and desorption time as determined by SIMS.

4. Summary

In summary, we have observed and quantified significant interactions in the growth of AlGaAs-InGaAs quantum-well structures. We have shown that statistical experimental design is an efficient means for optimizing growth conditions for a given device structure. In addition, we quantify, for the first time, the important interactions between effective substrate oxide desorption and AlGaAs and InGaAs growth conditions.

Acknowledgements

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References

MBE growth and physics of strongly coupled p-type double quantum wells showing correlated $v = 1$ quantum Hall state

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Abstract

We present a systematic magnetotransport study of high mobility (311)A strongly coupled p-type double quantum wells which demonstrate the existence of a correlated $v = 1$ quantum Hall state in the limit of weak tunnelling. Our results are in good agreement with the finite temperature phase transition predicted for this state.

Keywords: Two-dimensional hole gas; Molecular beam epitaxy; Gallium aluminium arsenide; Tunnelling

1. Introduction

Until recently, it has not been possible to realize a two-dimensional hole gas (2DHG) with high mobility using Be-doped GaAs/(AlGa)As heterostructures. Mobility enhancement of 2DHG has been observed by Henini et al. [1] in modulated p-type GaAs/(AlGa)As heterostructures on the (311)A GaAs surface using Si as the acceptor. A 2DHG with low temperature mobility as high as $1.2 \times 10^6 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$ was obtained. Such high-quality samples allowed them to study the fractional quantum Hall effect for holes, a fundamental physical phenomenon which has been intensively investigated in the two-dimensional electron gas (2DEG) in n-type GaAs/(AlGa)As heterostructures with mobilities exceeding $10^7 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$. Using the same growth procedure as those of Ref. [1] we have recently grown a series of very closely spaced p-type double quantum wells.

In this article we report the MBE growth of high-quality p-type modulated double quantum wells on (311)A using Si as the p-type dopant. We will show that the p-type double quantum wells exhibit interesting phenomena compared to the n-type double quantum well samples which have recently attracted considerable interest [2–4].

In single-layer, two-dimensional (2D) conductors the integer quantum Hall effect (IQHE) is a single particle phenomenon which arises when the Fermi energy lies in the region of localised states between Landau levels. The FQHE is a many-body phenomenon which arises from the intra-layer
Coulombic interaction. This leads to transitions into correlated liquid-like ground states when the Landau level filling factor, \( v \), has particular odd denominator fractional values. In these cases, there exists a finite gap separating the ground state from the lowest-lying excited state. When two such 2D layers are in close proximity, the inter-layer Coulombic interaction can lead to new FQHE states when the filling factor in each layer is an even denominator fraction [2].

In a double layer system the quantum Hall effect at a total Landau level filling factor \( v = 1 \) (i.e., \( v = \frac{1}{2} \) in each layer) can arise in two ways. If tunnelling is strong a single particle state can occur in which the energy gap is that separating the lowest symmetric and antisymmetric states of the system (\( \Delta_{\text{SAS}} \). A many body \( v = 1 \) state is also possible for strong inter-layer Coulombic interaction even in the absence of tunnelling [2]. The \( v = 1 \) state is expected to occur when the width of the barrier between the two quantum wells, \( d \), is comparable with the magnetic length, \( \ell_B \). An explanation to the origin of the \( v = 1 \) state has been treated theoretically [2] by considering that the carriers in the two quantum wells have a fictional spin of \( \pm \frac{1}{2} \).

The lowest energy charged excitations of the bi-layer correlated \( v = 1 \) states are merons which can be pictured as vortices in the spin orientation and which carry a charge of \( \pm e/2 \). At low enough temperatures, these objects are bound together in vortex–antivortex pairs with a total charge of 0 or \( \pm e \) (a closely bound meron pair carrying charge \( \pm e \) can be viewed as a skyrmion). At a finite temperature, \( T_{\text{KT}} \), a Kosterlitz–Thouless transition is predicted to occur. At \( T_{\text{KT}} \) the pairs unbind and the energy gap to charged excitations disappears. It is predicted [2] that the bi-layer \( v = 1 \) state should only be stable for \( d/\ell_B < 1.6 \), since \( T_{\text{KT}} \) tends to zero at this value.

In n-type GaAs/GaAlAs double layer systems at small layer separation, when the inter-layer Coulombic energy is large, tunnelling also tends to be strong due to the small electron effective mass. Due to the added effect of tunnelling of the carriers between the two quantum wells the assignment of the origin of the \( v = 1 \) state becomes difficult. States for \( v = 1 \) showing behaviour consistent with that predicted for the correlated state have been observed in a series of n-type double quantum wells which have \( \Delta_{\text{SAS}} \) values as low as 0.8 K [3]. However, in these samples tunnelling still plays an important role as is shown in tilted field experiments [3]. Complex behaviour has also been observed in very wide quantum wells which have some bi-layer character, but which also have very large \( \Delta_{\text{SAS}} \) [4]. It is clear from these studies that inter-layer correlations are important in stabilising the observed \( v = 1 \) states. However, the prediction of a \( v = 1 \) state in the limit of small \( \Delta_{\text{SAS}} \) has still not been directly demonstrated experimentally. In this paper we will concentrate on results for high mobility p-type double quantum wells for samples from one MBE grown wafer (NU1174), though the \( v = 1 \) state is observed in samples with appropriate densities from several wafers.

2. Experimental procedure

The structures were grown by MBE on 2-in liquid-encapsulated Czochralski semi-insulating substrates oriented within \( \pm 0.5^\circ \) of (3 1 1)A. The growth parameters were: substrate temperature of 630°C, beam equivalent pressure of \( \text{As}/\text{Ga} = 12 \), and a preferred growth rate of one monolayer per second and half a monolayer per second for GaAs and AlAs, respectively. The \( \text{As}_4 \) beam was generated using an Intevac Cracker Cracker cell filled with arsenic lumps from Preussag.

The preparation procedures of the MBE equipment broadly followed the method adopted by various groups and recommended by Intevac, and the growth parameters used above were those which gave us optimal electrical properties of (3 1 1)A heterostructures [1]. As we reported in Ref. [1] we used silicon as a p-type dopant for the (3 1 1)A orientation.

The modulation-doped samples consisted of a buffer layer followed by two GaAs quantum wells bounded on each side by an undoped \( \text{Al}_{0.35}\text{Ga}_{0.67}\text{As} \) spacer layer, and lightly silicon-doped \( \text{Al}_{0.33}\text{Ga}_{0.67}\text{As} \) layer. The whole structure was terminated by a 17 nm GaAs capping layer. The buffer layer consisted of 1 \( \mu \text{m} \) thick undoped GaAs followed by a superlattice \( 20 \times (2.5 \text{ nm} \text{ Al}_{0.33}\text{Ga}_{0.67}\text{As} + 2.5 \text{ nm} \text{ GaAs}) \) and 0.5 \( \mu \text{m} \) un-
doped GaAs. A two-dimensional hole gas (2DHG) is confined at each of the GaAs/Al0.33Ga0.67As interfaces. Each quantum well is 100 Å wide and they are separated by a 30 Å AlAs barrier as is shown in the insert of Fig. 2. The layer structure is almost symmetrical with Si-doped layers with the same doping density and the same thickness on each side of the quantum wells. However, it has been found necessary to make the lower spacer layer thicker (650 Å) than the upper spacer (600 Å) to compensate for the upward drift of the dopant with the growth surface.

The epitaxial layers were processed into Hall bars orientated along the [2 3 3] direction, by means of conventional photolithography and wet etching. The contacts consisted of 5 nm of Au, 10 nm of Zn, and 200 nm of Au annealed for sixty seconds at 510°C.

3. Results and discussion

The total carrier density and the relative densities in the two wells can be controlled using front and back gates. This is illustrated in Fig. 1 where the low field Shubnikov–de Hass oscillations for a series of different front gate biases are shown. Off balance a complex beating is seen while at balance one observes a single oscillatory period at the lowest fields before the spin-splitting is resolved at slightly higher fields. The density can be accurately determined from the Fourier transform of the low field Shubnikov–de Haas oscillations. As grown the carrier density in each well for NU1174 samples is within 5% of $1.1 \times 10^{15}$ m$^{-2}$ and the average mobility is 400 000 cm$^2$ V$^{-1}$ s$^{-1}$. The mobility is found to be a very weak function of carrier density as is the case for p-type heterostructures [1]. The potential and density distribution for this structure, shown in Fig. 2, was obtained by solving the Poisson equation and the one-electron Schrödinger equation self-consistently. A parabolic dispersion with a mass of 0.45$m_0$ was used and many-body effects were included in the local-density-approximation. This yields a weakly hole-density-dependent $\alpha_{\text{SAS}}$ of $\sim 70$ mK. This very small value is the result of the very weak tunnelling for the high mass holes.

Fig. 2 shows the typical behaviour of the longitudinal resistance, $R_{xx}$, and the Hall resistance,
Fig. 2. Typical $R_{xx}$ and $R_{xy}$ results (at 60 mK) for equal hole number densities of $1.1 \times 10^{11}$ cm$^{-2}$ in each well. The insert shows the calculated potential profile and hole density distribution.

$R_{xy}$ when the hole number densities in the two wells are balanced. As we will show, strong $\nu = 1$ quantum Hall states are observed which have activation energies of up fifty times $\Delta_{SAS}$. The states at $\nu = 3, 5, 7, \ldots$ are all absent as expected for weak tunnelling. This is direct unambiguous proof of the existence of the correlated $\nu = 1$ state in the limit of weak tunnelling.
Activation plots of $\ell n R_s$ against $1/T$ are given in Fig. 3. This shows very unusual behaviour since at low enough temperatures one observes normal activated behaviour indicating the presence of an energy gap, $\Delta E$. But as the temperature is raised the activated behaviour is terminated rather abruptly and no minimum in $R_s$ is observable above a temperature $T^*$ which is much smaller than $\Delta E$. This is in contrast to the behaviour of the adjacent $\nu = \frac{3}{2}$ state, which shows normal behaviour. This behaviour is predicted from the well-known Kosterlitz-Thouless transition in which one has the disappearance of the energy gap for charged excitations at a finite temperature, $T_{KT}$. It is worth noting that, surprisingly, similar behaviour has been seen [5] in very wide quantum wells despite the very strong tunnelling.

In Fig. 4 we present the dependence of the activation energy and our $T^*$ values as a function of $d/\ell_B$. $T^*$ steadily falls towards zero as $d/\ell_B$ approaches $\sim 1.7$, while $\Delta E$ falls rapidly to zero at this value. The predicted values [2] of $\Delta E$ are $\sim 10$ K. However, it is expected that finite thickness corrections, Landau level mixing and disorder will all tend to reduce this value. Our values are, therefore, not inconsistent with expectation. A more useful comparison can be made between our values of $T^*$ and the predicted behaviour of $T_{KT}$ since this transition temperature should be less influenced by disorder. To do so, we have taken the prediction of Ref. [1] for the behaviour at a fixed $\ell_B$ as a function of $d$ and scaled them by the intra-layer Coulombic energy. In Fig. 4 we have plotted the expected $T_{KT}$ against our upper ($d/\ell_B)_{eff}$ scale. Good agreement in functional form but a disagreement by a factor of $\sim 2$ in magnitude is found. Given the uncertainties in the calculated magnitude and in our scaling procedure, this disagreement in magnitude is not surprising.

4. Conclusion

We have shown that samples of Si-doped GaAs/AlGaAs coupled 2DHG structures with mobilities as high as 400 000 cm$^2$ V$^{-1}$ s$^{-1}$ can be achieved on the (3 1 1)A GaAs surface. Furthermore we have observed correlated $\nu = 1$ bi-layer states in the limit of weak tunnelling and find evidence of a finite temperature phase transition. The nature of the state is clear from the fact that it systematically weakens with increasing carrier density, and is destroyed for $d/\ell_B > 1.7$. Our results are in good agreement with recent theoretical predictions.

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References


Photoluminescence study of Si/SiGe multiple quantum wells grown by MBE

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Abstract

Pseudomorphic Si/SiGe multiple quantum well (MQW) structures were grown by solid source molecular beam epitaxy (MBE). The optical and structural properties of MQWs grown with and without bias applied to the substrate during the deposition were studied in detail. SiGe phonon resolved band-edge luminescence of as-grown samples is found only for MQWs deposited with a positive bias at the substrate. Subsequent rapid thermal annealing improves the photoluminescence properties of films grown with zero or positive bias, whereas for samples grown with a negative bias a broad deep luminescence band is observed. A strong correlation between the observation of the deep luminescence and the appearance of defects in the SiGe layers is established. It is most likely that these defects originate from ion bombardment of the growing film during e-beam evaporation in solid source MBE.

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Keywords: Quantum well structures; Photoluminescence; SiGe alloys; Molecular beam epitaxy

1. Introduction

Molecular beam epitaxy (MBE) is a key technology for the development of Si/Ge-based material systems. In fact it is the only epitaxial system which permits a controlled growth of structurally perfectionized Si/Ge superlattices (SL) [1] and more recently, of coupled Si_{1-x}Ge_xC_y/Si_{1-y}C_y quantum wells (QWs) [2]. Both structures are of particular interest for the development of an efficient silicon-based electrically pumped light emitter. For the SL structures intense luminescence is predicted due to zone folding effects and symmetry breaking of states at the interfaces [3], whereas for the coupled QWs the enhancement is attributed to spatially indirect (type II) transitions of electrons and holes confined in neighbouring quantum wells [2]. Besides a well-resolved band-edge-related luminescence spectra consisting of a no-phonon (NP) peak and phonon replicas, the luminescence of samples
grown by solid source MBE frequently show a deep luminescence, 80–120 meV below the band-edge related luminescence [4, 5]. In particular for SiGe QWs grown at low temperatures (< 700°C) this broad band is often observed, whereas band-edge luminescence is observed typically for samples either grown at higher temperatures [6–8] or subsequently annealed above 700°C [9]. The physical origin of this deep luminescence is still a matter of controversy. Several suggestions, including an optically active defect [5], donor acceptor pairs [10] and Ge-rich platelets [11], have been put forward to explain the phenomena. More recently, Sturm et al. [12] were able to create deep photoluminescence by ion implantation of Si into Si/SiGe QWs grown by rapid thermal chemical vapor deposition. They were able to recover the band-edge luminescence by annealing at 800°C. This result suggests that the deep luminescence of MBE grown samples is related to ion bombardment during growth. Some initial results using a substrate bias during the growth by MBE also point towards this model [13].

In this study we report on the optical properties (low-temperature photoluminescence (PL)) in relation with the structural properties, analysed by transmission electron microscopy and X-ray diffraction, of SiGe/Si multiple QWs. The deposition temperature, the substrate bias and the time and temperature of an subsequent rapid thermal annealing (RTA) of the QW samples were chosen as variables.

2. Experimental procedure

The SiGe/Si structures were grown in a solid source MBE chamber containing e-beam evaporators for Si and Ge, a pyrolitic graphite filament for C sublimation and effusion cells for B and Sb. A schematic view of the chamber is shown in Fig. 1. The distance between the e-beam evaporators and the substrate is approximately 0.5 m, which gives on the one hand good uniformity, but on the other hand requires quite high electron emission currents of the e-beam evaporators to achieve reasonable growth rates for the Si and SiGe layers (0.1–0.2 nm/s). In turn, this high emission current will also lead to enhanced generation of secondary and scattered electrons. For the Ge source a custom made Si crucible was used. The Si and the Ge source were shielded by Si walls to reduce the amount of scattered and secondary electrons traveling throughout the chamber. The flux of the e-beam evaporators is feedback controlled by a mass spectrometer.

The samples discussed were grown on undoped Si(1 0 0) substrates with a resistance of > 400 Ω cm at temperatures ranging from 550°C to 750°C. The temperature was measured by a thermocouple close to the filament, the actual temperature of the wafer was probably lower. The QW structures were grown on a 100 nm thick Si buffer layer. The QW structures contained 10 periods with 10 nm-wide Si barriers and SiGe wells of typically 2–8 nm and Ge concentrations between 5% and 30%. The composition and thicknesses of the wells were determined by X-ray diffraction and cross-sectional TEM.

Prior to growth the substrate bias was adjusted between −1500 and + 1500 V. After the e-beam evaporators were powered the + 1500 V bias dropped to about + 200 V. The currents between the substrate and the ground was measured to be 60–70 nA.

After the growth, the QW structures were thermally treated by a RTA process in the temperature range from 750–950°C for 240 s, in a N₂/H₂ gas mixture as well as in pure N₂. PL spectra were taken before and after the RTA treatment. The
spectra were recorded at 2.4 K using an Ar⁺ laser at 488 nm for excitation (excitation power: 6 mW) and a cooled Ge photoconductor for detection.

3. Results and discussion

The dependence of low-temperature (2.4 K) PL spectra on the substrate bias is shown in Fig. 2.

Fig. 2b depicts the PL spectra of a sample grown at 550°C with 0 V bias, i.e. with a floating substrate, and subsequently annealed at temperatures ranging from 750°C to 950°C. The as-grown sample shows beside the Si PL lines of the substrate a broad-band luminescence. This broad band gets even more pronounced when a substrate bias of −1500 V is applied to the sample as shown in Fig. 2a. The PL spectra of the sample shown in Fig. 2c is grown with a substrate bias of + 1500 V. In this case the broad-band luminescence is suppressed and band-edge luminescence of the SiGe QWs is visible. After the sample anneals at temperatures above 750°C the multiple quantum wells (MQWs) grown with positive and no bias show only band-edge luminescence and the broad-band luminescence vanishes. With increasing RTA temperatures the NP lines get more intense and the line width decreases. These results are observed independently of the gas ambient, pure N₂ or N₂/H₂ mixtures, indicating that the effect cannot be attributed to hydrogen passivation of defects in the QWs. It is remarkable that the PL lines of the MQW grown with the substrate bias of + 1500 V shift continuously with temperature to higher energies, indicative of Ge diffusion from the wells into the Si barriers. In contrast, the MQW grown with floating bias shows only very a small shift up to RTA temperatures of 900°C and then a significant shift at 950°C. The MQW grown with the negative bias (−1500 V) shows no band-edge luminescence after RTA processing and the broad-band luminescence remains quite significant.

A possible interpretation of the results is the following. During e-beam evaporation of Si and Ge, many secondary and scattered electrons hit the substrate thus charging it negatively. Due to this negative charge, positive Si and Ge ions are accelerated towards the substrate damaging the growing layer. By applying a positive bias to the substrate the effect is suppressed, whereas a negative bias enhances the damage caused by the ions.

In order to obtain more information about the structural properties of these defects, a careful analysis by TEM of as-grown MQW deposited with and without bias as well as after RTA processing has been done. Before discussing the results in detail, a few general comments about the TEM
analysis are warranted. Fig. 3a shows a cross-sectional bright field image of an as-grown MQW containing 5 nm wide SiGe quantum wells (dark layers in the image) with a Ge concentration of 21.8%. A bias of +1500 V was applied before starting the growth. The image reveals no extended defects, no interface contamination and smooth interfaces. Fig. 3b and Fig. 3c show high resolution images of the same sample investigated with (b) 300 keV and (c) 200 keV electrons in the TEM. In both images small defects show up as dark shadows, with extensions not larger than 5 nm. In the case of 300 keV conditions they show up in some cases with a stronger contrast. The defocus conditions optimized for these defects result in a rather weak contrast between Si and SiGe layers.

To insure that the defects are grown in defects, and are not created by the ion milling during the preparation of the TEM samples, or stem from electron bombardment during examination, the preparation and examination parameters were varied, and samples grown by other techniques, like atmospheric pressure chemical vapor deposition, were examined. The defects are found independent of the electron energies and of the dose, rendering damage during examination rather unlikely. However, the defects are only found in samples grown by solid source MBE, and not in samples grown by other techniques and prepared and examined with the same parameters. This gives strong indications that the defects are present in the as-grown film, although they might be point defects which get enlarged during preparation and examination.

In the next paragraph we demonstrate that these defects can be correlated with the PL features. As discussed in Fig. 2 the PL spectra of MBE grown Si/SiGe MQWs show band-edge luminescence after the RTA step at temperatures > 750°C or by applying a positive substrate bias. For this reason we carefully prepared TEM samples of MQWs which were grown under different bias conditions and with different subsequent annealing steps.

Fig. 4 compares TEM micrographs of a MQW with 5 nm wide Si$_{0.83}$Ge$_{0.17}$ wells separated by 10 nm-thick barriers grown at 550°C before and after annealing at 950°C and a MQW sample

![Cross-section TEM micrographs of a Si/SiGe MQW structure](image)

Fig. 3. Cross-section TEM micrographs of a Si/SiGe MQW structure, (a) bright field TEM image, (b) high-resolution TEM taken at 200 kV and (c) taken at 300 kV.
grown at 750°C. Whereas the TEM micrograph of the as-grown (550°C) sample (Fig. 4a) shows defects distributed throughout the wells and the barriers, the TEM micrograph of the annealed sample (Fig. 4b) exhibits defects only in the Si barriers. For clarity, the latter TEM micrograph is divided into two parts taken with different defocus conditions giving different contrasts for the defects. They are visible as dark spots lined up in the otherwise rather bright area of the Si wells. The SiGe layers are visible as gray bands of quite uniform color. For the sample grown at 750°C a similar behavior is
Fig. 5. High-resolution TEM micrographs of samples grown with (a) negative (-1500 V), (b) zero and (c) positive (+1500 V) substrate bias.
observed, the defects are lined up in the Si barriers. Notice that for the annealed sample, where the defects are found only in the Si barriers, intense phonon resolved luminescence of the SiGe wells is observed. In contrast, the as-grown sample reveals only a broad deep PL band, and at the same time the defects can be found in the SiGe as well as in the Si layers. This result gives clear proof of a correlation between the defects and the broad-band luminescence. Apparently the annealing leads to an outdiffusion of the defects from the SiGe layers into the Si. This diffusion process may be strain driven, i.e. the defects lower overall strain in the system by moving from the SiGe into the Si layer. The diffusion behavior agrees well with the PL results of Sturm et al. [12] for CVD-grown Si/SiGe heterostructures subsequently implanted with Si and annealed at 600–800°C. It is also noteworthy that defects are only obtained in the Si barriers even without subsequent annealing for the samples grown at elevated temperatures ( > 700°C). Typically, at these growth temperatures phonon resolved PL is observed for MBE-grown SiGe quantum wells [6].

Since band-edge luminescence is observed also for samples grown at low temperatures (550°C) which were deposited with a negative bias, it is interesting to do some careful TEM analysis with those samples. Fig. 5 compares TEM micrographs of MQWs grown with (a) positive bias (+1500 V), (b) zero bias and (c) negative bias (−1500 V). All samples have 17–21% Ge in the wells, a barrier width of 10 and well widths of 5 nm. The negative applied bias lead to a very defective sample throughout all layers. In addition, samples grown with a negative bias show contaminated substrate interfaces and rather large precipitate-like defects in the layers. Apparently the negatively biased substrate attracts positively charged particles in the MBE chamber.

The difference in the TEM cross-sectional micrographs for the positively and zero biased samples is not obvious. However, there are significantly less defects in the SiGe wells as in the Si barriers in the biased sample, whereas in the sample grown on the floating substrate the defects are more equally distributed. This result indicates that the activation energy to create a defect in the SiGe layer is lower than in a Si layer, which may again be an effect of the build in strain of the SiGe layers. This result agrees with the observation made for the photoluminescence of the samples. MQWs grown on floating substrates showed a broad deep luminescence, whereas the MQWs deposited on positively biased samples exhibit in no case broad luminescence but frequently phonon resolved band-edge luminescence.

4. Conclusions

Si/SiGe MQWs were grown by solid source MBE using different growth temperatures and substrate bias conditions. The samples were analyzed by TEM and PL before and after RTA at temperatures ranging from 750°C to 950°C. Our results give clear proof of an correlation between the appearance of defects in the SiGe wells and broad band PL. In addition, our results indicate that the defects are caused by ion bombardment during e-beam evaporation in solid source MBE. A positive substrate bias reduces the damage, particularly in the SiGe films. Subsequent annealing of the QW structures strongly improves the photoluminescence of the Si/SiGe MQWs by driving the defects out of the SiGe wells into the Si barriers.

References


Photoluminescence spectra of shadow masked multiple quantum wells

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Abstract

Lines of local epitaxy of GaAs/GaAlAs multiple quantum wells with linewidth of 7 and 5 μm having line separation of 250 μm were fabricated by molecular beam epitaxy (MBE) using shadow masking technique. Photoluminescence (PL) measurements at 10 K were performed by using an argon laser with cylindrical lens producing line focus and with circular lens producing point focus for the excitation. The focussed line and focussed point of laser beam were ~100 μm and could be aligned to the stripes of MQW. The PL spectrum characteristics reflect the quantum well structure as well as the crystal quality of shadow masked epitaxy layers. The PL spectrum shows double PL peaks at 758 and 774 nm with nearly equal intensity when the exciting beam lined parallel to the stripes of MQW have a 7 μm window and quantum well width of 30 monolayers. Overlapping PL peaks at 810 and 818 nm were observed from MQW having 5 μm window and quantum well width of 60 monolayers. Similar results were obtained by point focussed laser beam and by line focussed laser beam at perpendicular configuration. A stronger PL peak at 818 nm was clearly emphasized by strip-off MQW samples having 5 μm stripes. The PL peak from a shadow masked MQW sample was sharp having spectral FWHM of 14 nm and was relatively stronger than that obtained from conventional MQW samples. These experimental data indicated that MBE grown MQW mesa using shadow masking technique is suitable for quantum device fabrication.

Keywords: Photoluminescence; Shadow mask; Molecular beam epitaxy

1. Introduction

Mesa structures of GaAs/GaAlAs multiple quantum well (MQW) have been widely used in many quantum device fabrications. Generated or injected carriers could be easily confined in this mesa structure. Vertical switching device could be designed and realized by the mesa structuring technique. Photolithography and chemical etching are conventional processing steps of the mesa structure. However, chemical etching creates many defects on the etched surfaces affecting active MQW layers. These defects degrade device performance and cause low yield of device mass production. In addition, chemical etching does not allow a reproducible control of the layer thickness of an atomic scale which is needed for three dimensional integration of complex devices having MQW structure.

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Direct preparation of mesa structure by molecular beam epitaxy (MBE) growth is preferable [1] to avoid any chemical etching of active MQW layers. A novel method for mesa growth of epitaxial GaAs layers by MBE technique using patterned shadow masks was investigated [2] and was demonstrated in fabricating GaAlAs/GaAs hetero-nipi structure [3].

This research paper is an attempt to apply the shadow masking technique in growing GaAs/GaAlAs mesa epitaxy MQW by MBE technology. Shadow masked MQW samples were examined by photoluminescence (PL) measurement to confirm the quantum well phenomenon and its sharpness. The successful MBE grown mesa MQW using shadow masks gives stronger PL than a conventional MBE sample. An experiment of PL measurement on a strip-off sample having mesa MQW structure was also conducted to ensure the result. It is found that PL peaks from mesa MQW by shadow masking technique are three times stronger than a reference plain surface MQW and the PL spectrum has an acceptable sharpness of quantum effect (14 nm). This shadow masking technique is suitable for fabricating of MQW having complex structures.

2. Shadow mask preparation

The shadow masks consisted of MBE grown Ga$_{0.5}$Al$_{0.5}$As and GaAs layers. The Ga$_{0.5}$Al$_{0.5}$As layer, so-called mask spacer is grown on (1 0 0) semi-insulator GaAs substrate for 3.5 μm thickness. The GaAs layer is then consecutively grown over the top surface of the mask spacer for about 1 μm thickness. This GaAs layer is called a mask layer.

A shadow mask preparing process is adapted from Gulden et al. [2] by using $H_3PO_4$ : $H_2O_2$ : $H_2O = 1 : 1 : 10$ (by volume) to open the windows on the mask layer instead of $NH_4OH : H_2O_2 : H_2O$ solution for reducing shadow effect due to the space between an edge of the mask and the surface during the MBE growing process. Line shaped window patterns of 7 and 5 μm width and 250 μm separation were provided for the experiment. The micrograph cross-sectional view of shadow mask is shown in Fig. 1.

![Cross-sectional view of the shadow mask by an optical microscope.](image)

3. Growth of multiple quantum wells through shadow mask

There were three samples fabricated in this experiment.

- Sample #1: This is a reference sample to test the conditions of MQW growth. The MQW structure was grown on the plain (1 0 0) GaAs substrate.
- Sample #2: The sample which was prepared with 7 μm wide line patterns.
- Sample #3: The sample which was prepared with 5 μm wide line patterns.

A conventional MBE method was used to fabricate series of GaAs/Ga$_{0.8}$Al$_{0.2}$As to form a structure of 10 identical unit quantum wells. There are two different well widths, 60 monolayers and 30 monolayers. Samples #1 and #3 have 60 monolayers well width and sample #2 has 30 monolayers well width. The barrier thickness of all samples are 180 monolayers. Clean surfaces of GaAs mask layers and GaAs substrate under shadow masks is confirmed by RHEED pattern before each MBE growth. Substrate temperature during the growths was 650°C while the substrates were rotated at 4 rpm for uniformity. Figs. 2 and 3 show the cross-sectional views of shadow masked mesa MQW by an optical microscope and a scanning electron microscope, respectively.
Furthermore, after MBE growth process, a part of sample #3 was passed through a strip-off procedure by second reverse photolithography. The mesa-line structures were protected by photoresist after the photolithography process. Shadow masks and GaAs/Ga$_{0.8}$Al$_{0.2}$As quantum well structure grown over the masks were then partially removed with \( \text{NH}_4\text{OH} : \text{H}_2\text{O}_2 : \text{H}_2\text{O} = 3 : 1 : 15 \) (by volume). Fig. 4 shows a cross-sectional view of mesa-MQW after the strip-off to eliminate the shadow masks for effectiveness of MQW signal.

4. Photoluminescence results

The photoluminescence (PL) measurements were conducted by using 50 mW Ar$^{-1}$ laser as the excitation of all samples set up in 10 K cryogenic system. The laser beam was point focussed by a circular lens and was line focussed by a cylindrical lens onto the stripes of MQW. The line focus could be aligned parallel to the stripes (0°) and perpendicular to the stripes (90°) for analysis of the PL results. The focussing dimension was \( \sim 100 \) µm for spot and line. PL signals were collected by a set of collimated lens and passed through a monochromator. Spectral PL data were detected by a photomultiplier and recorded in a computer for analytical works.
Fig. 5 shows PL spectra of shadow masked mesa-MQW stripes with 7 μm width (Sample #2) when the sample was excited by point and line of laser beams both at 0° and 90° configurations. The dash lines in the figures were created by using the Multiple Gauss fitting utility in Origin™ plotting program. Double PL peaks at 758 and 774 nm were observed in all cases. The strongest PL peaks were obtained by point focus due to the most intense excitation laser beam. In case of line focus, at 0° configuration, the sample gave nearly equal intensity of double PL peaks but less strong signals due to shadow mask effect in optical alignment of the PL experimental setup. At 90° configuration, PL peaks become weaker due to the smaller effective area. The separation of these double PL peaks (about 16 nm) could be explained by the variation of narrow quantum well widths between MQW on shadow mask and mesa MQW under shadow mask. The quantized states of electrons in quantum wells are defined by well thickness [4].

Fig. 6 shows PL results from sample #3. An objective for preparing this sample is to provide a tunability of the quantized state in the shadow masked MQW. Overlapping of PL peaks at 810 and 818 nm were observed and referred to the previous experiment [5]. Small variations occur due to broader quantum well width. The strongest PL peaks were again observed by point focus due to the most intense excitation of laser beam. However, line focus at 0° configuration gave lowest PL peaks due to narrower window and this made PL alignment more difficult. When the sample was rotated to 90° configuration, PL alignment became easier and better PL signals were detected.

To ease the PL alignment, shadow masks of sample #3 were partially stripped-off and were studied under the same conditions. The only strongest PL peak was always at 818 nm which corresponded to the mesa-MQW stripe while the PL peak at 810 nm was comparatively much smaller. The best configuration for the stripped-off sample
5. Conclusions

The 10 MQW mesa structures were grown through shadow masks onto the GaAs substrates by MBE technology. The structures were self-organized by the mesa shapes grown. PL spectra indicate that the mesa structures consist of multiple quantum wells as in the previous study. PL peaks from the MQW can be tuned by varying the well layer thickness. The sharpness of PL peaks reflects the mesa crystal quality. The experimental results showed that the shadow mask technique was suitable for future patternized MQW device fabrications.

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References

Radiative decay in type-II GaP/AlP/GaP quantum wells

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Abstract

We have studied time-resolved photoluminescence (PL) of type-II GaP/AlP/GaP quantum wells with different well widths and acquired detailed information concerning the mechanisms of the radiative recombination. At low temperatures, the PL spectra consist mainly of a no-phonon line. The decay of the no-phonon line was slow and nonexponential. The decay curves could be fitted to a model which assumes that the radiative recombination occurs as the result of both incoherent and coherent Γ-X scattering due to disorder at the interface. The fit revealed that the dominant Γ-X mixing mechanism was random scattering caused by fluctuations in the potential at the interface. Based on simple perturbation theory, the observed increase in the radiative decay rate with decreasing well width was explained by the increase of overlap of the electron envelope wave functions as well as the decrease of Γ-X energy separation. The model suggests that the magnitude of the potential responsible for the Γ-X mixing is considerably greater than those observed in the type II GaAs/AlAs heterostructures.

Keywords: Time-resolved photoluminescence; Type-II quantum wells; Localized indirect excitons; Γ-X mixing

1. Introduction

Lattice matched GaP and AlP are both indirect bandgap compound semiconductors with conduction band minima at the X points where the band lineup is type-II at the AlP/GaP heterointerface [1]. The schematic band alignment of a GaP/AlP/GaP quantum well (QW) is shown in Fig. 1. In such type-II (staggered) band lineups, the AlP well layer only confines the electrons at the X minima; the holes lie on the band edge at the zone center in the GaP layers. Thus in the type-II configuration, electrons and holes are separated both in real and in k-space, and one can expect no oscillator strength for the optical transition if there are no momentum mixing mechanisms. In the type-II GaAs/AlAs heterostructures, however, there have been a number of studies showing the presence of a finite oscillator strength caused by Γ-X mixing effects [2–14]. The Γ-X mixing can arise purely from a potential step at the heterointerface without any zone folding effects associated with the periodicity of the superlattice [9].

This paper reports on our study of radiative decay in type-II GaP/AlP/GaP QWs with different well widths. Highly efficient continuous-wave (cw)
photoluminescence (PL) is obtained at low temperatures from GaP/AIP/GaP QW structures with simple type-II configurations [15]. Time-resolved PL has been employed to clarify the recombination mechanism. A slow and nonexponential decay of the luminescence was explained by the emission of the localized indirect excitons at the interface. We analyzed the decay data based on simple perturbation theory.

2. Experimental procedure

The GaP/AIP/GaP multiple QW (MQW) structures were grown on GaP (0 0 1) substrates by gas source MBE (VG Semicon V80H) using PH₃ and elemental Ga and Al. The MQW structures consist of 5 periods of AIP layers separated from each other by 30-nm thick GaP layers. The AIP layer thicknesses were varied with the samples (1, 2, 3, 5, and 8 monolayers (ML)). After thermal cleaning of the surface and subsequent growth of a 250-nm GaP buffer layer, the MQW structures were grown at 620°C. The growth rates of AIP and GaP were 0.25 and 0.5 ML/s, respectively. During the growth of the GaP and AIP layers, reflection high energy electron diffraction (RHEED) showed intense and sharp spots on the Laue rings and long lasting intensity oscillations in specular beams, indicating the progression of layer-by-layer growth [16]. Especially for the AIP layers, very precise ML growth control was performed by means of RHEED intensity oscillations so as to reduce the effect of the fluctuations of the Al molecular beams just after opening of the effusion cell shutter.

Cw PL spectra were excited with the 325 nm line from a He–Cd laser and detected via a spectrometer by a photomultiplier using lock-in techniques. The excitation power density was 0.3 W/cm². For time-resolved measurement, the excitation source was the second harmonic light of a Ti:Sapphire laser, pumped by a frequency-doubled Q-switched Nd : YAG laser. The repetition rate, pulse width and pulse energy of the excitation pulse were 3 kHz, 100 ns and 0.3 μJ, respectively. The luminescence was detected by a gated photon counter. The temporal resolution of the detector was ~5 ns.

3. Results and discussion

Fig. 2 shows the cw PL spectrum from the 2 ML AIP wells measured at 4.2 K. This spectrum is typical of the samples studied, consisting of a strong no-phonon line and a weak phonon sideband.
50 meV lower in energy, very close to the $\Gamma$-point LO phonon energy of GaP. The no-phonon line is derived from the transition between the confined AlP $X_2$ and the GaP $\Gamma$ state [15]. The presence of the strong no-phonon line and the GaP LO ($\Gamma$) phonon satellite suggests that there are significant momentum mixing mechanisms between the $X$ and $\Gamma$ electron states, which result in a finite oscillator strength. The possible sources are coherent and incoherent scatterings by the heterointerface. Coherent scattering is expected to occur due to breakdown of symmetry at the interfaces by the presence of ultra-thin AlP layers in the GaP bulk crystalline structure. Incoherent scattering is induced by roughness at the interface, which gives rise to random potential fluctuations.

The decay after pulsed excitation is slow and nonexponential, as shown for 1 ML AlP wells in Fig. 3. This temporal behavior can be explained by a model which assumes $\Gamma$–$X$ mixing mechanisms for localized indirect excitons by disorder at the interface [17, 18, 6]. According to the model, for localized indirect excitons at the zone boundary, whose decay is permitted by scattering to $\Gamma$ by a random potential, the time decay of the no-phonon line is given by

$$I(t)/I(0) = (1 + 2w_r t)^{-3/2},$$

where $w_r$ is the average radiative decay rate due to random potential scattering. In addition to random (incoherent) scattering, there is a non-random (coherent) contribution $w_n$ to the decay rate, Eq. (1) is modified [6] as

$$I(t)/I(0) = e^{-w_r t}[1 + w_n t/(1 + w_n w_r)](1 + 2w_r t)^{-3/2}.\tag{2}$$

The experimental data fitted the time dependence predicted from this model. The best fit was obtained with $w_r = 4.2 \times 10^5 \text{ s}^{-1}$ and $w_n = 0 \text{ s}^{-1}$ for 1 ML AlP wells, as shown in Fig. 2. The fitted values of $w_r$ and $w_n$ are given in Table 1. The parameter $w_r$ increased with decreasing AlP well width. In all the samples $w_n$ was considerably small and could be zero. The dominant mixing mechanism is therefore considered to be random scattering.

For the no-phonon transitions of indirect excitons, the $k$-selection rule requires there to be mixing between the $X$ and $\Gamma$ states. In the first-order perturbation theory for $\Gamma$–$X$ mixing by a potential $V$ [10], the matrix element for electric dipole transitions between conduction and valence bands is given by

$$M_{c,v} = \frac{\langle \psi_C^{AlP(X)}|V|\psi_C^{GaP(\Gamma)}\rangle \langle \psi_C^{GaP(\Gamma)}|p|\psi_C^{GaP(\Gamma)}\rangle}{\Delta},$$

Table 1

| AIP (ML) | $\Delta$ (eV) | $\langle \phi_C^{AlP(X)}|\phi_C^{GaP(\Gamma)}\rangle$ ($\times 10^{-2}$) | $w_r$ (10$^5$ s$^{-1}$) | $w_n$ (10$^5$ s$^{-1}$) |
|---------|---------------|-------------------------------------------------|-----------------|-----------------|
| 1       | 0.62          | 6.4                                             | 4.2             | < 0.01          |
| 2       | 0.70          | 4.3                                             | 0.38            | < 0.01          |
| 3       | 0.77          | 3.6                                             | 0.33            | < 0.001        |
| 5       | 0.85          | 3.0                                             | 0.28            | < 0.01          |
| 8       | 0.90          | 2.6                                             | 0.12            | < 0.001        |
where \( \psi_C \) and \( \psi_V \) are wave functions at the conduction and valence-band extrema, \( \Delta \) is the \( \Gamma - X \) splitting \( (E_T - E_X) \), and \( p \) is the dipole operator. The mixing is assumed to occur only between the AIP X states and the lowest \( \Gamma \) states in GaP. The oscillator strength for radiative transition is proportional to the square of the matrix element. The mixing factor in Eq. (3) can be transformed as follows [9]:

\[
\langle \psi_C^{\text{AIPX}} | V | \psi_C^{\text{GaP\Gamma}} \rangle = \langle \psi_C^{\text{AIPX}} | V | \psi_C^{\text{GaP\Gamma}} \rangle \langle \phi_C^{\text{AIPX}} | \phi_C^{\text{GaP\Gamma}} \rangle \Delta \%
\]

(4)

where \( u \) is the Bloch part of the wave function and \( \phi \) is the slowly varying envelope wave function. The radiative decay rate is thus given by

\[
w_{X,\Gamma} = w_{\text{f-R}} V_{\text{mix}} \frac{\langle \phi_C^{\text{AIPX}} | \phi_C^{\text{GaP\Gamma}} \rangle^2}{\Delta^2},
\]

(5)

where \( w_{\text{f-R}} \) is \( \langle \psi_C^{\text{GaP\Gamma}} | p | \psi_V^{\text{GaP\Gamma}} \rangle^2 \), and \( V_{\text{mix}} \) is \( \langle \psi_C^{\text{AIPX}} | V | \psi_C^{\text{GaP\Gamma}} \rangle^2 \). Table 1 gives the energy separation between the \( \Gamma \) and X electron states and the overlap integral of the electron envelope wave functions as calculated using a Kronig–Penney model. In spite of the large energy separation between the \( X \) and \( \Gamma \) electron states and the small overlap integral due to the sample structure, we observed finite radiative decay rates in the range of \( 10^8-10^9 \text{ s}^{-1} \). This suggests that the GaP/AIP heterostructure has a considerably large mixing potential \( V_{\text{mix}} \). In the direct bandgap semiconductors, the radiative decay rate is larger than \( 1 \times 10^9 \text{ s}^{-1} \), so we assume a value of \( 10^9-10^{10} \text{ s}^{-1} \) for \( w_{\text{f-R}} \). The resulting \( V_{\text{mix}} \) from Eq. (5) is of the order of \( 10-10^2 \text{ meV} \), which is far greater than those obtained in the type-II GaAs/AlAs system [9, 13].

Fig. 4 shows the radiative decay rate \( w \) as a function of \( \langle \phi_C^{\text{AIPX}} | \phi_C^{\text{GaP\Gamma}} \rangle / \Delta \). The linear relationship seen in Fig. 4 shows that the model can well explain the dependence of \( w \), on AIP well width. The increase in \( w \) with decreasing AIP well width results from an increase in the overlap integral of the electron envelope functions as well as a decrease in the energy splitting \( \Delta \). The mixing potential \( V_{\text{mix}} \) is almost completely independent of AIP well width.

One might expect a coherent contribution to the radiative decay rate due to the abrupt potential step along the growth direction at the heterointerface. The absence of such a coherent contribution is attributed to the small lateral scale of roughness at the interface. In the samples, the lateral scale of roughness is far smaller than the lateral size of excitons and the effective interface roughness is much less than one atomic layer [15]. In GaAs/AlAs QWs, the lateral size of roughness on AlAs-on-GaAs interfaces is close to or larger than exciton diameter, whereas on GaAs-on-AlAs interfaces, it is so small as compared with exciton diameter that they act as a smooth interface [19]. This is due to a far smaller diffusion length of the Al atoms along the AlAs surface in contrast to that of the Ga atoms along the GaAs surface. In the GaP/AIP system which is being grown in this study, both of the AIP-on-GaP and GaP-on-AIP interfaces have such a “pseudo-smooth” nature. Diffusion of both the Ga atoms along the GaP surface and the Al atoms along the AIP surface is expected to be very small. Since this pseudosmooth interface has a stochastic variation in the distribution of microsteps, excitons localized in different regions would experience different effective well widths. Thus the pseudo-smooth interface would act as an effective and random scattering source due to its abruptness and random fluctuations of average local potential.
4. Conclusions

We investigated radiative decay in type-II GaP/AlP/GaP quantum wells with thickness of 1, 2, 3, 5, and 8 ML. The no-phonon line at 2 K showed a slow and nonexponential decay. This temporal behavior was interpreted as the emission from the localized indirect excitons scattered by random fluctuations in potential at the GaP/AlP interface. The average random radiative decay rate increased with decreasing well width, which was interpreted to be the result of the increase of overlap between the X and Γ electron envelope wave functions as well as the decrease of Γ−X energy splitting. The simple perturbation theory suggested that the Γ−X mixing potential at the GaP/AlP heterointerface was far larger than that observed in the type-II GaAs/AlAs system.

References

Temperature-insensitive photoluminescence above 300 K in strained Ga\textsubscript{x}In\textsubscript{1-x} As multiple quantum wire heterostructures

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Abstract

Stable peak wavelength photoluminescence (PL) spectra has been observed in Ga\textsubscript{x}In\textsubscript{1-x} As strained multiple quantum wire (MQWR) heterostructures over a temperature range between 250 and 380 K. The MQWR samples were grown in situ by molecular beam epitaxy using the strain-induced lateral-layer ordering (SILO) process to create the quantum wires. The formation of the strained Ga\textsubscript{x}In\textsubscript{1-x} As quantum wires and the amount of the strain associated with the wires depends on the growth conditions and structural configuration of the MQWR active region. Samples grown near a substrate temperature of 500°C show a negligible net PL peak wavelength shift between 77 and 380 K. Red shifts and blue shifts of PL peak wavelengths between 77 and 380 K are observed in samples grown at temperatures less than and greater than 500°C, respectively. Due to the multi-axial strain fields induced by the SILO process in the MQWR active regions, the PL peak wavelength of all the MQWR samples stabilize at $\sim 1.61$ μm for temperatures above 250 K. The temperature dependent shift of peak PL wavelength between room temperature and 72°C for these MQWR heterostructures is as low as 0.2 K/°C.

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Keywords: QWR; PL; PPL; Multi-axial strain; MBE; SILO

1. Introduction

The well documented improvements in performance of quantum well (QW) lasers over double heterostructure lasers serve as motivation to develop quantum wire (QWR) lasers in the hope of attaining further improvements. The superior characteristics of QW devices such as lower threshold currents, decreased temperature sensitivity, higher differential gain, and a narrower gain spectrum are mainly due to the localized step-like nature of the density of states (DOS) function for QWs [1]. Since the DOS function for QWRs is even more localized, QWR lasers are expected to out perform their QW
counterparts. However, due to limitations in current QWR fabrication technology it is very difficult to physically realize nanometer-size wires with uniform cross-sections. As a consequence of wire imperfection, the DOS function is QW-like making the theoretically predicted advantages of QWRs not noticeable [2]. Nevertheless, many interesting properties have already been observed in these non-ideal QWR device structures [3–5].

Recently, we discovered a unique temperature insensitive photoluminescence (PL) behavior in strained Ga_{1–x}In_xAs/InP multiple quantum wire (MQWR) heterostructures prepared by the strain induced lateral-layer ordering (SILO) process [6]. The PL peak wavelength would either not shift between 77 K and room temperature or would blue shift with increasing temperature depending on the MQWR layer structure. Furthermore, the peak PL wavelength converges to about 1.61 µm at 300 K for every sample. In order to utilize such strained MQWR structures in high speed fiber optical communication systems, the PL peak wavelength needs to be tuned to 1.55 µm and maintained at that value for temperatures greater than 300 K. It is plausible to meet these requirements based on the results presented in Ref. [6] since the PL wavelength shift was found to be a function of the strain distribution within the MQWR active region where the multi-axial strain itself is controlled by the layer structure in the active region and the growth conditions [6, 7]. Therefore, to further pursue strained Ga_{1–x}In_xAs MQWR structures as stable emitters for high speed fiber optical communication systems we have investigated the multi-axial strain which exists in the MQWR active regions. Specifically, in this study, we report on the growth temperature effect on the PL peak wavelength shift as well as the high temperature (∼100°C) PL behavior in strained Ga_{1–x}In_xAs MQWR structures.

2. Experimental procedure

Strained Ga_{1–x}In_xAs MQWR heterostructures have been grown in situ on (0 0 1) on-axis InP substrates by MBE using the SILO process. The details of the growth conditions have been described in a previous publication [7]. To have precise control of the layer composition, the Ga, In, and Al fluxes were calibrated using reflection high energy electron diffraction intensity oscillations. Different growth temperatures were examined as well as different structural configurations to see what effect they had on wire formation and the corresponding PL spectra. Typical structures examined here contain a 1200 Å Al_{0.48}In_{0.52}As buffer, and a 1200 Å Al_{0.25}Ga_{0.24}In_{0.52}As cladding region, followed by the strained Ga_{1–x}In_xAs MQWR active region, another 1200 Å Al_{0.24}Ga_{0.24}In_{0.52}As cladding region, and a 500 Å Al_{0.48}In_{0.52}As cap. The active region consists of one to five 100 Å conventional QWs separated by 70–150 Å Al_{0.24}Ga_{0.24}In_{0.52}As barriers. A schematic of a typical MQWR structure with 5 QWs is shown in Fig. 1. Each QW contains 8 pairs of (GaAs)$_m$/(InAs)$_n$ short period superlattice (SPS) grown at a fixed ratio of $m:n \approx 2:2.2$ where $m$ and $n$ are the number of monolayers of GaAs and InAs, respectively. Within these QWs a composition modulation perpendicular to the growth direction is induced by the strain associated with the small non-zero values of mismatch $\Delta T = \Delta T(m, n, a_{GaAs}, a_{InAs})$ between the SPS layers and the substrate, and global average strain $\Delta S = \Delta S(m, n, a_{GaAs}, a_{InAs})$ within the SPS [3, 8]. The quantities $a_{GaAs}$ and $a_{InAs}$ correspond to the

![Fig. 1. The schematic diagram of a typical Ga_{1–x}In_xAs strained MQWR heterostructure used in this study.](image-url)
lattice constants of InP, GaAs, and InAs, respectively. Since the lateral composition modulation is formed within conventional QWs in the [1 0 0] direction, two-dimensional quantum confinement is attained resulting in QWRs.

To investigate the effect of strain on the MQWR region and the resulting PL spectra, samples with differing amounts of strain were studied by preparing samples at different substrate temperatures $T_s$ using the desorption temperature of InP, $\sim 540^\circ$C, as a reference. For comparison, a multiple quantum well (MQW) sample with latticed matched GaInAs QWs was prepared.

The luminescence properties of the strained MQWR samples were studied using standard PL measurement techniques. The samples were mounted in a liquid nitrogen cooled dewar for measurements between 77 and 300 K. To measure the PL spectrum of the sample for temperatures greater than 300 K, a hot plate was used to heat the sample and a thermocouple monitored the temperature of the crystal. The heterostructures were excited using the 5145 Å line of an Ar$^+$ laser with an excitation intensity of approximately 100 W/cm$^2$. The collected light was dispersed in a 0.5 m focal length grating spectrometer and detected with a liquid nitrogen cooled Ge detector using the lock-in technique. To examine the wire formation of the heterostructures, polarized photoluminescence (PPL) spectra were taken. This was done using the same setup as for PL measurements but with a polarization analyzer located in front of the spectrometer and orienting it either parallel or perpendicular to the [1 1 0] direction (e.g., the QWR axis) of the sample [3, 7]. A depolarizer is placed in front of the spectrometer to remove any possible grating polarizations.

3. Results and discussion

The existence of the lateral composition modulation in the strained Ga$_x$In$_{1-x}$As MQWR structures was examined by cross-sectional transmission electron microscopy (TEM). The results indicate that a composition modulation consisting of alternating larger band gap Ga-rich regions and smaller band gap In-rich regions on the order of 100 Å has been formed [7, 9]. In addition to TEM, confirmation of wire formation is provided by the observation of PPL spectra. It is expected that the PL emitted from the [1 0 0] direction will appear to be polarized when the polarization analyzer located in front of the spectrometer is oriented parallel and perpendicular to the wires. The polarization effect reflects the existence of QWRs formed by the SILO process and the strain fields associated with the lateral composition modulation. The magnitude of polarization is proportional to the degree of the lateral composition modulation [3]. Fig. 2 shows the PPL spectra of a strained Ga$_x$In$_{1-x}$As MQWR heterostructure at 77 K. The peak intensities at 1.61 μm are highly polarized whereas the PL from the AlGaInAs cladding and barrier layers display the usual small preferential polarization for bulk crystals with a zinc-blende type cubic symmetry [10]. It is thus concluded that the peak PL wavelength at 1.61 μm emanates from the MQWR active region within the sample.

In a previous study, the influence of QWRs on the PL behavior of strained Ga$_x$In$_{1-x}$As MQWR heterostructures measured at temperatures between 77 and 300 K was reported [6]. It was found that the temperature dependence of the peak PL wavelength for samples with 2 QWs in the MQWR region differed from that of samples with 5 QWs.
This structural difference manifests itself as differing strain fields in the MQWR active region. Since the strain induced by the SILO process is a cumulative effect [7], the highly strained structures with 5 QWs will have a stronger lateral composition modulation than those with 2 QWs. This is supported by the fact that the PL emitted from the 5 QW sample has a stronger polarization than that observed from the 2 QW sample. The samples with 2 QWs show no net shift in peak PL wavelength for temperature ranging from 77 to 300 K, whereas the 5 QW sample has a net blue shift of peak PL wavelength with increasing temperature up until 300 K. This effect was attributed to the multi-axial strain distribution which exists in the MQWR active region [6].

To further examine this effect, the PL behavior of strained Ga$_{x}$In$_{1-x}$As MQWR heterostructures with 5 QWs grown under different substrate temperatures $T_s$ is examined in this study. The PL spectrum of these samples was observed to be polarized indicating QWR formation in each sample. In Fig. 3 the resulting trend in temperature dependence of peak PL wavelength shift between 77 and 300 K for various values of $T_s$ is given. It has been shown that the lateral composition modulation is optimized in Ga$_{x}$In$_{1-x}$As MQWRs grown at a temperature above and near 500°C [7]. At growth temperatures lower than 500°C, a less regular weaker lateral composition modulation occurs resulting in irregular QWRs, whereas samples grown above and near 500°C have a stronger composition modulation which allows for the formation of regular arrays of QWRs. Thus, Fig. 3 indicates that strained Ga$_{x}$In$_{1-x}$As MQWR samples with a stronger lateral composition modulation have a larger blue shift with respect to increasing cryostat temperatures. However, for samples grown at lower values of $T_s$, the magnitude of the blue shift decreases until the shift of peak PL wavelengths approaches the expected red shift value of approximately 1000 Å for the lattice matched GaInAs MQW control sample.

The high temperature properties of the strained Ga$_{x}$In$_{1-x}$As MQWR samples were studied using PL spectroscopy. Fig. 4 compares the temperature dependence of peak PL wavelengths of strained MQWR heterostructures and the MQW control sample from 77 to 380 K. Two types of strained MQWR samples are shown here, those that are strongly strained and those that are moderately strained. The strongly strained samples correspond to samples with five conventional QWs grown above and near 500°C. Moderately strained samples refer to samples with 2 QWs or samples with 5 QWs prepared at temperatures less than 500°C.
Two very interesting features can be seen in Fig. 4. First of all, the overall shift in wavelength of the strained MQWR samples is less than the MQW control sample. In fact, the moderately strained MQWR samples tend to have no net effective temperature dependence where the strongly strained MQWR samples have a net blue shift with increasing temperature. Both of these trends are in stark contrast to the expected net red shift of 1000 Å depicted by the MQW control sample. Most importantly, the peak PL wavelengths of both strained MQWR heterostructures are relatively stable at 1.61 µm for temperatures between 250 and 380 K. In the temperature range of room temperature to 72°C, the wavelength shifts with a dependence as low as 0.2 Å / °C. This thermal induced wavelength drift is only 4% of that in conventional MQWs (~5 Å / °C). Furthermore, we conclude that the stable peaks originate from the MQWR region since the PL at 1.61 µm is coincident with the polarized peaks seen in Fig. 2.

The samples studied in Fig. 4 correspond to samples grown under different substrate temperatures and to samples with different structural configurations. Thus, it is clear that by varying $T_s$ and by making structural modifications, the strain in the Ga$_x$In$_{1-x}$As MQWR active region can be manipulated. The resultant effect is apparent in the temperature behavior of the peak PL wavelength where, most importantly, it is stable above room temperature.

4. Conclusions

It has been shown that the peak PL wavelength of strained Ga$_x$In$_{1-x}$As MQWR heterostructures is stable for temperatures ranging from 250 to 380K. It has a temperature dependence as low as 0.2 Å / °C between room temperature and 72°C. This is over a 25 time improvement over the conventional MQW structures. The PL behavior of these samples between 77 and 300 K varies depending on the amount of strain induced in the MQWR active region by the SILO process. The strain in the MQWR regions, which directly affects the PL peak wavelength shift, can be controlled by the growth temperature and structural configuration. Between 250 and 380 K, all samples show a nearly stable PL peak wavelength near 1.61 µm. The fact that the wavelength is stable above 300 K is of paramount importance to the continued pursuit of QWR technology since QWRs with a stable emission wavelength could be used as the active region of lasers, thus, improving the temperature characteristics of emitters for communication and information systems.

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References

Growth-mode-induced surface morphology and its relation to optical properties of GaAs single quantum wells

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Abstract
Quantum wells with short period superlattice barriers are grown by molecular beam epitaxy in growth modes which range from two-dimensional island nucleation to step flow. These different growth modes result in different surface morphologies as imaged by atomic force microscopy. Micro-photoluminescence of the quantum wells and of their barriers reveal spectra with a rich fine structure. These narrow lines in the micro-photoluminescence spectra are attributed to exciton localization by potential fluctuations at the interface. The formation of a pronounced roughening of step edges for samples grown either at high As4-partial pressure or on misoriented substrates with As-terminated steps is observed, which is accompanied by the appearance of a pronounced fine structure in photoluminescence and a broadening of its distribution.

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Keywords: Superlattices; Quantum wells; Molecular beam epitaxy; AlAs/GaAs

1. Introduction
The electrical transport and the optical properties of novel devices with reduced dimensionality are sensitively influenced by the defect structure of the interfaces including the geometrical and the chemical profile across the interfaces [1] because the magnitude of these defects is on a length scale comparable to, e.g., the extension of the confined wave function of electrons. However, it is not clear which kind of imperfections govern the local physical properties—defects on an atomic scale as revealed by scanning tunneling microscopy [2] or those on a submicron scale already visualized by atomic force microscopy [3]. For a better understanding of these phenomena, a detailed characterization with high spatial resolution is required. Spatially resolved photoluminescence studies of thin quantum wells (QW) have revealed that the radiation emitted from localized areas [4, 5] reflects potential fluctuations due to well thickness inhomogeneities and compositional transients at

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interfaces. Furthermore, the interfaces of a quantum well (normal and inverted) are inequivalent as manifested by, e.g., a unidirectional segregation [6].

In this paper we present results of micro-photoluminescence (μ-PL) and atomic force microscopy (AFM) investigations of a series of QWs. They were prepared under extremely different growth conditions in order to examine the influence of growth-mode-induced morphologies on the optical properties of these structures.

2. Experimental procedure

GaAs single quantum wells with a thickness of 16 monolayers (ML) embedded in short period superlattice (SPSL) barriers consisting of 8 ML GaAs/4 ML AlAs are grown by solid-source molecular beam epitaxy. The temperatures and As$_4$-to-Ga beam-equivalent pressure (BEP) ratios are varied in a range from 550°C to 640°C and 8 to 30, respectively. Almost exactly oriented GaAs(0 0 1) substrates and those with misorientations up to 2° towards (1 1 1) Ga and 2° towards (1 1 1) As are used. The upper barrier is capped with GaAs of the same thickness as the QW. For growth rates of about 300 nm/h the above-mentioned growth conditions allow the adjustment of the (2 x 4) or (3 x 1) surface reconstructions and growth modes ranging from two-dimensional island nucleation to step flow. Growth is interrupted for 40 s after the deposition of the 16 ML QW. The samples are cooled down at a rate of 20°C/min in an As$_4$ atmosphere, thereby maintaining the (2 x 4) surface reconstruction. AFM investigations are performed in ambient in the constant force contact mode. Low-temperature μ-PL studies (8 K) are performed with a confocal imaging system using a microscope objective lens together with a fixed diaphragm. The area of excitation and detection is confined to about 2 μm$^2$.

3. Results and discussions

The use of an air AFM for the study of surface morphologies in the GaAs/AlAs material system is restricted to the steady-state GaAs surfaces. For comparability of surface and interface morphologies we ensured that under identical growth conditions the buffer layer, the QW and the cap layer surface show identical terrace width distribution and step edge profiles which remain unchanged within a few minutes, the time needed to quench the surface. Fig. 1 displays AFM micrographs of homoepitaxial GaAs layers annealed under As$_4$ atmosphere at the growth temperature ($T_g = 610°C$). A well-developed step array corresponding to the 0.2° misorientation is maintained after 2 min annealing (Fig. 1a) which is nearly identical to that obtained for unannealed samples (not shown). After 5 min of annealing the surface starts to change by the onset of increased step bunching (Fig. 1b)

Fig. 1. AFM micrographs (3 μm x 3 μm) of homoepitaxial layers grown on 0.2° misoriented (0 0 1) GaAs after annealing for 2 min (a), 5 min (b) and 20 min (c) at $T_g = 610°C$ in As$_4$ ambient. The initially well-ordered monolayer step system is progressively disturbed by increased step bunching.
which is initiated predominantly at multikinks in the step edges. Prolonged annealing totally changes the initial appearance of the surface (Fig. 1c) by the formation of very narrow terraces or even multi-layer steps [7] with abruptly changing step edge directions.

Fig. 2 shows AFM micrographs of the cap layer surface of QW structures grown in the two-dimensional island nucleation mode (Fig. 2a), predominantly in the step flow mode (Fig. 2b), in the pure step flow mode (Fig. 2c) and in a modified step flow mode (incomplete condensation at high growth temperature) (Fig. 2d). The surfaces display a (2×4)x (Fig. 2a and Fig. 2b) or a weak (3×1) (Fig. 2c and Fig. 2d) reconstruction during growth.

Details of growth parameters are listed in Table 1. Caused by an unintentional miscut of the substrates and due to the distinct growth conditions, the step systems exhibit very different terrace and step edge profiles. High arsenic partial pressures \( p_{\text{As}} \) and high substrate temperature \( T_s \) lead to a pronounced step edge roughening by the formation of larger Ga- and As-terminated step segments (Fig. 2b). Thus, the total length of the step edge segments between a fixed distance is maximized compared to the case of reduced arsenic flux which results in less roughened step edges (Fig. 2c). At \( T_s = 640^\circ\text{C} \), the surface of the layer contains a high number of holes and islands on the terraces (Fig. 2d) indicating a collapse of the pure step flow

![AFM micrographs](image)

**Fig. 2.** AFM micrographs (2 μm × 2 μm) of the cap layer surfaces of QW structures prepared under different growth conditions. \( T_s \) and \( \text{As}_{\text{ex/Ga}}\)-BEP ratios were 550°C/10 (#1) (a), 610°C/30 (#3) (b), 610°C/10 (#4) (c) and 640°C/10 (#6) (d). Substrates are unintentionally misoriented by about 0.03°–0.05° towards (0 1 1). Growth is performed in the two-dimensional island nucleation mode (a), predominantly in the step flow mode (b), pure step flow mode (c) and step flow mode with incomplete condensation (d).

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* Denotes predominantly step flow mode.

b Step flow mode at incomplete condensation, 2DN the two-dimensional island nucleation mode.
Fig. 3. Micro-PL spectra of the QW emission (1.66–1.68 eV) and SPSL emission (1.85–1.885 eV) of samples grown on different misoriented substrates (2° B, 2° A, 0°) denote miscut towards (1 1 1)As (#8), (1 1 1)Ga (#7) and no intentional miscut (#4) (a), with the (2 x 4) surface reconstruction at $T_s = 550^\circ$C in the two-dimensional island nucleation mode (#1) and $T_s = 610^\circ$C in the step flow mode (#3) (b) and in the step flow mode at $T_s = 610^\circ$C with different surface reconstructions (#3, 4) (c).
mode due to an incomplete GaAs condensation. At lower \( T_s \), layers grow in the 2D-nucleation mode (Fig. 2a).

The optical characterization of the layer system is performed by \( \mu \)-PL. The spectra of all QW and all SPSL emission show a rich fine structure and the envelopes of which correspond to the conventional PL spectra with full-width at half-maximum (FWHM) values of 6–8 and 8–14 meV, respectively, (Fig. 3a and Fig. 3c). The spikes in the \( \mu \)-PL spectra are very narrow (FWHM \( \leq 0.1 \) meV).

Fig. 3a shows \( \mu \)-PL spectra from the QW and the SPSL of samples all grown in the step flow mode at the same \( T_s \) but on differently misoriented substrates. The reconstruction during growth for larger miscuts is \( (1 \times 1) \), whereas for small miscuts an asymmetrical \( (3 \times 1) \) is maintained. For misorientations between 0° and 2° towards \( (1 1 1) \) Ga, no quantitative variation in the PL spectra is observed while misorientation towards \( (1 1 1) \) As results in a broadening of the spike distribution. In Fig. 3b we compare PL spectra of two samples grown both with a \( (2 \times 4) \) surface reconstruction but at different \( T_s \) and \( p_{As} \). Obviously, with increasing arsenic partial pressure the fine structures in the QW and SPSL spectra become more pronounced. The SPSL emission is asymmetrical for the sample grown at lower \( T_s \) and narrower compared to the one grown under high \( p_{As} \). This asymmetry reflects exciton transfer into local potential minima at the interface defects. Fig. 3c shows PL spectra of two samples grown in the same growth mode and at the same \( T_s \) but with different reconstructions for different \( p_{As} \). Again we observe a remarkable broadening of the SPSL emission in the case of excessive As incidence.

Growth mode, growth rate, \( T_s \), \( p_{As} \), and surface reconstruction are interrelated, and thus, they cannot be adjusted independently. Even when taking into account this difficulty, we neither recognize a systematic influence of the growth mode nor the reconstruction on the PL properties (Fig. 3b and Fig. 3c). However, a high \( p_{As} \) leads to a remarkable broadening of the PL spectra. The same broadening is observed for samples grown on vicinal substrates with As-terminated steps. By comparing this finding with AFM images (Fig. 2b) we conclude that the total length of step edge segments is maximized for the above mentioned growth conditions which seems to assist the formation of localization sites. The generally observed undulation of the surface or interface in the 1 \( \mu \)m range [2] is not responsible for the occurrence of numerous spikes because of the limited probe area (1.5 \( \mu \)m in diameter).

The spikes in the PL spectra are attributed to the localization of excitons at potential fluctuations due to topological defects at step edges or compositional gradients across the interfaces. From present results it is not justified to correlate exciton localization sites to morphological features. However, it is reasonable to assume that local potential minima for excitons are preferentially formed by intermixing which is mainly caused by Ga segregation at step edges or boundaries of the surface reconstruction domains on the terraces. Therefore, strong step edge roughening favors the formation of localization sites.

4. Summary

A series of GaAs single quantum wells are prepared on almost exactly and on misoriented GaAs \((0 0 1)\) substrates by solid-source MBE with the aim to examine the impact of surface morphologies on the optical properties of the layer system. The samples are grown under controlled conditions giving rise to distinct surface reconstructions and growth modes and studied in combination of AFM and \( \mu \)-PL. For all structures the optical emission from the QW and from the AlAs/GaAs SPSL barriers show a pronounced fine structure in the \( \mu \)-PL spectra which is attributed to exciton localization. Direct correlations to either growth mode or surface reconstruction to PL properties are not found. However, As-rich growth conditions lead to enforced roughening of step edges which is concomitant with a broadening of the PL emission distribution. The latter is also observed when As-terminated misorientations steps are involved. It is supposed that this step edge roughening increases the probability of local intermixing, and thus the formation of additional exciton localization sites.
Acknowledgements

We appreciate the experience and assistance of M. Höricke in MBE growth.

References


Photoluminescence of low-temperature AlGaAs/GaAs multiple quantum wells

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Abstract

The photoluminescence (PL) of low-temperature-grown AlGaAs/GaAs multiple quantum wells (LT-MQWs) has been investigated and compared with a normal-temperature-grown AlGaAs/GaAs MQW structure (NT-MQWs) implanted with protons. The as-grown LT-MQWs show much more intensive PL than the as-implanted NT-MQWs. Upon anneal, the PL from the LT-MQWs and the NT-MQWs exhibits different behaviors. For the NT-MQWs, the PL intensity increases monotonously with rising annealing temperature due to the decrease of implantation-induced defects. By contrast, the PL from the LT-MQWs is drastically quenched, and the PL intensity drops nearly three orders of magnitude after anneal at 600°C. The annealing behavior of the PL from the LT-MQWs is attributed to the formation of As clusters that act as deep trap centers for the photoexcited carriers and cause the quenching of the PL intensity. After anneal, enhanced interface intermixing and roughening have been observed in the LT-MQWs.

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Keywords: Molecular beam epitaxy; Photoluminescence; Multiple quantum wells

1. Introduction

GaAs grown by molecular beam epitaxy (MBE) at low temperatures (LT-GaAs) has generated great scientific and technological interest since the report of Smith et al. [1]. LT-GaAs is highly non-stoichiometric, containing up to 1–2% excess As in the form of As antisite (AsGa), As interstitial and Ga vacancy [2–5]. It exhibits extremely short carrier lifetimes [6, 7], and when annealed, very high resistivity. Upon anneal, the point defect densities in LT-GaAs are dramatically reduced, and the excess As collects in As clusters. Now, there exist two models explaining the electrical properties of LT-GaAs, i.e., the “antisite defect compensation” model and the “buried Schottky barrier” model. The former explains the electrical properties of LT-GaAs by the role of AsGa defects in compensation and hopping conduction [3], while the latter attributes the semi-insulating properties to overlapping depletion regions around As clusters [8].

On the other hand, ion implantation is a well-established technique to introduce traps or recombination centers in semiconductors. Very high
resistivity and subpicosecond carrier lifetimes can also be realized by ion implantation. In this work, we investigated the effect of annealing on the PL of the LT-MQWs and the proton-implanted NT-MQWs. The variations of PL reflect the changes of defect concentrations in samples upon annealing. By comparing the annealing temperature dependence of PL of the LT-MQWs and the proton implanted NT-MQWs, anomalous annealing behavior of PL for the LT-MQWs was found, and explained by the formation of As clusters after annealing.

2. Experimental procedure

The samples were grown on semi-insulating GaAs(100) substrates in a V80H MBE machine with uncracked As$_4$. During growth, the As$_4$/Ga beam equivalent pressure (BEP) ratio was kept at 10. A standard MQW sample consists of 1 μm of GaAs buffer layer, 50 Å of AlAs$_{0.5}$Ga$_{0.7}$As, and 75 periods of Al$_{0.3}$Ga$_{0.7}$As (100 Å)/GaAs (70 Å), as illustrated in Fig. 1. Two MQW samples were grown in this study, one at 310°C (referred to as LT-MQWs) and the other at normal temperature (referred to as NT-MQWs). After growth, the NT-MQWs was implanted with protons at 180 keV to a dosage of $10^{14}$ cm$^{-2}$. Such an implantation will create $10^{15}$ defects per cm$^3$ in the sample. Two samples were then cut into pieces and each piece was subjected to 30 s isochronic annealing at 500–900°C. The PL measurements were made at 77 K. The 5145 Å line of an Ar$^+$ laser was used as an excitation optical source. The PL from the samples was dispersed by a focus length $f = 0.5$ m monochromater and detected by a photomultiplier tube coupled to a lock-in amplifier.

3. Results and discussion

Shown in Fig. 2 are the PL spectra for two MQW samples at several annealing temperatures. The PL peaks are due to the $n = 1$ heavy-hole transitions in the AlGaAs/GaAs MQWs. As is seen, the PL spectra of the LT-MQWs and the NT-MQWs exhibit different dependences on annealing temperature. The integrated PL intensity as a function of annealing temperature is given in Fig. 3. For the NT-MQWs, the PL intensity increases monotonously with increasing annealing temperature,
due to the gradual decrease of concentration of defects created by implantation which act as non-radiative recombination centers for photoexcited carriers. For annealing temperatures above 600°C, the PL intensity becomes saturated, indicating a nearly complete recovery of implantation-induced damage. By contrast, the relatively strong PL for the as-grown LT-MQWs is dramatically quenched after annealing at 600°C. The PL intensity first decreases nearly three orders of magnitude, and then recovers gradually as the annealing temperature is raised above 600°C. At 900°C anneal, the PL intensity of the LT-MQWs becomes comparable with that of the NT-MQWs. There have been many reports that annealing at 600°C or below can greatly reduce point defect densities in LT-GaAs [3, 9]. Since the deep defects can act as nonradiative recombination centers, the reduction in defect densities after annealing should lead to an enhancement in the PL intensity, as is observed for the NT-MQWs. However, we observe an opposite trend in the PL spectra of the LT-MQWs annealed at 600°C and below. This anomalous annealing behavior of PL intensity for the LT-MQWs is believed to be related to the formation of As clusters, and will be discussed later.

The PL peak energy shift versus annealing temperature is shown in Fig. 4. No change in the peak energy is observed for the NT-MQWs with annealing up to 900°C. The PL peak energy of the LT-MQWs also shows no visible change with annealing below 600°C. For annealing above 600°C, the PL peak begins to move toward higher energy. The energy shift reaches 30 meV with annealing at 900°C. The observed energy shift for the LT-MQWs is attributed to Al-Ga interdiffusion at AlGaAs/GaAs interfaces. The Al–Ga intermixing changes square wells into parabolic ones, and raises the subband energy levels in the wells. The large energy shifts indicate that strong Al–Ga interdiffusion occurs at the AlGaAs/GaAs interfaces. The enhancement of intermixing can be ascribed to the high density of V_{Ga} vacancies in the LT-MQWs, which assist the diffusion of atoms on the Ga sub-lattice [10]. The Al–Ga interdiffusion also results in interface roughening that broadens the PL peaks. Fig. 5 shows the full width at half maximum (FWHM) of PL peak as a function of annealing temperature. The increasing PL linewidths indicate that the interfaces are heavily roughened for the LT-MQWs after annealing.

There are two possible explanations for the sharp decrease in the PL intensity for the LT-MQWs annealed at 600°C or below. One is that after annealing, some new point defects and/or defect complexes appear which act as more effective non-radiative recombination centers than those in the as-grown sample. From a detailed study on the annealing-temperature dependence of PL intensity for LT samples grown at various temperatures, we
found that the annealing temperature at which the PL intensity of a LT sample is quenched the most is dependent on its growth temperature. The higher the growth temperature, the higher the annealing temperature required to quench the PL intensity. For example, the annealing temperature corresponding to the minimum PL intensity is 800°C for the samples grown at ~350°C, and 550°C for the samples grown at ~300°C or below. These facts imply that the supposed new defects should appear for annealing temperatures ranging from 550°C to 800°C, i.e., they can withstand annealing at temperatures up to 800°C. However, it is seen in Fig. 2 that the PL intensity increases by about four orders of magnitude from 600°C to 800°C. Such an increase in PL intensity indicates that the new defects, if they exist, cannot withstand annealing at temperatures up to 800°C. Therefore, the quenching of PL intensity cannot be explained by the formation of new point defects (or defect complexes).

The second explanation is evidently related to the formation of As clusters in the annealed LT-MQWs. The As/GaAs interfaces exhibit very high nonradiative recombination rates [6]. The photoexcited carriers are fast trapped by As clusters. Thus, the radiative recombination probabilities and carrier lifetimes are drastically reduced, causing the quenching of PL intensity. With further increase of annealing temperature, the As cluster coarsening process results in a lower cluster density and larger spacing between As clusters. The trapping probabilities for photoexcited carriers at the As clusters decrease. Therefore, the radiative recombination probabilities and thereby the PL intensities increase. The above-mentioned annealing-temperature dependence of the PL intensity on growth temperature can be explained as follows: Under the same As/Ga BEP ratio, less excess As will be incorporated into the LT sample grown at higher temperature, and therefore annealing at higher temperature is needed to form sizable As clusters.

Upon annealing, the line shapes of PL spectra of the LT-MQWs also show different changes than the NT-MQWs. From Fig. 2, it is seen that the PL peaks for the NT-MQWs keep Gaussian line shapes with annealing at 500–900°C. On the other hand, the PL line shapes for the LT-MQWs become strongly asymmetric upon annealing. The higher energy tail gets more prominent with increasing annealing temperature up to 700°C. With further increase of annealing temperature from

![Diagram](image_url)
700°C to 900°C, a Gaussian line shape gradually recovers. The changes in PL line shape have relation with the quenching of excitons in the LT-MQWs, as can be seen from the optical absorption shown in Fig. 6. The excitonic features trail off with increasing annealing temperature, and become fully quenched at 700°C. For annealing temperatures 700–900°C, only the interference fringes can be seen in the absorption spectra. For annealing temperatures below 700°C, PL of the LT-MQWs changes from exciton-dominated to band-to-band-recombination-dominated, resulting in asymmetric PL line shapes. For annealing temperatures higher than 700°C, strong interface intermixing and roughening occur. The increasing statistical broadening in the LT-MQWs makes the band-to-band-recombination line shape become more and more a Gaussian [11].

4. Summary

In summary, the effects of thermal annealing on the optical emission properties of the LT-MQWs and the NT-MQWs have been studied and compared. Upon annealing, the PL of the LT-MQWs and the NT-MQWs show different annealing behaviors. This difference clearly reveals out the role of As clusters in determining the optical properties of the LT-MQWs.

Acknowledgements

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References

Surface chemistry during metalorganic molecular beam epitaxy studied by pulsed molecular beam scattering

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Abstract

The surface chemistry during metalorganic molecular beam epitaxy (MOMBE) is discussed on the basis of the time-of-flight (time-of-arrival) distributions of trimethylgallium (TMG) molecules scattered from structure-controlled GaAs(1 0 0), GaAs(1 1 0) and GaAs(1 1 1)B surfaces as well as oxidized GaAs surfaces. The scattering (adsorption and desorption) of TMG from GaAs surfaces, the initial stage of the surface chemistry, is interpreted within the framework of precursor-mediated chemisorption, where the precursor state is deepened by the charge distribution of the relaxed or reconstructed GaAs surface and the dissociative chemisorption is suppressed on the surface with the highly stabilized structure. The growth suppression on oxidized GaAs surface is due to the absence of the deep precursor states, implying the mechanism of selective area growth. The growth suppression on an As-excess GaAs(1 1 1)B surface, by which the GaAs lateral growth is realized on a grooved GaAs(1 1 1)B substrate, is explained by neutral As trimers which geometrically hinder the TMG trapping in the precursor state. It is also found that the TMG scattering varies with the surface preparation method even if electron diffraction shows similar reconstructions, which suggests that TMG molecules diffuse over the wide area during trapping in the precursor state.

1. Introduction

It is well known that the growth rate of metalorganic molecular beam epitaxy (MOMBE) drastically varies with the growth condition. By utilizing this behavior of MOMBE, we can three-dimensionally control the epitaxial growth. For example, GaAs can be selectively grown on window areas opened on mask materials, such as oxides and nitrides, where MOMBE growth is highly suppressed [1–3]. Atomic layer epitaxy (ALE) is realized by alternately supplying source gases, since the MOMBE growth is limited to one atomic monolayer under single source gas supply [4, 5]. Furthermore, using the behavior that the growth rate is very low on the GaAs(1 1 1)B surface under a high As flux [6, 7], we can laterally grow GaAs on a grooved GaAs(1 1 1)B substrate [8]. However, the mechanism governing the drastically varying
MOMBE growth rate has not been well understood so far.

In order to clarify the mechanism, the growth rate as a function of growth condition has been investigated in detail [9–12]. Several analytical methods such as reflection difference spectroscopy [13], surface photo-absorption measurement [14], grazing incidence X-ray diffraction [15], have been applied. Mass spectrometry of desorbed species enable us to in situ observe the surface chemistry during MOMBE [16–19]. These studies implied the mechanism in ALE. Furthermore, it is demonstrated that the GaAs growth selectivity in selective area growth is attributed to the difference in the decomposition rate of such Ga source gases as trimethylgallium (TMG) and triethylgallium (TEG) from the mass spectrometric analysis of the desorbed species [2]. The decomposition suppression of source gases on the As-rich GaAs(111)B surface was also found by mass spectrometry [6], which is against the expectation that the As-containing species promote TMG decomposition as is known in the vapor-phase reaction [20]. Also on the GaAs(100) surface, we found that the surface reaction of TMG is not influenced by the As density on surface but by the surface structure; TMG decomposition is suppressed on a GaAs(100)-(2×4) surface with a specific As coverage, which has a highly stabilized structure [21].

As mentioned above, great efforts have been done to understand the mechanism of the surface reaction. However, within static behaviors of MOMBE growth, the mechanism of the surface-dependent reaction has not been clear. In order to extract the essential factors in the surface-dependent reaction, the information beyond the static behaviors has been required.

Pulsed molecular beam scattering is a powerful technique to observe the dynamical behavior of the surface reaction, especially of the adsorption and desorption of gases [22]. Recently, we demonstrate that the surface residence time and the energy relaxation of TMG during scattering can be directly evaluated from the time-of-arrival distributions of TMG scattered from well-defined GaAs surfaces and that the surface scattering of TMG is interpreted within the framework of precursor-mediated chemisorption mechanism [23–25].

In this paper, we discuss the surface chemistry during MOMBE on the basis of the results of the pulsed TMG beam scattering from the well-defined GaAs surfaces which carefully prepared by epitaxial growth and subsequent treatments, focusing on the mechanism of selective area growth and lateral growth of GaAs. Furthermore, we examine the TMG scatterings from GaAs surfaces with similar reconstructions which are prepared by different methods.

2. Experimental procedure

The experiments were carried out in a multi-chamber system comprising an MOMBE growth facility, a pulsed molecular beam scattering chamber, and a surface treatment chamber, which have already been described elsewhere [23–25].

We generated pulsed TMG beams using a rotating chopper with two slits of 2 mm in width and a synchronized high-speed pulsed valve with a 0.5 mm nozzle. The pulse shape of incident TMG beams was well reproduced by the velocity distribution of translationally shifted Maxwell–Boltzmann with a translational shift velocity of 457 m/s and a velocity spread of 103 m/s. Using a cryoshrouded quadrupole mass spectrometer, we measured the time-of-arrival (time-of-flight) distributions of TMG molecules scattered from GaAs surfaces prepared in the MOMBE growth facility and the surface treatment chamber.

GaAs surface has variously reconstructed or relaxed structures, depending on the surface stoichiometry as well as the surface orientation. Furthermore, surface properties are known to vary with the surface preparation method even if the reconstructions are similar [26]. In this study, we compared the TMG scatterings from differently prepared GaAs surfaces, containing epitaxially prepared GaAs(100), GaAs(110), and GaAs(111)B surfaces with controlled structures as well as As-desorption-prepared GaAs surfaces. We also examined the scattering from the oxidized GaAs surface, where TMG decomposition is known to be suppressed. The oxidized GaAs surfaces were prepared by oxygen exposure under no light irradiation.
3. Results and discussion

3.1. Scattering of TMG from oxidized GaAs surfaces

First, we examined the scattering of TMG molecules from oxidized GaAs surfaces in order to study the mechanism of selective area growth of GaAs.

The time-of-arrival distributions of TMG molecules scattered from oxidized GaAs surfaces were found to be broader than those expected from the velocity distribution of the incident TMG beams. This broadening is due to the relaxation of translational energy of TMG molecules during scattering. From the analysis of the time-of-arrival distributions, we can evaluate the relaxation of the translational energy of TMG molecules. These time-of-arrival distributions were well reproduced by the convolution of the velocity distributions of incident and scattered TMG, where the velocity distribution of scattered TMG molecules was assumed to be of shifted Maxwell–Boltzmann, while the velocity distribution of incident TMG molecules are directly measured [24].

From the curve-fitting of the time-of-arrival distributions, we extracted two parameters in the velocity distribution of scattered TMG molecules, velocity spread and shift velocity. Fig. 1 shows these parameters as functions of the surface temperature.

If there is no energy relaxation during scattering, the velocity distribution of the scattered TMG molecules would be equal to that of the incident TMG molecules. On the other hand, if TMG scattering occurs under thermal equilibrium at the surface temperature, the velocity spread would be equal to the solid line and the shift velocity would be zero. It is noticed from Fig. 1 that the parameters extracted from time-of-arrival distributions trace the values corresponding to those under thermal equilibrium. According to the mass spectrometric study of desorbed species, TMG molecules do not decompose on this oxidized GaAs surface [27]. Therefore, it is considered that TMG molecules are not decomposed even when TMG molecules obtain thermal energy from surface during scattering under thermal equilibrium. It should be noted that TMG

![Fig. 1. (a) Velocity spread and (b) shift velocity in velocity distribution of TMG scattered from oxidized GaAs surface as functions of the surface temperature. These values were obtained by curve-fitting of time-of-arrival distributions of scattered TMG to the convolution formula of velocity distributions of incident and scattered TMG. The velocity distributions are assumed to be of shifted Maxwell–Boltzmann. If there is no energy relaxation during scattering, the velocity distribution of the scattered TMG molecules would be equal to that of the incident TMG molecules; that is, velocity spread and shift velocity would be 457 and 103 m/s, respectively. On the other hand, if TMG molecules are completely thermalized by the substrate surface, the velocity spread would be equal to the solid line ($\sqrt{2kT}$ s/m) and the shift velocity would be zero.](image-url)
molecules are efficiently decomposed on the GaAs surface at the same temperature. The result suggests that the difference in the TMG decomposition rate is not attributed to the difference in the amount of the energy transfer from GaAs surface to TMG molecules during scattering.

3.2. Scattering of TMG from GaAs surfaces

According to the TMG decomposition rate on GaAs(1 0 0) surface as a function of surface As coverage, the TMG decomposition is anomalously suppressed on a specific GaAs(1 0 0)-(2 × 4) surface (θAs = 0.75), which is reproducibly obtained by a controlled cooling process after GaAs epitaxy [21]. Since the structure of this surface is known to be highly stabilized, we presume that TMG decomposition is suppressed when the structure stability is sufficiently high.

In the case of the TMG scattering from the specific GaAs(1 0 0)-(2 × 4) surface, the time-of-arrival distribution was found to be much broader and cannot be interpreted only by the translational energy relaxation. This distribution was well reproduced by adding a component for scattering without a trapping during scattering for such a long period that we can directly observe within the time resolution of the system we used. Fig. 2 shows the result of the curve-fitting of the time-of-arrival distribution of TMG to the sum of the components for scatterings with and without surface residence. This indicates that there exists a precursor state, where TMG molecules are temporarily trapped in the initial stage of scattering. From the temperature dependence of the reciprocal surface residence time (TMG escape rate from the precursor state to vacuum), the depth of the precursor state is estimated to be 0.85 eV [28]. Although TMG molecules are dominantly trapped in the deep precursor state, almost all the TMG molecules are desorbed without a permanent sticking (the sticking probability is nearly zero).

This behavior was also observed on the GaAs(1 1 0) and GaAs(1 1 1)B-(√19 × √19) surfaces. It has been established that the GaAs(1 1 0) surface is stabilized [29]. From the temperature programmed desorption study, the stability of the GaAs(1 1 1)B-(√19 × √19) structure is considered to be high [25]. Therefore, we conclude that this behavior is common for the scattering of TMG from the highly stabilized GaAs surfaces.

From the temperature dependence of the surface residence time during scattering, the depths of the precursor states for GaAs(1 1 0) and GaAs(1 1 1)B-(√19 × √19) surfaces are estimated to be 0.32 and 0.35 eV, respectively [30].

In order to explain the depth of precursor states, we simply examine the charge distribution of the relaxed or reconstructed GaAs surface structures. The electrons of dangling states in Ga are completely moved to As to maintain a semiconducting surface [31]. Therefore, in the buckled picture of the GaAs(1 1 0) surface [29], the topmost Ga–As pair generates a sufficiently large electrostatic dipole moment outside the crystal, which can polarize TMG molecules coming near to the surface, resulting in the TMG trapping in the deep precursor states observed here [25]. In the case of the GaAs(1 1 1)B-(√19 × √19) surface [32], the electrostatic behavior is considered to be similar to that for the (1 1 0) surface. The deeper precursor state for the GaAs(1 0 0)-(2 × 4) surface is explained by the higher density of the Ga–As pairs in the proposed (2 × 4) structure [33]. Therefore, the surface electrostatic behavior explains well the depth of the precursor state [30].
In the case of the scattering from less stabilized GaAs surfaces such as GaAs(1 0 0)-c(4×4), GaAs(1 0 0)-(1×6), and GaAs(1 1 1)B-(1×1) surfaces, where high TMG decomposition rate is observed [6, 21], the time-of-arrival distributions were observed to have no component for the scattering with a long surface residence, as shown in Fig. 3. The small amount of scattered TMG corresponds to the large amount of TMG dissociatively chemisorbed [25]. We consider that there exists a precursor state also on the GaAs surface with less stability, since the surfaces have As-Ga pairs generating the large electric dipole, and that TMG trapped in the precursor state is efficiently decomposed, providing such a high TMG decomposition rate as about 0.5.

From the comparison of the scatterings from clean GaAs and oxidized GaAs surfaces, it is concluded that the TMG decomposition suppression on the oxide is attributed to the lack of the precursor state [24].

3.3. TMG decomposition suppression by excess As

TMG decomposition is highly suppressed on the As-excess GaAs(1 1 1)B surface with (2×2) reconstruction, resulting in the lateral growth of GaAs on a grooved GaAs(1 1 1)B substrate by controlling As flux [8]. We examined pulsed TMG beam scattering also on the GaAs(1 1 1)B-(2×2) surface. The TMG scattering was found to be very similar to that from the oxidized GaAs surfaces; that is, all the TMG molecules are scattered without trapping in a deep precursor state although the energy relaxation occurs.

This feature is interpreted by the reconstructed surface structure. The GaAs(1 1 1)B-(2×2) surface consists of As-trimers on As-terminated GaAs(1 1 1)B surface [32]. According to the simple estimation of electron distribution similar to that for the above-mentioned GaAs surfaces, the topmost As trimers on As-terminated surface are neutral [30]. The As–Ga pairs in the second and third atomic layers, where electrons are biased, certainly generate large electric dipole moments. However, the space between As trimers where TMG molecules can approach is small compared with the TMG molecular size. We consider that TMG trapping is hindered geometrically by such neutral adatoms as topmost As-trimers on the As surface. This mechanism is consistent with the results on the super-As-rich GaAs(1 0 0)-c(4×4) that TMG sticking probability decreases as the As coverage further increases [25].

3.4. Surface preparation dependence of the TMG scattering

The above-mentioned GaAs(1 0 0)-(2×4) surface, where TMG decomposition is suppressed, was prepared by controlling As flux during the cooling process after GaAs epitaxial growth (referred to as epitaxially prepared surface). A GaAs(1 0 0) surface showing (2×4) reconstruction can also be obtained by another method such as a process containing As desorption from an As-covered GaAs surface (referred to as As-desorption-prepared surface). It has been reported that the work function of the epitaxially prepared GaAs(1 0 0)-(2×4) surface is largely different from that of the As-desorption prepared GaAs(1 0 0)-(2×4) surface [26]. This demonstrates that these surfaces are not the same. Here, we examine the scatterings of pulsed TMG beam from these surfaces.

Fig. 4 shows the temperature-programmed desorption (TPD) spectrum of As$_2$ from an As-rich GaAs(1 0 0)-c(4×4) surface, in which observed reconstructions are also given. The ramp rate is set at 0.1°C/s. It is considered that the bottom of the deep valley at 480°C in Fig. 4 corresponds to the
GaAs(1 0 0) surface with the highest stability among the As-desorption-prepared (2 x 4)-reconstructed surfaces.

For TMG scattering experiments, we prepared GaAs(1 0 0) surfaces by heating the As-rich GaAs(1 0 0)-(4 x 4) surface up to different treatment temperatures (T_c) at a ramp rate of 0.1 °C/s and quenching it. The time-of-arrival distributions of TMG molecules scattered from these surfaces were analyzed by curve-fitting to the sum of the components for scatters with and without surface residence during scattering in the same manner as in the cases of the Fig. 2.

Fig. 5 shows the ratio (R_a) of the amount of TMG molecules in the component for scattering with a long surface residence to the total amount of scattered TMG as a function of T_c. The R_a value was evaluated by integrating the fitted time-of-arrival distribution. The dashed line in Fig. 5 indicates the R_a value for the scattering from the epitaxially prepared GaAs(1 0 0)-(2 x 4) surface (Fig. 2).

We found in Fig. 5 that R_a’s for the As-desorption-prepared surfaces are much smaller than that for the epitaxially prepared surface. This result indicates that TMG surface chemistry varies with the surface preparation even if the reconstructions observed by low-energy electron diffraction (LEED) or reflection high-energy electron diffraction (RHEED) are similar. The component for the scattering with the surface residence corresponds to the TMG molecules which survive dissociative chemisorption in the precursor states for a long period. The dissociative chemisorption on GaAs surface is considered to occur at the surface sites with less stabilized structure (referred to as active sites). Therefore, high R_a indicates a low density of active sites. This result suggests that the density of active sites on As-desorption-prepared surface is higher than that on the epitaxially prepared surface.

As can be seen in Fig. 5, R_a is relatively high when the surface shows (2 x 4) reconstruction. However, we cannot observe a tendency that R_a become larger at the treatment temperature corresponding to the valley bottom in the TPD spectrum. Here the difference in As desorption rate in the TPD spectrum suggests the difference in the stability of the surface. This discrepancy is explained by taking into account the surface diffusion, which enables TMG efficiently to reach to active sites even if the number of the active site is very small. This behavior is consistent with the As coverage dependence of the TMG decomposition rate [21].

The GaAs surface during MOMBE growth is considered to be different from the static surfaces
we examined in this study, even if the surface reconstructions are observed to be similar. During growth, surface atoms are not necessarily in the most stable sites but moving dynamically. Then, the density of active sites is probably higher. Therefore, it is expected that MOMBE growth is not so strongly suppressed by the surface stability. In fact, it was observed that the As-coverage dependence of MOMBE growth rate on GaAs(1 0 0) is not so drastic as that of TMG scattering in this study [10], although the growth rate decrease due to the high surface stability, which is well explained by the mechanism proposed in this study, is clearly observed.

4. Summary

We measured the time-of-arrival distributions of TMG scattered from variously reconstructed or relaxed GaAs(1 0 0), GaAs(1 1 0) and GaAs(1 1 1)B surfaces as well as oxidized GaAs surface. The initial stage of the TMG surface chemistry on GaAs is interpreted within the framework of precursor-mediated chemisorption. We discussed the TMG decomposition suppression in the cases of the selective area growth and the lateral growth of GaAs. The growth suppression on oxidized GaAs surfaces during the selective area epitaxy of GaAs is due to the absence of the deep precursor states, rather than smaller energy that TMG obtains during scattering from substrate surface. On the other hand, the growth suppression on the As-excess GaAs(1 1 1)B surface during the lateral growth of GaAs is originated from neutral As-trimers geometrically hindering TMG trapping. It was also found that TMG scattering was so sensitive that it varies with the surface preparation method even if the diffraction patterns are similar. This suggests that TMG diffuse over wide area during trapping in the precursor states.

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References

Lateral coupling of InP/GaInAsP/InP structures by selective area MOMBE

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Abstract

Lateral couplings of InP/GaInAsP/InP structures selectively grown by metalorganic molecular beam epitaxy (MOMBE) are presented. The heterostructures were grown by either using the hydrides AsH\textsubscript{3} and PH\textsubscript{3} or tertiarybutylphosphine and tertiarybutylarsine as group V precursors in a prototype multiwafer (MOMBE) system. The base heterostructures of the first epitaxy were patterned with SiO\textsubscript{2} mask stripes and trenches were reactive-ion etched. Heterostructures were selectively filled in by a second growth run forming lateral heterojunctions of different quaternary materials. The structures exhibit a bright photoluminescence and a sharp transition at the boundary of the locally grown material indicating a high crystal quality up to the lateral junction with only a minor change in the PL wavelength (< 2 nm). These optimized lateral couplings were applied to laser/waveguide butt-couplings. Optical coupling losses were determined by means of reactive-ion etched waveguides across cascades of coupled GaInAsP layers with an emission wavelength of $\lambda_G \approx 1050$ nm. Values as low as 0.12 dB/coupling were determined in cut-back measurements.

1. Introduction

The realization of photonic integrated circuits (PICS) is a field of growing interest [1–3]. The monolithic integration of devices like lasers, waveguides and photodetectors require some lateral coupling of heterostructures. Lateral couplings can be achieved by overgrowth of patterned semiconductor surfaces [4, 5]. A more attractive pathway for the production of PICS with lateral couplings is the embedded selective-area epitaxy (SAE), due to the planarity of the structures and a higher degree of freedom in the design of the device structures [1, 2, 6].

The true SAE of InP/GaInAsP heterostructures is only achieved in metalorganic growth systems [2, 7]. The lateral couplings of heterostructures by SAE have been realized in metalorganic vapour phase epitaxy (MOVPE) [1, 8–10] and in metalorganic molecular beam epitaxy (MOMBE) [3, 11].

In MOVPE the gas-phase-concentration gradients and diffusion processes lead to a growth-rate enhancement and to a change of the material composition starting at a distance larger than 50 $\mu$m from the masked area. This behaviour is a function of the growth parameters and depends on the mask conditions.
geometry and the aspect ratio of the masked to the unmasked area [2, 12–14].

In MOMBE the growth is determined by the adsorption, diffusion and desorption processes of the molecules on the semiconductor surface only. The variation of the material composition on the (1 0 0) surface at a transition of a growth to a non-growth area is not significantly affected [6] and is rather independent of the aspect ratio [2, 15]. In the case of embedded SAE, the growth starts in a tub at various crystal planes. The growth rate on these side walls has to be optimized for planar infills. The published couplings achieved by selective area MOMBE so far were all grown by using a tilted molecular beam geometry for the group III precursors [3, 11]. Due to shadowing and side wall growth effects there is only a small growth parameter window to achieve a good lateral contact without gaps or planar layers with no growth rate enhancement at the transition area for the use of a tilted beam geometry. Therefore, we have here applied a perpendicular molecular beam geometry for reducing the lateral growth rate [16].

In terms of production, the replacement of hydrides by less dangerous precursors like tertiarybutylarsine (TBAs) and tertiarybutylphosphine (TBP) is of growing interest [17, 18]. High-quality heterostructure couplings are obtained with both classes of precursors. The optimization of the embedded SAE-process yields extremely low-loss waveguide couplings applicable in PICs.

In the first epitaxy an InP/GaInAsP/InP double hetero(DH)structure was grown on a 2 inch InP wafer misoriented 2° towards the nearest (1 1 0) plane. This base structure was covered by different 120 nm-thick SiO2 masks. The mask consists either of SiO2 stripes with a width between 4 and 100 μm and a periodicity of 500 μm or of 15 μm wide SiO2 twin stripes separated by a gap ranging from 1.5 to 5 μm with a periodicity of 250 μm. Trenches up to 2.7 μm in depth were reactively ion etched into the base structure of the first epitaxy. Prior to the second SAE growth run, the samples were etched in H2SO4 : H2O2 : H2O (3 : 1 : 6 in volume ratio) for 5 to 20 s just before introduction into the loading chamber. DH structures were filled in by the second growth run with a typical V/III input ratio of 10, a growth temperature of 495°C and a vertical growth rate of 0.5 μm/h.

We have investigated lateral couplings between InP/GaInAsP/InP base structures with an emission wavelength of λG ≈ 1550 or 1050 nm and a filled in DH structure with λG ≈ 1550 or 1050 nm. All the couplings were oriented in the [0 1 1] or [0 1 1] direction, which is the standard orientation for laser cavities in the production.

3. Results and discussion

The test structure for the integration of lasers and waveguides is the coupling of a DH structure with λG ≈ 1550 nm and the infill DH structure with λG ≈ 1050 nm.

The scanning electron microscopy (SEM) cross-section view in Fig. 1a shows a 5 μm wide selective infill (λG = 1048 nm) in between two 15 μm wide masked DH base structures (λG = 1510 nm). The SEM investigations reveal planar infill-layers and smooth contacts with lateral heterojunctions of the quaternary layer. The selective infill was grown under conditions where the (0 1 1) and (0 1 1) planes are kinetically stable in planar SAE [7, 16, 19]. The perpendicular molecular beam geometry ensures a low lateral growth rate. In the case of a high lateral growth rate, the infilled layers are curved and the vertical growth rate enhancement in the vicinity of the junction facet leads to the formation of the significant “ears” in
the transition area [2]. Anyhow the lateral growth rate has to be high enough to avoid gaps between the infill and the side wall of the masked base structure. The planarity of the SAE layers shown is independent of the aspect ratio (here investigated from 2 to 22%) of the masked to the unmasked surface and of the width of the masked ridges or the width of the etched trenches.

During the growth of the InP buffer layer a vertical InP separation layer was grown at the side wall of the tub (see Fig. 1a). This InP separation layer is adjustable in the thickness in the range from 10%
to 90% of the InP buffer thickness by changing the growth parameters of the buffer layer and the molecular beam geometry. In the micrograph shown in Fig. 1a this lateral thickness is 50%. The thickness of the InP separation layer is reduced for a lower vertical growth rate and a lower growth temperature. For comparison, when using the tilted injection geometry the thickness of the InP separation layer is increased up to a factor of 2.

An efficient butt-coupling of two DH structures requires a constant material composition of the quaternary infill up to the mask edge. We have

Fig. 2. Lateral junctions between selective infill [InP(d = 0.15 μm)/GaInAsP(d = 0.4 μm, λ₀ = 1047 nm)/InP(d = 0.15 μm)] and masked DH structure [InP(d = 0.8 μm)/GaInAsP(d = 0.2 μm, λ₀ = 1540 nm)/InP(d = 0.2 μm)] grown by using TBAs and TBP. (a) SEM cross-section view; (b) Spatially resolved PL measurements at 300 K. Linescans of the PL intensity at the PL wavelength of the maximum and PL wavelength versus the lateral position of the laser spot.
investigated the material quality at the transition area by means of spatially-photoluminescence (μ-PL) measurements at room temperature. The plot in Fig. 1b shows linescans of the PL intensity dependent on the lateral position of the excitation laser spot (spot size 5 μm in diameter) on the (1 0 0) surface of the sample. The SAE quaternary layer exhibits a bright luminescence and a sharp decay at the boundary of the locally grown structure. This decay of the PL intensity at \( \lambda_G = 1048 \text{ nm} \) coincides spatially with the increase at \( \lambda_G = 1510 \text{ nm} \) and vice versa. The measured decay/increase length of 4–5 μm coincides with the spot size of the excitation laser beam. Consequently, a sharp transition boundary is realized. The measured maximum PL wavelength at the lateral heterojunction in the coupling zone is very stable. The small deviation of ±2 nm is in the order of the wavelength error using 300 K PL for the rapidly decreasing intensity at the junction. At a distance of 5 μm, the PL wavelength of the infill is identical to the emission wavelength of a large-area-grown reference sample emphasizing the independence of the aspect ratio of the mask.

Previous investigations by spatially resolved X-ray diffraction of heterostructures selectively grown by MOMBE have shown a change of the lattice mismatch of less than \( 1.0 \times 10^{-4} \) in a transition width of 6 μm. μ-PL measurements at these samples have demonstrated low variations in the PL wavelength too [20, 21]. We conclude from this knowledge and our PL results that the quaternary material composition with a high crystal quality is not affected up to the lateral junction and that the composition of the infill is identical to the reference sample.

In comparison to the sample presented in the SEM micrograph in Fig. 1a, the filled in DH structure (\( \lambda_G = 1047 \text{ nm} \)) in Fig. 2a was grown using TBAs and TBP as group V precursors. This infill shows a slight ear formation. The linescans of the PL intensity in the plot in Fig. 2b exhibits again a sharp transition of the quaternary materials with emission wavelengths of \( \lambda_G = 1047 \text{ (infill)} \) and 1540 nm (base structure). The measured wavelength shift of ±4 nm of the quaternary infill at the junction is slightly higher for the infill grown with TBAs and TBP. This example was grown with a vertical growth rate of 1 μm/h instead of the standard growth rate for SAE of 0.5 μm/h.

Further experiments with the growth rate of 0.5 μm/h have shown a slight growth rate enhancement at the mask boundary too. This overgrowth and wavelength shift is probably caused by different surface diffusion processes, so that an individual optimization for the TBP/TBAs-process is required.

The results of the optimization in the hydride process of the selective infill were applied for laser/waveguide butt-couplings. Fig. 3 presents a cross-section of a laser/waveguide butt-coupling grown with the conventional hydrides. There is a smooth contact between the strained multiquantum-well laser structure with two separate confinement layers (2 SCH MQW) [22] and the waveguide infill [23]. The infill with a height of 2.5 μm shows neither any overgrowth over the mask nor ears at the junctions. During the growth of the entire selective infill the growth rate, the growth temperature and the V/III ratio was kept constant. This convenient growth procedure is only applicable in the perpendicular geometry [16, 24].

Masked islands of 100 μm \( \times 600 \mu \text{m} \) were embedded by the waveguide DH structure with \( \lambda_G = 1050 \text{ nm} \) forming lateral couplings. Laser-waveguide chips were cleaved. The lasers (400 μm in length) are surrounded on three sides by the waveguide DH structure. The threshold current
density of a laser/waveguide combination with a 300 μm-long waveguide-DH structure is $j_{th} = 1.28 \text{ kA/cm}^2$. The threshold current density decreases to $j_{th} = 880 \text{ A/cm}^2$ if the waveguide-DH structure in the direction of the laser cavity is cleaved off. This threshold current density is identical with the value of a reference sample before selective-area-growth processing [22, 23]. Consequently, the etching and SAE growth of the waveguide have not degraded the laser performance. In order to determine quantitatively the optical losses at such lateral couplings a waveguide/waveguide integration was processed.

The SEM cross-section view in Fig. 4a gives an example of a coupling of an InP/GaInAsP($\lambda_G = 1058 \text{ nm}$)/InP and an InP/GaInAsP($\lambda_G = 1100 \text{ nm}$)/GaInAsP/InP. This structure was cleaved off to reveal the selective mask (InP/InP) and the selective fill (InP/GaInAsP/InP).
1100 nm)/InP. In this example the masked structure was grown with TBAs and TBP.

Such a structure allows for the investigation of the material composition at the transition areas by μ-PL due to a difference in $\lambda_\text{c}$ higher than the full width of maximum of the PL peak at 300 K and for the precise loss measurements at the butt-couplings due to an approximately identical refractive index of both quaternary materials. This SAE infill exhibits absolutely planar layers, a thin InP separation layer (10% of the buffer layer thickness) and a smooth contact to the masked quaternary material. The μ-PL data in Fig. 4b are comparable to the data of the example in Fig. 1b, except for the lateral
transition of an emission wavelength of \( \lambda_G = 1058 \) to 1100 nm. Again, there is only a shift of the PL wavelength at the junction of less than \( \pm 2 \) nm. These properties promise highly efficient butt-couplings.

Fig. 5a gives a Nomarski interference micrograph of a top view on a series of reactive-ion-etched-rib waveguides perpendicular across the lateral heterojunctions. The horizontal 8 and 12 \( \mu \)m-wide lines represent the heterostructure with \( \lambda_G = 1058 \) nm from the first grown structure. In between, there is an infill of the DH structure with \( \lambda_G = 1100 \) nm. We have etched two different types of rib waveguides as shown in the inset of Fig. 5b. In type 1 the wave is guided by a 3 \( \mu \)m-wide reactive-ion-etched ridge in the quaternary material. The second type of waveguide is obtained by a 4.5 \( \mu \)m-wide ridge etched in the InP cap layer.

The precise determination of the loss of the waveguide/waveguide butt-couplings was achieved with the cut-back method using the TE polarized laser light with a wavelength of 1540 nm. In Fig. 5b, the coupling loss of these different types of waveguides versus the number of butt-couplings is plotted. The small spread of data points for the two different waveguides of the same type in the case of type 1 and three in the case of type 2 over 12 cutbacks illustrates the high uniformity of the couplings. Some scatter of cleaved edge surface gives the major source of error in the measurement. Type 1 shows an optical loss of \( 0.22 \pm 0.05 \) dB/coupling in the cut-back measurements for a series of 24 butt-couplings and type 2 the best value of \( 0.12 \pm 0.04 \) dB/coupling for a series of 22 butt-couplings. The propagation loss of identical waveguides on a reference sample without any couplings was measured to be \( 0.59 \pm 0.06 \) dB/cm by the Fabry–Perot resonance method [25]. This low propagation loss of the waveguides is comparable to the best published data [26, 27] and is still included in the cut-back measurements.

4. Conclusion

The embedded SAE grown lateral couplings of InP/GaInAsP/InP-DH structures have a constant material composition up to the lateral contact. The PL intensity is high up to the junction with a rapid decay at the junction given by the resolution of the laserspot of about 5 \( \mu \)m. We have presented the first lateral coupling of GaInAsP/InP heterostructures selectively grown by using the replacement precursors TBAs and TBP.

The reactive ion etching process and the selective regrowth of waveguides in MOMBE allow for a high quality butt-coupling of laser and waveguides and does not degrade the threshold current densities of the MQW lasers.

The butt-coupling efficiency of 97% measured at a cascade of 22 couplings on a length over 5.5 mm, which is in the order of the dimensions of chips incorporating PICs, is very attractive for applications in the integration of photonic devices.

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References

New semiconductors TlInGaP and their gas source MBE growth

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Abstract

New III–V compound semiconductors Tl\textsubscript{x}In\textsubscript{1-x}Ga\textsubscript{y}P (thallium indium gallium phosphide), which we proposed recently for 0.9 \textmu m to over 10 \textmu m laser diodes, are grown by gas source molecular beam epitaxy on InP and GaAs substrates for the first time. They have a potential to exhibit a temperature-independent band-gap energy. Grown layers exhibit (2 x 4) surface reconstruction. X-ray diffraction measurements show the successful growth of TlInP, TlGaP and TlInGaP, although phase separation is observed in TlGaP grown on GaAs. Photoluminescence emission is observed for TlInP and TlInGaP grown on InP. Hall measurements show n-type conduction with a room temperature electron concentration of 6.3 x 10\textsuperscript{15} cm\textsuperscript{-3} and an electron mobility of 2500 cm\textsuperscript{2}/V·s.

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Keywords: Thallium alloy; TlInP; TlGaP; TlInGaP; Infrared; Mid-infrared; Temperature-independent band gap; Gas source MBE

1. Introduction

Recently, we have proposed the new III–V compound semiconductors Tl\textsubscript{x}In\textsubscript{1-x}Ga\textsubscript{y}P (thallium indium gallium phosphide) as an active layer for 0.9 \textmu m to over 10 \textmu m wavelength range laser diodes [1]. TlInGaP can be lattice-matched to InP as well as GaAs. This material system can cover the wavelength range from 0.92 \textmu m (InP) to over 10 \textmu m (TlInP) and is suitable for optical devices, especially for 1 \textmu m range laser diodes (LDs) for optical fiber communications and mid-infrared (longer than 2 \textmu m) LDs. Furthermore, these LDs can have the potential to operate without changing the wavelength with ambient temperature variation. TlInGaP can also be lattice-matched to GaAs, covering wavelengths from 0.65 \textmu m (InGaP) to about 1.24 \textmu m (TlGaP).

The estimated band gap energy variation with alloy composition is shown in Fig. 1 [1]. The line

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for InGaP is based on experimental data, the line for TIlnP is from theoretical calculation [2], while the line for TIGaP is only estimated. These semiconductor materials are alloys consisting of the semiconduc-
tor InGaP and the semimetal TIP. This alloy system has only one group V atom and the type-I band lineup having a wider conduction band discontinuity than that of valence band is expected. This characteristics is very suitable for the optical devices. Furthermore, the semiconductors whose band gap does not change with ambient temperature are expected because of the alloy of semiconductor and semimetal similar to Hg$_5$Cd$_{1-x}$Te, where the temperature-independent band-gap energy at a Hg composition of 0.4 was observed [3]. This characteristics is very promising to fabricate semiconductor lasers whose wavelength does not change with ambient temperature variation, which is very important for the advance in WDM (wavelength division multiplexing) optical fiber communication, because one problem in using InGaAsP/InP LDs in WDM system is that the lasing wavelength fluctuates with ambient temperature variation due to the temperature dependence of band gap energy and LDs in WDM system must be equipped with Peltier elements that works to stabilize LD temperature.

In this paper, we report the first successful growth of this material system by using gas source molecular beam epitaxy (MBE) on both InP and GaAs substrates. Preliminary results on optical and electrical properties are also described.

2. Experimental procedure

The growth was conducted in a gas source MBE apparatus (ANELVA GBE-830), which was evacuated by an oil diffusion pump with liquid-nitrogen trap [4]. Elemental Tl (5N), In (7N) and Ga (7N) were used as group III sources and thermally cracked PH$_3$ was used as a group V source. The substrates used were Fe-doped (100) InP and Cr-doped (1 0 0) GaAs. The substrate temperature was monitored with an optical pyrometer calibrated by the melting point of InSb at 525°C. PH$_3$ flow rate was 0.5–1.5 SCCM. The vapor pressure of Tl is similar to that of Sb (antimony) at low cell temperatures and lower at high cell temperatures, and Tl vaporizes from melt. Therefore, the Tl flux controllability is much better than that of Sb. The toxicity of Tl is known to be higher than that of As, so that extreme care during treatment must be taken. Tl or Tl contaminated surfaces should not be touched with bare hands and Tl flakes or vapors should not be inhaled: lethal intake is documented to be at 600 mg.

Grown surfaces were studied by reflection high energy electron diffraction (RHEED). Crystallographic quality and mole fraction of layers were evaluated by X-ray diffraction. Electrical properties were studied by van der Pauw Hall measurement. Photoluminescence (PL) measurement was conducted with an Ar ion laser (488 nm) as an excitation source.

3. Results and discussion

3.1. Growth on InP

The InP substrate surface was thermally cleaned at 480°C under a PH$_3$ flow rate of 1.0 SCCM. After
thermal cleaning, an InP buffer layer (0.1 μm thick) was grown at a substrate temperature of 350–450 °C with a PH₃ flow rate of 0.3–0.5 SCCM. RHEED patterns showed (2×4) phosphorus-stabilized reconstruction. On these InP buffer layers, TIP binary, TIlnP ternary and TIlnGaP quaternary layers were grown (thickness: about 0.5 μm). Tl composition of TIlnP was 0–1, while Tl and Ga compositions of TIlnGaP were about 0.05–0.2 and 0.1–0.2, respectively.

RHEED patterns for TIP and TIlnP revealed (2×4) reconstruction at substrate temperatures of 400–450 °C showing phosphorus-stabilized surfaces. The surface exhibited mirror-like. During the growth of TIlnP, RHEED intensity oscillation was observed as shown in Fig. 2. By the addition of Tl flux during the growth of InP, the oscillation period was decreased indicating the growth of TIlnP. At lower temperatures RHEED pattern showed phosphorus-excess (2×2) reconstruction.

Fig. 3 shows the double crystal (monochromatic) X-ray diffraction rocking curves for TIP/InP and TIlnP/InP samples. In addition to the peak from InP, those from TIP and TIlnP are clearly visible, indicating the successful growth of TIP and TIlnP. TIlnP alloys with various Tl compositions were also grown by varying Tl flux during growth and the alloy composition of TIlnP was found to vary with Tl flux [5]. Therefore, it was shown that the entire range of TIlnP can be grown by gas-source MBE. Recently, phase separation was reported in the MOVPE growth of TIlnSb (thallium indium antimonide) [6]. However, in the present study we observed no phase separation in TIlnP grown on InP by gas-source MBE, though phase separation was observed in the TIGaP growth as described below.

PL emission was detected from the TIlnP layers grown on InP substrates. Fig. 4 shows the 77 K PL spectra from TIlnP and InP. The PL shift to lower energy is small in Fig. 4 because of low Tl composition. PL study on TIlnP with larger Tl composition is in progress, and results will be reported in the separate paper. Preliminary Hall measurement data on TIlnP showed n-type conduction with an electron concentration of $6.3 \times 10^{15}$ (3.9 $\times 10^{17}$) cm$^{-3}$ and an electron mobility of 2500 (22000) cm$^2$/V·s at room temperature (77 K) for TIlnP with a Tl composition of 5%.

Fig. 5 shows the double crystal X-ray diffraction rocking curves for the TIlnGaP quaternary layer grown on InP. The peaks from InP and TIlnGaP are clearly observed. The shifts to the lower diffraction angle for the TIlnGaP peak with increasing Tl flux and to the higher angle with increasing Ga flux were observed. These shifts agree with the increase of Tl mole fraction and Ga mole fraction, respectively. This indicates that quaternary TIlnGaP
layers can also be grown by gas-source MBE. Reduction of full width at half maximum (FWHM) was also observed by decreasing the degree of lattice-mismatch for TlInGaP as can be seen in Fig. 5. No phase separation was also observed. This result indicates that the TlInGaP alloy on InP substrate is promising from the viewpoint of crystal growth. We have observed the PL (not shown) only from the TlInGaP with a composition close to InP at present.

![Image of photoluminescence spectra from InP and TlInP grown on InP.](image1)

**Fig. 4.** Photoluminescence spectra from InP and TlInP grown on InP.

![Image of double crystal X-ray diffraction rocking curves for TlInGaP on InP.](image2)

**Fig. 5.** Double crystal X-ray diffraction rocking curves for TlInGaP on InP.

![Image of double crystal (monochromatic) X-ray diffraction rocking curve for TlGaP on GaAs around the angle of GaAs. Peak from TlGaP is observed as well as that of GaAs.](image3)

**Fig. 6.** (a) Double crystal (monochromatic) X-ray diffraction rocking curve for TlGaP on GaAs around the angle of GaAs. Peak from TlGaP is observed as well as that of GaAs. (b) Powder (unmonochromatic) X-ray diffraction rocking curve for the same sample. Phase separation is observed.
3.2. Growth on GaAs

We have also grown TiGaP layers on GaAs substrates. TiGaP layers (thickness: about 0.5 μm) were grown after the growth of GaAs buffer layer. Fig. 6 shows (a) double crystal (monochromatic) X-ray diffraction rocking curve and (b) conventional powder (unmonochromatic) X-ray diffraction rocking curve from TiGaP/GaAs. Two peaks are observed in Fig. 6a. Lower and higher angle peaks are confirmed to come from GaAs substrate and TiGaP layer, respectively, by selective etching of TiGaP layer with HCl. This means that the nearly lattice-matched TiGaP was grown here. However, as shown in Fig. 6b, phase separation was also observed. In addition to the diffraction peaks from GaAs and nearly lattice-matched TiGaP at around 33.06° where the Kz1 and Kz2 peaks are resolved, the broad peak at the angle for GaP (34.45°) is also observed. The broad peak is also visible at the angle for TIP (31°), though weak for this sample. This observation indicates the occurrence of phase separation and is contrast to the result for TInnP on InP. This difference seems to be due to the difference in lattice-mismatch between the two systems. Tl composition of phase separated TiGaP having lattice constant close to GaAs ranged from about 0.47–0.53 in the present study. More detailed results will be reported elsewhere.

4. Summary

We have, for the first time, studied the growth of new semiconductor TInGaP, which we recently proposed as an active layer of 0.9 μm to over 10 μm laser diodes, by gas source MBE on (1 0 0) InP and GaAs substrates. Grown layers exhibited (2 × 4) surface reconstruction. RHEED intensity oscillation was observed during the growth of TInnP as well as InP indicating the layer by layer growth. X-ray diffraction measurements showed the successful growth of TInnP, TiGaP and TInGaP. No phase separation was observed in TInnP and TInGaP grown on InP substrate, while the phase separation was observed in TiGaP grown on GaAs. PL emission was observed for TInnP and TInGaP grown on InP, though composition was close to InP. The layers showed n-type conduction with an electron concentration of \(6.3 \times 10^{15} (3.9 \times 10^{15}) \) cm\(^{-3}\) and an electron mobility of 2500 (22 000) cm\(^2\)/V·s at room temperature (77 K) for TInnP with a Tl composition of 5%. It is concluded that the gas source MBE is a promising method for the growth of TInGaP.

References

CBE of 1.55 µm (GaIn)(AsP) lasers for monolithic integration

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Abstract

Chemical beam epitaxy is well suited for monolithic integration due to its mask selective growth. We have grown embedded GaInAsP/GaInAsP SCH MQW laser structures in grooves in InP substrates etched by ECR-RIE using CH\textsubscript{3}/H\textsubscript{2} with Si\textsubscript{3}N\textsubscript{4} masks, both, for etching and selective growth. Our laser structure consisted of six compressively strained quaternary MQWs, quaternary barriers (λ = 1.2 µm) and confinement layers (λ = 1.1 µm). Selectively grown stripe lasers in 4 µm wide grooves showed threshold currents of 16 mA for a length of 280 µm. No change in composition of the quaternary core appeared for lasers in 7 and 30 µm wide grooves, when compared with lasers on planar substrates. Quantum wells grown in narrow 4 µm wide grooves, on the other hand showed wavelength shifts depending on substrate misorientation. A 10 nm redshift is observed on (1 0 0) substrates oriented 2° off towards the next [1 1 0] direction, whereas a slight blueshift on (1 0 0) exact oriented substrates occurred.

1. Introduction

The monolithic integration of differing photonic devices requires selective area epitaxy. Chemical beam epitaxy (CBE) allows selective growth of (GaIn)(AsP) layers on structured and masked substrates, thus the integration of lasers, waveguides, etc. become possible. The growth is restricted to non-masked areas, as no nucleation takes place on dielectric masks, e.g. Si\textsubscript{3}N\textsubscript{4}. We describe the realisation of strained 1.55 µm laser structures for monolithic integration. Filling the laser into a groove on the substrate reduces the number of regrowths, lowering the thermal stress on the laser structure.

Best device performance is usually obtained with compressively strained GaInAsP/GaInAsP SCH MQW structures [1, 2]. The use of quaternary wells gives an additional degree of freedom, allowing a variation of strain at a fixed well width to obtain the required wavelength of 1.53 µm. It is essential in view of monolithic integration to grow these complicated laser structures selectively on Si\textsubscript{3}N\textsubscript{4} masked and patterned InP substrates.

2. CBE of high quality GaInAsP

Epitaxy was performed on sulfur-doped (1 0 0) InP substrates oriented 2° off towards the nearest
[110] direction. The precursors were pure trimethylindium and triethylgallium as group-III sources. A high temperature (1050°C) injector decomposed arsine and phosphine as group-V sources. We investigated the epitaxial growth of lattice matched quaternary compounds and found only very narrow parameter windows suited for high crystal quality, essentially if growth temperatures are kept constant to avoid growth interruptions at the interfaces of the MQW structure.

Fig. 1 emphasizes the viable V/III\textsubscript{BEP} ratios for our rather high growth temperature of 543°C as a function of the emission wavelength, characterizing the composition. Increasing In and P content, i.e. decreasing wavelength, requires higher supply of hydrides, changing from a V/III\textsubscript{BEP} ratio of 3 for long wavelength GaInAsP to 8.5 and 10 for compositions corresponding to wavelengths of 1.2 and 1.1 μm, respectively.

GaInAsP layers (300 nm thick), lattice matched to InP, grown with the above parameters, always had high luminescence yield and mirror-like surface morphology. For long wavelength GaInAsP the typical 4.2 K photoluminescence linewidth was 3–4 meV. The compositions used for confinement and barriers showed slightly higher values of 5 meV. The linewidths were nearly independent of the strain, ranging from (Δa/a)\textsubscript{relax} = −1.5 × 10\textsuperscript{−3} to 1.5 × 10\textsuperscript{−3}. Photoluminescence linewidths obtained at 300 K are typically 40 meV. These results emphasize the high quality of GaInAsP in the narrow growth window.

3. Planar growth of quaternary lasers

A typical MQW structure consisted of 6 compressively strained GaInAsP quantum wells and barriers with a material composition of λ = 1.2 μm. The MQW was embedded in-between two 1.2 and 1.1 μm GaInAsP separate confinement heterostructure layers with thicknesses of 35 and 100 nm, respectively. The n and p cladding layers with thicknesses of 1000 and 1600 nm were doped with elemental Si and Be.

These all quaternary MQWs showed an intense and narrow photoluminescence at 300 K (E\textsubscript{PL} = 0.81 eV for laser emission at 0.80 eV) and 4.2 K, with FWHM values of 24 and 6.4 meV, respectively. The well width was 5.4 nm and the strain of the QW was (Δa/a)\textsubscript{relax} = 6.5 × 10\textsuperscript{−3}, determined by XRD measurements. All other quaternary layers were lattice matched within |Δa/a|\textsubscript{relax} < 0.25 × 10\textsuperscript{−3}. Broad area laser diodes processed from these structures [3] had extrapolated threshold current densities of 410 A cm\textsuperscript{−2} and internal losses of 12 cm\textsuperscript{−1}. These extremely low threshold current densities of 68 A cm\textsuperscript{−2} per quantum well clearly demonstrate the high quality of quaternary laser structures grown with CBE. Ternary lasers with equal well widths and half strain for the identical emission wavelength showed higher values of 78 A cm\textsuperscript{−2}, indicating the influence of higher strain for the reduction of threshold current densities.

4. Selective growth of laser structures

Having established optimized growth and material parameters on planar, i.e. nonstructured substrates, we used these growth conditions for the epitaxy of SCH MQW structures in grooves, 1.5 μm deep with varying widths. To obtain planar growth at the bottom of the groove, we modified
layers (0.4 nm s⁻¹), small {1 1 1} facets at the edge of the SCH MQW layer appeared. Growth of 1.4 μm thick InP:Be cladding layers with \( \alpha_{(100)} = 0.4 \text{ nm s}^{-1} \) to a height above the groove edge, resulted in a faceted ridge. This guaranteed an excellent coverage with the evaporated metallisation for contacting. Laser facets were obtained by cleaving perpendicular to the grooves. Fig. 3 shows the light power output-current characteristics for a 280 μm long, 4 μm broad device. The lasers show a low threshold current of 16 mA and an external quantum efficiency of 47% for both facets.

5. Selective growth on 2° misoriented substrates

Compositional changes of homogeneous thick GaInAsP layers and quaternary SCH MQW structures were investigated with cathodoluminescence for different groove widths. The laser core and thick GaInAsP layers did not change in composition in 7 and 30 μm wide grooves, compared to the planar growth. In narrow, 4 μm wide grooves the energy shift was \(- 5 \text{ meV} \) for quaternary and \(- 13 \text{ meV} \) for ternary quantum wells (see Fig. 4).

The decrease in emission energy of 1.2 μm wavelength, thick GaInAsP was found to be 10 meV. For GaInAs the energy shift indicates an increase in compressive strain. The relative

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**Fig. 2.** SEM picture of the facet of a selectively grown MCRW Laser.

**Fig. 3.** Light power output characteristic of a (GaIn)(As)P SCH MQW laser selectively grown in an InP groove. The identical threshold current is 16 mA, for pulsed and cw operation.

**Fig. 4.** Cathodoluminescence measurements for different groove widths. For 4 μm wide grooves on (1 0 0) 2° misoriented substrates a redshift is observed. For the same grooves on exactly oriented (1 0 0) substrates a blueshift occurs.
incorporation ratio In/Ga during planar growth for quaternary and ternary compositions showed a constant value of 2.5, independent of composition. Due to the observed equal incorporation ratio In/Ga at planar growth for quaternary and ternary compositions the energy shift indicated a lower Ga incorporation for GaInAs and for GaInAsP. In the case of the GaInAsP (\(\lambda = 1.2 \mu m\)) the energy shift linked with a reduced phosphorus incorporation.

With decreasing misorientation of the substrate, i.e. lower surface step density, the gallium and the phosphorus incorporation of GaInAsP (\(\lambda = 1.05 \mu m\)) were found to decrease [4]. In our case, we observed a tilted surface in the 4 \(\mu m\) wide grooves, corresponding to the misorientation of the substrate. These phenomena did not occur when using exactly oriented (1 0 0) InP substrates as shown in Fig. 5. The SEM picture shows the facet of a selectively grown laser in a groove on a substrate with an exact (1 0 0) surface orientation. Here the energy shifted to slightly higher values (+3 meV) as shown in Fig. 4.

Growth of laser structures yielded higher compressive strain when using (1 0 0) oriented substrates in comparison to structures grown at the same temperature and material fluxes on 2° misoriented substrates. XRD spectra show this clearly in Fig. 6. The upper spectrum is that of a SCH MQW structure on a misoriented substrate, the lower spectrum shows the same structure grown on a (1 0 0) exactly oriented substrate. Especially, the peaks attributed to the confinement layers shifted to lower angles indicating a change in lattice mismatch of \(\Delta(\Delta a/a) = 3 \times 10^{-3}\) for \(\lambda = 1.1 \mu m\) and \(1.2 \mu m\) GaInAsP. For the long wavelength GaInAsP of the quantum wells the change of mismatch was half with \(\Delta(\Delta a/a) = 1.5 \times 10^{-3}\). The photoluminescence energy rose by 25 meV on the misoriented substrate, while the superlattice period was constant within 0.2 nm, indicating unchanged growth rates.

The energy shift to lower values in narrow grooves during selective growth on 2° misoriented substrates was due to lower step densities, by reconstruction of the (1 0 0) plane in the groove. Investigations of wide area selective growth on (1 0 0) oriented substrates substantiated this result.

6. Summary

The presented results show the capability of chemical beam epitaxy for monolithic integration of optoelectronic devices. The optimized growth parameters of planar grown laser structures allow the selective growth of high quality lasers with threshold currents as low as 16 mA. The redshift of the emission wavelength for narrow 4 \(\mu m\) wide devices in grooves on 2° misoriented substrates indicate a reconstruction of the (1 0 0) plane during selective growth.
For geometrical reasons (lateral symmetries of infilled structure) the growth on exactly oriented substrates has to be preferred, especially as the 4.2 K photoluminescence linewidth of 5 meV was the best one achieved during this work. On the other hand it is easy to suppress \{111\} facets in grooves on misoriented substrates. However, growth parameters on exact oriented substrates are more sensitive, especially control of InP growth is difficult at our rather high growth temperatures.

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References

MOMBE growth of highly tensile-strained InGaAsP MQWs and their applications to 1.3-μm wavelength low threshold current lasers

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Abstract

We have studied metalorganic molecular beam epitaxy (MOMBE) of InGaAsP multi-quantum wells (MQWs) with 0.5–1.8% tensile-strained well layers. The MQW photoluminescence intensity depends significantly on the barrier layer strain to compensate well layer strain. Optimizing the strain compensation conditions, i.e., keeping the net strain less than 0.3%, makes it possible to grow high optical quality MQWs with 1.8%-strained, 10-well layers. A critical thickness-net strain ($t_c$) curve for the tensile-strained MQWs has been experimentally determined by analyzing more than 50 samples with various strains and total thicknesses. The $t_c$ curve fairly agrees with the one calculated by the Matthews formula for more 0.5% net strain, but the $t_c$ values are much higher than the expected ones below that range. For 6-well MQW lasers, the threshold current densities, $J_{th}$s, decrease with increasing tensile strain; the minimum $J_{th}$ obtained is 0.6 kA/cm$^2$ at 1.3% strain.

1. Introduction

Strained multi-quantum well (MQW) lasers emitting at 1.3 μm wavelength are of interest for optical communication and optical interconnection systems [1]. Theoretical studies [2, 3] predict that tensile-strained MQW lasers have even better performance than compressive ones. Little work, however, has been carried out for tensile-strained MQWs owing to the difficulty of growing them.

Theoretical studies show that, in order to extract the beneficial effects of tensile-strained MQWs, the well layer must be more than 100 Å thick and have more than 1% strain. The stacking of such thick and highly strained layers easily generate misfit dislocations. For example, photoluminescence intensity of 1.1% tensile-strained MQWs grown by metalorganic chemical vapor deposition (MOCVD) remains strong for up to three wells but drastically degrades for four wells. The threshold current density of the MQW lasers decreases with increasing tensile strain in less than the 1% range as predicted, but it increases with over 1% strain [4, 5]. Increasing the strain and well number is expected to improve the laser performance.

The growth of strained MQWs with high optical quality requires a knowledge of the critical
thickness over which misfit dislocation are generated. Houghton et al. [6] derived a theoretical formula for MQWs using the Matthews model [7]: the Matthews formula holds for MQWs by replacing strain and layer thickness with net strain [8] and MQW total thickness. Many researchers have reported that strain compensation to reduce net strain is effective for growing highly strained MQWs and improving laser performance [5, 9]. However, there has been no systematic study on the experimentally determined critical thickness for tensile-strained MQWs.

Recently, one of the authors reported that metalorganic molecular beam epitaxy (MOMBE) enables the growth of 1.45%, 6-well strained MQWs [10] without generating misfit dislocations. In this letter, we describe how to grow tensile-strained InGaAsP MQWs with many well layers and show what parameters determine critical thickness of the strained MQWs grown by MOMBE. The strain effect on the MQW laser performance is also mentioned.

2. Experimental procedure

We used the Vacuum Generators V-400 CBE system. Triethylgallium (TEG) and trimethylindium (TMI) were the group III sources, and AsH₃ and PH₃ the group V sources. The hydride gases were decomposed in a low-pressure crack reactor cell heated at 905°C. Substrate temperature was kept at 520°C for InGaAsP films. The temperature was calibrated with the InSb melting point. Solid Be and Sn were used as p- and n-type dopants, respectively. Pre-etched (1 0 0)-oriented substrates purchased from Sumitomo were used without any chemical treatment.

Typical MQWs consisted of tensile-strained InGaAsP (equivalent wavelength \( \lambda = 1.5 \) μm) as well layers and compressive-strained InGaAsP (\( \lambda = 1.13 \) μm) as barrier layers. The well layer thickness \( L_w \) and the barrier thickness \( L_b \) were kept at 100 and 140 Å, respectively. Double-crystal X-ray diffraction spectra of the MQWs were analyzed with a kinematic step model to determine the lattice constants of the unrelaxed well and barrier layers. The strain values used in this paper represent the lattice mismatch to InP. The tensile strain \( \varepsilon_w \) in the wells was varied from 0.5% to 1.8% while the compressive strain \( \varepsilon_b \) in the barriers was varied from 0% to 0.5%. Each value contains an ambiguity of \( \pm 0.03\% \). Hereafter, we refer to the MQWs with 1.3% tensile strained layer simply as 1.3% strained MQWs. A plan-view transmission electron microscopy (TEM) study was carried out to observe directly misfit dislocations. Microphotoluminescence (μ-PL) [11] was measured with a Waterloo Scientific SPM-2000 to observe misfit dislocations in a \( 2 \times 2 \) mm² area. Conventional PL measurements were performed at room temperature using a YAG laser as an excitation source.

The MQW lasers had separate confinement double heterostructures (SCH) with MQWs. Using MOMBE, we grew a structure consisting of a 5000 Å thick n-type InP buffer layer, 1000 Å thick undoped InGaAsP (\( \lambda = 1.13 \) μm), the undoped MQWs mentioned above, 1000 Å thick undoped InGaAsP (\( \lambda = 1.13 \) μm), a p-type 0.5 μm thick InP, and an InGaAs cap layer. After removing the cap layer, a p-type 2 μm thick InP layer and a p-type InGaAs contact layer were sequentially overgrown at 620°C by MOCVD. The SCH structure was processed into broad lasers with a 40 μm wide stripe.

3. Results

Fig. 1 plots the MQW PL intensity against barrier layer compressive strain as a parameter of well tensile strain. The well number was fixed at six. The tensile strain values were 1.3%, 1.45%, and 1.6% while the barrier strain was varied from 0% to 0.5%. The figure indicates two important features. First, the PL intensity of the MQWs remarkably improves by increasing the barrier strain. Though not shown for 1.8% tensile strain, a similar curve was obtained. This means that the strain compensation technique is still effective in the growth of MQWs consisting of such highly strained and thick well layers. Second, the barrier strain needed for obtaining strong PL intensity must be increased for larger tensile strain.

We have studied the MQWs in Fig. 1 from structural property point of view. Cross-sectional TEM study showed that the thickness modulation of the well layers due to spinodal decomposition was about 10 Å which did not depend on barrier strain.
values. Plain-view TEM and μ-PL measurement revealed that the low PL intensity MQWs had misfit dislocation as shown in Fig. 2a and Fig. 2b, respectively. The misfit dislocation generation is the cause of reducing the PL intensity [12]. By using μ-PL, we classified the samples used in Fig. 1 into two groups, i.e., with and without misfit dislocations. The results are summarized in Fig. 3. The vertical axis shows the net strain ε* calculated using the formula of ε* = (L_w ε_w + L_o ε_o)/(L_w + L_o). The figure clearly shows that misfit dislocations are generated when the net strain exceeds 0.43% irrespective of well tensile strain. This fact strongly suggests that MQW strain condition reflects net strain, not well layer strain. Since the MQWs have a total thickness of 1440 Å, we can say that the critical thickness t_c for 0.43% net strain is 1440 Å.

In order to determine experimentally the critical thicknesses of MQWs in a wide net strain range, we grew more than 50 MQWs with various total thicknesses and net strains. The results are summarized in Fig. 4, where the data are classified into two groups, i.e., with and without misfit dislocations. The solid squares represent the MQWs with tensile-strained InGaAsP well layers in which misfit dislocations are generated while the open squares represent those in which dislocations are not generated. The solid circles represent the MQWs with compressively strained InAsP well layers containing misfit dislocations while the open circles are for those not containing dislocations [13]. The solid line is the border dividing the two classes of the MQWs. In other words, the solid line is the experimentally determined critical thickness–net strain curve. The figure shows that the curve is almost the same for the MQWs with either tensile- or compressive-strained well layers. The gray line in the figure is a theoretical curve calculated using the Matthews formula applied for MQWs [6]. Comparing the two curves, we can extract two

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**Fig. 1.** Dependence of MQW PL intensity on barrier layer compressive strain as a parameter of well layer tensile strain. The barrier strain needed for obtaining strong PL intensity must be increased for larger tensile strain.

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**Fig. 2.** (a) and (b) Misfit dislocation patterns of a tensile strained MQW obtained by plan-view TEM (a) and μ-PL (b). The misfit dislocations orient to [110]. The μ-PL observed area was 2×2 mm².
Fig. 3. Relation between the net strain and well layer strain of the MQWs with respect to misfit dislocations. The samples are used in Fig. 1. Misfit dislocations are generated for net strains over 0.43%.

Fig. 4. Map of MQWs for various MQW total thicknesses and net strains in which misfit dislocations are and are not generated. The solid and gray lines are respectively the experimental and theoretical critical thickness–net strain curves.

Fig. 5. Map of the MQWs for various well numbers and tensile strains in which misfit dislocations are and are not generated. Each MQW had a net strain between 0.2 and 0.3 except for the 1.45% strained MQWs.

According to Refs. [4, 5], the critical thickness at 0.47% net strain for the tensile strained MQWs is about 330 Å. This value is much smaller than our value of 1440 Å and is rather close to the expected value calculated by the Matthews formula (see Fig. 4). The critical thickness difference can be explained in terms of the growth temperature difference between the two growth methods; 650°C for MOCVD and 520°C for MOMBE. Using a modified Matthews model, Price [14] theoretically indicated that the critical thickness can be increased when temperature is lowered. Low-temperature growth might prevent the motion of dislocation due to low lattice vibration.

We conducted an experiment to see how many well layers can be grown for tensile-strained MQWs without generating misfit dislocations. The results are summarized in Fig. 5 as a map of MQWs for various well numbers and tensile strains. Solid and open circles are for the MQWs with and without misfit dislocations, respectively. The well layer tensile strain was varied from 0.5% to 1.8%. Each MQW was designed to have a net strain between 0.2 and 0.3 except for the 1.45% MQWs, whose net strain was 0.37%. From the figure, the maximum well number below which dislocations are not generated is more than ten throughout the tensile strain range from 0.5% to 1.8%. This value is more than twice that obtained in MOCVD growth.

Important features. First, the tendency for critical thickness to decrease with increasing net strain is the same for the two curves. Second, the experimentally determined critical thickness is much larger than the theoretical values in the low net strain range, i.e., less than 0.5%. This latter result suggests that the Matthews formula needs modification in the low net strain range.

Here, we compare our critical thickness results with those for MQWs grown by MOCVD.
Finally, we describe the laser characteristics of the tensile strained MQWs emitting at 1.3 μm wavelength. Misfit-dislocations free MQWs were processed into broad area lasers with a cavity length of 300 μm. All the lasers operated in TM mode. For a fixed strain value of 1.45%, the threshold current density \( J_{th} \) decreased from 2 to 0.7 kA/cm\(^2\) by increasing the well number from 2 to 6. Fig. 6 shows the dependence of the threshold current density on the well strain for the 6-well MQW lasers. The \( J_{th} \)'s decrease with increasing tensile strain from 0.5% to 1.3%, but they slightly increase or remain rather constant in the 1.45–1.8% strain range; the minimum \( J_{th} \) is 0.6 kA/cm\(^2\) at 1.3%. This value is the smallest ever reported for tensile strained 1.3 μm wavelength lasers for such a cavity length. The optimum strain of 1.3% is slightly different from the reported values of about 1% in MOCVD [4, 5].

The reduction of \( J_{th} \) with the larger strain can be explained in terms of a greater separation between the light-hole and the heavy-hole energy band. The saturation of the \( J_{th} \) in the 1.45 –1.8% strain range may be related to the well layer microstructure caused by spinodal decomposition, which was revealed by cross-sectional TEM microscopy [15].

4. Summary

We have systematically studied strain compensation conditions for the MOMBE-grown tensile strained MQWs. The net strain must be kept less than 0.3% in order to obtain MQWs with strong PL intensity. As a result, a 10-well, 1.8% tensile-strain MQW can be grown without generating misfit dislocations. The experimentally determined critical thickness–net strain curve for the strained MQWs agrees with the one calculated by the Mathews formula except in the net strain range of less than 0.5%, which suggests that modification of the formula is required in the low strain range. The threshold current density of the 6-well MQW lasers decreases with increasing well layer strain; the lowest \( J_{th} \) is 0.6 kA/cm\(^2\) at 1.3% strain.

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Growth and transformation of ultra-thin InAs/InP layers obtained by chemical beam epitaxy

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Abstract

We have studied the morphology of InAs layers grown on InP epitaxial surfaces and their evolution under cracked arsine anneal using atomic force microscopy. On an exactly oriented (0 0 1) surface, a 2 ML InAs layer annealed under arsine breaks into elongated dots oriented along the [1 1 0] direction. During annealing, the dot density first increases, then decreases, pointing to stages dominated by nucleation and by coalescence, respectively. Nucleation and filamentary growth of InAs is observed along the steps of a step-bunched InP surface, with an average misorientation of 0.8° towards [1 1 0], while no localization can be achieved on surfaces tilted towards [1 1 0]. A more homogeneous population of isotropic InAs dots with a base size dispersion as low as ±10% is obtained when a silicon-doped InP buffer layer is used.

Keywords: InAs, InP, Quantum dots, Self-organization, Chemical beam epitaxy

1. Introduction

InGaAs/GaAs and InGaAs/InP low-dimensional nanostructures, particularly quantum wires and dots, have interesting optical properties which could make them good candidates for use in novel optoelectronic devices, for instance as the active layer of improved lasers [1]. However, these applications require a high density of very small and regular wires or dots. An efficient way to fabricate such dot structures is to take advantage of the Stranski–Krastanow (SK) growth mode in lattice-mismatched systems like InAs/GaAs [2, 3].

More recently, the compressively strained InAs/InP system has been drawing attention. A thin InAs layer grows in a two-dimensional (2D) way, but a few seconds anneal under cracked arsine triggers an evolution from a two to a three-dimensional (3D) surface as seen by reflection high energy electron diffraction [4, 5]. Transmission electron microscopy confirmed the formation of InAs dots following this procedure [6]. However, no direct observation of this surface was carried out so far.

In the present paper, we report the atomic force microscopy (AFM) investigation of the surface of InP and InAs/InP epilayers grown by chemical beam epitaxy (CBE). We demonstrate the influence of the substrate miscut and its doping on the surface structure and its evolution under arsine anneal.

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2. Experimental procedure

The samples discussed here were grown by CBE using pure trimethylindium, cracked arsine, cracked phosphine and a solid Si source for n-type doping. We used exactly oriented (0 0 1)InP substrates as well as misoriented ones with various miscuts in both [1 1 0] and [1 1 0] directions (with an orientation tolerance smaller than ± 0.1°). The layer structures consisted of a 4000 Å InP buffer grown at a rate of 1 µm/h at a temperature of 500°C for exactly oriented substrates and 520°C for misoriented ones, and subsequently annealed for 5 min under phosphine flux. After optimization of the InP layer morphology, we have grown between 0.3 and 2 monolayers (ML) InAs, at a growth rate of 0.3 ML s⁻¹ and a temperature of 500°C. Next, an anneal under arsine for a few seconds and an anneal under phosphine to quench the process were performed. The samples were finally investigated at atmospheric pressure using a digital instrument nanoscope III AFM.

3. Exactly oriented surfaces

After growth of a 4000 Å thick InP layer at 1 µm/h and 500°C, the surface shows a fragmented topography with a patchwork of 2D islands 5000 Å wide and 1 ML high, as expected under these conventional CBE growth conditions for InP [7]. Having this surface anneal under phosphine was found to smoothen it and create large atomically flat terraces (Fig. 1).

Fig. 2 is an AFM picture corresponding to the case of a 2 ML thick InAs layer annealed for 1 min under arsine. It reveals a high density (4.6 × 10⁹ cm⁻²) of dots oriented along the [T 1 0] direction. A cross-section study reveals a grossly pyramidal dot shape; their main sidewall facets are close to [1 1 5] planes. Their average length, width and thickness are 1165, 367 and 53 Å, respectively, with dot-to-dot variations of less than ± 20% for the lateral dimensions, but as high as ± 30% for their height. The main difference between these dots and the InAs dots obtained by gas source molecular beam epitaxy and arsine anneal [6] is in the stronger anisotropy observed in our case and

Fig. 1. AFM image of a 4000 Å InP buffer grown at 500°C on an exactly oriented substrate and annealed under cracked phosphine for 5 min.
also observed for InAs dots obtained by the SK growth mode on InP by molecular beam epitaxy [8].

The effect of annealing time $t_a$ is shown in Fig. 3 which presents the evolution of the dots density ($D$), average dot volume and shape versus $t_a$. For $15 \text{ s} \leq t_a \leq 300 \text{ s}$, the dots density lies in the $1 \times 10^2$–$7 \times 10^5 \text{ cm}^{-2}$ range (Fig. 3a). First, $D$ increases to a maximum value of $6.75 \times 10^5 \text{ cm}^{-2}$ corresponding to $t_a = 30 \text{ s}$, then it monotonically decreases showing dot gathering. Fig. 3a also shows the dots volume evolution which presents an almost monotonous increase with $t_a$. The dot shape variations are shown in Fig. 3b which presents the width, length and height evolution with $t_a$. For $t_a = 15 \text{ s}$, the dots are small and well oriented along the [1 1 0] direction. When $t_a$ increases to 300 s, the average dot length hardly changes, while both their width and height slightly increase. The dot size regularity is not varying when $t_a \leq 60 \text{ s}$ and is about $\pm 20\%$ for the lateral dimensions and $\pm 30\%$ for the height. For $t_a > 60 \text{ s}$ the regularity of dot size deteriorates drastically (the height variations are more than $\pm 50\%$ for $t_a = 300 \text{ s}$). This is probably the result of random coalescence.

These results show several regimes in the dot evolution with $t_a$. At the beginning of the anneal under arsine, small, regular and very elongated dots are formed. The orientation, along the [1 1 0] direction, could be explained by the surface dimerisation which induces a higher binding energy in this direction [9, 10]. The total volume of these dots ($V_{3D}$) is less than the total deposited InAs volume which means, since In recrystallization is negligible at 500°C, that there is still a 2D InAs layer under the dots. When the annealing time increases, the dot size and particularly their density increase to reach a maximum. This might be called the nucleation step. During the anneal under arsine, there is a surface reorganization which creates spatial fluctuations of the 2D layer thickness. The critical thickness is thus locally exceeded and some dots nucleate [11]. On the other hand, there is a minimum distance between the dots which is a consequence of the repulsion between them [12] and the dot density increases until this minimum is reached.
During this step, $V_{3D}$ increases but is still less than the total InAs volume. When $t_1$ increases further, the dot density decreases and the dot size increases; the dots touch each other and start to gather in a regular way (their size regularity does not deteriorate). During this growth stage, however, they...
become less elongated and thicker. It is important to notice that the dot size variations do not increase during the nucleation and growth stages because their gathering is regulated by the consumption of the 2D layer. Finally, when \( t_n > 60 \text{ s} \) the dots exhibit a random coalescence step which deteriorates their size regularity (the 2D layer is already consumed).

We want to stress that our nominal InAs layer is below the critical thickness (at which the 2D growth is replaced by a 3D one) which is 3 ML for the InAs/InP system [13] and that the formation of our dots is clearly different from the SK mode. However, the evolution with annealing time under arsine shows features which are similar to the formation of the InAs/GaAs dots by the SK growth mode [3, 12].

4. Vicinal surfaces

As we have seen, the size variations of our dots are larger than for InAs/GaAs dots [14]. To further investigate this point, we have studied the substrate influence on the dot formation and size regularity. As we know that the dot density and positioning play a key role in self-assembly processes [3] and that defects on the surface drive the dot nucleation [15, 16], we optimized the morphology of the InP layer (underneath the dots) creating purposely “organized” surface defects. For this purpose, we have used misoriented substrates with different miscuts and various growth parameters in order to obtain multilaminate steps using the step bunching phenomenon [17–20]. We have systematically studied the morphology of InP layers grown on substrates tilted towards [1 1 0] (A) and [1 1 0] (B), with the growth temperature in the range of 490–520°C. The layers grown on the A steps were more regular than those grown on the B steps. The 0.8°-tilt surface was finally selected because the steps were the most regular at a temperature of 520°C and we obtained 5 Å high double-steps separated by 440 Å wide terraces, oriented along [1 1 0]. On these optimized surfaces, we have grown a thin InAs layer with different thickness from 0.2 ML to 0.6 ML and then annealed them under arsine for 1 min.

Our main results are shown in Fig. 4. For a 0.3 ML InAs thick layer and on the exactly oriented substrate there is no dot formation (Fig. 4a). The InAs film can be distinguished from the InP background as it is composed of small 2D islands which are typical of an incomplete layer [12]. On the 0.8°-off (0 0 1) misoriented surface (Fig. 4b), elongated dots oriented along the step direction are formed on the numerous steps which serve as preferential nucleation sites [21]. This means that for such a thin InAs layer and during this short annealing time, kinetics does not allow the formation of dots on a surface without defect, but the multisteps are efficient nucleation sites which favor dot formation. When the InAs layer thickness increases the dots become wider, higher but especially longer along the steps, becoming short wires (Fig. 4c).

It is interesting to note that for similar experiments on a surface with B steps, the multisteps along the [1 1 0] direction do not influence much the dot shape which is still elongated and oriented along [1 1 0].

5. Influence of silicon doping

Another approach for creating surface defects for dot nucleation is to dope the buffer layer. We have studied by AFM the morphology of a Si-doped InP layer with a doping concentration of \( 3 \times 10^{18} \text{ cm}^{-3} \) grown on (0 0 1) InP surface. It reveals that, in addition to large terraces present on the undoped InP layer surface, there are numerous 2D islands. As we can see in Fig. 5, the Si-doping of the InP buffer influences the InAs dots morphology (cf. with Fig. 2). For the same InAs layer thickness (2 ML), the dots are not elongated anymore, but are pyramid or cone shaped. In addition, they exhibit a higher density: \( 1.6 \times 10^{10} \text{ cm}^{-2} \) instead of \( 4.6 \times 10^9 \text{ cm}^{-2} \) and they are bigger (87Å high and 687 Å wide on average). Moreover, the Si presence significantly improves the dot size regularity, ±10% instead of ±20% for the width, ±20% instead of ±30% for the height. This size dispersion is as good as that observed for the InAs/GaAs system which is twice as much strained as the InAs/InP one.
Concerning the isotropic shape of the InAs dots, it is well known that the presence of Si in a GaAs layer can modify the surface structure, even leading to Ga rich reconstruction [22]. As we have seen previously, the elongated shape of the dots nucleated on an undoped InP buffer can be correlated with the surface reconstruction. If this reconstruction changes, the dot shape could change as well.

Since Si doping of the InP buffer was found to strongly influence the dot formation on exactly oriented substrates, we also extended this experiment to vicinal surfaces. However, there was no dramatic influence on the features of the step-bunched InP surface.

6. Summary

The direct observation by AFM of the surface of thin InAs films grown as 2D layers on InP surfaces confirms that arsine anneal induces a surface reorganization leading to the formation of InAs dots.
This behavior can be modified by introducing perturbations on the surface. For instance, a significantly better size uniformity and an isotropic island shape are observed on a Si-doped surface as compared with an undoped layer. Moreover, nucleation and growth along steps on vicinal surfaces tilted toward $\langle 110 \rangle$ was observed, while such preferential growth could not be obtained when the tilt was toward $\langle 110 \rangle$. Further investigations are needed in order to improve the size distribution of such dots for applications in long wavelengths optoelectronic devices.

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References

Transient surface states during the CBE growth of GaAs

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Abstract

We report the occurrence of a transient surface state during the initial stages of CBE GaAs(0 0 1) growth. The state was detected in real-time reflectance (R) and reflectance anisotropy spectroscopy (RAS) growth monitoring. At low growth rates, less than 1 μm/h, beam equivalent pressure (BEP) of triethylgallium (TEG) < 2.5 \times 10^{-5} \text{ mbar} there was no change in R and the RAS signal changed from its pre-growth value under arsenic stabilisation at the growth temperature to its "during growth" value upon admission of the TEG, with the familiar monolayer oscillations. At higher TEG BEPs there was a rapid increase in R at all monitoring wavelengths, followed by a monotonic decay to its pre-growth value. This transient increase in R was accompanied by a change in the RAS signal, the magnitude and sign of which varied with wavelength. The initial increase in R is shown to be associated with the development of a metallic-like surface whereas the changes in the RAS signal are consistent with the formation of Ga dimers.

Keywords: CBE growth; GaAs; RAS

1. Introduction

Optical monitoring of epitaxial growth is gaining wide-spread acceptance as a valuable aid in understanding growth processes. Optical techniques are relatively insensitive to the growth pressure, and whilst this is not particularly important in MBE growth, it is of tremendous benefit for higher-pressure growth environments such as MOVPE. One such optical technique is reflectance anisotropy spectroscopy (RAS) \cite{1, 2}, which measures $\Delta R/R$, the ratio of the difference, $\Delta R$, in normal incidence reflectance for light linearly polarised along two orthogonal directions in a sample surface to the average normal incidence reflectance, $R$, as a function of photon energy. RAS is sensitive to short-range order (bond energies and directions) in the material under investigation. For a cubic material such as GaAs with isotropic bulk optical properties, RAS gives a direct measure of surface optical anisotropy, primarily arising from the surface reconstructions. Indeed, RAS has been applied with considerable success in the study of MOVPE growth, in particular, in mapping out surface reconstructions as a function of III/V ratio and temperature for MOVPE grown GaAs \cite{3, 4} from correlations of simultaneous RAS and RHEED measurements on MBE \cite{5, 6} and CBE \cite{3} grown GaAs.
Like RHEED, RAS has also been used to study the dynamics of the growth. For such measurements the monitoring is undertaken at a fixed photon energy, the energy often being that where the difference in the pre-growth RAS and the RAS determined during growth is maximum [7]. For GaAs this energy is 2.6 eV. However, this is not a necessary condition for a dynamic study of GaAs growth using RAS; such studies have been reported at photon energies down to 1.85 eV [8]. Monolayer period oscillations are readily observed in the dynamic RA traces of GaAs and AlAs CBE growth [9] at relatively low group III BEPs corresponding to growth rates up to \( \sim 1 \) ML/s. The purpose of this study was to use RAS to probe the growth dynamics at the higher group III BEPs normally used in our CBE GaAs growth.

2. Experimental procedure

The CBE growth experiments were carried out in a VG80H growth system fitted with gas lines which has been described elsewhere [10]. The precursors were thermally pre-cracked arsine (AsH3) and triethylgallium (TEG). The real-time RAS monitoring system was based on the photo-elastic modulator design reported by Aspnes [2].

The RAS measurements are calculated and displayed in real time as \( \Delta R/R \). Thus, any temporal changes in the surface anisotropy (\( \Delta R \)) can be correlated with changes in \( R \).

Typically, a spectrum (1.5–4.0 eV) with data points at every 0.02 eV took 2 min to acquire. In the single wavelength dynamic mode, measurements were taken and displayed in real time at intervals down to 20 ms.

The experiments were carried out at temperatures where the CBE growth rate of GaAs is maximum for a given group III beam equivalent pressure (BEP), that is over the range 510–560°C [11]. Spectral and dynamic \( R \) and RAS measurements were made with the group III and group V BEPs over the ranges \( 3.0 \times 10^{-5}–4.5 \times 10^{-5} \) mbar and \( 10^{-4}–4 \times 10^{-4} \) mbar, respectively, where the GaAs growth rate is in the range 1.5–2 ML/s.

3. Results and discussion

RAS spectra over the range 1.5–4.0 eV for temperatures of 510°C and 535°C are shown in Fig. 1 in the pre-growth (PG) condition (group III BEP = 0) and during growth (DG) with a group III BEP of \( 3 \times 10^{-5} \) mbar; the group V BEP was \( 10^{-4} \) mbar in all cases. From correlations with previous simultaneous RHEED and RAS measurements on our CBE growth equipment [3] these spectral RAS traces show the 510°C pre-growth surface reconstruction to be \( \sqrt{2} \langle 4 \times 4 \rangle \), changing to \( 2 \times 4 \) during growth, whilst the 535°C pre-growth reconstruction can be approximately represented as a linear combination of \( \sqrt{2} \langle 4 \times 4 \rangle \) and \( 2 \times 4 \), also changing to \( 2 \times 4 \) during growth. During growth at higher temperatures and for pre-growth the RAS signal is that of the \( 2 \times 4 \) surface [3].

Upon admission of TEG at BEPs up to \( 2.5 \times 10^{-5} \) mbar, there was no change in \( R \) and the dynamic RAS trace changed from the pre-growth.

![Fig. 1. RA spectra of GaAs at different temperatures in the pre-growth and during growth conditions.](image-url)
Fig. 2. Transients in \( R \) and \( \Delta R/R \) as a function of monitoring energy and group III BEP.

to the “during growth” value with the familiar oscillatory behaviour. The period of the oscillations corresponded to the growth of 1 ML of GaAs and the oscillations damped out to the during growth RAS value. When the group III BEP was higher than \( \sim 3 \times 10^{-5} \) mbar, immediately upon admission of the TEG, there was a rapid increase in the value of \( R \) at all monitoring energies followed by a decay to the pre-growth value of \( R \). At these higher BEPs the oscillatory behaviour of the dynamic RAS trace disappeared and transients were observed in the RAS trace in the interval between the pre-growth and the during-growth conditions. Examples of the transients are shown in Fig. 2, in which the RAS traces have been shifted in the \( \Delta R/R \) axis but the scale has been preserved. Likewise, the reflectance traces have been shifted whilst maintaining the scale and have been normalised to the pre-growth value, \( R_{pe} \); they therefore appear as a percentage change.

In all transients the maximum in \( R \) occurred some 0.8–1.0 s after the admission of the TEG. However, the decay time, back to the pre-growth value, increased with increasing group III BEP (for a constant group V BEP). The magnitude of the increase in \( R \) depended on both energy and group III BEP (again at a constant group V BEP). Fig. 2 shows that the increase in \( R \) was 27% at 1.8 eV falling through 12% at 2.6 eV to 9% at 3.5 eV when the group III and group V BEPs were \( 4 \times 10^{-5} \) and \( 10^{-4} \) mbar, respectively; reducing the group III BEP to \( 3.5 \times 10^{-5} \) mbar had the effect of reducing the increase in \( R \) to 6%.

Using a bilayer model (ambient/addlayer/bulk GaAs) the change in \( R \) with thickness can be estimated, provided the optical constants (refractive index, \( n \), and extinction coefficient, \( k \)) of the addlayer and GaAs are known. Over the spectral range of interest, the refractive index of GaAs, \( n_{GaAs} \), is \( \sim 4 \) and the extinction coefficient, \( k_{GaAs} \), is in the range 0.2–2.5 [12]. Over this short period of time the maximum amount of material that may have been added to the substrate is \( \sim 2 \) ML and thus the increase in \( R \) requires either \( n \gg n_{GaAs} \) or \( k \gg k_{GaAs} \) or both, where \( n \) and \( k \) are the optical constants of the surface addlayer. The refractive index of GaAs is high and it is unlikely that \( n \) can be much greater than \( n_{GaAs} \). Large values of the extinction coefficient \( (k \gg 1) \) occur in metals, the magnitude falling with energy over the spectral range of this study. A bilayer simulation assuming the addlayer to have a thickness of 2 monolayers for the group III BEP of \( 4 \times 10^{-5} \) mbar and taking a range of values of \( n \) from 1 to \( n_{GaAs} \) at the monitoring wavelength indicated that \( k \) would have the values \( 16 \pm 2 \) at 1.8 eV, \( 9 \pm 1 \) at 2.6 eV and \( 6 \pm 1 \) at 3.5 eV. Thus, the increase in \( R \) upon admission of the TEG is consistent with the addlayer being metallic in terms of its optical behaviour. The bilayer model predicts an oscillatory behaviour in \( R \) with thickness if the addlayer remained as deposited. The monotonic decay back to the original value of \( R \) and the absence of oscillations indicates that the addlayer becomes incorporated into the growing GaAs.

Fig. 2 demonstrates that the transient in \( R \) was always accompanied by a transient in the dynamic RAS trace, the duration of the two transients being similar. As the RAS signal is presented as \( \Delta R/R \) then any changes in \( R \) will automatically be manifest as a change in \( \Delta R/R \). A given percentage increase in \( R \) will cause the magnitude of \( \Delta R/R \) to
fall by a corresponding fraction. As can be seen from Fig. 2 the magnitude of the changes in $\Delta R/R$ does not correspond in percentage terms with the fractional increase in $R$; depending on the monitoring energy $\Delta R/R$ may increase or decrease during the transient. This demonstrates that there is a change in the surface optical anisotropy over the duration of the transient.

The changing surface optical anisotropy was investigated by monitoring the period immediately after the admission of TEG with fixed photon energies over the range 1.5 to 4.0 eV. Examples of the dynamic RAS traces are given in Fig. 3 for BEPs of $3.8 \times 10^{-5}$ mbar (group III) and $10^{-4}$ mbar (group V) at 510°C. This enabled RAS spectra to be derived at various times during the dynamic RAS monitoring. The results are shown in Fig. 4 for the pre-growth and steady-state during growth conditions along with two points in the transient. The spectra indicated as T1 in Fig. 4, which are similar at each temperature, correspond to the peak in the reflectance transient whilst those marked T2 correspond to the feature at 3.6 s in Fig. 3. The similarity in the derived PG and DG spectra (Fig. 4) and the directly determined PG and steady state DG spectra (Fig. 1) is evident.

We have established that at point T1 in the transient the surface is metallic, the metallic nature probably arising from a high Ga content in the surface addlayer. To generate a steady-state Ga rich surface, the temperature was reduced to 300°C with the group III BEP = 0, and the group V supply switched off. RAS spectra were taken as the group V BEP was reduced from $10^{-5}$ to $10^{-8}$ mbar, Fig. 5. At $10^{-5}$ mbar, the RAS spectrum is typical of the $d(4 \times 4)$ reconstruction, indicating that the surface is still As rich. However, with a group V BEP in the range $10^{-7}$–$10^{-8}$ mbar the spectra (Fig. 5) show the development of a trough below 2 eV which deepens and shifts to lower energies as the group V BEP falls, consistent with the development of the Ga rich $4 \times 2$ reconstruction [5]. The T1 spectra of Fig. 4 exhibit some common features...
Fig. 5. RA spectra of GaAs as a function of group V BEP at 300°C with the group III BEP = 0.

with the lower temperature Ga rich spectra (Fig. 5) especially the large trough in the region below 2.0 eV, where the Ga dimers are known to dominate the RAS of GaAs, and the peak developing at \( \sim 2.4 \) eV.

The dynamic RAS measurements suggest that the transient addlayer initially comprises Ga dimers, whereas the reflectivity measurements demonstrate it is initially metallic, at least in terms of its optical properties. It has been shown [13] that when metal atoms are deposited on semiconductor surfaces, metallisation may occur with coverage of between 0.5 and 1 ML. Indeed, there are calculations which indicate that the deposition of \( \sim 1 \) ML of alkali metals on GaAs(0 0 1) produces a metallic surface [14] and that the As saturated GaAs(0 0 1) (2 \times 4) surface, in which six As atoms form three dimers in the unit cell and there are no Ga atoms in the outermost cell, is also metallic [15]. However, we have not found any theoretical work on the nature of the Ga rich GaAs(0 0 1) 4 \times 2 surface. Our results suggest that this surface is metallic also.

It therefore appears that the initial part (\( t < 1.0 \) s) of the transient is due to the rapid dissociation of the TEG molecule leaving Ga on the surface, which is yet to be incorporated as GaAs. Thus, with the group III BEP greater than \( 3.5 \times 10^{-5} \) mbar the net deposition rate of Ga is initially too high, in relation to the As flux, to be incorporated into the growing structure. Decreasing the III/V ratio has the effect of reducing the overall duration of the transient by accelerating the rate of incorporation. Transitory behaviour has been observed in a RHEED study of the MOMBE growth of GaAs with TEG and As \(_2\) as precursors [16] with a high TEG flux. The reconstruction of the transients was Ga rich 4 \times 2 and the authors [16] suggested that Ga was initially deposited as clusters subsequently being incorporated on the arrival of the As. This suggests that there may be a different growth rate over the duration of the transient than in the steady state when equilibrium is established. It was further reported that the growth rate was lower during the transient [16]. By observing the transient in both RAS and R during the growth of GaAs on a GaAs(0 0 1) substrate incorporating a thin layer of AlGaAs, we were able to study the growth rate over the duration of the transient via multiple reflection interference [17]. The results [18] indicate that the growth is slower over the transient in agreement with Ref. [16].

4. Conclusion

At group III BEPs greater than \( 3.5 \times 10^{-5} \) mbar there exists a transient surface state in the initial stages of the CBE growth of GaAs. The surface reconstruction changes over the duration of the transient. Initially it appears to be 4 \times 2 Ga rich and metallic and is due to the deposition of Ga on to the surface faster than it can be incorporated as growth of GaAs is initiated. After about 1 s the Ga begins to be incorporated and after a further few seconds, depending on the III/V ratio, the growth continues as normal.
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References

Be redistribution in InGaAs and InP grown by gas source molecular beam epitaxy

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Abstract

The redistribution of Be during the growth of InGaAs and InGaAs/InP heterostructures by gas source molecular beam epitaxy has been studied using secondary ion mass spectrometry. Be deposition to the InP layer grown next to the highly Be-doped InGaAs layer has been shown to cause significant Be segregation. Although the growth temperature and AsH3 flux dependence of Be redistribution in InGaAs layers can be explained by the substitutional–interstitial (S–I) diffusion mechanism, no enhancement of Be redistribution in InGaAs is observed by n+ doping in nearby layers, inconsistent with previous results of Zn diffusion. The temperature dependence of Be profiles in InP layers follows the S–I diffusion mechanism, however the PH3 flow rate dependence is inconsistent with the model.

1. Introduction

To achieve the full potential of heterojunction bipolar transistors (HBTs) in terms of high-speed performance, rigorous control of doping profiles in the base region is necessary. Specifically, device performance can be improved through heavy p-type doping of the transistor base region to reduce base resistance while maintaining narrow abrupt base-doping profiles to keep the minority-carrier transit time through this region short. For the acceptor-doped base layer, beryllium (Be) and zinc (Zn) have been the conventional dopants used in molecular beam epitaxy (MBE) and organometallic vapor-phase epitaxy (OMVPE) because of their high doping efficiency, good hole transport properties, and acceptable diffusion rate at moderate doping levels. However, for doping levels in the 10¹⁹ cm⁻³ range, concentration-dependent diffusion has been reported to be a serious problem [1]. Be and Zn diffusion in GaAs has been extensively studied for many years [1–6]. A model of diffusing interstitial impurity which is captured on a group-III lattice site has proved successful in explaining results. The substitutional–interstitial (S–I) diffusion mechanism [6, 7] is based on the reaction equation

\[ A^m_+ + V_{Ga}^0 \leftrightarrow A^- + (m + 1)h_0 \]  

(1)

where A stands for a Be or Zn atom with A⁻ being the singly ionized substitutional species and \( A^{m+} \) the interstitial species with charge \( m^+ \). Here, \( V_{Ga}^0 \) is the neutral Ga vacancy, and \( h_0 \) is the hole.

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The diffusivity of interstitial Be or Zn is assumed to be much larger than that of substitutional Be or Zn or of a lattice vacancy.

It is also well known that the diffusion of Zn is enhanced by nearby highly n-type layers, as in HBT structures \([4, 5, 8]\), and this was explained by Deppe \([9]\) in terms of nonequilibrium point defects induced by a combination of surface pinning of the Fermi level and n\(^+\) doping. On the other hand, Be diffusion is suppressed by the presence of nearby n\(^+\) layer [10–12]. A satisfactory model of Zn and Be diffusion should be able to account for these experimental characteristics. The above-described objective is yet to be attained.

Up to now, Be diffusion has been extensively studied in GaAs/AlGaAs systems. There are few reports on Be redistribution in InGaAs/InP.

In this article, to gain a better understanding of the processes involved in Be redistribution in InGaAs and InGaAs/InP heterostructures, we have investigated the effects of growth temperature, group V flux, and n-type doping concentrations in nearby layers by studying secondary ion mass spectrometry (SIMS) depth profiles.

2. Experimental procedure

All studied layers were grown in a gas source molecular beam epitaxy (GSMBE) system. Details of this system were reported in Ref. \([13]\). Elemental Ga and In were used for the group-III growth species derived from effusion cells. 100% AsH\(_3\) and PH\(_3\) were decomposed in a low-pressure thermal cracker cell maintained at 900°C. The substrate temperature was determined using an Incon, Model-V, pyrometer.

In order to investigate the effects of growth temperature, group-V flux, and n-type doping to nearby layer, we grew three types of samples as shown in Fig. 1. Fig. 1a and Fig. 1b show an InGaAs n-i-p-n structure and a n-InP/i-InGaAs/p-InGaAs/n-InP heterostructure. Be- and Si-doping concentrations for these samples were set around \(1 \times 10^{19}\) and \(1 \times 10^{17}\) cm\(^{-3}\), respectively. In the course of growing samples (a) and (b), 400 s growth interruptions were introduced prior to growing the intentionally Be-doped InGaAs layer to raise Be cell temperature. After the Be-doped InGaAs layer was grown, Be cell temperature was lowered by 200 to 650°C and no growth interruption was introduced. Fig. 1c shows the InGaAs pn-junction structure used to study the effect of n-type doping in nearby layers. In this sample Be concentration was set around \(2 \times 10^{19}\) cm\(^{-3}\) and Si concentration was varied from \(7 \times 10^{17}\) to \(3 \times 10^{19}\) cm\(^{-3}\). A 30 nm low Si-doped InGaAs layer was introduced between each highly Be- and Si-doped layer. 200 s growth interruptions were introduced to set the Si and Be temperatures. After

![Fig. 1. Sample structures. (a) InGaAs pn-junction structure. (b) InGaAs/InP double heterostructures. (c) InGaAs stacked pn-junction structure.](image-url)
growing Be-doped layers, no growth interruptions were introduced, while Be cell temperature was lowered by 200°C.

SIMS measurements were provided by Evans East from their Physical Electronics 6600 SIMS system. Cesium ions and oxygen ions were used as primary ions and the primary ion acceleration potential used was 3 and 6 kV, respectively.

3. Results and discussion

3.1. Effects of Be-cell temperature profile

Anomalous Be redistribution towards the surface was observed in samples grown at 460°C by keeping the Be cell temperature constant throughout the growth (samples (a) and (b) in Fig. 2). We found that the extended growth of high-concentration Be-doping layers causes Be deposition even when the Be shutter is closed, as was previously observed by Panish et al. [14], and this unintentional Be doping causes significant Be segregation into the Si-doped InP-layer grown next to them. The term “segregation” will be used for dopants moving predominantly towards the surface during crystal growth. The unintentional Be-doping level is around $1 \times 10^{17}$ cm$^{-3}$, as is seen in the undoped InGaAs layer in Fig. 2. Be accumulation during growth interruption is also observed (sample (b)). The full-width at half-maximum (FWHM) of the Be peak in sample (a) is about 58 nm. Considering the SIMS depth resolution, the measured FWHM is in good agreement with the intended thickness of 50 nm. Therefore, the unintentional Be-doping seems to cause no effect on the top profile of the Be peak. This is probably because the unintentional doping density is more than two-orders of magnitude smaller than the peak concentration. The origin of the unintentional Be-doping is probably the re-evaporation of Be from the shutter.

Therefore, in the course of the experiments, Be cell temperature was lowered by 200 to 650°C before and after the Be-doping layers were grown. When the Be temperature was lowered, a sharp Be profile was obtained on both sides of the intentionally Be-doped region (sample (c)). The asymmetry found in the Be profile (c) is probably due to the well known knock-on effect.

Schubert et al. [15] explained Be surface segregation in MBE grown GaAs by taking into account Fermi-level pinning at the semiconductor–vacuum interface. Using this model, the segregation velocity of the dopants is given by

$$v_d = \frac{D_e e N_A^{2D}}{kT} \frac{1}{e},$$

where $v_d$ is the growth rate and $N_A^{2D}$ is two-dimensional doping density. Suppose the diffusion coefficient of InGaAs is the same as that of GaAs, the diffusion coefficient $D < 10^{-16}$ cm$^2$/s yields a negligible segregation velocity at 460°C. Therefore, for samples grown at low growth temperature, segregation does not dominate as was pointed out by these researchers.

3.2. InGaAs pn-junction structures

The Be-profiles of InGaAs pn-junction samples (Fig. 1a) are shown in Figs. 3 and 4. The influence of growth temperature on Be redistribution in InGaAs pn-junction structures grown using an AsH$_3$ flow rate of 10 sccm is given in Fig. 3. The Si-SIMS profile is also shown as a layer marker. When the growth temperature is set between 400–460°C, sharp Be profiles are obtained on both sides of the doping peaks. When the AsH$_3$ flow rate is reduced to 1.5 sccm, although the substrate side of the Be profile is the same as samples grown with a higher AsH$_3$ flow rate, the surface side tail of the profile shows clear Be-segregation (Fig. 4).

AsH$_3$ flow rate dependence can be explained on the basis of the S–I diffusion mechanism, in which a positively ionized Be interstitial (Be$_i^{+}$) rapidly diffuses until it encounters a group III vacancy site ($V_{m}$), which it may then occupy, thereby becoming negatively charged (Be$_m^{-}$). For an interstitial diffusion controlled process, the criterion of the process is $(m + 1)D_i C_i \gg D_e C_e$, where $m$ is the charge state of the interstitials, $D$ is the diffusion coefficient, $C$ is the diffusion source concentration, and $i$ and $s$ stand for interstitial and substitutional species. The difference in the diffusivities can be as much as six orders of magnitude faster than known
Fig. 2. Be cell temperature dependence of Be-SIMS profiles in gas source MBE grown InGaAs/InP heterostructures. (a) and (b) are samples grown by keeping Be cell temperature high. Growth interruption of 200 s is introduced during InP layer growth of sample (b). (c) A sample grown by setting Be cell temperature 200°C lower to 650°C before and after the Be-doping layer was grown. G.I. stands for growth interruption.

Fig. 3. Growth temperature dependence of Be-SIMS profiles in InGaAs pn-junction structures grown with an AsH₃ flow rate of 10 sccm.
substitutional impurities, therefore this difference in diffusion coefficients is expected to be able to offset the concentration difference which is estimated to be two to three orders of magnitude [15]. This mechanism implies that Be diffusion can be enhanced by decreasing the number of $V_{\text{III}}$. A decrease in $V_{\text{III}}$ can be accomplished by decreasing the $V/\text{III}$ flux ratio during growth, or by raising the growth temperature while keeping the arsenic flux constant.

3.3. InGaAs/InP heterostructures

Figs. 5–7 show the Be-profiles in the InGaAs/InP double-heterostructures shown in Fig. 1b. When the flow rates of AsH$_3$ and PH$_3$ are 6 sccm (Fig. 5), although the Be profiles in InGaAs show no growth temperature dependence, the profile in the InP layers of the sample grown at 490°C is considerably broader. The FWHM of the Be peak in sample grown at 400°C is about 58 nm, which is almost coinciding with the intended thickness of 52 nm. Therefore, it can be concluded that, when the AsH$_3$ flow rate is 6 sccm, the Be-profile can be tailored in InGaAs layers even at higher growth temperature, as is shown in Figs. 3–5, while significant redistribution of Be occurs in InP layers for sample grown at 490°C. When the PH$_3$ flow rate is reduced to 4 sccm while keeping AsH$_3$ flow rate constant, Be redistribution observed in the tail region of sample grown at 490°C is reduced (Fig. 6). The substrate side of the profiles show no temperature dependence. Fig. 7 shows the PH$_3$ flow rate dependence for samples grown at 490°C. In samples grown with PH$_3$ flow rates of 6 and 8 sccm, the Be profiles in the InP layers broaden inconsistently with the above-mentioned S–I diffusion mechanism. From Figs. 5–7, it is confirmed that, for Be redistribution in InP, the PH$_3$ flow rate dependence is inconsistent with the proposed S–I diffusion model, although the temperature dependence follows the S–I diffusion mechanism. From the facts that the thermal cleaning of InP in GSMBE under high PH$_3$ flow rate results in roughened surface, and exposure to H$_2$ plasma leads to the roughened surface by preferential desorption of P [16], the PH$_3$ flow rate dependence of Be redistribution seems to be due to hydrogen assisted preferential desorption of P at the InP surface. Further investigations, particularly PH$_3$ flow rate dependence of the InP surface structure is necessary to understand the Be redistribution characteristics in InP.

3.4. n-type doping dependence

It has been reported that an abnormal Zn redistribution into the AlGaAs emitter and the GaAs collector in the HBT structure takes place when all the layers except the base are doped with Si [4, 5]. In the case of Zn diffusion in InGaAs/InP heterostructures grown by OMVPE, Kurushima et al. [17] pointed out that, at lower growth temperature, high n$^+$ doping in the subcollector layer in a heterojunction bipolar transistor structure is the main cause of enhanced Zn diffusion. The basis for explaining Zn diffusion is that a high concentration of nonequilibrium group-III interstitials generated in the n$^+$ layer move toward both the surface and the underlying layers, and kick-out the Zn from substitutional to interstitial sites via the reaction

$$Zn_{\text{III}} + \text{IIIi} + 2h^+ \leftrightarrow Zn_{\text{I}}^+.$$ 

The interstitial Zn then rapidly diffuses towards both the surface and the underlying layers.

We have examined the effects of Si-doping in adjoining layers to Be-doped layers in the InGaAs stacked pn-junction structure shown in Fig. 1c. Si doping density was varied from $7 \times 10^{17}$ to $3 \times 10^{19}$ cm$^{-3}$. No Be diffusion was observed even for layers with the highest Si doping concentrations (Fig. 8). The Fermi level in InGaAs is reported to be pinned close to the conduction band [18]. On the basis of Deppe's model [9], it is clear that if the Fermi level is not pinned close to midgap at the surface, then the number of group-III interstitials in n-type layer ($N_{\text{I}_3}^+$) is same as that in intrinsic layer ($N_{\text{I}_3}$). Therefore, the number of interstitial Be is reduced and Be redistribution should be suppressed.

4. Summary

In conclusion, we have shown that unintentional Be doping to the InP layer grown next to the highly
Fig. 4. AsH$_3$ flow rate dependence of Be-SIMS profiles in InGaAs pn-junction structures grown at around 440°C.

Fig. 5. Growth temperature dependence of Be-SIMS profiles in InGaAs/InP double heterostructures grown with AsH$_3$ and PH$_3$ flow rates of 6 sccm.
Fig. 6. Growth temperature dependence of Be-SIMS profiles in InGaAs/InP double heterostructures grown with AsH₃ and PH₃ flow rate of 6 and 4 sccm.

Fig. 7. PH₃ flow rate dependence of Be-SIMS profiles in InGaAs/InP double heterostructures grown at 490°C. AsH₃ flow rate was set at 6 sccm. (a), (b) and (c) are samples grown with PH₃ flow rates of 4, 6 and 8 sccm, respectively. The Si profile shown in the figure is taken from the sample grown with a PH₃ flow rate of 4 sccm.
Be-doped InGaAs layer causes significant Be redistribution, and we have also shown that the growth temperature and AsH$_3$ flux dependence of Be redistribution in InGaAs layers can be explained by the S–I diffusion mechanism. However, in InP/InGaAs heterostructures, although the temperature dependence of the Be profile follows the S–I diffusion mechanism, the PH$_3$ flow rate dependence is inconsistent with the model. Regarding the effects of n$^+$ doping in nearby layers, no enhancement of Be redistribution in InGaAs was observed, which is inconsistent with previous results. This can be explained in the framework of Depp’s model if the Fermi level of InGaAs is pinned close to the conduction band.

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References

Growth temperature-dependent conduction-type inversion of C-doped InGaAs grown by chemical beam epitaxy

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Abstract

We report for the first time on the growth temperature-dependent conduction-type transition of carbon-doped InGaAs grown by chemical beam epitaxy (CBE) using triethylgallium (TEGa), trimethylindium (TMIn), precracked arsine (AsH\textsubscript{3}), and unprecracked monoethylarsine (MEA). The hole concentration of InGaAs layers decreased with increasing growth temperature and conductivity changed from p to n at around 450°C. The conduction-type inversion of InGaAs layer is attributed to the surface segregation of indium and decomposition and incorporation kinetics of group-III source materials on the surface depending on the growth temperatures. The variations of indium composition and the V/III ratio did not affect the growth temperature-dependent conduction-type inversion of carbon-doped InGaAs. This behavior is attributed to the effect of surface segregation of indium on the carbon incorporation during the growth of InGaAs.

1. Introduction

The behavior of dopants in semiconductor is of basic interest and its understanding is fundamental for device applications. Electrical and optical device performances are influenced by properties of dopants, such as the incorporation efficiency or activation energy of free carriers. Carbon, a p-type dopant in the GaAs/AlGaAs material system, has been of great interest and widely investigated [1–4]. GaAs films with carrier concentrations of $1 \times 10^{20}$ and $1 \times 10^{21}$ cm$^{-3}$ have been demonstrated by metalorganic chemical vapor deposition (MOCVD) [5] and metalorganic molecular beam epitaxy (MOMBE) [4] or chemical beam epitaxy (CBE), respectively. The main advantage of carbon in these materials is the low diffusivity [6,7] that accompanies the high dopant activity, permitting thin, highly conducting layers to be used in devices such as the heterojunction bipolar transistors (HBTs). However, the behavior of carbon as an intentional dopant in III–V materials has not been sufficiently understood. Until recently, carbon had been regarded as an undesirable residual impurity in III–V materials for the MOMBE, CBE, and MOCVD methods.

InGaAs is a promising material for high-speed device applications owing to its high electron
mobility and high peak electron drift velocity. The behavior of carbon in InGaAs is not well characterized at present, but the use of a p-type carbon-doped InGaAs base in InP/InGaAs HBTs would be of great interest if high doping levels can be obtained, and if carbon similarly shows low diffusivity in InGaAs. However, it is more difficult to achieve a high p-type doping level in InGaAs with carbon, because the carbon incorporation efficiency decreases with an increase in indium mole fraction [8, 9], and the amphoteric behavior of carbon produces n-type or p-type InGaAs, depending on growth conditions [10]. Up to now, p-type In_{x}Ga_{1-x}As was obtained only when trimethylgallium (TMGa) together with solid In and solid As, CCl_{4}, or CBr_{4} was used for carbon dopant sources. Also, the p-n conversion occurs at x = 0.8 for InGaAs films grown by MOMBE [11], whereas it occurs at x = 0.6 for those grown by conventional molecular beam epitaxy (MBE) with a heated graphite filament as a carbon source [10]. This paper reports the growth temperature-dependent conduction-type transition of carbon-doped InGaAs on GaAs grown by CBE using trimethylindium (TMIn), triethylgallium (TEGa), precracked arsine (AsH_{3}), and unprecocracked monoethylarsine (MEAs). To the best of our knowledge, no work has been known to report on the growth temperature-dependent conduction-type inversion of carbon-doped InGaAs. In our previous work, p-type carbon-doped InGaAs was successfully grown at low growth temperature by CBE using TEGa, TMIn, and MEAs [12]. In this work, we extended our investigation on the effect of hydride source gases, indium compositions, and V/III ratios on the growth temperature-dependent conduction-type inversion of carbon-doped InGaAs layers.

2. Experimental procedure

Growth of InGaAs was done in a modified CBE apparatus which consists of an ultrahigh vacuum growth chamber and a gas source control system [13]. TMIn, TEGa, precracked AsH_{3}, and unprecocracked MEAs were used as the source materials. All growths of InGaAs were carried out on Cr-doped GaAs(100) with 2° off toward (110). Prior to the growth, the substrate was heated to remove adsorbed impurity and surface oxide layers. The growth temperature of InGaAs was in the range of 340–550°C, as measured by optical pyrometer. The In compositions of grown layers, determined by photoluminescence (PL) and double-crystal X-ray diffraction (DCXD), agreed to within 5%. Layer thicknesses were determined by scanning electron microscope (SEM) on stained and etched cleavage planes. The epilayer thicknesses used in the experiment were 0.5 to 1.5 μm. Some of the samples had some surface morphology due to the lattice mismatch. The carrier concentration and mobility were determined by van der Pauw–Hall effect measurement at room temperature. The Hall samples were cleaved 6 × 6 mm² squares with indium ohmic contacts at the sample corners.

3. Results

Fig. 1 shows the growth temperature-dependent carrier concentration of InGaAs layers obtained from van der Pauw method. The In composition of InGaAs layers were about 0.2. The hole concentration of InGaAs using TEGa, TMIn, and MEAs decreased with increasing growth temperature and the conductivity type changed from p to n at around 450°C. Above 460°C, the conduction type of grown layer shows n and the electron concentration of InGaAs layer decreases with increasing growth temperature. As shown in Fig. 1, the growth temperature-dependent type transition of InGaAs layer with precracked arsine is similar to that of InGaAs with MEAs. This suggests that group-III source material is influencing the growth temperature-dependent type transition rather than group-V source gases in the growing film. In Fig. 1, two different characteristics of carbon dopant consistently explains the behavior of carbon incorporation in InGaAs. The average bond strengths for Ga–C_{2}H_{6}, As–CH_{3}, and In–CH_{3} are 57, 55, and 47 kcal/mol, respectively [14]. Although these values do not represent actual binding energies for precursors on the epilayer surface during growth, the relative amplitudes are considered to be still...
valid in the comparison of the binding energies between carbon and metals. Below 450°C, the observed p-type conduction is due to the carbon which may be incorporated as a Ga–C bond into the growing layer and can be activated effectively at As site. The binding energy for Ga–C is larger than that for As–C. This may be attributed to the fact that carbon preferably occupy As sites. Above 460°C, a conductivity inversion from p- to n-type was observed without a much variation of indium composition in InGaAs grown layers. It was mentioned in a previous work [15] that the dissociation kinetics of precursors is not changed during growth of InGaAs at these temperatures. It is well known that even small amounts of lattice indium have such a pronounced effect on the growth characteristics of InGaAs. This appears to stem from the occurrence of surface segregation of indium. Although indium is fairly immobile in the lattice at low temperatures, rapid segregation from bulk to surface occurs as the substrate temperature is raised above 450°C [16]. This phenomenon results in an increase in the surface concentration of indium, until it decreases at much higher temperatures as a result of indium evaporation [17]. As a result, the surface adatom population of indium is increased with growth temperature. As mentioned above, the binding energy of In–C is smaller than that of As–C. This should cause carbon atoms to preferably occupy indium sites at the growth front of InGaAs layer. If the conduction-type inversion is originated from the self-compensation of carbon in InGaAs, a reduction of carrier concentration and mobility should be observed around the conduction-type inversion border. But, a significant reduction of carrier concentration was not observed at this growth temperature region, where the conduction-type inversion occurs, as shown in Fig. 1. These results imply that the growth temperature-dependent conduction-type inversion is not due to the self-compensation of carbon in this material. Our result suggests that the dissociation kinetics of precursors and the surface segregation of indium critically change the lattice site of carbon in the InGaAs material system, depending on the growth temperature.

Fig. 2 shows the dependence of the indium composition in InGaAs layer on the growth temperature. The InGaAs layers were grown on the GaAs substrate with beam equivalent pressure flux ratios of TEGa to TMIn of 7 : 3. Unlike the MOMBE-grown samples [11], severe temperature-dependent variations of the indium composition were not observed in CBE-grown InGaAs using unprecracked MEAs. It may be due to the different growth mechanism of InGaAs using unprecracked MEAs compared to that of elemental arsenic. Here, the surface segregation of indium depending on the growth
temperature does not seem to significantly influence the bulk composition of indium in InGaAs grown layers. These results indicate that the inversion of conductivity type with increasing growth temperature as shown in Fig. 1 is not resulted from the variation of indium composition.

Fig. 3 shows the dependence of free carrier concentration of carbon-doped In$_x$Ga$_{1-x}$As on the InAs mole fraction grown at the temperature of 400°C. The beam equivalent pressure flux ratio of TEGa to TMIn was varied from 0 to 1 with fixed MEAs beam flux and substrate temperature. The conductivity-type changes from p to n between $x = 0.3$ and 0.45. The p-n inversion occurs between $x = 0.3$ and 0.45 for the In$_x$Ga$_{1-x}$As layers grown by CBE using unprercracked MEAs, whereas it occurs at $x = 0.6$ and 0.8 for those grown by MOMBE [11] and solid-source MBE with heated graphite filament [10], respectively. It is worth noting that the hole concentration of InGaAs layers grown by CBE using TEGa, TMIn, and unprercracked MEAs is comparable to that of the carbon-doped InGaAs layers grown by conventional MBE with heated graphite filament. The conduction-type inversion with indium composition usually accompanies the decrease of free carrier concentration and mobility [10]. As carbon can occupy both donor and acceptor sites, strong carrier compensation causes a large increase in ionized impurity scattering. A significant reduction of carrier concentration is shown around the border as shown in Fig. 3. These results imply that the indium composition-dependent conduction-type inversion is due to the self-compensation of carbon in InGaAs. Therefore, the mechanism of conduction-type inversion with indium composition is different from that with growth temperature. A reduction of electron mobility was observed in InAs samples. We believe that it results from the defects occurring in InAs due to the large lattice mismatch between GaAs and InAs. The growth temperature-dependent conduction-type inversion of InGaAs layers occurred at the indium compositions less than 0.3, as shown in Figs. 1 and 2. Therefore, the growth temperature-dependent conduction-type transition is not attributed to an increase of the indium composition with growth temperature.

Fig. 4 shows the dependence of carrier concentration of InGaAs epilayers on V/III ratio obtained at the growth temperature of 400°C. All samples grown under the conditions of various V/III ratios and low growth temperature show p-type conduction, which is consistent with the results shown in Fig. 1. The surface segregation of indium during the growth of InGaAs is suppressed by increasing the arsenic pressure [17]. As shown in Fig. 4, samples under the condition of low V/III ratios were
measured as p-type conduction. The elementary process of carbon incorporation at low temperatures is governed by dissociation kinetics of source materials on the surface and the effect of surface segregation of indium on the carbon incorporation is negligible in this region. Therefore, the growth temperature-dependent conduction-type transition is not attributed to the variation of V/III ratio. Furthermore, the carrier concentration and mobility of grown layer depended distinctly on the partial pressure of MEAs. Therefore, it is believed that the presence of partially cracked arsine radical and hydrogen atom dissociated from MEAs on the growing surface can inhibit the incorporation of hydrocarbon, since the surface can play a key role in inhibiting hydrocarbon pyrolysis due to an available surface population of arsine radical and atomic hydrogen.

4. Summary

We have examined for the first time the growth temperature-dependent conduction-type inversion of carbon-doped InGaAs on GaAs grown by CBE using TMIn, TEGA, precracked AsH3, and unprecked MEAs. A conduction-type inversion from p to n is found to occur in the InGaAs layers with increasing growth temperature. This phenomenon can be understood by considering surface segregation of indium during InGaAs growth at high growth temperatures and different decomposition and incorporation kinetics of carbon-containing source materials on the surface at low growth temperatures. From the investigation on the effects of V/III ratio and indium composition, the growth temperature-dependent conduction-type inversion is not attributed to the influence of V/III ratio and the variation of indium composition. From this work, it is suggested that the surface segregation of indium has a significant effect on the dopant incorporation and activation behavior in the growth of indium-containing compound semiconductor, for example InGaAs or InAlAs. These results suggest that p–n junction of indium-containing materials can be fabricated by the control of growth temperature without introduction of dopants.

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References

Selective area etching of III–V semiconductors using TDMAAs and TDMASb in metalorganic molecular beam epitaxy chamber

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Abstract

Selective area etching of SiO₂-masked GaAs using trisdimethylaminoarsenic (TDMAAs) is studied. GaAs substrates are partly masked with stripe shaped SiO₂ films along the [0 1 1] and [0 1 1] directions. Cross-sectional scanning electron microscopy observation shows that the etched shape depends on both the direction of the mask and substrate temperature, and is almost independent of the TDMAAs flow rate. At low substrate temperatures, the (1 1 1)B surface is preferentially formed and a V-shaped groove is formed for the [0 1 1] mask direction. With increasing substrate temperature, the V-groove changes to the channel with vertical side walls due to the drastic increase of the etching rate of (1 1 1)B. For the [0 1 1] mask direction, the vertical (0 1 1) plane is formed at all temperatures. It is found that etched shapes are determined by the difference of etching rates and that etching rates are influenced by the crystal structure on the surface. On the other hand, the etching rate of GaSb by TDMASb is independent of substrate surface orientation. This is because the bond strength of GaSb is weak and the etching proceeds along the direction of molecular beams. Enhanced etching rate is observed for InP by TDMAAs, which is explained by the formation of thin InAs layer on InP due to the irradiation of As released from TDMAAs.

1. Introduction

Selective area etching and regrowth using patterned substrates are emerging as a very promising technique for the fabrication of nanoscale structures, such as quantum dots or wires [1]. Considering these multiple processes, the clean environment is required since the optical and electrical properties of devices will be affected by a contamination due to an air exposure or surface residues. Furthermore, the accurate control of etched profiles with a minimal damage is necessary. For this purpose, metalorganic molecular beam epitaxy (MOMBE) is a useful method because the etching and regrowth can be conducted in the same vacuum chamber by switching from etching to growth and vice versa as described below.

An As precursor, trisdimethylaminoarsenic [As[N(CH₃)]₃, TDMAAs], has been proposed as an alternative candidate in the MOMBE growth of
GaAs and AlGaAs [2–9] instead of the extremely toxic and high thermally stable hydride-like AsH₃. Since TDMAs releases As atoms even at low temperatures (300–450°C) compared with other precursors [5, 6], thermal precracking in a gas cell before the supply on the substrate surface is not needed. This leads to a low carbon incorporation by TDMAs in the growth of GaAs with both triethylgallium (TEGa) and trimethylgallium (TMGAs) [3, 7].

It was also reported that GaAs and GaSb substrates are etched by TDMAs and by tris-dimethylaminoantimony [Sb[N(CH₃)₂]₃, TDMASb] [9–12]. The etching by these metalorganics was considered to be due to the reaction of surface Ga atoms with TDMAs and TDMASb related species (probably, amine species) resulting in the formation of volatile Ga species although the exact mechanism has not been understood yet. The etching proceeds in the monolayer-by-monolayer mode, which was confirmed by the reflection high-energy electron diffraction (RHEED) intensity oscillations [12].

For the regrowth, TDMAs is also a suitable As source because the selective area epitaxy of GaAs was easily achieved by the combination of TEGa and TDMAs at lower temperatures than by the use of TEGa and solid As [8].

In this paper, we report on the metalorganic molecular beam etching of SiO₂-masked GaAs substrates by TDMAs and discuss the mechanism of the formation of etched shapes. In addition, the dependence of etching characteristics on etched substrate (GaSb, GaAs, GaP, InP) and metalorganic molecules (TDMAs, TDMASb) is also described.

2. Experimental procedure

The etching experiments were conducted using a modified ULVAC MBE 300 apparatus equipped with gas cells for TDMAs and TDMASb. The growth chamber was evacuated by an oil diffusion pump with a liquid nitrogen trap, yielding a pressure of less than 5 × 10⁻⁶ Torr during etching. Non-precracked TDMAs and TDMASb were directly introduced into the growth chamber through mass flow controllers (MFCs) without any carrier gases. Source cylinder, gas line and gas cell were heated at proper temperatures in order to avoid their condensation and to get reasonable gas flow rates. The substrates used were (1 0 0)-GaAs partly masked with SiO₂ films for selective area etching, and non-masked (1 0 0) and (n 1 1) \( \{ n = 5, 4, 3, 2, \frac{5}{2}, 1 \} \), GaAs and GaSb. Stripe mask patterns aligned along [0 1 1] and [0 1 1] directions were formed by a conventional photolithography and a wet chemical etching. The substrate temperature \( T_{sub} \) was monitored by an optical pyrometer calibrated by the melting point of InSb at 525°C. The substrate holder was rotated intermittently during etching. Cross-sectional etched profiles were studied by the scanning electron microscopy (SEM).

3. Results and discussion

3.1. Selective area etching on SiO₂-masked GaAs substrate

Selective area etching using TDMAs was performed on SiO₂-masked GaAs substrates. Fig. 1 shows typical cross-sectional SEM photographs of etched substrates with a stripe mask aligned along the [0 1 1] and [0 1 1] directions. It was found that etched shapes were influenced by the direction of a stripe mask and \( T_{sub} \). At \( T_{sub} \) lower than 550°C, for the [0 1 1] mask direction, only (1 1 1)B surfaces were observed and the V-shaped groove was formed (Fig. 1a). With elevating \( T_{sub} \) up to 600°C, (1 1 1)B surfaces disappeared and only vertical (0 1 1) side walls emerged (Fig. 1b). On the other hand, for the [0 1 1] mask direction, the etched side walls were a combination of (1 1 1)A and vertical (0 1 1) at low temperatures (Fig. 1c), and in addition to these surfaces (4 1 1)A facets emerged at \( T_{sub} \) higher than 600°C (Fig. 1d). But etched shapes were almost independent of a TDMAs flow rate. These results indicate the possibility of controlling the etched profiles by selecting the etching condition.

In order to study the mechanism for the formation of etched side walls, the ER variations with \( T_{sub} \) were measured for (1 0 0), (4 1 1), and (1 1 1)
surfaces as shown in Fig. 2. These (1 1 1) substrates were mounted on a same Mo holder, and the etched thickness (or etching rate (ER)) was measured by the step height between the etched area and the masked one with a small Ta metal strip. As can be seen, ERs were increased with $T_{\text{sub}}$ for all surfaces. Our previous work [12] showed that the etching occurs when amino-species react with surface Ga atoms, which are formed by the desorption of outermost As atoms from the surface. This suggests that the ER increases at higher $T_{\text{sub}}$ due to the the low surface coverage of group V atoms, where the desorption of As is enhanced.

Now, we consider the dependence of the ER on the substrate orientation. At low $T_{\text{sub}}$, the ER of the (1 1 1)B was smallest, indicating that the (1 1 1)B is
difficult to be etched. However, by increasing $T_{sub}$ it drastically increased and approached those for other surfaces. On the other hand, that of the (1 1 1)A was slightly smaller than that of the (1 0 0) at all $T_{sub}$, but did not increase above 600°C. These differences in ERs are explained by taking into account the crystal structure of three surfaces, i.e., (1 1 1)A, (1 0 0) and (1 1 1)B. Although the ERs for other (n 1 1) surfaces were not considered here, they are assumed to be similar to those for the (100) and (1 1 1) surfaces because the (n 1 1) surfaces are composed of (1 0 0) and (1 1 1) surfaces. The (1 1 1)A and (1 1 1)B are terminated by Ga and As atoms, respectively. That is, As atoms on the (1 1 1)B surface are strongly bound with three back-bonds to the underlying Ga atom, whereas those on the (1 0 0) and (1 1 1)A have two and only one back-bonds, respectively. At low $T_{sub}$ the surface coverage of As on the (1 1 1)B surface is highest due to the strong strength of As-bonds and a little desorption of As, resulting in the slow ER. From the above discussion, the surface coverage of As on the (1 1 1)A is lowest and the ER is considered to be the highest. However, the ER of (1 1 1)A surface was lower than that of the (1 0 0) surface. This discrepancy might be due to the strong bonding of Ga atoms on the (1 1 1)A surface, where Ga atoms have three back-bonds to the underlying As atoms. At higher $T_{sub}$ above 600°C, the RHEED pattern indicated that the desorption of As molecules is enhanced and that the surface is terminated by the Ga surface or covered with Ga droplets. Therefore, ERs are independent of the surface coverage of As and depend only on the strength of Ga atoms. It is also understood that the ER of (1 1 1)B, where Ga atoms are bound to surface As atoms with only one bond, drastically increases with increasing $T_{sub}$ and that the ER of (1 1 1)A, where Ga atoms with three back-bonds, is lower than those of the other surfaces.

Now, we discuss the dependence of etched profiles on the etching condition (Fig. 1). For the [0 1 1] mask direction, the (1 1 1)B surface are selectively formed due to the low ER of the (1 1 1)B at low $T_{sub}$. However, as the $T_{sub}$ increases, ER for the (1 1 1)B plane increases rapidly and approaches those for the other planes. Therefore, at higher $T_{sub}$, vertical (0 1 1) side walls are formed because the etching proceeds along the direction of molecular beams which is perpendicular to the surface. On the other hand, for the [0 1 1] mask direction, the ER of the (1 1 1)A plane is slightly smaller than those for the other surfaces at all $T_{sub}$. Therefore, vertical etching is mainly expected, but (1 1 1)A planes are also formed at the bottom region of the etched walls due to the low etching rate of (1 1 1)A. But (4 1 1)A planes also appeared at the base of the walls. This is not clear, but might be related to the reflection of TDMAAs molecules from the (1 1 1)A planes inside the etched channel or to the higher stability of the (4 1 1)A surface [13].

3.2. Comparison of etching rates between GaAs and GaSb

We investigated the etching process on GaAs and GaSb substrates. The ER of GaAs by TDMAAs increased with $T_{sub}$ for all surface orientations. In addition, the (1 0 0) surface was etched with the highest ER and the ER was decreased in accordance with an increase of the tilt angle $((1 0 0) > (4 1 1) > (1 1 1))$ as shown in Fig. 2. From the decomposition study of TDMAAs, it is completely pyrolyzed above 350°C [5, 6]. Since etching experiments here were conducted at $T_{sub}$ higher than 450°C, the decomposition efficiency is almost unity, producing no orientation dependence. Therefore, ERs on high index substrates are considered to be determined by the crystal structure of the surface.

Fig. 3 shows the substrate orientation (tilt angle) dependence of the ER of GaSb by TDMAAs as a function of $T_{sub}$. As can be seen in this figure, the ER is almost independent of the tilt angle. Solid circles in Fig. 4 show the ER of (1 0 0)-GaSb by TDMAAs. The ER of GaSb saturated at $T_{sub}$ 550°C, whereas that of GaAs by TDMAAs increased with $T_{sub}$ as shown in Fig. 4. This saturation of the ER is caused by the weak bond of GaSb compared with that of GaAs. In the case of GaSb, Sb atoms are easily desorbed from the top surface and Ga droplets exist on the Sb sublayer at $T_{sub}$ higher than 500°C [14]. On the surface with Ga droplets, the ER depends neither on the surface coverage of Sb, nor on the tilt angle because TDMAAs easily reacts with Ga droplets on the surface under the Sb free condition. Furthermore,
the ER is limited by the supply of TDMAb, resulting in the saturation of the ER.

3.3. Etching of different substrates by TDMAAs and TDMAb

We studied the influence of the ER on the etched material, (1 0 0)-GaSb, GaAs, GaP and InP substrates. Fig. 4 shows the ER dependence of various substrates etched by TDMAb on the Tsub. All substrates were etched and the order of the ER is GaSb > InP > GaAs > GaP. In order to explain this result, we consider the difference of bond strength. The bond of GaSb (45.9 kcal/mol) is the weakest among them, resulting in the highest etching rate, while the etching rate of GaP is relatively low because of the strongest bond strength (54.9 kcal/mol) [15]. Therefore, the order of bond strength, which is Ga–Sb < In–P (47.3 kcal/mol) < Ga–As (50.1 kcal/mol) < Ga–P, corresponds to the order of the ER. However, the ER of InP is higher than that of GaSb when TDMAAs was used for etching instead of TDMAb as shown in Fig. 5. The SEM observation indicated that surface morphologies of InP were rough after etching. In solid source MBE, it has been known that the surface of InP is replaced by thin InAs layers by the irradiation of As beam due to the high vapor pressure of P [16]. Hence, the surface of InP is exchanged into InAs by As atoms released from TDMAAs, followed by etching by amino-groups. Therefore, the ER of InP substrates increases due to the etching of InAs instead of InP.

4. Summary

Selective area etching of SiO2-masked GaAs using TDMAAs was studied. Cross-sectional SEM
observation showed that the etched shape depends on both the direction of the mask and the substrate temperature, and is almost independent of the TDMAAs flow rate. It was found that these etched shapes are determined by the ER for each surface and that the ER is influenced by the crystal structure on the surface. Therefore, it is possible to control the etched profiles by choosing etching conditions. TDMAAs is not only a promising arsenic source for growth but also is a quite useful in situ etching gas source. In contrast, the ER of GaSb etched by TDMAAsb was independent of the crystal structure and was similar for all high index substrates. This is due to the weak bond strength of GaSb. When the InP substrate was etched by TDMAAs, the surface of InP was replaced by thin InAs layers due to the irradiation of As released from TDMAAs and the growth rate of InP was enhanced.

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References

Indium surface segregation during chemical beam epitaxy of Ga$_{1-x}$In$_x$As/GaAs and Ga$_{1-x}$In$_x$P/GaAs heterostructures

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Abstract

We report on the investigation of In surface segregation in GaInP/GaAs and GaInAs/GaAs heterostructures grown by chemical beam epitaxy (CBE). Owing to the peculiarities of CBE growth, it is shown that In segregation can be quantitatively evaluated in real-time by using reflection high-energy electron diffraction (RHEED). A segregation coefficient and its variation with the growth temperature is extracted from the RHEED data. It is used to determine the In composition profiles at the interfaces of GaInAs/GaAs quantum well (QW) structures as a function of the growth temperature. A good agreement is found between optical transition energies calculated from these profiles and the experimental photoluminescence (PL) energies. PL energy shifts observed in GaInP/GaAs QW's as a function of the growth temperature are qualitatively explained by In segregation.

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Keywords: Chemical beam epitaxy; Surface segregation; Photoluminescence; Ga$_{1-x}$In$_x$P; In$_x$Ga$_{1-x}$As

1. Introduction

It is now recognized that surface segregation is a severe limitation to the building of perfectly abrupt III–V semiconductor interfaces. However, most of the available data are mainly concerned with Ga$_{1-x}$In$_x$As/GaAs grown by standard solid source molecular beam epitaxy (MBE) and only a few results have been reported for other III–V materials [1] as well as for different growth methods such as chemical beam epitaxy (CBE) [2–4]. Also, only scarce reports concern real-time investigations during growth [5]. In a previous paper, we have reported on the study of In surface segregation during CBE growth of Ga$_{1-x}$In$_x$P/GaAs heterostructures [6]. We have demonstrated that In segregation at both normal and inverted interfaces can be evidenced in real-time by reflection high-energy electron diffraction (RHEED) via growth rate variations observed in the first stages of the growth. In the present paper, we investigate in the same way the temperature dependence of the In segregation in GaAs. The validity of our method of in situ determination of In

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segregation is demonstrated by correctly predicting from the RHEED data the transition energies experimentally observed by photoluminescence (PL) in GaInAs/GaAs QWs grown at various temperatures. Finally, the variation of PL energies of GaAs/GaInP QWs as a function of growth temperature is discussed in the framework of In segregation.

2. Experimental details

Growth was carried out in an ultra-high vacuum reactor especially designed for CBE (RIBER-EPINEAT) and which has been described elsewhere [7]. Triethylgallium (TEG) and trimethylindium (TMI) were used as group III-element precursors. Group V-element fluxes were provided by high temperature cracking (950°C) of AsH₃ and PH₃. All the samples were grown on GaAs (0 0 1) substrates. The temperature was controlled using an infrared pyrometer calibrated with the melting point of InSb (525°C). For the GaInP/GaAs heterostructures, GaInP was grown at 550°C with a growth rate of 0.92 monolayers per second (ML/s), while GaAs was grown at various temperatures with a constant growth rate of 0.82 ML/s. The temperature was varied during the growth interruptions at the direct and inverted interfaces, while the surface was stabilized under cracked PH₃ and AsH₃ for GaInP and GaAs, respectively. The growth of GaInP on GaAs was performed by switching all the precursors in the same time, in order to minimize the formation of GaP at this interface [8]. At the GaAs on GaInP interface, the growth was interrupted for 1 s under cracked AsH₃ leading to a partial substitution of the phosphorus atoms of the surface layer by arsenic ones [9]. However, it has to be noted here that different growth interruption sequences were investigated at the beginning of this study in order to check that the growth interruption procedure has no influence on the growth rate of the following layers. Growth at the interfaces of GaInAs/GaAs heterostructures is less critical, so standard growth interruptions were performed (30 s growth interruption under As₂ flux at both interfaces) in order to smooth the interfaces between the two materials. Photoluminescence was excited with a 4880 Å line of an argon laser and measured at 9 K in a closed-cycle He cryostat.

3. Results

3.1. Correlation between growth rate transient and In surface segregation

During the early stages of the growth of GaInP on GaAs, the growth rate deduced from RHEED specular beam intensity variation decreases before reaching a steady state value (Fig. 1a). With the assumption that the decomposition of TEG, and as a consequence the incorporation rate of Ga, is decreased by the presence of In on the growth front, this particular feature of the RHEED oscillatory period can be explained by a gradual increase of the In composition at the beginning of the growth up to a steady state value reached after few monolayers [2, 6]. The observed growth rate transient is in fact the signature of In surface segregation. At the inverted interface, i.e. GaAs on GaInP (Fig. 1b), an increase of the growth rate is now observed at the beginning of the growth. It can be explained in the same way by considering now the incorporation of the surface segregated excess In atoms at the GaInP surface into the GaAs growing layer [6].

The In profile calculation from these data is straightforward assuming that the growth rate

![Fig. 1. Growth rate variations deduced from RHEED specular beam intensity oscillations of (a) Ga₀.₃₅In₀.₆₅P on GaAs and (b) GaAs on Ga₀.₃₅In₀.₆₅P. The growth temperature is 550°C.](image-url)
difference between one of the layers in the transient regime (n-th layer) and the steady state growth rate is proportional to the difference of In surface population of the (n-1)th layer and layers at the steady state. The simple model of Muraki et al. [10] is then used to quantify the segregation process. This model needs the determination of two experimental data, \( x_0 \) and \( x_{ss} \), which are respectively the In bulk and surface layer compositions when the steady state of GaInP growth is reached. The \( x_{ss} \) value is assumed to be given by the monolayer fraction of InP which is necessary to predeposit onto GaAs in order to eliminate the growth rate transient of GaInP [5, 11, 6]. It allows us to determine the exchange coefficient \( R \) in GaInP, which is the fraction of In atoms on the topmost layer segregating to the next layer. It can be obtained following the expression:

\[
\frac{\Delta r_n}{\Delta r_{n-1}} = \left( \frac{x_{ss} - x_0 \sum_{i=0}^{n-1} R^i}{x_{ss} - x_0 \sum_{i=0}^{n-1} R^i} \right)
\]

where \( \Delta r_n \) is the growth rate difference between the \( n \)-th layer and layers at the steady state.

With the previous assumptions, \( R \) for GaAs is extracted from the growth rate variation \( \Delta r_n \) between the \( n \)-th layer and pure GaAs: \( R = (\Delta r_n/\Delta r_{n-1})^{1 - n'} \).

We have applied the same method for the In segregation in GaInAs/GaAs by considering the growth rate transient observed at both interfaces through the RHEED specular beam intensity oscillations.

### 3.2. Influence of the growth temperature

In this part, we first investigate the growth temperature \( T_g \) dependence of the In surface segregation in GaAs grown on Ga\(_{0.45}\)In\(_{0.55}\)P. For this study, the GaInP growth conditions were kept constant (550°C and 0.92 ML/s) and only the GaAs growth temperature was varied. The \( R \) values corresponding to the In surface segregation in GaAs grown on GaInP as a function of \( T_g \) were determined as discussed above and are shown in Fig. 2 (closed squares). Between 500°C and 560°C, these values are well fitted by an Arrhenius law, indicating that the segregation process has a temperature dependent behavior with an activation energy of \( \approx 0.23 \) eV. In the low temperature range (below 500°C), \( R \) becomes more sensitive to the growth temperature. This behavior is similar to the one observed by limura et al. [2] for the GaInAs growth rate variation as a function of the temperature. It can be explained by an increasing coverage of the growth front by alkyl species when the growth temperature is decreased. The growth front is in fact encapsulated by the remaining alkyl radicals which may finally act as surfactant-like species and decrease the In surface segregation phenomenon [12, 13].

The \( R \) values have been determined as a function of the growth temperature for the growth of GaAs on Ga\(_{0.45}\)In\(_{0.55}\)As and are also reported in Fig. 2 (closed circles). This figure clearly shows that, as expected, there is no difference in the segregation process of In when growing GaAs on top of GaInP or on top of GaInAs layers.

The knowledge of the temperature dependence of \( R \) can be very useful to predict the transition energy of QWs. In fact, the best way to demonstrate this, but also to validate our in situ method for
Fig. 3. Experimental PL transition energies of a 7 ML wide Ga$_{0.65}$In$_{0.35}$As/ GaAs QW grown by CBE as a function of growth temperature (closed triangles). Open circles correspond to transition energies calculated using $R$ values from the data of Fig. 2 (straight lines). The inset is the In composition profile determined using $R$ values (Fig. 2 data) at 545°C (full line) and at 475°C (darkest line). The dashed line corresponds to the ideal QW profile (without segregation).

determining $R$, is to use Fig. 2 data (straight lines) to calculate the optical transition energies of QWs in the well known GaInAs/GaAs system and to compare them with experimental results. With this aim, a sample containing three 7 MLs wide QWs of Ga$_{0.65}$In$_{0.35}$As separated by 500 Å GaAs barriers was grown with decreasing the temperature, in such a way that the three QWs correspond to a growth temperature of 545°C, 510°C and 475°C, respectively. The $R$ segregation coefficients, deduced from the fit of Fig. 2 data, are 0.81, 0.70 and 0.45, respectively. In Fig. 3, are reported the theoretical transition energies obtained using the potential profiles determined with these $R$ coefficients (open symbols) and the experimental transition energies obtained at 9 K by PL (closed symbols) (the insert shows the In composition profiles deduced from the $R$ values at 475°C and 545°C). A very good agreement is obtained between both sets of data. On the other hand, the QW PL energy can also be fitted by using $R$ as an adjustable parameter [14, 15]. The corresponding $R$ values are reported as crosses in Fig. 2. It can be seen that they agree well with the data obtained through RHEED real-
time measurements, confirming once more the validity of this new in situ method for determining In segregation coefficient. We can then compare with confidence the $R$ values determined in this study with the values reported in the literature for MBE growth of GaInAs/GaAs heterostructures (plotted in Fig. 2 with open symbols) [10, 14, 16, 17]. It should be emphasized that for similar growth parameters, the $R$ values corresponding to CBE growth are always lower than that generally reported for MBE experiments, as previously noted by Leymarie et al. [15] from optical spectroscopy results. This can be tentatively attributed to the effect of a kinetic limitation specific to the CBE growth environment (due, for example, to the hydrogen or other radical adsorption at the growing surface).

Finally, in order to complete the comparison between GaInAs/GaAs and GaAs/GaInP heterostructures, we have grown four samples, each containing two GaAs/GaInP QWs of 25 and 6 ML, respectively, separated by 500-Å-thick Ga$_{0.51}$In$_{0.49}$P barriers (the well thicknesses are determined by RHEED oscillations and therefore take into account the growth rate transient). For each sample, the 6 ML-thick QW is grown at various temperatures (respectively 480°C, 505°C, 525°C and 550°C), while the wide QW (25 ML), which is first grown, corresponds to a fixed temperature of

Fig. 4. Photoluminescence (9 K) of Ga$_{0.51}$In$_{0.49}$P:GaAs QW structures containing a wide QW (25 ML) grown at 550°C (reference QW) and a 6 ML QW grown at different temperatures.
550°C in order to check the reproducibility of the growth conditions from sample to sample. The growth temperature for the GaInP barriers is kept constant (550°C). Fig. 4 displays the PL spectra of these four samples. The first point to note is that the energy shift of the 6 ML GaAs QW as a function of the growth temperature is less important than the shift observed for GaInAs (7 ML)/GaAs QW's (Fig. 3). Actually, a significant red shift occurs when increasing the growth temperature from 480°C to 505°C and then saturates (note that the slope of the R variation with temperature decreases drastically above 505°C, Fig. 2). This red shift can be, at least partly, explained by considering that In is incorporated in the GaAs QW. In fact, above a R (growth temperature) value of ≈ 0.7 (≈ 510°C), this red shift saturates as In escapes from the thin GaAs well at the GaInP on GaAs interface. Nevertheless, a quantitative description of this phenomenon is a complex problem due to additional effect of As–P intermixing at the interfaces which is poorly known but has a dramatic influence on the optical properties of the QWs. Further investigations are therefore necessary to correctly understand the energy shifts observed in GaAs/GaInP QW structures.

4. Conclusions

In conclusion, it is shown that In surface segregation in III–V compounds, such as GaInP and GaInAs, can be quantitatively evaluated in real-time during CBE growth. This is possible owing to the growth rate transients due to the variation of the In surface composition in the first stages of the growth, which can be measured through RHEED intensity oscillations. Following this, we have determined the temperature variation of the segregation coefficient of In in GaAs. The validity of our approach is demonstrated by correctly predicting the photoluminescence energies of GaInAs/GaAs QWs, grown at different temperatures, using In composition profiles deduced from the in situ determined segregation coefficient. The influence of In segregation on optical transition energies in the GaInP/GaAs system has also been evidenced, though the quantitative interpretation of the results is not as straightforward as in the GaInAs/GaAs system, presumably due to the group V element interfacial intermixing.

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References

Full gaseous source growth of separate confinement MQW 1.55 μm laser structures in a production MOMBE

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Abstract

The feasibility of metalorganic molecular beam epitaxy (MOMBE or CBE) as a production process is studied with respect to flexibility, uniformity, long term stability and device quality. Gaseous doping sources were used for growth of GaInAsP/InP device structures. For n and p type doping disilane and diethylzinc (DEZn), respectively, were injected. The importance of precracking for efficient dopant incorporation is described. For uniform growth across larger areas the dependence of layer composition on growth temperature is critical. Therefore this effect along with the effect of V/III ratio have been studied in detail, and the optimum growth conditions are discussed. It is demonstrated that composition uniformity and long term stability can be obtained leading to variations of emission wavelength and lattice mismatch below ±1.5 nm and ±1.5×10⁻⁴, respectively. These material properties are found across an area of about 90% of a 2 in wafer revealing the high yield of this process. As a device test vehicle, strained layer MQW laser structures were grown, and data on broad area threshold current densities are compared with state of the art results from structures grown by metalorganic vapour phase epitaxy (MOVPE).

1. Introduction

It has been demonstrated that metalorganic molecular beam epitaxy has the potential to deliver state-of-the-art InP-based device structures for the fabrication of lasers, photodetectors, modulators and HBTs [1–4]. The flexibility of MOMBE is in addition enhanced by the unique material qualities obtained in the selective area epitaxy process (SAE) which enables a high quality of device integration like lasers and waveguides [5, 6]. First reports on the development of production type MOMBE setups are recently presented [7, 8]. However, the growth chemistry of the GaInAsP on InP is ruled by a complex dependence on growth temperature, V/III ratio and absolute injection fluxes [9–12]. In order to obtain an excellent lateral and vertical uniformity, an optimization process concerning the temperature sensitivity for all quaternaries having a gap wavelength between 1.05 and 1.65 μm is performed and an accurate temperature control with time and across the wafers is absolutely required [8].
There are instabilities due to passivation of the solid dopant Be and Si in MOMBE. The implementation of cooled baffles can suppress this aging process of the dopant [13], but this hinders an uniform upscaling for large area growth or multi-wafer production. As a consequence and in order to move all sources for growth and doping out of the growth chamber several doping gases for donor and acceptor incorporation have been investigated [14-19].

In this communication we concentrate on the important issues for the upscaling of the MOMBE process. The quaternary growth in the wavelength range from 1.05 to 1.65 μm is investigated targeting for stable growth conditions with optimized parameter dependencies. The doping is achieved by using the identical doping gases DEZn and disilane as in MOVPE. This selection of dopants enables steep dopant profiles in InP-based heterostructures and is of major importance for the acceptance in the production. The stability and flux uniformity of this full gaseous MOMBE process is finally tested by the fabrication of strained MQW lasers with double separate confinement layers. This test bed for our production type MOMBE incorporates as well several quaternary layers as p- and n-type doped films.

2. Experimental procedure

The experiments were carried out in a 3 × 2 in multi-wafer production type MOMBE system. As precursors trimethylindium (TMI), triethylgallium (TEG), AsH₃, PH₃, diethylzinc and disilane were used. The system is equipped with two equivalent high temperature cells each with one AsH₃ and PH₃ line and a third high temperature cell for the disilane injection. In this investigation only one high temperature cell for the group V elements was used. The group III elements were injected via a low temperature central- and a low temperature off axis cell, each connected to a TEG and TMIII line. For Zn-doping the system was equipped additionally with a low temperature dopant cell [19]. Details about system layout and gas injection were given in ref. [8, 19, 20].

InP: Fe and InP: S-doped (1 0 0)-substrates with 2° misorientation towards the (1 1 0) direction were used. A growth rate between 1.0 and 1.3 μm/h was selected depending on the material. The quaternary layers were mainly grown at a constant V/III ratio of 3.7, in selected cases the V/III ratio was increased to 6.8 for optimized surface smoothening. The layers were characterized by room temperature photoluminescence (PL) and high resolution X-ray-diffraction. The growth temperature was accurately (±1°C) controlled in the center of the 2 in wafer via a two channel optical infrared pyrometer in the range between 450°C and 530°C. The Si and Zn doped InP layers were characterized by Hall measurement and secondary ion mass spectrometry (SIMS).

Separate confinement MQW laser structures were grown at an optimized growth temperature of 510°C for the active region. The Zn doped cladding layers were grown at an optimized growth temperature of 450°C [19]. The Zn doping starts after a recess in the InP cladding layer of 120 nm undoped InP. The test structures were characterized by PL, X-ray diffraction transmission electron microscopy (TEM) and SIMS. The emission wavelength yields 1.55 μm. The layer sequence was processed as broad area laser (50/100 × 400 μm²).

For the ohmic contact a sequence of Au/Zn/Au was used on the p-side, the n-contact is made of Au/AuSn/Au.

3. Results and discussion

3.1. Growth of GaInAsP

To investigate the temperature dependence of the composition in the GaInAsP-alloys, single quaternary GaInAsP layers in the wavelength ranges of 1.05 (Q-1.05), 1.25 (Q-1.25), 1.55 (Q-1.55) and 1.65 μm (GaInAs) were grown in the temperature range of 460–530°C with a thickness of 0.3–1.3 μm. Generally we used an input V/III ratio of 3.7 at a constant growth rate. The TEG, TMI, AsH₃ and PH₃ fluxes were precisely kept constant for each wavelength range. The material composition in GaInAsP was calculated according the references [21, 22]. At lower temperatures (< 490°C) the
thermal dissociation of the TEG molecules is the limiting factor whereas at higher temperatures (> 490°C) a Ga molecule loss is observed [11, 23, 24]. This leads to the typical u-curve in the plot of the measured lattice mismatch Δd/d versus growth temperature (Fig. 1) in the wavelength range of 1.05–1.65 μm. The minimum of these u-curves is observed at 490°C for GaInAs, at 505°C for Q-1.55 layers, at 513°C for Q-1.25 layers and then 500°C for Q-1.05 layers. The Q-1.05 process has the strongest temperature dependence, whereas GaInAs reveals the weakest dependence in the temperature range of 460–530°C.

Around the most practical growth temperature of 510°C the optimum room temperature photoluminescence was achieved for all quaternary materials. At this growth temperature we observe a change in lattice mismatch Δa/a – which is half of the measured value Δd/d – of 95 ppm/°C for GaInAs, for Q-1.55 material this coefficient is reduced to 85 ppm/°C and for the Q-1.25 material this coefficient is negative (−130 ppm/°C). The coefficient for Q-1.05 layers is positive again (150 ppm/°C). Consequently, this optimum temperature enables coefficients of < 150 ppm/°C for all GaInAsP compositions. The maximum of Ga incorporation is observed at 490°C for GaInAs and at 500°C for the other quaternary materials, which is represented in the Ga/In ratio in Fig. 2a. Above a temperature of 490°C we observe an increasing P uptake in GaInAsP (see Fig. 2b). The shift of the minimum in lattice mismatch for the Q-1.25 material towards higher temperatures (see Fig. 1) is a consequence of the strong increase of the P incorporation as shown on Fig. 2b. The comparison of the data from Fig. 2 reveals that the increase in the P uptake at higher temperatures (Fig. 2b) is always accompanied by a decrease of the Ga uptake at higher temperatures. Changing the composition from GaInAs to InP at the constant temperature of 510°C a drastically decrease of the Ga and P incorporation efficiencies is observed. This can be evaluated from the data in Fig. 2 by plotting the Ga/In (P/As) ratio in the solid layer versus the Ga/In (P/As) ratio in the beam flux (not shown here). A similar correlation was found earlier for the GaInAsP growth on vicinal surfaces [10]. It was speculated that the Ga molecule loss is due to a mutual effect between the higher P content and the Ga molecules.

This is in line with the results from Refs. [10, 25], but in contrast to the data from Ref. [9], where a higher Ga incorporation was reported for quaternary materials near InP. This discrepancy is not clear at the moment, but the difference might be originating from the fact that in Ref. [9] the layers for each material were grown in a stack of quaternary layers grown at different growth temperatures. Our procedure using single layers allows for a detailed and independent determination of the absolute Ga and P incorporation in GaInAsP by PL and X-ray-diffraction.

3.2. Uniformity and reproducibility

For device structure production vertical as well as lateral uniformity over a 2 in wafer is required. Therefore single quaternary layers Q-1.55 were grown with a thickness of 1.3 μm. Uniformity of the layer in vertical direction was achieved using an automated optical pyrometer system [8] for the
growth temperature measurement. Fig. 3 gives the lateral dependence of the variation and the variation in lattice mismatch \( \Delta a/a \) across a single 2 in wafer. The wavelength uniformity is better than \( \pm 1.5 \) nm and in lattice mismatch a uniformity of \( \pm 150 \) ppm is achieved. On the basis of the data in Figs. 1 and 2 it can be calculated that the growth temperature at the edge of the 2 in wafer is 3°C higher than in the center. Notice that more than 90% of the wafer area is suitable for device fabrication in this production type MOMBE system. The reproducibility measured for 10 samples during a period of 1\( \frac{1}{2} \) month was \( \pm 1.5 \) nm in wavelength and \( \pm 150 \) ppm in lattice mismatch for the Q-1.55 material. These stability data are comparable to those reported earlier by Heinecke et. al. [4] and within the scatter of the 2 in uniformity shown in Fig. 3.

3.3. Doping using disilane

As in MOVPE Si\( _5 \)H\(_6\) for the n doping of InP layers has been used. For the application in device structures a sufficient uniformity in the doping process is required. In order to avoid clogging a high conductance, high temperature cell was used. The injector is comparable to the hydride cracker yielding a good 5 in uniformity. The growth temperature was set to 510°C. The layer thickness was 1.3 \( \mu m \) (growth rate 1.3 \( \mu m/h \), V/III = 4.5). Fig. 4 gives the dependence of the precracking temperature on the net electron concentration in InP using 5% disilane diluted in H\(_2\) and pure disilane. The dopant gas flux was held constant. The Si incorporation in InP starts at 600°C precracking temperature. A maximum of the net electron concentration is observed for 5% disilane at 900°C.
precracking temperature and for undiluted disilane at 1000°C precracking temperature. Above 1000°C in the cracker the electron concentration in the layer is reduced. We assume, that at lower cracking temperatures (600–1000°C) the disilane is not completely precracked (SiH₂ formation supports the Si-incorporation in InP [26]). Above 1000°C the disilane decomposes and Si deposits in the HTD-injector, but the vapour pressure of Si at this temperature is too low to achieve significant dopant fluxes. The electrical activation calculated by the comparison of SIMS and Hall data is independent of the precracking temperature between 400 and 1100°C and reaches 85–90%. At a growth temperature of 450°C the electrical activation is reduced to 46%. Furthermore, as expected a linear dependence of the electron concentration on the Si₂H₆ flux is measured in the range of 10¹⁶–7×10¹⁷/cm³.

3.4. Separate confinement MQW-laser structures

The basic layer sequence for the laser structures was 1000 nm Si-doped InP cladding layer/100 nm Si doped Q-1.15 confinement/30 nm Q-1.25/MQW with 5 wells/30 nm Q-1.25/100 nm Q-1.15/120 nm undoped InP spacer layer/1600 nm Zn-doped InP-cladding/200 nm Zn-doped GaInAs. In the MQW the lattice matched Q-1.25-material was used as barrier (thickness 10 nm) and unstrained GaInAs or 0.6%/0.8%/1.0% compressively strained quaternary layers as wells (thickness 6 nm). Fig. 5 presents the SIMS-measurement of a double separate confinement MQW laser structure. On the basis of the results achieved earlier [19] the spacer thickness of 120 nm at the p-side cladding layer is sufficient to avoid Zn diffusion into the quaternary MQW region, which is located 2 μm from the surface on the SIMS data and bounded by the slight decrease in the Ga and the P signals. The decrease of the Zn signal in the SCH layers is more rapid than 30nm/decade indicative of minimal Zn diffusion. Table 1 shows the results of the threshold current densities using GaInAs or 0.6%/0.8%/1.0% compressive strained quaternary wells. The lowest threshold current densities with 100 × 400 μm² broad area test structures are observed with 1.0% strained wells. Under optimized growth conditions we achieve 640 A/cm² for the 400 μm long devices, which is comparable with lasers grown by CBE and MOVPE [1, 27].
The uniformity of the threshold current densities across a 2 in wafer varies in the range of 750 ± 110 A/cm² with the lowest threshold current densities at the outer diameters of the wafer (Table 2).

4. Conclusions

This study has shown that for growth of InP based device material even for n and p type doping the identical precursors can be used in MOMBE as in MOVPE. The gaseous dopant sources disilane and diethylzinc, which are qualified precursors in MOVPE production lines, were successfully applied in a MOMBE process. It has been demonstrated that for doping with disilane a high temperature cracking cell is required, whereas diethylzinc can be injected without precracking. With both precursors steep dopant profiles were achieved.

From a detailed study of the effect of growth temperature on quaternary layer composition the optimum temperature range for good layer uniformity has been elaborated. A temperature around 510°C should be chosen to maximize the useful wafer area (yield = 90%) for device applications with respect to composition uniformity and high optical quality.

The recipe of a constant growth temperature along with a fixed V/III-ratio reduces extensive optimization work for the growth of quaternary layers of any desired composition.

As a device test vehicle the strained layer MQW laser was chosen. Our MOMBE results from the full 2 in wafer process are comparable with state-of-the-art MOVPE data. This investigation clearly reveals that the MOMBE process has taken a further step on its way towards production maturity with the option of multi-wafer processing.

Acknowledgements

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Table 1
Threshold current densities of the 1.55 μm broad area lasers (50/100 μm × 400 μm)

<table>
<thead>
<tr>
<th>Quantum wells</th>
<th>$j_\text{th}$ (A/cm²)</th>
</tr>
</thead>
<tbody>
<tr>
<td>(a) unstrained-GaInAs</td>
<td>1600</td>
</tr>
<tr>
<td>(b) 0.6% strained-GaInAsP</td>
<td>1040</td>
</tr>
<tr>
<td>(c) 0.8% strained-GaInAsP</td>
<td>1040</td>
</tr>
<tr>
<td>(d) 1.0% strained-GaInAsP</td>
<td>880</td>
</tr>
<tr>
<td>(e) 1.0% strained-GaInAsP optimized</td>
<td>640</td>
</tr>
</tbody>
</table>

Table 2
Uniformity of threshold current density across a 2 in. wafer (from (d) in Table 1)

<table>
<thead>
<tr>
<th>Distance from center (mm)</th>
<th>$j_\text{th}$ (A/cm²)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>880</td>
</tr>
<tr>
<td>10</td>
<td>800/780/680</td>
</tr>
<tr>
<td>18</td>
<td>760/880/800/640</td>
</tr>
<tr>
<td>24</td>
<td>670/670</td>
</tr>
</tbody>
</table>
References

GSMBE growth and characterization of In$_x$Ga$_{1-x}$As/InP strained-layer MQWs in a P-i-N Configuration

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Abstract

In$_x$Ga$_{1-x}$As/InP (0.39 ≤ x ≤ 0.68) strained-layer quantum wells having 20 wells with thickness of 50 Å in a P-i-N configuration were grown by gas source molecular beam epitaxy (GSMBE). High-resolution X-ray diffraction rocking curves show the presence of up to seven orders of sharp and intense satellite reflection, indicative of the structural perfection of the samples. Low-temperature photoluminescence and low-temperature absorption spectra were used to determine the exciton transition energies as a function of strain. Good agreement is achieved between exciton transition energies obtained experimentally at low temperature with those calculated using the deformation potential theory.

PACS: 68.65.+g; 68.55.Bd; 78.66.Fd; 73.20.Dx

Keywords: Quantum wells; Molecular beam epitaxy; InGaAs/InP; Optical properties

1. Introduction

Strained layer quantum wells (QWs) are of considerable interest due to the added flexibility in controlling their optical and electrical transport properties. The biaxial in-plane strain caused by lattice constant mismatch has been used to optimize device performance such as reduced threshold current densities in strained quantum well lasers [1–3] and reduced switching times of high electron mobility devices [4]. The In$_x$Ga$_{1-x}$As/InP (0 ≤ x ≤ 1) quantum well structure, with relative lattice mismatch −3.8% for x = 0 to +3.2% for x = 1, is one of the most important material systems for device applications such as long wavelength lasers [1, 2], high electron mobility transistors [5] and electro-optic modulators and switching devices [6–9]. Although many investigations on this structure have been conducted [10–15], optical absorption characteristics [12–15] have not sufficiently been clarified, in particular, for strained QWs.

In this work, we report the growth of the strained layer InGaAs/InP multiple quantum wells (MQWs) with P-i-N configuration using a home-made GSMBE system and the use of optical techniques to.
study on the excitonic transition energies in the InGaAs/InP MQWs as a function of strain.

2. Experimental procedure

The epitaxial growth was carried out in a homemade GSMBE system equipped with a low-pressure hydride gas-cracking cell for As$_2$ and P$_2$ molecular beams. These were supplied as 100% arsine and phosphine. Control of flow rates was achieved using mass flow controllers. The Group-III elements were produced from two indium and one gallium conventional effusion cells. The pressure in the growth chamber during growth was usually between $5 \times 10^{-5}$ and $1 \times 10^{-4}$ Torr. The growth rates of GaAs, InP and InGaAs were calibrated with in situ observations of the intensity oscillations of reflection high-energy electron diffraction (RHEED). The growth temperature was determined by a thermocouple located just behind the molybdenum sample holder and calibrated by an pyrometer.

To optimize the growth parameters such as growth temperature and V/III ratio, two series of InGaAs epilayers were grown, one with different growth temperatures, and the other with different AsH$_3$ flow rates. The growth rate for all samples was 1 ML/s. The double-crystal X-ray diffractions were used to determine the InGaAs lattice-mismatch relative to InP substrates, and the results show that the lattice mismatch for all samples was small than $1 \times 10^{-3}$. The low-temperature (10 K) photoluminescence spectra (PL) measurements were carried out for InGaAs/InP heterostructure samples. The results show that when the growth rate is 1 ML/s, the optimized growth conditions were such that the AsH$_3$ flow rate was 8–10 sccm and the growth temperature was about 500°C, i.e., when the growth rate is 1 ML/s, the arsine flow rate is between 8 and 10 sccm, and the growth temperature is about 500°C, the grown InGaAs epilayer has the most intense and narrowest PL peak. The obtained optimized growth parameters were used to grow InGaAs/InP strained MQWs in a P-i-N configuration.

The structures grown in our experiments are described in Fig. 1. The InGaAs/InP MQWs were grown on (0 0 1)-oriented InP substrates in a P-i-N configuration. The undoped MQWs in the i-region was prepared on a 0.2 μm-thick unintentionally doped InP layer under which a 0.2 μm n-InP buffer layer was first grown. It consists of 20 InGaAs quantum wells with thickness of 50 Å separated by 200 Å-thick InP barrier. On the MQWs a 0.2 μm thick undoped InP and then a 0.5 μm-thick p-InP were deposited. Five samples were grown and studied, one lattice matched with $x = 0.53$ (sample C), two under compression with $x = 0.63$ (sample D) and $x = 0.68$ (sample E), and the reminder under tension with $x = 0.39$ (sample A) and $x = 0.50$ (sample B). The indium concentration $x$ was varied from $x = 0.39$ to 0.68.

The gas switching procedure used for growing MQWs is shown on Fig. 2. Two indium conventional effusion cells were used, one (denoted by In(1)) for growing InP barriers, the other (denoted by In(2)) for InGaAs wells. To improve interface quality, the growth interruption was used at each interface. At the InP/InGaAs interface, the surface was first stabilized under cracked PH$_3$ (mainly P$_2$) during time $T_1$, then under cracked AsH$_3$ (mainly As$_2$) during time $T_2$. At the InGaAs/InP interface, the reverse procedure was employed using times

<table>
<thead>
<tr>
<th>Material</th>
<th>Layer Thickness</th>
</tr>
</thead>
<tbody>
<tr>
<td>InP:Be</td>
<td>5000 Å</td>
</tr>
<tr>
<td>i-InP</td>
<td>2000 Å</td>
</tr>
<tr>
<td>MQWs</td>
<td>20 Periods</td>
</tr>
<tr>
<td></td>
<td>well=50 Å</td>
</tr>
<tr>
<td></td>
<td>Barrier=200 Å</td>
</tr>
<tr>
<td>i-InP</td>
<td>2000 Å</td>
</tr>
<tr>
<td>InP:Si</td>
<td>2000 Å</td>
</tr>
<tr>
<td>N-InP</td>
<td>Substrate</td>
</tr>
</tbody>
</table>

Fig. 1. Schematic diagram of the strained layer InGaAs/InP multiple quantum wells in a P-i-N configuration.
3. Results and discussion

Fig. 3 presents the high-resolution CuKα1 X-ray diffraction (HRXRD) scans of five samples with different quantum well compositions (therefore with different strains). The rocking curves show the presence of up to seven orders of satellite reflection, indicative of the structural perfection of the samples. The satellite reflections of the lattice-matched sample (sample C) show a highly symmetric intensity distribution on both sides of the (0 0 4) InP peak as expected from strain-free layers. In comparison, the asymmetric shape of the two upper traces (sample D and E) with their enhanced left-hand-side diffraction intensity, is indicative of in-plane compressive strain, resulting in lattice tension in the growth direction. The opposite sign of the strain (in-plane tensile strain) results in the enhanced right-hand-side diffraction intensity of the two lower traces (sample A and B). Despite the lattice mismatch, however, the strained samples retain their structural integrity, as judged by the sharpness and intensity of the satellite reflection.

To determine the $n = 1$ exciton recombination energies in the InGaAs MQWs, the 10 K photoluminescence measurements were performed. For each of the five samples, we observed a sharp and intense PL peak, which is due to the excitonic recombination of electron and hole from the $n = 1$ subband level. The peak energies obtained are shown in Fig. 5 by triangle data points.

Fig. 4 represents the 10 K optical absorption spectra for the five samples A, B, C, D and E. The number indicates sublevels, and the letters H and L denote an electron to heavy-hole and electron to light-hole transition, respectively. From Fig. 4 we can see that the samples exhibit well-shaped excitonic transitions. The characteristic step-like structure, which reflects their two-dimension nature, are clearly shown with the exception of sample A. For sample B, C, D, and E, the spectra show two clear excitonic absorption peaks at the absorption edge, one corresponding to the $n = 1$ heavy-hole excitonic transition (1 1 H), the other corresponding to the $n = 1$ light-hole excitonic transition (1 1 L). For sample A, the spectrum show only one broader absorption peak, which is due to the $n = 1$ light-hole excitonic transition. The above assignment has
been made on the basis of an energy shift calculation using the deformation potential model [15]. In the calculation, the heavy- and light-hole energy split at a Γ point due to the strain effect has been taken into account. The numerical parameters used in the calculation of the band structure are taken from Ref. [15]. The parameters of InGaAs are calculated using the well-known Vegard’s law between GaAs and InAs. For all five samples, due to the InGaAs quantum wells are thin (50 Å), only the first level in the conduction band is allowed. Therefore, no other absorption peak can be seen as in the Fig. 4.

The curve shown in Fig. 5 is the calculated \( n = 1 \) excitonic transition energies as a function of InGaAs quantum well indium composition using the deformation potential model [15]. The data points on the same plot are the experimental results obtained by PL and absorption measurements for four strained and one lattice-matched InGaAs/InP MQWs in a P-i-N configuration. From Fig. 5, we can see that the experimental results agree well with those of calculation. As indium composition \( x \) increases, the transition energies corresponding to \( 1\!\!1H \) and \( 1\!\!1L \) decreases and the separation between \( 1\!\!1H \) and \( 1\!\!1L \) absorption peaks vary. When \( x < 0.42 \), the \( 1\!\!1H \) energy is larger than \( 1\!\!1L \) energy for a given indium composition \( x \), indicating that the first light-hole sublevel is above the first heavy-hole sublevel due to biaxial in-plane tensile strain. When \( x = 0.42 \), the \( 1\!\!1H \) energy is equal to the \( 1\!\!1L \) energy, showing that the first light-hole sublevel and the first heavy-hole sublevel degenerate at this quantum well composition. When \( x > 0.42 \), the \( 1\!\!1L \) energy is larger than \( 1\!\!1H \) energy, indicating that the first light-hole sublevel is below the first heavy-hole sublevel for a given \( x \) due to quantum size and strain effects.

4. Summary

We have grown a series of In\(_x\)Ga\(_{1-x}\)As/InP strained quantum wells with \( x \) varying from \( x = 0.39 \) to 0.68 using a home-made GSMBE system. High-resolution X-ray diffraction patterns show the presence of up to seven orders of sharp and intense satellite reflection, indicative of the structural perfection of the samples. The low-temperature PL and absorption spectrum are used to determine the exciton transition energies as a function of strain. The agreement between transition
energies obtained experimentally at low temperature with calculated results using the deformation potential theory is good.

References

Symmetric triangular-barrier optoelectronic switch (S-TOPS) by gas source MBE

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Abstract

We report a novel optical functional device, a symmetric triangular-barrier optoelectronic switch (S-TOPS), which consists of a symmetric n⁺–i–δp⁺–i–n⁺ structure of In₀.₅₅Ga₀.₄₅As, grown by a gas source molecular beam epitaxy (GSMBE). We fabricated the S-TOPS by changing the sheet-carrier concentration of the δp⁺ gate layer to introduce an avalanche multiplication for positive feedback in the operation. In the current vs. voltage characteristics of the S-TOPS under illumination of 1.55 μm wavelength light, we successfully confirmed a bipolar S-shaped negative differential resistance (NDR) characteristics on a certain range of sheet-carrier concentration of the δp⁺ gate layer. Clear latch characteristics in the input-light power vs. current characteristics were also obtained in both the positive and negative biased conditions. We found that it is very important to optimize the sheet-carrier concentration to fabricate the S-TOPS.

PACS: 73.40.Kf; 85.60.Dw

Keywords: Optical functional device; Gas source MBE; Negative differential resistance; Avalanche multiplication

1. Introduction

For optical signal processing, various kinds of functions are needed for optical devices such as logic, memory, and so on. For these purposes, an optoelectronic switch showing nonlinear optical response is quite promising, and several kinds of devices have been studied to date [1–6]. Especially the devices which use S-shaped negative differential resistance (NDR) are very attractive [1–4]. Among them devices such as the p–n–p–n structure [1, 2], p⁺–n⁺–i–δp⁺–i–n⁺ structure [3], and the heterojunction bipolar phototransistor (HPT) with n⁺–p⁺–i–n⁺ structure [4] have been studied. These devices utilize the avalanche multiplication phenomena in the reverse-biased condition for switching.

We have already proposed a novel device, triangular-barrier optoelectronic switch (TOPS) [7].
It consisted of an n⁺−i−δp⁻−i−n⁺ doping profile in In₀.₅₅Ga₀.₄₇As/In₀.₅₂Al₀.₄₈As heterostructure and showed clear S-shaped NDR. By utilizing this S-shaped NDR, we successfully obtained differential gain, bistable and latch characteristics by simply changing the bias voltages [8]. This device also showed good characteristics such as high-sensitivity, high-gain and high-contrast outputs. TOPS was grown by a gas-source molecular beam epitaxy (GSMBE), which is a promising technique for realizing sharp interfaces and good δ-doped layers [9]. We confirmed optoelectronic logic operations such as AND and OR operations by a single TOPS, and NOT, NAND, NOR, XOR operations by dual TOPS [10]. It is noted that these logic gates can be implemented by using only one or two devices, and that these operations can be realized without any optical or electrical reset signals. However, TOPS has only unipolar S-shaped NDR due to its asymmetric heterostructure. Flexibility of the circuit structure of TOPS is limited by its unipolarity when it is used in the arrayed structure. If TOPS has a bipolar S-shaped NDR, we can use the functions of TOPS at both polarities. Therefore, it is expected to extend the flexibility of the circuit structure of TOPS in the arrayed structure by using bipolar type.

In this paper, we propose a novel symmetric structure triangular-barrier optoelectronic switch (S-TOPS) for 1 μm wavelength-range operation by GSMBE, and demonstrate the optically controllable bipolar S-shaped NDRs. To our knowledge, this is the first demonstration of optically controllable bipolar S-shaped NDRs using n⁺−i−δp⁻−i−n⁺ structure.

2. Growth and measurement

The devices were grown by GSMBE. Group III elements of In and Ga and respective p- and n-type dopants of Be and Si were used. As for the group V sources, 100% AsH₃ and PH₃ with cracking at 1000°C were used. The flow rates of AsH₃ and PH₃ were 4.0 and 10.0 sccm, respectively. The background pressures during the growth were 6 × 10⁻⁵ Torr for AsH₃ and 2 × 10⁻⁴ Torr for PH₃. PH₃ was used for InP buffer layer. The device was grown on an (1 0 0) semi-insulating InP substrate. The typical growth temperature and the growth rate were 420°C and 0.4 μm/h, respectively, for InGaAs. We obtained the proper sheet-carrier concentration of the δp⁺ layer by precise control of Be cell temperature. The devices were formed into a mesa structure with 60 μm diameter. Contact with the top layer was achieved by evaporating Au/Sn to form an open-ring structure using the lift-off technique.

The input light was introduced at the top of the device with a lensed fiber, and the wavelength of the input light was 1.55 μm. I–V characteristics were measured under positive and negative bias of the substrate side, separately.

3. Results and discussion

To clear the difference between a unipolar NDR and a bipolar NDR characteristics, a brief summary of the TOPS is described first [7]. Fig. 1 shows a band diagram of the TOPS, in which the bias voltage is applied. It had n⁺−i−δp⁺−i−n⁺ structure of In₀.₅₅Ga₀.₄₇As/In₀.₅₂Al₀.₄₈As for 1.5 μm wavelength-range operation. The sheet-carrier concentration of δp⁺-In₀.₅₅Ga₀.₄₇As layer was 9.8 × 10¹¹ cm⁻². The avalanche multiplication occurs in the drain region as the bias voltage is increased. As a result, electrons and holes are generated in the drain, and the holes move to the gate layer and accumulate in it. Resultant lowering
of the gate-potential barriers increases the majority-carrier flow of electrons over the potential barrier from the source to the drain region with a thermionic emission. Multiplied holes lower the potential barrier further, so the majority-carrier flow increases. This positive feedback phenomenon gives rise to S-shaped NDR. It should be noted that the avalanche multiplication can be stopped by decreasing the input-light power, and the various functions can be realized by only changing bias voltages. Experimental $I-V$ characteristics of the TOPS at different input-light powers observed at room temperature are shown in Fig. 2. Here, a significant S-shaped negative differential resistance (NDR) was observed in forward bias when the input-light power was less than 2 $\mu$W. However, in the reverse bias condition no NDR characteristics were observed, though a small change in $I-V$ characteristics with low input-light power.

To realize a bipolar NDR characteristics, we propose a novel device, S-TOPS. Fig. 3 shows the device structure of the S-TOPS. The device structure consisted of $n^+\text{-InP}$ buffer layer (0.64 $\mu$m, $1.9 \times 10^{18}$ cm$^{-3}$), $n^+\text{-In}_{0.53}\text{Ga}_{0.47}\text{As}$ (0.098 $\mu$m, $2.2 \times 10^{18}$ cm$^{-3}$), $i\text{-In}_{0.55}\text{Ga}_{0.45}\text{As}$ (0.39 $\mu$m), $\delta p^+\text{-In}_{0.53}\text{Ga}_{0.47}\text{As}$ (98 Å, $n_{sp+}$), $i\text{-In}_{0.53}\text{Ga}_{0.47}\text{As}$ (0.39 $\mu$m) and $n^+\text{-In}_{0.53}\text{Ga}_{0.47}\text{As}$ (0.098 $\mu$m, $2.2 \times 10^{18}$ cm$^{-3}$). Here, $n_{sp+}$ is the sheet-carrier concentration of the $\delta p^+$ layer. The $\delta p^+$-InGaAs region was designated as the gate and both the $i$-InGaAs layers were designated as the source or drain region determined by the polarity of the bias voltage. An S-shaped NDR is expected to occur due to avalanche multiplication in both $i$-InGaAs drain regions. The parameters of the $\delta p^+$ layer were critical to attain the NDR operation, which was caused by the avalanche multiplication near the $\delta p^+$ layer. We fabricated the S-TOPS devices which have the same sheet-carrier concentration as

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**Fig. 2.** $I-V$ characteristics at different input-light powers of the TOPS.

**Fig. 3.** Device structure of the S-TOPS.
that of the $\delta p^+$ gate layer. Fig. 4 shows a band diagram of the S-TOPS under biased condition. The major difference of the S-TOPS from a conventional TOPS is the symmetric structure.

Experimental $I$–$V$ characteristics of S-TOPS at different input-light powers are shown in Fig. 5a–Fig. 5c, in which the sheet-carrier concentration $n_{sp^+}$ of $\delta p^+$ layer were varied as follows:

(a) $n_{sp^+} = 3.8 \times 10^{11}$ cm$^{-2}$,
(b) $n_{sp^+} = 1.4 \times 10^{12}$ cm$^{-2}$,
(c) $n_{sp^+} = 1.1 \times 10^{13}$ cm$^{-2}$.

Fig. 5. $I$–$V$ characteristics of the S-TOPS at different input-light powers by changing the sheet carrier concentration $n_{sp^+}$.

(a) $n_{sp^+} = 3.8 \times 10^{11}$ cm$^{-2}$, (b) $n_{sp^+} = 1.4 \times 10^{12}$ cm$^{-2}$, (c) $n_{sp^+} = 1.1 \times 10^{13}$ cm$^{-2}$. 
In Fig. 5a, we obtained only gain characteristic because electrons in i-layer can easily flow over the potential barrier when the photo-generated hole accumulate in the gate layer and lower the potential barrier. The avalanche multiplication, however, does not occur in this case due to low-potential height made by the $\delta p^+$ doping. On the other hand, we obtained clear bipolar S-shaped NDRs and gain characteristic simultaneously in Fig. 5b. It is because of the positive feedback due to the avalanche multiplication and gain characteristic. In the case of further high doping such as in Fig. 5c, we could not obtain the gain characteristic any more. Photocurrent was only obtained on this condition that electrons could not flow over the potential barrier.

It is noted that there is a certain range of sheet-carrier concentration $n_{sp+}$ of the $\delta p^+$ gate layer for positive feedback to occur. The reason is described as follows. The potential-barrier height depends on the thickness and the sheet-carrier concentration of the $\delta p^+$ layer, and the widths of the source and the drain. Since the parameters except the sheet-carrier concentration $n_{sp+}$ of these samples were not changed, the barrier height depends only on the sheet-carrier concentration. When the sheet-carrier concentration $n_{sp+}$ is large enough, the electric field in the drain becomes large enough for the avalanche multiplication. Here it is important that the majority-carriers, electrons, flow over the potential barrier when the bias voltage and/or the input-light power is increased. When the sheet-carrier concentration $n_{sp+}$ is extremely large, electrons cannot easily flow over the potential barrier. If the electrons do not flow from the source at all, positive feedback does not occur in the drain. When the positive feedback does not occur in the drain, it is possible to have two types of operation. One is the phototransistor type which has a gain without NDR in Fig. 5a, the other is the photodiode type which has only photocurrent without a gain and NDR in Fig. 5c. Therefore, we must determine the potential-barrier height to satisfy both, electron flow and avalanche multiplication. For example, the parameters of $n_{sp+} = 1.4 \times 10^{12} \text{ cm}^{-2}$, in which bipolar S-shaped NDR was obtained as shown in Fig. 5b, corresponds to about 3.9 eV for the barrier height, and $1.0 \times 10^{4} \text{ (V/cm)}$ for the electric field. We can also observe a slight asymmetric NDR characteristics in Fig. 5b. It is likely that the quantity of the majority carriers, which are generated in the i-InGaAs source layer by the absorption of the input-light, is different by the polarity of the bias voltage because the input-light is incident on the surface of the devices.

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Fig. 6. Input-light power versus output-current characteristics of the S-TOPS by changing the bias voltages $V_b$. (a) $V_b = +3.0 \text{ V}$. (b) $V_b = -3.0 \text{ V}$. 
Fig. 6a and Fig. 6b shows input-light power versus current characteristics by changing the bias voltages $V_b$ in the device of Fig. 5b. We obtained latch characteristic with positive bias condition in Fig. 6a. We also obtained nearly the same latch characteristics with negative bias in Fig. 6b, therefore, we can use this device as a bipolar optoelectronic switch. We can also observe the difference in the input-light power threshold between the polarities of the bias voltage. This, as also the small difference in $I-V$ characteristics mentioned above, is due to the difference in the number of the photo-generated carriers in the i-InGaAs source layers. From these results, we confirmed that it is important to optimize the sheet-carrier concentration of $\delta$-doped layer for the S-TOPS operations.

4. Conclusions

We fabricated and examined a novel symmetric triangular-barrier optoelectronic switch (S-TOPS) by GSMBE, which consisted of a symmetric n$^+$$-\delta$p$^+$$-i$-n$^+$ structure. Bipolar S-shaped NDR and clear latch characteristics were obtained in both positive and negative biased condition. We also found that it was very important to optimize the sheet-carrier concentration of the $\delta$p$^+$ gate layer to realize the S-TOPS operation. These characteristics also show the high potential of GSMBE for fabricating well-controlled $\delta$-doped structure.

We can apply this S-TOPS device to optical functional devices such as optical logic devices and optical memories as well as two-dimensional optical signal processing.

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References

Chemical beam epitaxy growth of tensile-strained GaAsP/GaAlAs quantum well heterostructures for laser application

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Abstract

We report on the chemical beam epitaxy growth conditions of high-quality Ga\textsubscript{1-x}Al\textsubscript{x}As (x: 0.2–0.6) and tensile-strained GaAs\textsubscript{1-x},P\textsubscript{x} quantum wells (y: 0.05–0.29) on GaAs(0 0 1) surfaces using trimethylamine alane and alternative precursors of group V elements (tertiarybutylarsine (TBAs) and tertiarybutylphosphine (TBP)). Cracking conditions of TBAs and TBP have been improved in order to reduce the carbon incorporation. Low levels of carbon and oxygen were detected by SIMS analysis of AlGaAs layers, independent of the aluminum concentration used ([O] ≈ 6–8 × 10\textsuperscript{17} cm\textsuperscript{-3}, [C] ≈ 4–5 × 10\textsuperscript{17} cm\textsuperscript{-3}). GaAsP quantum wells have been grown exhibiting high luminescence efficiency and full-width at half-maximum of 10 meV for 30 Å thick quantum well. The QW photoluminescence of E\textsubscript{1}-HH\textsubscript{1}, E\textsubscript{1}-LH\textsubscript{1} energy transitions as a function of P mole fraction is well accounted for by envelope function calculations including strain effects. Preliminary results concerning laser diodes are discussed in the light of laser structure designs.

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1. Introduction

AlGaAs-based high-power laser diodes (LDs) emitting in the 0.8 µm range have key applications in the field of optical information processing and solid state laser pumping. Up to now, these laser diodes have been fabricated using metal-organic chemical-vapor deposition (MOCVD) [1, 2] or molecular beam epitaxy (MBE) [3]. Although chemical beam epitaxy (CBE) is comparatively a recently developed technique, it is very promising for low-cost mass production devices with a reduced environmental impact compared to MOCVD. On the other hand, the use of gases as material sources adds versatility to the conventional solid sources MBE technology. Using this technique, it is, for example, easy to incorporate tensile strain in the active region of AlGaAs-based LD structures by inserting GaAsP pseudomorphic quantum well (QW) layers. This is an important point since

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tensile strain has been shown to improve LDs characteristics [4–6]. However, the growth of high-quality AlGaAs material by CBE has been hindered for years by the lack of a suitable metalorganic (MO) Al starting source. Owing to the recent progress in the development and synthesis of Al containing MO precursors, such as trimethylamine-alane (TMAAl) or dimethylethylamine-alane (DMEAAI), it is now possible to obtain high purity CBE AlGaAs material [7]. However, these advances have been mainly used to realize bipolar or two-dimensional electron gas transistors [8, 9] and there is only a few reported results on the optical properties of CBE grown AlGaAs-based QW structures (see Ref. [10] and references therein).

The aim of this paper is to report on the CBE growth conditions of high-quality LD structures combining AlGaAs (x = 0.2–0.6) and tensile strain GaAsP QWs (y = 0.05–0.25) using TMAAl and low toxicity alternative precursors of group V elements (tertiarybutylarsine and tertiarybutylphosphine).

2. Experimental procedure

The experiments have been carried out in a all gaseous sources 3X2° VG90 machine. The gas source configuration is the following: group III atoms are provided by organosiliconates (triethylgallium (TEGa) and trimethylamine-alane (TMAAl)); precracked tertiarybutylphosphine (TBP), tertiarybutylarsine (TBAs) and uncracked tridimethylaminarsenic (tDMAAs) are the group V starting sources. Uncracked hydrogen sulfide (H2S) and trimethylgallium (TMGa) were used for n- and p-type doping, respectively. All the gas sources are regulated using a two valves pressure control system which permits a very high level of reliability but also of flux reproducibility. As an example, over a period of 6 months, Al compositions variation did not exceed ±1% for fixed TEGa, TMAAl setpoint pressures and substrate temperature.

Thanks to the relatively high vapor pressure of TBAs and TBP, it has been possible to use a high-pressure type cracker cell design previously developed for arsine and phosphine. The optimum cracking temperature for TBAs is 760°C leading to a complete decomposition of TBAs into As2, CH4 and butene radicals. Increasing the cracking temperature results in an increase of carbon incorporation in GaAs up to 1018 cm−3 (to compare with 1016 cm−3 under optimal conditions). The cracking of phosphorus containing products is more difficult and less efficient. A complete cracking of the TBP molecule without production of PH3, which represents the limiting step of the reaction, is obtained at a cracker temperature of 1000°C. However, this temperature results in high levels of carbon incorporation, in particular for ternary Ga0.5In0.5P compounds, degrading the growth front and then the surface morphology. In fact, the optimum cracking temperature in terms of epilayer structural quality and carbon incorporation is found to be 950°C. Under these conditions, the cracking of the TBP molecule is incomplete [11] but allows the growth of phosphorus-based compounds (InP, GaInAsP) at a growth rate of about 1 μm/h.

3. Assessment of high purity GaAlAs materials

To our knowledge, no informations have been reported, up to now, on the growth of GaAlAs by CBE using the TMAAl, TBAs source combination. Smooth surface morphology and low defect densities (<10 def/cm²) have been obtained in the 520–600°C growth temperature range even for aluminum concentration exceeding 55%. The use of tDMAAs as a starting source leads to rough GaAlAs surface mainly due to the relatively poor purity (solvent traces) of the source resulting in high oxygen incorporation (>5 × 1018 cm−3). Carbon and oxygen incorporation have been characterized as a function of growth temperature and V/III ratio. Growth temperature has been found to have little effect on both oxygen and carbon incorporation in the epilayers. The carbon concentration determined by SIMS versus the V/III ratio is shown in Fig. 1. The carbon concentration is reduced to 4 × 1017 cm−3 with increasing the V/III ratio (i.e., TBAs flux) with no significant variation of the oxygen doping level lying at 6 × 1017 cm−3. These values are not influenced by the aluminum concentration up to 55%. Residual doping level for
bulk $\text{Ga}_{0.65}\text{Al}_{0.35}\text{As}$ is p-type in the low $10^{16}$ cm$^{-3}$ range.

4. GaAsP phosphorus composition control

GaAsP bulk layers can be grown in the full temperature range 500–600°C using TBAs and TBP group V starting sources. The P mole fraction has been tentatively determined by using group V induced RHEED intensity oscillations [12] as proposed by Hou et al. [13]. Indeed, these authors have shown that the phosphorus composition deduced under these conditions (excess of Ga) is the same as that obtained under a V/III ratio higher than 1. Consequently, they concluded that, even under a V/III ratio < 1, the incorporation of phosphorus is partially hindered by the presence of As. Fig. 2 shows the phosphorus concentration in GaAs$_{1-x}$P$_x$ deduced from the measurement of arsenic and (As + P) incorporation rate under V/III ratio less than 1 compared to phosphorus concentration in the solid determined from ex situ X-ray diffraction measurements. In the same plot, P solid concentrations in GaInAsP/GaAs [14] and GaInAsP/InP [15] versus the gas phase P/(As + P) ratio are reported. The same behavior for the incorporation of the group V elements is observed when comparing GaAsP and quaternary GaInAsP alloys lattice matched on InP and GaAs. It can then be suggested that group III elements play a minor role in the incorporation mechanisms of As and P.

Besides that, it is clear that the P compositions determined from RHEED intensity oscillations under Ga excess conditions do not reveal the real solid P composition. Then, it can be concluded that the interaction between As and P must be negligible under these conditions and consequently, the phosphorus composition deduced from group V induced RHEED oscillations represents, in fact, the phosphorus to arsenic flux ratio P/(As + P) in the gas phase. These results give evidence of the important parameter which is the displacement of phosphorus by arsenic and vice versa under group V rich growth conditions. Nevertheless, it should be pointed out that, if both arsenic and phosphorus fluxes are significantly lower than the Ga one, the condition V/III > 1 being filled by the sum of As and P fluxes, the in situ P determination would be valid [16]. However, in case where As or P individual flux is able to maintain a group V rich growth front, again, the in situ measurements diverge from ex situ composition values.

5. Optical properties of GaAsP QWs

Optical properties of separate confinement heterostructure GaAsP/GaAlAs QWs have been characterized by room temperature photolumine-
Table 1
Parameters for GaAs_{0.56}Al_{0.44}As [18]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\alpha (A^2)$</td>
<td>5.6533 − 0.2021y</td>
</tr>
<tr>
<td>$m^*_0/m_0$</td>
<td>0.069 + 0.023y</td>
</tr>
<tr>
<td>$\gamma_1$</td>
<td>6.85 − 2.75y</td>
</tr>
<tr>
<td>$\gamma_2$</td>
<td>2.1 − 1.61y</td>
</tr>
<tr>
<td>$\gamma_3$</td>
<td>2.9 − 1.65y</td>
</tr>
<tr>
<td>$E_p/(eV)$ at 300 K</td>
<td>$1.424 + 1.174y + 0.186y^2$</td>
</tr>
<tr>
<td>$C_{11} (10^{11} \text{ dyn/cm}^2)$</td>
<td>11.88 + 2.24y</td>
</tr>
<tr>
<td>$C_{12} (10^{11} \text{ dyn/cm}^2)$</td>
<td>5.38 + 0.87y</td>
</tr>
<tr>
<td>$a (eV)$</td>
<td>9.77 + 0.47y</td>
</tr>
<tr>
<td>$b (eV)$</td>
<td>1.7 + 0.2y</td>
</tr>
<tr>
<td>$\Delta E_c$</td>
<td>100% $\Delta E_{\text{strain}}$</td>
</tr>
</tbody>
</table>

The structure is composed of two cladding 1.2 μm thick Ga$_{0.56}$Al$_{0.44}$As layers, an optical Ga$_{0.75}$Al$_{0.25}$As cavity of 2000 Å with a single 100 Å GaAsP quantum well, the P composition being varied from 0 to 30%. Double X-ray measurements have been used to determine the phosphorus concentration with a precision of ±1%. For phosphorus concentrations higher than 10% two well-defined luminescence peaks have been observed corresponding to the heavy hole and light hole transitions. These measurements indicated a lowest HH1 transition energy for the low P compositions and a lowest LH1 transition energy for the high P compositions. The subband edges were calculated using a standard transfer matrix method [17]. The parameters [18] taken for the simulation are listed in Table 1. Due to the tensile strain in the GaAsP layer the HH1 and LH1 levels are split by the shear energy which increases with increasing the phosphorus concentration. For the lower P composition, the HH1 level is more confined than the LH1 level due to the difference in effective masses. For higher P composition the shear energy overcomes this effect so that we observe a higher transition energy for E1–HH1 transition than for E1–LH1 transition as soon as phosphorus composition is greater than 5%. Fig. 3 shows the variation of E1–HH1 and E1–LH1 energy transitions as a function of phosphorus composition for 100 Å thick quantum wells. A good agreement is obtained between experimental and calculated results. Note that we used a band offset between GaAsP and GaAlAs which affects 100% of the unstrained gap difference to the valence band discontinuity. The inset in Fig. 3 shows that above 30% of phosphorus, the difference (E1–LH1) − (E1–HH1) decreases and nearly vanishes above 44%. This reflects the shrinkage of the carriers confinement in the well and their progressive delocalisation in the whole cavity.

To complete this study, we report preliminary results concerning tensile-strained separate confinement heterostructure double quantum well (SCH-DQW) lasers. The P composition in the GaAsP quantum wells is 14% leading to an emission wavelength of 780 nm, the Al composition is 25% in the barriers and 44% in the cladding regions. Broad area lasers operate with a threshold current density of 1.6 kA/cm$^2$ and an external quantum efficiency 0.45 W/A for 0.7 mm long devices. The internal waveguide losses are of about 6 cm$^{-1}$ which are typical for this kind of device. These results compare favourably with laser performances on single quantum well structures reported
by Agahi et al. [19] except for the threshold current density. These laser structures [19] were grown by MOVPE but with an Al content in the barrier of 34%. The lower threshold current density reported by Agahi et al. can be explained by an increase of Al content in the barrier which decreases both population of hole and conduction electron in the cavity and so the current losses due to radiative and non-radiative recombinations in the cavity. Note that the population inversion depends mainly on the properties of the QW layers, so the gain threshold condition is unchanged for both cases. In conclusion, the Al fraction in the cavity material has to be increased up to 30% while the GaAlAs confinement barrier has to be consequently increased to at least 55% in order to maintain a significant optical confinement.

6. Conclusion

In summary, we have demonstrated the possibility of using alternative group V organometallic precursors (TBAs, TBP) for the growth by CBE of GaAlAs/GaAsP quantum well structures. Carbon and oxygen concentrations in GaAlAs layers are in the mid-$10^{17}$ cm$^{-3}$ range whatever the aluminum composition used ($<$55%). In situ measurements of P and As incorporation rates under Ga excess conditions did not appear suitable for the determination of group V alloy composition. Finally, GaAsP QW structures have been grown exhibiting high luminescence efficiency for P compositions ranging from 5 to 28%. The QW photoluminescence energy as a function of P mole fraction is found to be well accounted for by envelope function calculations including strain effects. At least for $y_P \leq 25\%$, a good agreement is obtained between experimental and calculated E1–HH1 and E1–LH1 energy transitions. These calculations have been used to optimize GaAlAs/GaAsP laser structure designs.

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References

Contamination in molecular beam epitaxy: the role of arsenic drag effect

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Abstract

We have measured levels of Al, In and Si in nominally pure GaAs layers grown by molecular beam epitaxy as a function of arsenic flux for a number of different combinations of idling temperatures of Al, In and Si cells, using either As\textsubscript{2} or As\textsubscript{4} arsenic molecules. We find that a considerable number of atoms find their way to the growing layer in spite of closed shutters and no direct line of sight from their effusion cell to the wafer surface. The number of these atoms is proportional to the arsenic flux used and to the equilibrium vapor pressure over the considered element. We present arguments for the existence of the “arsenic drag” effect which deflects a fraction of the atoms that by-passed their effusion cell shutter towards the wafer.

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\textit{Keywords:} MBE; Contamination; Molecular scattering

1. Introduction

Much progress has been made in the field of purification of elements used for molecular beam epitaxy (MBE) with the result that in many areas quality of sources available in the market is no longer the limiting factor. In spite of that, large gaps still exist between the quality of the best and “typical” layers grown in any given class. In order to bridge this gap, a better understanding of the remaining factors is needed in areas such as MBE system design, its preparation, and growth procedures. In the present work, we examine a contamination source which for the past four years has been controversial, namely unintentional doping with elements from idling effusion cells. Proposed mechanisms ranged from suboxide transport [1] to direct re-evaporation from coated shutters and inadequately cooled cell ports [2]. Although plausible under some circumstances, none of the proposed mechanisms explains the growing evidence.

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that such contamination is quite common, even in the absence of either of the conditions considered. In the present paper, we demonstrate that this contamination is linked to the direct interaction between molecular beams traversing the chamber, a process which has been regarded so far as negligible in the typical MBE environment.

2. Experimental procedure

The experiment was performed using a modified [3] V80H VG-Semicon MBE machine with effusion cells configuration as shown in Fig. 1a. The wafer temperature in the growth chamber is monitored using an infrared pyrometer calibrated against melting point of InSb. Molecules desorbing from the central 2 cm\(^2\) of the wafer can be monitored with the narrow-acceptance angle, ion-counting Quadrupole Mass Spectrometer (QMS) located in the vicinity of the In cell port. The shutters in Fig. 1a are shown in the positions used for the “collecting planes” (to be defined later) and the “dark” ports indicate cells not used in the present experiment. Fig. 1b shows the 3D perspective view of the shutters within the cell ports. The system of tantalum panels shown, which isolates the effusion cells, is in good thermal contact with the bottom of the cryoshroud. This minimizes potential thermal cross-talk and cross-contamination between the cells. With shutters in closed position, no point on a 3" substrate lies in a direct line of sight from the emitting surfaces of any effusion cells. The shutters, however, do not provide a tight seal of the cell compartment, leaving considerable gaps through which some of the flux can enter the growth chamber. Examples of such peripheral molecular beams are shown in Fig. 1c–Fig. 1e. Fig. 1c shows the approximate intersection of the As beam with the peripheral In beam. A high-capacity EPI valved cracker cell was used as a source for both As\(_2\) and As\(_4\). The composition of the flux leaving this cell was determined by the temperature of its Rhenium cracking zone, giving predominantly As\(_2\) flux at 950°C, and predominantly As\(_4\) flux at 400°C. To quantify the arsenic flux incident on the wafer surface, we have calibrated the background pressure readings of the ion gauge placed between the growth chamber and one of the two cryopumps. A reading during typical growth should be proportional to the arsenic flux used. For the calibration purpose, while growing GaAs at 2 Å/s and 600°C, the arsenic flux (As\(_2\) or As\(_4\)) was decreased gradually until the transition to 4 × 2 Ga-stabilized surface reconstruction had been detected with RHEED. This transition occurs very abruptly and can be clearly detected even with wafer rotation. At this point [4] sticking coefficients are equal to 1 and 0.5 for As\(_2\) and As\(_4\), respectively. With GaAs grown at the rate of 2 Å/s, the gallium flux at the surface is 4.42 × 10\(^{14}\) atoms/cm\(^2\) s. This offers reference points at molecular fluxes of 2.21 × 10\(^{14}\) molecules/cm\(^2\) s for both As\(_2\) and As\(_4\). The background pressures we registered for such conditions were 5.5 × 10\(^{-8}\) and 2.0 × 10\(^{-7}\) mbar for As\(_2\) and As\(_4\), respectively. Using this calibration we converted the background pressure \(P_{\text{bkg}}\) to the As flux \(\phi\) incident on the wafer using the following formula:

\[
\phi = \begin{cases} 
2.21 \times 10^{14} \left( \frac{P_{\text{bkg}}}{5.5 \times 10^{-8}} \right) \text{ molecules/cm}^2 \text{s} & \text{for As}_2, \\
2.21 \times 10^{14} \left( \frac{P_{\text{bkg}}}{2.0 \times 10^{-7}} \right) \text{ molecules/cm}^2 \text{s} & \text{for As}_4.
\end{cases}
\]

Fig. 2 shows the functional diagram of the growth recipe designed to test Al, Si, and In cells for their ability to deliver their respective elements to the wafer surface in spite of the closed shutters. The schematic of the layer grown is shown on the right-hand side of the figure (growth progresses in the bottom-top direction). The information to the left of the layer schematic shows, for different segments of the layer deposited, the type of molecules leaving the arsenic cell, its valve setting (3, 2, 1 squares\(^1\) and closed), as well as the Al, Si and In nominal\(^2\) fluxes. To overcome SIMS sensitivity problems and to minimize the sputtering time, the previously

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\(^1\) We use here the “square” unit for simplicity. For the actual valve controller the 3, 2, 1 squares and closed translate to 180, 120, 60 and 0 setting, respectively. The arsenic flux leaving the cell is approximately proportional to these settings. 

\(^2\) By “nominal” we mean here the flux which would be incident on the wafer surface if the cell shutter was open.
Fig. 1. Effusion cell configurations in the growth chamber: (a) Relative placement of the effusion cells with respect to each other. The shutters are drawn in the positions used for the "collecting planes" and the "dark" ports indicate cells not used in the present experiment. (b) 3D perspective view of the shutters within the cell ports. The system of tantalum panels shown isolates the effusion cells from each other. (c)-(e) Cross sections through the growth chamber showing the cell designs and their positioning with respect to the shutters as well as the peripheral beams which enter the chamber via the gaps between the shutters and the panels.
mentioned collecting planes were introduced. At this stage GaAs growth was interrupted for 30 min (the planes are indicated with the dashed lines). The arsenic valve settings shown are the settings used during corresponding growth interrupts. For the 1000 Å GaAs spacing layers, an arsenic valve setting of 0.5 (in the “square” units) was used (not indicated in Fig. 2). The nominal fluxes for Al, Si and In were calculated from their respective cell temperatures using typical calibration procedures. Note that the Al flux was reduced in steps for the P_9 and P_10 collecting planes (keeping the As_4 flux constant). This was done to measure the influence of the nominal Al flux value on the contamination of the layer with Al atoms. The vertical line that extends down from the In-related flux marker indicates that, even for this over 100 times lower setting (flux of 2.9 \times 10^{-2} \text{ 1/cm}^2\text{s}), considerable In contamination was subsequently measured with SIMS.

The growth was performed on N + (001)GaAs 2° epi-ready substrates. The wafer was mounted in 3° molybdenum holder using an In-free mounting technique. Prior to transferring into the growth chamber, the wafer was outgassed for 2 h in the preparation chamber at 600°C, as indicated by the outgassing stage thermocouple (the actual wafer temperature is 50–70°C lower). In order to remove possible remains of oxide, the wafer was further annealed for 30 min at 650°C in the growth chamber under arsenic flux. Subsequently, the substrate temperature was lowered to 600°C and a buffer layer of 500 Å GaAs was grown followed by smoothing superlattice of ten periods of 20 Å GaAs/20 Å AlAs and 2000 Å GaAs layer. During the growth of the 2000 Å GaAs layer the substrate temperature was lowered to 500°C in order to limit diffusion and/or segregation of the contaminants deposited on the collecting planes.

SIMS profiles of Al, In and Si were performed ex-situ with Cameca IMS 4f secondary ion mass spectrometer. Si was detected as the negative ion with 14.5 keV Cs⁺ primary beam, while In and Al were both detected as positive ions with 2.75 keV O₂⁺ primary beam. The primary beam was rastered over 350 × 350 μm area and a 63 × 63 μm was analyzed. The calibration of the absolute Si concentration was done with an implant standard. The In standard was obtained by profiling In_{0.17}Ga_{0.83}As/ GaAs multiple quantum well (3 periods of 430 Å GaAs/70 Å InGaAs) where the total amount of In was derived independently from X-ray rocking curve measurement. To calibrate the Al contamination level, we have normalized the SIMS counts to the total amount of Al expected in the buffer AlAs/GaAs superlattice of the structure studied.

The top three panels in Fig. 3 show the results of SIMS profiles for Al, Si and In plotted using linear scale. Note that the profiles are plotted with the substrate on the left-hand side of the plot. The amounts of respective elements deposited on each collecting plane are proportional to the shaded
are plotted versus the distance from the surface. In spite of this disparity, the periodic nature of the structure (1800 s interrupt/500 s of GaAs growth) allowed us to align the SIMS peaks with corresponding arsenic flux traces. The strong correlations between the amount of contamination on each collecting plane with the product of the arsenic flux and the respective element nominal flux, can be seen in the figure without any numerical analysis. As mentioned earlier, traces of In are found on all collection planes, even those where In was idling at over 200°C lower than the nominal setting of 932°C used to test its contaminating potential. On the other hand, the QMS counts of the atomic In, showing clear correlation with the arsenic flux, are seen only for the high nominal In flux. In this case, however, we are detecting only In atoms which by-passed the cell shutter and were back-scattered into the narrow acceptance angle of QMS before reaching the LN₂-cooled shroud (see Fig. 1c).

3. Analysis and discussion

Fig. 4 shows the results of analysis of the experimental data discussed above. To facilitate comparison between the three elements studied, the data are plotted on a log–log scale. On the X-axes we plot the average As flux impinging the wafer surface during the given growth interrupt. The closed symbols represent the data obtained with As₂, while the open symbols correspond to the data obtained with As₄. The Y coordinates were obtained in the following way. First the surface density of the element on the collecting plane was calculated by integrating the area under the SIMS profile of the corresponding region. This quantity was then converted into the effective contamination flux by dividing the calculated density over the 1800 s it took to accumulate it. The flux leakage ratio plotted is the ratio of this contamination flux to the nominal flux of the element (flux which would be arriving at the surface if the shutter was opened). In much simplified but practical terms this ratio may be looked upon as the shutter-effective transparency. The lines plotted through the points (note the log–log scale) are of the form \( Y = zX \), where the coefficient \( z \) was fitted using the least-squares algorithm applied to the areas under the corresponding peaks. Dark-shaded banners included in each panel indicate the nominal flux for the given element (proportional to the banner height) as used in Fig. 2. Below the SIMS profiles are shown the arsenic fluxes incident on each collecting plane. The fluxes were calculated directly from the recordings of the background pressure in the growth chamber using Eq. (1). The arsenic fluxes are also plotted using linear scale, and the 30 min growth interrupts are marked by the shading under corresponding stretches of the recordings. The bottom panel in Fig. 3 shows the QMS readings at a.m.u. of 115, which corresponds to single ionized atomic In. Note that the last two panels are plotted versus time, while the first three
experimental data separately for every element, but combining As$_2$ and As$_4$ points. It is clear that such a simple relationship describes experimental results very well, proving that for all the cells investigated, the leakage rates are proportional to the arsenic flux used. The most complete set of data is available for Al, which follows such proportional behavior for over two orders of magnitude change in the incident As flux. Fig. 5 shows the Al contamination flux for three different settings of the Al cell temperature using the same As$_4$ flux of $3.1 \times 10^{15}$ molecules/cm$^2$s. It is clear that the Al contamination is proportional to the Al nominal flux, indicating that no intermediate steps in the Al transport are present (such as re-evaporation from the shutter edges). Qualitatively, we see the same behavior for In and Si.

The experimental results discussed above provide a solid evidence that the direct interaction between the peripheral molecular beams from Al, In and Si cells with the arsenic molecules traversing the growth chamber is responsible for deflecting trajectories of these atoms towards the substrate and towards the source flange. Since the correlation was established by monitoring the As flux using the background pressure, the actual proportion of scattering events caused by the direct arsenic beam and those caused by the background arsenic molecules remains speculative. The direct arsenic molecular beam likely plays the dominant role in deflecting other atoms towards the substrate, because within its range (i) the beam-equivalent pressure is
considerably higher than that of the background arsenic and (ii) the beam is very directional, thus providing efficient momentum transfer in the direction of the substrate. This resembles the mechanism of a UHV diffusion pump [5], where heated vapor is injected into high-vacuum region through a system of narrow nozzles. Rapid expansion gives rise to the supersonic jets, that provide enough downward molecular “drag” to sustain a very large differential pressure between the high- and low-vacuum regions within the pump. This resemblance is particularly striking for the case of the cracker design used in our system, where the arsenic molecules are rapidly decompressed from relatively high-pressure bulk evaporator region (pressure approaching 1 mbar) into the UHV chamber through a very narrow (a couple of millimeters in diameter) nozzle. On the other hand, for the case of the atoms deflected towards the source flange (as demonstrated by correlations of the QMS reading at 115 a.m.u. with the arsenic background pressure), significant role is likely played by other arsenic molecules bouncing around the chamber, particularly the excess arsenic re-evaporated from the substrate. The process of redirecting the atoms back to the source flange has been used in the past to study flux transients of group III atoms with QMS [6]; however, the mechanism responsible was not identified at that time. The same phenomenon explains also many of the elusive memory effects and cross-contamination problems in MBE systems. Indeed, with the aid of arsenic scattering, material from one cell can contaminate another cell. The level of such contamination can build up over time and will be particularly aggravated if high arsenic fluxes are used and many shutters are opened at the same time.

Another issue which cannot be conclusively resolved at present is the relative contributions of the nonreactive (elastic and inelastic) scattering events to the reactive processes, where as a result of collision a new molecule is formed (say As$_2$ + In $\rightarrow$ As$_3$In). The dominant role of the nonreactive scattering in the observed wafer contamination finds indirect support in the very similar values for the scattering efficiencies measured for As$_2$ and As$_4$ molecular beams (see Fig. 4). Indeed, the arsenic dimer and tetramer show very different behavior in coating MBE chamber parts, as well as interact in a different way with the growing layer surface [7, 8]. Thus, it is reasonable to expect that they should also show much different reactivity when it comes to bonding single group IV or group III atom in the process of collision. A simple estimate shows that in our experiments the average momentum of As$_2$ molecule is very close to that of As$_4$ (As$_3$ is half the mass, but it leaves the cracking zone with roughly twice the speed). Since the total momentum is conserved in the reactive collision, it would require very similar reactivity of As$_2$ and As$_4$ to explain the similarity of measured dependences. However, since the complexity of atomic scattering mechanisms is considerable [9], a dedicated study on crossed molecular beams would have to be performed to shed more light on this issue.

Note that In leakage rate is about 40 times larger than that measured for Al, while Si is in between the two. Such large differences in the flux leakage rates measured for different effusion cells can be understood qualitatively by examining the individual cell designs and locations (see Fig. 1c and Fig. 1d). Indeed, the direction of the “stray” In beam and its overlap with the direct As flux ensures the maximum momentum transfer between the two. Also, the In cell is a dual filament “hot-lip” design which will give relatively strong flux at large angles to the cell axis. In contrast, the Al cell is of a “cold lip” design which minimizes the risk of Al “creeping out” of the crucible. In our case the “cold lip” protection is supplemented by the dual crucible arrangement which minimizes the risk to the cell in the event of the crucible cracking. The final result is that the top portions of the walls for Al crucible emits very little, and the rings of Al which accumulate there further reduce the flux leaving the cell at large angles, thus strongly reducing the Al peripheral beam. The Si cell is most conveniently compared to the Al cell. Indeed, both cells are located symmetrically on each side of the arsenic cracker cell; thus, one would expect similar effective intersection of their respective “stray” beams with the arsenic beam. The Si cell, however, is located higher up in the source flange and its 5 cm$^3$ conical crucible is loaded with strips of cleaved Si wafer. The orientation of the strips within the crucible is such that its emission profile should be much closer to
that characteristic of the hot-lip cell, giving a considerably stronger peripheral beam than that for Al cell.

In summary, we find that hot effusion cells can contaminate the wafer surface with their respective elements in spite of the closed shutter and no direct line-of-sight from the crucible to the wafer. The amount of such contamination is proportional to both the arsenic flux used and the element vapor pressure over the crucible. The efficiency of such process is similar for As$_2$ and As$_4$ molecules, regardless of the contaminating material. All these observations are qualitatively consistent with the model where the peripheral atomic beams traversing the growth chamber are scattered towards the wafer surface by the arsenic molecular flux. The scattering efficiency is sufficient to redirect a fraction of the atoms back to source flange before they are chemisorbed at the shroud walls. We present arguments in support of a model in which the process is dominated by the nonreactive scattering; however, this particular issue can be resolved only through further dedicated experiments. Most of the problems related to the effects described are best taken care of by proper MBE system design. However, with existing systems many of the negative consequences can be avoided by altering certain growth procedures or simple modification of the flux blocking structures adjacent to the cell ports.\footnote{For instance, by employing appropriate growth procedures, we have been able to grow reproducibly in our system AlGaAs/GaAs ultra-high mobility two-dimensional electron gas structures (peak mobilities up to 6400000 cm$^2$/V s)\cite{10} while the same structures grown without the corrective measures are significantly worse (peak mobilities up to 1000000 cm$^2$/V s).}

Although the experimental results presented pertain to scattering by arsenic molecules, the physics behind this phenomenon is very general. The importance of the effect for other material systems and molecular beams will depend on the growth conditions and design of the apparatus used.

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References

Photoluminescence and X-ray characterization of relaxed \( \text{Si}_{1-x}\text{Ge}_x \) alloys grown on silicon on insulator (SOI) and implanted SOI substrates

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Abstract

Boron and oxygen were implanted into the \( \text{SiO}_2 \) region of a thin silicon on insulator (SOI) substrate to form borosilicate glass and reduce the reflow temperature needed for compliant substrates. The effect of lowering the reflow temperature was examined by characterizing a strained \( \text{Si}_{0.7}\text{Ge}_{0.3} \) layer which was grown on the implanted compliant substrate by molecular beam epitaxy. Photoluminescence (PL) and triple-axis X-ray diffraction were used to characterize film quality, Ge concentration and percent relaxation. A marked decrease in the annealing temperature required (roughly a difference of 300°C) to relax the \( \text{Si}_{0.7}\text{Ge}_{0.3} \) layer was fully observed. The PL spectra show a broad band around 800 meV that is believed to be related to the relaxation of the SiGe layer. The intensity of this peak first increases and then decreases dramatically when the annealing temperature reaches the point where the SiGe layer begins to relax or the underlying glass begins to flow. For the sample grown on the boron and oxygen implanted BESOI, this low-energy peak is shown to disappear completely at an annealing temperature of 900°C. Above 900°C, we begin to see the evolution of near band-gap luminescence for the \( \text{Si}_{0.7}\text{Ge}_{0.3} \) layer grown on implanted BESOI. An energy peak which shifts to higher energies, from 970 to 1025 meV, with greater relaxation is only observed for the sample grown on the implanted substrate. It is believed that the observation of near band-gap luminescence is primarily due to the reduced reflow temperature and a low-defect dislocation concentration in the \( \text{Si}_{0.7}\text{Ge}_{0.3} \) film.

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1. Introduction

Recently, there has been an interest in achieving high-mobility n-channel field-effect transistors by using tensilely strained silicon for the electron channel [1, 2]. Electron mobilities greater than 520,000 cm²/V s at 4.2 K have been reported [2] for tensilely strained silicon on relaxed Si₁₋ₓGeₓ. Another application of such a high-quality relaxed SiGe buffer layer is for optical devices where dislocations are of important concern [3, 4]. Normally, relaxed Si₁₋ₓGeₓ films have been grown on abrupt or more recently graded buffer layers [5]. The major drawbacks have been high dislocation densities for the former and extremely large buffer thicknesses for the latter.

In 1994, Powell et al. [6] proposed a novel approach in which a very thin 'free-floating' buffer layer is allowed to absorb most of the strain of a SiGe epilayer grown on top of it. The dislocations develop in the buffer layer and allow the epilayer to relax its strain without formation of dislocations. The 'compliant substrate' may consist of a thin monocrystalline Si layer on top of a SiO₂ layer on top of a silicon substrate. In this approach, the surmised mechanism leading to the reduced dislocation density is the viscous flow of the buried SiO₂ layer, which allows the Si to float and the strain in the SiGe layer to be relaxed by the introduction of dislocations primarily in the thin Si layer. Recently, we have investigated the quality of annealed Si₉₀.₈₆Ge₀.₁₄ films grown on thin Bond- Etchback silicon on insulator (BESOI) wafers using photoluminescence and X-ray diffraction techniques [7]. We presented evidence for partial strain relaxation of these layers, resulting in higher film quality than for the same layers grown on bulk Si. However, the temperatures required for strain relaxation of the Si₉₀.₈₆Ge₀.₁₄ film were high (in the range of 950°C–1100°C). Such a high temperature would cause the strained epitaxial film to develop dislocations before the substrate becomes 'free-floating'. In this paper, we present an approach using low-temperature glasses formed by ion implantation to lower the reflow temperature of the buried oxide layer such that the strain may be relaxed before unwanted dislocations develop in the epitaxial layer.

2. Experimental procedure

Phosphosilicate (PSG) or borosilicate (BSG) is known to have a lower reflow temperature than SiO₂ [8]. By implanting SiO₂ with phosphorous and a subsequent implant of oxygen to activate the phosphorous, these glasses can be formed [9]. Likewise, borosilicate glass can be formed in a similar manner. The main advantage for the latter is that the boron ion is smaller than the phosphorous ion and hence will result in smaller surface damage. Thus, annealing or recrystallization will be easier from the surface, in view of the fact that there is the absence of a bulk 'seed'. This lower reflow temperature offers several important advantages. First, it allows strain relaxation at lower temperatures which would precede mechanisms likely to occur at higher temperatures, like Ge segregation, surface roughening, and dislocation formation. Second, it allows for the possibility of in-situ reflow annealing and thus allows for better control over the growth conditions of SiGe epilayers.

The BESOI substrates in this work were produced by Sibond, Inc. [10]. The silicon and silicon dioxide layers were initially 200 and 230 nm thick, respectively. The silicon film was subsequently etched using an anisotropic etchant, tetra-methyl ammonium hydroxide (TMAH) to roughly 60 nm. One of the BESOI substrates was then implanted first with boron and then oxygen. The implant fluences were determined by the stoichiometrical ratio of B₂O₃ and the targeted concentration of 5% by weight. We determined the implanting energies by setting the projected range, Rₚ, to approximately the center of the SiO₂ layer. The actual energies and fluences were 40 keV and 9.1 × 10¹⁵ cm⁻² for boron and 50 keV and 1.4 × 10¹⁶ cm⁻² for oxygen, respectively, [11]. After the implant and a 800°C anneal for 30 min, the top Si layer was thinned down further to the desired thickness (~ 20 nm). This extra step not only allows the top thin Si layer to be strained or relaxed easily but also allows us to etch off some of the Si surface damaged by the implantation process. The final thicknesses of the Si and the SiO₂ were confirmed using an ellipsometer. The surface roughness was characterized by an atomic force microscope and the root mean squared roughness was measured to be about 11 Å.
After the substrates were cleaned using a modified Ishizaka and Shiraki cleaning procedure [12], they were attached to a single holder and loaded into a Perkin-Elmer Model 430S MBE system. The substrates consisted of a p-Si substrate, an unimplanted BESOI substrate, and a B and O implanted BESOI substrate. Following a 10 min. thermal cleaning at 900°C, 10 nm of Si, 120 nm of Si_{0.7}Ge_{0.3} and 5 nm of Si were deposited consecutively at 475°C. All photoluminescence (PL) measurements were measured at 4.2 K with an excitation wavelength of 488 nm from an Ar laser. The actual Ge concentration was verified by high resolution triple-axis X-ray diffraction (TAD) using both symmetric [0 0 4] and asymmetric [2 2 4] scans.

3. Results and discussion

Figs. 1–3 show PL spectra as a function of annealing temperature for the Si_{0.7}Ge_{0.3} layer grown on the plain p-silicon substrate (Sample A), unimplanted BESOI (Sample B), and B and O implanted BESOI (Sample C), respectively. From the as-grown spectra in all cases Fig. 1a, Fig. 2a and Fig. 3a, we can see luminescence attributed to the substrates which are the peaks above 1 eV and a lower-energy broad peak sometimes seen on MBE grown samples [13–15]. For Sample A, as we slowly increase the annealing temperatures from 800°C, we see that the luminescence of the broad peak increases in intensity relative to the p-substrate luminescence in Fig. 1, which is to be expected since this broad energy peak is usually associated with dislocations. For sample B, the layers grown on the unimplanted BESOI substrate, we see an increase and then a decrease of the broad energy peak intensity. The intensity begins to decrease around 1100°C. This low-energy broad-band luminescence appears to be related to the T-band luminescence as seen by others [15] since it has the same temperature and band-gap dependences as the T-band. Its relationship to D1 or D2, whose energies are similar to this broad luminescence, is unknown at this time. For Sample C, the broad peak is more noticeable after a 800°C annealing. However, at temperatures above 900°C the broad peak disappears and a higher-energy peak begins to appear (900 meV–1.04 eV).

![Fig. 1. PL Spectra of Sample A as a function of annealing temperature and time — Si_{0.7}Ge_{0.3} alloy layer on a p-Si substrate; (a) as grown, (b) 800°C–30 min, (c) 800°C–60 min, (d) 900°C–30 min, (e) 900°C–60 min, (f) 1000°C–30 min, (g) 1000°C–60 min, (h) 1100°C–30 min, (i) 1100°C–60 min.](image)

As confirmed by high-resolution triple axis X-ray diffraction (TAD), the evolution of the low-energy peak in the luminescence demonstrates the relaxation mechanism. It can be seen for Sample C that the temperature at which this band reaches the maximum intensity occurs when annealed around 800–850°C for 30 min. Whereas for the unimplanted Sample B, this temperature and time are around 1100°C for 60 min. For Sample A, which is grown on a conventional p-silicon substrate, the peak never decreases in the annealing temperature range examined. It is assumed that if it does begin to decrease, it is obviously at a higher temperature and may be attributed to other mechanisms due to its extreme temperature. Hence, we can see that we have effectively lowered the reflow temperature by about 300°C by the use of implantation. This lower energy broad band also appears to correspond to
the mosaic spread in the high-resolution X-ray spectra observed from the sample (not shown) and in Ref. [7]. The decrease in the mosaic spread in the TAD spectrum can be explained by noting that when the strain of the SiGe layer is transferred to the thin Si layer fully or partially (with and without accompanying dislocations), the Si$_{0.7}$Ge$_{0.3}$ layer is ‘smoother’. The disappearance of this broad low-energy luminescence peak can be directly attributed to the relaxation of the Si$_{0.7}$Ge$_{0.3}$ without accompanying dislocations and to the subsequent transfer of the strain to the thin silicon layer fully or partially. Assuming no dislocation formation, the equilibrium relaxation of the finite bilayer system should be roughly 75% using the formulation of Huang et al. [16]. This 75% relaxation (confirmed by TAD) of the top SiGe layer, for example in the case of Sample C, occurs at an annealing temperature of 850°C when the broad low energy luminescence peak reaches a maximum. However, after further annealing (1000°C or above), the layer fully relaxes (95%). This can be attributed to the fact that the bottom thin silicon layer relaxes by way of dislocations. At higher temperatures, the thickness of the thin Si layer (~ 20 nm), will surpass the critical thickness determined by the Si$_{0.7}$Ge$_{0.3}$ ‘substrate’ and the thin Si film will be dislocated to relieve the strain. Hence, the Si$_{0.7}$Ge$_{0.3}$ layer is free to relax without formation of misfit dislocations in the Si$_{0.7}$Ge$_{0.3}$ layer itself. The disappearance of the broad low-energy luminescence peak also supports the notion that this peak does not arise from the dislocations themselves, but rather from the strain field around the SiGe/Si interface. This is especially
true in the case of the compliant substrates since it would not be expected that the dislocations will disappear at higher temperatures. Hence, in the case of the compliant substrates, the low-energy luminescence may arise due to the strain field around the interface.

Fig. 3 also shows the evolution of several higher energy peaks (900 meV–1040 meV) for Sample C at annealing temperatures of 900°C and above. We attribute these peaks to band-gap related luminescence both the NP and $\text{TO}_{\text{Si-Si}}$ phonon replica. The energy values of 1.015 and 0.935 eV from Fig. 3(i) correspond well to published reports [17, 18], particularly if we use the 95% relaxation value from TAD. The temperature dependence of these peaks follows the same temperature dependence as reported by others for SiGe near band-gap luminescence [19]. The 1.015 eV peak disappears around 55 K and shifts to lower energies with increasing temperature. The strain relaxation dependence discussed previously also supports our assignment of these peaks since we expect that at higher annealing temperatures the SiGe layer will relax more and that the SiGe band-gap will shift to higher energies as illustrated in Fig. 3.

The energy peaks around 900–1040 meV in Fig. 3 are shown to broaden as the annealing temperature increases beyond 900°C. The reason for the broadening is not fully understood at this time but may be related to Ge segregation at higher temperatures or more likely some relaxation distribution in the SiGe layer. Etching of the SiGe layer confirms that the near band-gap luminescence does indeed come from the SiGe layer and not from the thin silicon layer, nor the substrate. The fact that this near band-gap luminescence is not evident in the PL spectra of Samples A and B, may be attributed to the presence of dislocations or other mechanisms which compete for the recombination of the same carriers. We should also note that since the $\text{Si}_{0.7}\text{Ge}_{0.3}$ layer is 1200 Å, it is larger than the conventional critical thickness for a pseudomorphic growth of $\text{Si}_{0.7}\text{Ge}_{0.3}$ on Si. Augmentation of the pseudomorphic thickness may be expected since, for lower reflow temperatures, the thin silicon layer of the BESOI substrate should accommodate some of the strain within the thin silicon layer [16].

4. Conclusions

We have successfully demonstrated the lowering of the reflow temperature of BESOI by about 300°C by implanting the SiO$_2$ layer with both boron and oxygen. Near band-gap luminescence from the relaxed $\text{Si}_{0.7}\text{Ge}_{0.3}$ layers grown on implanted BESOI was observed. In addition, we showed the decrease and disappearance of a low energy ($\sim$ 0.80 eV) broad feature in the PL spectra. This low energy feature was shown to be directly related to the relaxation of the $\text{Si}_{0.7}\text{Ge}_{0.3}$ layer and possibly the generation of dislocations in the underlying Si layer. It may be possible to further lower the reflow temperature of the SiO$_2$ layer by implanting the region with a higher concentration of boron and oxygen. The approach described in this paper may be used to grow high-quality strained Si/SiGe on SOI. Likewise, this approach may be extended to other strained layer systems, such as GaAs on Si, etc.

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References

Modulated-beam studies of the layer-by-layer etching of GaAs(0 0 1) using AsBr₃: identification of the reaction mechanism

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Abstract

Modulated-beam mass spectroscopy (MBMS) has been used to study the reaction mechanism of the layer-by-layer etching of GaAs(0 0 1) using AsBr₃ under molecular beam epitaxy conditions. It is shown that GaBr is the main etching product and its “delay” time with respect to the incident AsBr₃ flux exhibits a strong dependence on substrate temperature, changing from 12 ms in the reaction limited regime at 387°C to less than 0.5 ms at 560°C. Desorbing As-containing species appear to have very short surface lifetimes throughout the temperature range investigated and the additional As₂ flux supplied from a solid source has no effect on the etching rate. The results suggest a reaction pathway where the rate limiting step is the formation or desorption of GaBr and not the decomposition of AsBr₃.

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1. Introduction

It has recently been demonstrated that III–V semiconductor materials such as GaAs and InP can be chemically etched in an ultrahigh vacuum (UHV) chamber using molecular beams of group-V halides, e.g. AsCl₃ [1, 2], PCl₃ [3] and AsBr₃ [4]. The etching process on GaAs(0 0 1) has been monitored by reflection high-energy electron diffraction (RHEED) [1, 4]. Specular beam RHEED intensity oscillations were observed over a wide temperature range indicating that the etching process occurs on a layer-by-layer basis, similar to the corresponding growth processes that are well known on this surface [5]. The combination of atomic controlled etching and growth in the same UHV chamber offers considerable promise for the fabrication of novel low-dimensional semiconductor heterostructures [6].
Kinetic information regarding the etching rates using AsBr₃ and their dependence on flow rate and substrate temperature has been obtained from RHEED intensity oscillation measurements on GaAs(0 0 1) [4, 7]. Oscillations were observed over a wide range of substrate temperatures, between 350 and 650°C, and the etching rate was found to be constant above 460°C but decreased significantly at lower temperatures. Two etching regimes were therefore identified, a supply-rate-limited regime at high temperatures ( > 460°C) and a reaction-rate-limited regime at low temperatures (<460°C) [4]. Furthermore, measurements have been made across AlGaAs/GaAs heterojunctions and the etching rate was found to be determined by the surface composition of the group-III elements [7].

Although RHEED intensity oscillation measurements can provide information regarding the rates of the etching process, it is not a species-specific technique and provides no direct information on reaction pathways. In contrast, modulated beam mass spectroscopy (MBMS) is ideally suited for detailed kinetic studies and it has been used successfully as a probe of reaction mechanisms in the growth of GaAs by molecular beam epitaxy (MBE) [8, 9] and metalorganic molecular beam epitaxy (MOMBE) [10, 11]. In common with other modulation-based techniques, MBMS has the advantage that it allows the detection of small signals over a large background. Modulation of the desorbing beam enables the species leaving the surface to be identified, whereas modulation of the incident beam allows the determination of reaction orders and surface residence times [8].

In this paper, we present MBMS measurements for the in situ etching of GaAs(0 0 1) using AsBr₃. We find the dominant etching product to be GaBr and its delay time with respect to the incident AsBr₃ is measured over a wide range of temperature. Our results suggest a reaction pathway where the rate-limiting step is the formation or desorption of GaBr and not the decomposition of AsBr₃ at the surface.

2. Experimental procedure

The experiments were performed in a purpose built metalorganic MBE system with additional group-V halide gas source and solid group III sources [12]. The growth chamber is also equipped with MBMS and RHEED systems. Epi-ready, singular GaAs(0 0 1) substrates were used and the initial oxide was removed in the growth chamber at 610°C in the presence of an As₂ flux supplied from a solid source cracker cell. Buffer layers of GaAs were grown using solid sources at 580°C to ensure that each subsequent etching experiment would not remove material and expose the underlying interface. The AsBr₃ gas source was maintained at 80°C to avoid condensation in the gas line and the AsBr₃ flow rate was 0.03 SCCM. A concomitant As₂ flux was used throughout all etching experiments to maintain an As-rich surface condition.

The MBMS system consists of a quadrupole mass spectrometer with a cross beam ioniser and two modulators. The ioniser of the mass spectrometer was surrounded by a liquid nitrogen tank, except for a solid angle directly facing the sample. The modulators were positioned in front of the entrance to the ioniser and the gas cell, to provide modulation of the molecular (atomic) fragments desorbing from the sample into the ioniser, or AsBr₃ molecules incident on the sample. A multichannel scaler (MCS) was used to collect the modulated signal which allows digital processing of the signal. This is a significant advantage over the conventional lock-in amplifier instrumentation as a number of parameters can be obtained using digital methods in addition to the phase angle and amplitude. It also allows signal averaging as the spectra is obtained through accumulation over many scans, thus improving the signal to noise ratio. The methods of obtaining these additional parameters, such as orders of reaction, can be found in the work of Foxon and Joyce [8, 9].

3. Results and discussion

The reaction products were identified by monitoring the desorbing beam during etching of GaAs(0 0 1) with AsBr₃ in both the supply- and reaction-limited regimes, as indicated by RHEED intensity oscillation studies. Modulated signals were detected in both regimes at mass-charge ratios (m/e) of 69 (Ga⁺), 148 (GaBr⁺), but no signals
were identified at 158 (Br$_3^+$) and 227 (GaBr$_3^+$). The specific mass charge ratios were chosen after consideration of the isotope distribution and the need to separate each species in the mass spectrometer. The intensity of the modulated signals at 69 and 148, determined from the amplitude at the fundamental modulation frequency, exhibited the same temperature dependence as the etching rate. Since the vapour pressure of Ga over GaAs is very low, it can be concluded that the main desorbing neutral species must be GaBr, and that the modulated signal at 69 derives purely from the cracking pattern of GaBr. There is, however, a possibility that the ionisation fragmentation patterns of GaBr$_x$ (x > 1) may not give a significant contribution to the signal at m/e = 227.

Modulation of the incident AsBr$_3$ flux allows determination of a number of kinetic parameters. Fourier analysis of modulated signals obtained at m/e = 69, 75 (As$^+$), 148, 150 (As$_2^+$) and 237 (AsBr$_2^+$) show characteristics of a linear response (zero even harmonics in Fourier analysis of the spectra) with respect to the incident AsBr$_3$ flux [8] in both supply-rate and reaction-rate limited regimes. This is consistent with the linear analysis using the convolution between the odd drive function and the impulse response, indicating that the apparent etching reaction is first order. This is in agreement with the etching-rate measurements obtained by RHEED in the supply-rate-limited regime due to the linear dependence of the etching rate on the incident AsBr$_3$ flux. However, rate measurements (using RHEED) in the reaction-rate-limited regime show a negligible dependence on the incident flux, i.e. the reaction is pseudo-zeroth order with respect to the AsBr$_3$ flux. The apparent discrepancy between results obtained from the two techniques can be explained as follows. A zeroth order reaction would not generate a modulated signal in an MBMS experiment. The zeroth order shown in rate measurements (RHEED) reflects saturation of the surface by AsBr$_3$. This would not be true if the AsBr$_3$ flux is reduced below a threshold value. Under the conditions used for MBMS, the AsBr$_3$ flux is modulated between zero and a designated flux which is higher than the threshold value. The part below this threshold value would give rise to the modulated signal detected in MBMS. Therefore, the first order indicated by the MBMS results reflects the condition when there is no saturation of AsBr$_3$ on the surface.

The surface delay time (lifetime) of the detected species can be obtained using a number of methods [8]. Fig. 1 shows the MCS spectrum obtained at an m/e ratio of 69 and at a substrate temperature of 387°C. Since the signal at m/e = 69 was due to the fragmentation of GaBr in the ioniser, the information obtained arises solely from GaBr. For a first-order reaction, the impulse response is an exponentially decaying function. Also shown in Fig. 1 are several examples of simulated spectra taking into account the flight time of the incident (mass = 312 amu) and desorbing neutrals (mass = 148 amu) and impulse response functions with different decay constants. Comparison of the simulated and experimental data indicates a surface delay time for GaBr of approximately 12 ms. This decreases with increasing temperature to a value of about 2 ms at 526°C. At a temperature of 560°C, the surface delay time is below the resolution of the instrument (0.5 ms) for the modulation frequency used.

The surface lifetime of the As-containing species are all very short compared to the resolution of the instrument at the modulation frequency used. Fig. 2 shows the temperature dependence of the etching rate at two different As$_2$ fluxes supplied from a solid source. The etching rate is identical under the two conditions indicating that As$_2$ has a negligible effect on the rate-limiting step of the etching reaction (at least within the range of As$_2$ flux used). The very short surface lifetime of the As-containing species over the temperature range studied suggests that dissociation of AsBr$_3$ is unlikely to be the rate-limiting step in the etching reaction. In contrast, the surface delay time of the etching product GaBr changes drastically as a function of substrate temperature in the reaction-limited regime. It is clear that either the formation or desorption of GaBr must be the rate-limiting step in the etching reaction. This result is entirely in agreement with previous RHEED studies which have looked at the effect on the etching rate of the surface composition [12], and results obtained from etching across AlGaAs/GaAs heterojunctions [7], where the etching rate was determined by the
Fig. 1. MCS spectrum obtained at 387°C for m/e = 69 and various simulated curves with increasing delay time.

Fig. 2. Etch rate as a function of substrate temperature at two different As$_2$ flux.
surface composition of the group-III elements. The As$_2$ flux supplied from the solid source has no effect on the etching rate or the position of the transition region between the supply-rate and reaction-limited regime. Clearly, the rate-limiting step is unlikely to depend on the surface As concentration.

4. Conclusions

MBMS has been used to obtain important kinetic information, complimentary to that obtained from RHEED intensity oscillations, concerning the layer-by-layer etching reaction of GaAs(0 0 1) using AsBr$_3$ in an MBE growth system. The underlying etching reaction is first order with respect to the incident AsBr$_3$ flux. The main reaction product has been identified as GaBr and the rate-limiting step in the etching reaction involves either the formation of GaBr or its desorption from the surface. A concomitant As$_2$ flux supplied from a solid source to maintain an As-rich reconstruction has little effect on the etching rate and suggests that the rate-limiting step does not depend on the surface As concentration.

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References

Morphology of MBE grown InAs films studied by atomic force microscope

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Abstract

The morphology of MBE grown InAs films on GaSb buffered GaAs(0 0 1) substrates with miscuts ranging between 0.2° and 4° has been studied by atomic force microscope (AFM). Step flow growth mode on nominal singular (1 0 0) substrates has been observed for samples grown at substrate temperatures higher than 500°C and for samples on miscut substrates grown at temperatures between 400°C and 530°C. Unstable growth mode has been observed on nominal singular (0 0 1) substrates when substrate temperatures are lower than 450°C. Relatively high density pits have also been observed on InAs surfaces with nucleation temperatures above 450°C.

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Molecular beam epitaxy (MBE) is a widely employed technique for growing III–V compound semiconductor materials and heterostructures. Because of its technical and scientific importance, MBE grown surfaces have been widely studied by a variety of techniques, such as reflection high energy electron deflection (RHEED), scanning tunneling microscope (STM), and atomic force microscope (AFM). RHEED has proven to be a powerful tool providing a lot of real time information, including growth mode transition, surface diffusion length, and surface reconstruction during the film deposition. However, RHEED can only provide averaged coherent diffraction information, failing to give local topographic information. More recently, STM and AFM have been employed to study the MBE grown films at various points during the evolution of the surfaces [1, 2], which have

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provided more detailed information about local noncoherent structures. However, among a large number of publications on STM and AFM studies on MBE growth of III–V semiconductors, most of them concentrate on GaAs [3, 4]. InAs is another III–V semiconductor with relatively narrow band gap (0.354 eV at room temperature). The novel broken-gap alignment with GaSb and small lattice mismatch between InAs and GaSb make the material very promising in infrared applications. In this paper we will report our AFM results of morphology of MBE grown InAs films on GaSb buffered GaAs substrates.

Our samples were grown on (1 0 0)GaAs substrates with a variety of miscuts ranging between 0.2° (nominally singular) and 4°. The As$_2$ was used as the As source. The typical As/In pressure ratio is about 15 for the sample grown at temperatures higher than the surface phase transition temperature [5], and 10 for the samples grown at relatively lower temperatures, the growth rate was approximately 0.3 μm/h. In order to make the substrate temperature measurements more reliable, the surface phase transition temperature during growth was employed as a calibration point. If the As/In beam flux ratio is high enough, the hysteretic range for this phase transition can be smaller than 2°C. The substrate temperatures during the growth was varied between 530 and 430°C. The samples consist of a 500 nm GaSb buffer layer grown at 500°C on GaAs substrates, a 10 nm InAs nucleation layer grown at either 430°C or higher temperatures, and an InAs layer with a thickness varying from 200 to 500 nm grown at different temperatures. Strong RHEED intensity oscillations were observed for the InAs films grown at temperatures higher than 500°C. However, these oscillations were absent for samples grown at temperatures lower than 460°C, even after 5 min annealing at 520°C.

The AFM measurements were conducted under ambient conditions. Although a native oxide layer has been formed, which is believed to be a few nm thick, the monolayer steps and terraces remain clearly observable. The sample grown on nominal singular (0 0 1) substrates show very large differences in surface morphology for different growth temperatures. As shown in Fig. 1a, for samples grown at 520°C, the surfaces are quite smooth and regular monolayer step edges with terrace widths of about 80 nm have been measured. However, in comparison with the samples grown at temperatures higher than 500°C, the surfaces of samples grown at temperatures below 450°C are relatively rough, as shown in Fig. 1b. Detailed examination of the surface reveals that the surface is formed by approximately circular shaped mounds with different sizes and heights. The very interesting feature of these mounds is that their slopes are approximately identical. The typical sizes of mounds are between a few hundred nm to a few μm. Their slope is
slightly larger than 1°. We have studied films grown at the same substrate temperature but with different thickness of InAs from 200–500 nm. The feature of the surfaces is almost the same as what has been shown in Fig. 1b. This means that the surfaces have reached a steady state after 200 nm growth at this substrate temperature.

This type of surface feature represents an unstable growth mode. It means that the growth on the singular surface is unstable. The growth will evolve into step flow growth mode through kinetic roughening. This kind unstable growth mode occurs when the competition between step flow and nucleation favors the nucleation of new islands. This happens when the terrace width is larger than the typical length of the adatoms travel before nucleating an island. However, this is not a sufficient condition for this unstable mode growth. For two dimension islands turning into multilayer mounds, the adatoms need to stay at the top of the islands rather than to jump off the terraces. Therefore, in order to sustain the unstable mode growth, the material should exhibit an energy barrier (Schwoebel barrier [6]) to prevent the adatoms from jumping off the terraces.

On a (0 0 1) singular GaSb surface which was grown on a GaAs(0 0 1) substrate, as shown in Fig. 2, the typical terraces width are larger than the distance of In adatoms traveling before nucleating an island on InAs surface. Therefore, it is very likely that the first few InAs layers take a two dimensional island nucleation growth mode. Once the islands have grown large enough, Schwoebele barrier will be formed to prevent the adatoms jumping off the terraces. The adatoms will be bounced back from the step edges, which will initiate the unstable growth.

In comparison with GaAs unstable mode growth, unstable mode growth on InAs surfaces exhibits a different feature. In the case of GaAs growth, the mounds are observed to be anisotropic [1, 7]. The STM and laser light scattering experiments show that the elongated mounds tend to take their main axis along the [1 1 0] direction. In their experiments, the ratio between the length and the width of the mounds is about 4. In our case for InAs growth, the mounds exhibit an isotropic feature. This means that the Schwoebel barrier, sticking probabilities, and adatom diffusion on InAs(0 0 1) surface are isotropic. On the other hand, some of them are anisotropic on a GaAs(0 0 1) surface.

In order to further confirm the above argument, we have also studied the surface morphology of InAs films grown on miscut substrates at different substrate temperatures. Fig. 3 shows the morphology of InAs films on 3° off GaSb buffered GaAs substrates. The films grown at temperatures of 430 and 520°C exhibit the similar morphology with high density of monolayer step edges. The film growth takes the step flow growth mode. The tilting angle of the miscut substrates is larger than the slope of the mounds which have been observed on the films grown on singular substrates. In other words, the average terrace width is smaller than the distance of the adatoms traveling before the nucleate islands. Therefore, the films growth favorite step flow growth mode.

The influence of the nucleation temperature on InAs growing on GaSb has also been studied. Fig. 4 shows a typical AFM image of the surface of
Fig. 3. AFM images of InAs films grown on miscut substrates: (a) grown at 510°C and (b) grown at 430°C.

Fig. 4. AFM image of an InAs surface grown at 460°C with a nucleation temperature of 460°C.

an InAs film with a thickness of 500 nm grown at 460°C. The InAs nucleation temperature is also at 460°C. The surfaces are relatively rough and with a relatively high density of deep pits (more than 200 nm in depth) surrounded by terraces (about 40 nm high). This phenomenon is consistent with that observed in the case of unstable growth. These facts can be explained on the basis of (1) insufficient and anisotropic In diffusions due to the Schwoebel barrier at the step edges, and (2) step pinning by surface defects, such as highly strained GaAs clusters which we believe to be generated on the GaSb surface by exposing the surface to As$_2$ beam flux during the nucleation process of InAs layer.

In summary, the surface morphology of InAs films grown on GaSb buffered GaAs substrates has been studied. Unstable mode growth has been observed on films grown on singular substrates at relatively low substrate temperatures. Unlike what has been observed on MBE grown GaAs films, the mounds on an InAs singular surface which formed due to growth instability were found to be isotropic. This means that the adatom diffusion, sticking probabilities, and Schwoebel barriers on InAs(0 0 1) surface are isotropic. The higher nucleation temperature (above 460°C) will result in relatively high density pits which we believe are due to the surface defects, such as highly strained GaAs clusters generated on the GaSb surface by exposing the surface to As$_2$ beam flux during the nucleation process of InAs layer.
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References


Luminescence anomaly in band gap tailored
In$_{0.53}$(Ga$_x$Al$_{1-x}$)$_{0.47}$As quaternary alloy grown
by molecular beam epitaxy

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Abstract

In$_{0.53}$(Ga$_x$Al$_{1-x}$)$_{0.47}$As epilayers with band gap energy varying from 0.78 to 1.42 eV are grown lattice matched to InP substrate by MBE technique. The lattice mismatch and crystalline quality of the layers are assessed by the double crystal X-ray diffraction and the lattice vibrations due to the three binary compounds in the alloy are observed in the Raman spectra. Photoluminescence studies in the range 4-100 K showed an anomalous “inverted S” shaped variation in the PL energy versus temperature plot for epilayers with Al mole fraction less than 0.2. For mole fraction greater than 0.2, the characteristic dip in the energy values disappears and the normal band gap energy versus temperature relation is restored. Strong localization of excitons is associated with such a behavior in Al rich InGaAlAs epilayers.

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Keywords: Molecular beam epitaxy; Photoluminescence; Exciton; Quaternary alloy; Lattice matching

1. Introduction

The III–V group of semiconductors are of potential importance in high-speed electronics and optoelectronics applications. The “arsenide” based ternaries (Al$_{0.48}$Ga$_{0.52}$As and Ga$_{0.47}$In$_{0.53}$As) and quaternary (In$_{0.53}$(Ga$_{x}$Al$_{1-x}$)$_{0.47}$As) alloy materials are an attractive alternative to the phosphide based (InGaAsP) material system covering the wavelength range of 1.1–1.7 μm. The band gap of InGaAlAs alloy can be engineered between the emission wavelengths of the two bounding ternary alloys namely: In$_{0.53}$Ga$_{0.47}$As (0.76 eV, 1.63 μm) and In$_{0.52}$Al$_{0.48}$As (1.46 eV, 0.85 μm), and can be an alternative to the InGaAsP quaternary material. With the conventional solid source MBE technique, the growth of InGaAlAs alloy is relatively easier than InGaAsP due to the following reasons: (a) only one group V element (As) is incorporated, avoiding the problem of As/P ratio control, (b) composition of the layer is controlled by each constituent element’s flux intensity, as noted by Ohno.
et al. [1] and (c) near-unity sticking coefficients of the three group III elements facilitate reproducibility of composition. The growth of lattice matched InGaAlAs alloy on InP substrates by MBE technique was first reported by Barnard et al. [2] and subsequently, the material was improved in its optical and crystal properties by other workers [3–6]. Olego et al. [7] developed an empirical expression between the band gap and composition of the alloy, suggesting that the alloy is a pseudo-binary of the two ternary materials. Most of the previous works on the MBE growths, incorporated pulse techniques [8] or two In/two Al cells [9, 10]. In our present work we have used one cell for each element and grown state-of-the art epilayers of InGaAlAs, as indicated by photoluminescence (PL) and double crystal X-ray diffraction (DCXD) results. These well characterized epilayers are studied under a temperature variation of 4–300 K at varying excitation power levels in PL experiments.

2. Experimental procedure

The quaternary alloy epilayers of In$_{0.53}$Ga$_x$Al$_{1-x}$As$_{0.47}$As were grown in a solid source MBE system (RIBER 32P model), with source elements of purity greater than 5 N. The In Beam Equivalent Pressure (BEP) was kept constant at 3.3 × 10$^{-7}$ mbar and Ga/Al BEP ratio was varied such that the band gap energy of the layers could be tailored, whilst keeping them lattice matched to the InP substrate. All the epilayers are grown to a constant thickness of 1.5 μm and at a constant substrate temperature of 510°C. The layers were characterized for lattice mismatch and crystalline quality on a double crystal X-ray diffractometer and the Raman spectra were observed on a Renishaw micro Raman system fitted with a high sensitive Charge Coupled Device (CCD) detector. The PL studies were made in a low-temperature closed loop He cryostat with argon laser as the exciting source and with Photo-Multiplier Tube (PMT), lock-in amplifier comprising the measurement setup for detecting the PL signal.

3. Results

In$_{0.53}$Ga$_x$Al$_{1-x}$As$_{0.47}$As epilayers with Al composition varying from 0.05 to 0.45 were grown lattice matched to InP substrate under a normalized In flux of 0.46. The normalized In flux is defined as the ratio of In flux to the total group III fluxes. The lattice mismatch obtained for the independently grown InGaAlAs epilayers is within 0.03%, with FWHMs in the range 20–35 arsec. The band gap of the material was estimated from the RT photoluminescence (PL) peaks, as shown in Fig. 1, and were observed to vary from 0.78 eV to 1.42 eV with FWHMs in the range 30–100 meV. Peaks at 1.35 and 1.42 eV, however, are not shown in the figure. The FWHMs. Table 1 shows the various growth parameters used for forming the varying band gap energy InGaAlAs epilayers.

The results obtained by DCXD and PL characterization indicate high crystalline and optical quality of the In$_{0.53}$Ga$_x$Al$_{1-x}$As$_{0.47}$As epilayers. Raman spectra of the samples showed distinct phonon vibrational mode peaks of the three binary (InAs, GaAs and AlAs) compounds with the intensity ratios being equivalent to the mole fraction of the elements in the alloy. Fig. 2. shows the Raman spectra of various epilayers with a noticeable shift in the peak positions due to the variation in alloy composition.

![Fig. 1. Room temperature photoluminescence peaks of band gap engineered samples. The peak values vary from 0.78 to 1.26 eV.](image-url)
In the PL spectra, an interesting phenomenon of “inverted S” shaped behavior is observed in the variation of PL energy with temperature in the range 4–100 K. Initially as the temperature is raised from 4 to 30 K, a red shift is observed in the energy peaks. Between 30 and 80 K a blue shift occurs and beyond that the normal red shift takes over following the Varshni’s equation [11] for band-to-band transition. This anomalous temperature dependence was first observed by Kondow and Minagawa [12] in Ga$_{0.5}$In$_{0.5}$P/GaAs samples and the probable cause was related to the crystal defects in the epilayer. Olshoorn et al. [13] and Ferguson et al. [14] observed similar phenomena in lattice matched InAlAs material and called the anomaly as “inverted S” shaped dependence. However, in this case the observations were related to the localization of excitons.

For sample #543 with Al = 0.42 and Ga = 0.05, the 4K PL spectrum shows a strong peak due to the bound exciton transition, at 1.495 eV with a FWHM of 18 meV. Deconvolution of the single peak suggests that in addition to the bound exciton transition, impurity levels, free exciton and band-to-band transitions are also built in. With increase in excitation power the peak intensities rise with the peak position remaining at a constant value. However, at high power levels a small peak develops at the high-energy side of the spectrum due to free exciton or band-to-band transitions with the bound exciton intensity tending to saturate. In Fig. 3, the PL spectra of sample #543 at variable temperatures are shown. On varying the temperature from 4 to 40 K the peak energy position is observed to shift to lower energy values. At 40 K

<table>
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<tr>
<th>Sample</th>
<th>Ga</th>
<th>Al</th>
<th>In</th>
<th>$J_{Ga}/J_{Al}$</th>
<th>$E_e$ (eV)</th>
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</thead>
<tbody>
<tr>
<td>#543</td>
<td>1.3 x 10^-8</td>
<td>1.0 x 10^-7</td>
<td>4.3 x 10^-7</td>
<td>0.07</td>
<td>1.42</td>
</tr>
<tr>
<td>#364</td>
<td>4.0 x 10^-8</td>
<td>9.4 x 10^-8</td>
<td>4.3 x 10^-7</td>
<td>0.23</td>
<td>1.34</td>
</tr>
<tr>
<td>#537</td>
<td>5.2 x 10^-8</td>
<td>8.0 x 10^-8</td>
<td>4.3 x 10^-7</td>
<td>0.35</td>
<td>1.26</td>
</tr>
<tr>
<td>#538</td>
<td>1.2 x 10^-7</td>
<td>5.0 x 10^-8</td>
<td>4.3 x 10^-7</td>
<td>1.28</td>
<td>1.02</td>
</tr>
<tr>
<td>#539</td>
<td>1.3 x 10^-7</td>
<td>3.9 x 10^-8</td>
<td>4.3 x 10^-7</td>
<td>1.80</td>
<td>0.94</td>
</tr>
<tr>
<td>#540</td>
<td>1.6 x 10^-7</td>
<td>2.6 x 10^-8</td>
<td>4.3 x 10^-7</td>
<td>3.20</td>
<td>0.87</td>
</tr>
<tr>
<td>#542</td>
<td>1.4 x 10^-7</td>
<td>9.0 x 10^-9</td>
<td>3.3 x 10^-7</td>
<td>8.40</td>
<td>0.78</td>
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the bound exciton partially converts to free exciton and a sudden shift towards the higher energy is observed. During this temperature range the free and bound excitons co-exist and two distinct peaks can be visualized. Up to about 80 K the free exciton dominates over the bound exciton and finally the latter quenches off at about 100 K. Beyond this temperature up to 300 K the band-to-band transition takes over and the Varshni’s equation is obeyed.

The difference in energies at 4 K, between the observed values and that obtained by Varshni’s equation is about 28.3 meV. The binding energies calculated by the expressions given by Goetz et al. [15] can account for only 11 meV ($E_b = 4.2$ meV, $E_D = 6.8$ meV). The excess energy of 17.3 meV is thus associated with the localization of excitons to some unknown energy levels. Fig. 4 shows the superposition of PL spectrum and the PLE spectrum which is detected at the emission peak of bound exciton. The absorption threshold in the PLE spectrum is attributed to the band edge transition which corresponds to the high energy peak of the deconvoluted PL spectrum. In sample 538 (Al = 0.2, Ga = 0.26) the energy peak observed at 4 K is due to the free (carrier) exciton and does not show any dip in energy values with the variation of temperature up to 300 K. At low excitation power levels the bound and free excitation transitions co-exist, but at high power levels the peak observed is completely dominated by free exciton transition. In Fig. 5 the PL spectra of 538 at different excitation power levels are shown. Finally in sample 539 (Al = 0.17, Ga = 0.30), the peak observed at 4 K is due to the band-to-band transition and hence in the temperature variation up to 300 K, the trend follows the Varshni’s band gap versus temperature relation. Fig. 6 shows the combined plot of all the samples in the PL energy versus temperature. For high Al content layers, the tendency to localize exciton strongly to an energy level below the excitonic levels is high due to the highly reactive nature of Al. It is estimated that for Al < 0.2 the characteristic dip in energy totally disappears.
4. Conclusions

InGaAlAs epilayers are grown lattice matched to InP substrate with band gap tailored between 0.78 and 1.42 eV. For Al rich layers strong localization of excitons is proposed since the energy discrepancy at 4 K between the observed values and that predicted by the unknown excitonic energies is large. Due to the highly reactive nature of Al, certain energy levels are formed in Al rich InGaAlAs layers which tend to localize the excitons.

References

Full wafer optical characterisation of resonant tunnelling structures using photoluminescence excitation spectroscopy

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Abstract

We report on a novel optical technique for accurately characterising unprocessed GaAs/AlGaAs tunnelling structures having two or more thin barriers and thick, heavily doped, contact layers. The results from photoluminescence excitation spectroscopy measurements are presented in which the intensity of the photoluminescence from the heavily doped contact layers rather than photoluminescence from the quantum wells is monitored. This method gives essential information for growth and quality control of “as-grown” layers prior to device fabrication. The usefulness and precision of this technique is illustrated by the measurement of a 0.3% variation in layer thickness from the centre to the edge of an as-grown wafer, and a 1% wafer to wafer variation in layer thickness in a series of triple barrier resonant tunnelling structures that have been designed for use at THz frequencies.

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1. Introduction

The capability of molecular beam epitaxy (MBE) for high precision growth of thin film structures has enabled the fabrication of many types of complex tunnelling structures. The ease with which such structures can be exploited commercially will be improved when it is possible for the as-grown wafers to be accurately and non-destructively characterised, without expensive, time-consuming processing and electrical measurements. As-grown wafer characterisation should provide essential feedback as rapidly as possible in order to control the growth of subsequent wafers. In particular, the accurate determination of quantum well thicknesses is extremely important, as device designs seek to exploit the full capability of MBE growth \cite{1,2}. Low-temperature photoluminescence spectroscopy has been shown to be a powerful tool for studying various characteristics of resonant tunnelling devices \cite{3–6}, but the problem of measuring as-grown (no applied bias) wafer material remains problematic. In general at low temperatures, for as-grown structures which contain a small number

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of quantum wells and thin barriers, very little or no photoluminescence is observed from the quantum well regions. This is because photo-injected carriers rapidly escape from the quantum well before any significant recombination can occur. For example, the electron escape time for a 70 Å GaAs quantum well confined by 50 Å Al$_{0.3}$Ga$_{0.7}$As barriers is of the order of 1 ps, whereas the exciton recombination time is of the order of 1 ns [7].

This paper presents results of a novel variant of photoluminescence excitation spectroscopy for characterising as-grown wafers that are to be processed into triple barrier tunnelling devices. The technique involves monitoring the luminescence intensity from the contact layers of the wafer instead of from the well regions. Strong well-resolved exciton transitions were observed from the quantum wells. These spectra were used to determine quantum well widths prior to processing and to monitor the uniformity of quantum well thickness across 50 mm diameter wafers.

2. Experimental procedure

The samples studied came from a set of closely related triple-barrier resonant tunnelling structures that have been designed for use as ultra high speed devices (THz) which demand very high precision growth for successful operation [2]. The electron energy subband spacing in these devices, for THz operation, must be the order of only 4 meV. Therefore, sharply defined energy states with minimal broadening are critical. The structures were grown without interruption at 0.85 µm/h on 50 mm semi-insulating GaAs substrates in a VG Semicon V90H system using conventional Ga and Al cells and a solid source cracker producing As$_2$. The layer sequence was as follows: (i) 1 µm n$^+$ ($N_{Si} = 3 \times 10^{18}$ cm$^{-3}$) GaAs lower contact layer (ii) 100 Å n($N_{Si} = 3 \times 10^{17}$ cm$^{-3}$) GaAs layer (iii) 100 Å undoped GaAs spacer (iv) 16 monolayer (ML) Al$_{0.3}$Ga$_{0.7}$As barrier (v) undoped wide GaAs quantum well (WW) (vi) 19 ML Al$_{0.3}$Ga$_{0.7}$As barrier (vii) undoped GaAs quantum well with either the same thickness as the WW or a narrower quantum well (NW) (viii) 16 ML Al$_{0.3}$Ga$_{0.7}$As barrier (ix) 200 Å GaAs spacer layer (x) $5 \times 10^{12}$ cm$^{-2}$ Si $d$-doped GaAs (xi) 0.5 µm n$^+$ ($N_{Si} = 7 \times 10^{18}$ cm$^{-3}$) upper contact layer. For the purposes of this discussion, the structures grown with both quantum wells of the same nominal thickness are referred to as symmetric structures, while those with quantum wells of different thicknesses are referred to as anti-symmetric structures. The growth temperature was 580°C for layers (i)–(ix) and was reduced to 540°C for layers (x) and (xi). A schematic of the band alignment for these structures is shown in Fig. 1, which includes the $n = 1$ electron, heavy hole and light hole confined states, and an indication of the heavily doped contact regions.

For the optical measurements the as-grown wafers were mounted on the cold finger of a variable temperature closed cycle helium cryostat. Light from a chopped Ar$^+$ pumped Ti:Sapphire laser was focused onto the wafer (spot size ~100 µm diameter) with typical power densities of 50 W/cm$^2$. The power density was small enough that no sample heating effects were observed, nor was there any change in the current through 80 µm square mesa devices with 50 µm square windows. The resulting luminescence was dispersed by a 0.75 m double grating spectrometer (SPEX 1404) and
detected by a thermoelectrically cooled GaAs photomultiplier. The signal was then processed by a conventional lock-in amplifier.

3. Results and discussion

In Fig. 2a is shown a photoluminescence spectrum from an as-grown wafer containing a symmetric well triple barrier structure with both quantum wells having a nominal thickness of 23 ML. We would expect the WW quantum well emission to occur at an energy around 1.580 eV. As can be seen from the spectrum we observe no emission from the quantum well due to the fast carrier escape discussed earlier but we observe strong emission bands at 1.530 and 1.490 eV. The broad emission at 1.530 eV is identified as being due to band-to-band recombination from the heavily doped n-type contact layers, with a clearly defined Fermi-energy labelled $E_f$. The emission process at 1.495 eV is identified as band-to-acceptor recombination. When excitation spectroscopy measurements are made on this sample over an energy range 1.550–1.660 eV, while detecting the band-to-band recombination from the contact layers, we observe the spectrum as shown in Fig. 2(b). The form of this spectra is not sensitive to the detection energy across the whole broad band–band luminescence feature. The two groups of lines around 1.578 and 1.600 eV are identified as involving $n = 1$ electron to $n = 1$ heavy hole transitions and $n = 1$ electron to $n = 1$ light hole transitions, respectively. The mechanism for the observation of these signals is that photogenerated electron/hole pairs and excitons in the quantum wells escape rapidly to the doped contact layers, resulting in an increase of the luminescence from the heavily doped contact region. The precise nature of the splitting of the two groups in this symmetric well sample is not yet clear. The splitting could be associated with symmetric and antisymmetric states of the $n = 1$ electron state or could be due to the two quantum wells having an effective well thickness different by 2.3 Å. The precise identification of these features is not as important as the main issue of this paper, which is that although it is not possible to detect photoluminescence from the quantum wells in an as-grown wafer, it is possible to measure quantum well optical transitions by using excitation spectroscopy and detecting luminescence from the contact layers. The usefulness of this characterisation technique is shown in Fig. 3. In this figure are shown a series of eleven PLE spectra taken in a uniformly spaced scan at radii from 5 to 22 mm on a 50 mm diameter wafer incorporating an asymmetric well triple barrier tunnelling structure. The peaks correspond to the $n = 1$ electron-heavy hole exciton transitions in each of the wells whose widths are calculated to be 69.1 Å and 66.2 Å. Further peaks are observed, attributed to $n = 1$ electron-light hole exciton transitions, but are not shown in Fig. 3. The extreme variation in the position of the lowest energy peak over the scan is 0.3 meV (average line width of 1.05 meV). This corresponds to a variation in the effective width of the quantum wells of only 0.3%. These data demonstrate the high uniformity afforded by the V90H MBE system, originally designed for multiple 2" or single 4" wafers, in growing uniform structures over large areas. The uniformity demonstrated (0.3%) is to an extent well below what is required by device processors (typically ± 1%). For a series of eight TBRTS, where the WW quantum well was maintained at a nominal 23 ML and the NW quantum well ranged from 23 ML down to 16 ML the wafer-to-wafer variation in WW width was only 1%.
To emphasise the significance of this data it demonstrates:

(1) The uniformity of the quantum well thickness across the majority of the slice.

(2) The usefulness of the technique in that accurate characterisation of multi barrier tunnelling structures can be applied to whole wafers without any sample processing.

This technique has been successfully used for a variety of triple barrier resonant tunnelling structures with differing well widths ranging from 24 ML (68 Å) down to as low as 9 ML (25 Å), and differing barrier aluminium concentrations of 33%, 29.5% and 26%. In the case of these highly asymmetrical narrow well structures, \( n = 1 \) electron-heavy hole transitions are observed at energies up to 1.725 eV.

4. Conclusion

We have demonstrated a high-resolution method for characterising multiple barrier resonant tunnelling structure wafer material. By monitoring the photoluminescence intensity from the heavily doped contact region we have used PLE on wafers to determine the quantum well transition energies. The important point to emphasise is that without any sample processing it is not possible to detect any photoluminescence from the quantum wells of such structures. The sensitivity of this technique has been demonstrated by conducting a detailed characterisation of a series of unprocessed wafers which include high-quality triple barrier resonant tunnelling devices. The measurements have been used to give valuable information about wafer uniformity and quality together with an accurate measurement of the lowest lying electron-heavy hole and electron-light hole transition energies.

References


Scanning tunneling spectroscopy and first-principles investigation on GaAs(0 0 1)(2 × 6)-S surface formed by molecular beam epitaxy

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Abstract

The surface reconstruction of, in situ prepared, S-terminated GaAs (0 0 1) is studied by scanning tunneling microscopy and a first-principles pseudopotential method. According to an electron counting model, in order to form this (2 × 6)-S on GaAs (0 0 1) surface, one electron needs to be transferred from each S dimer to the Ga dangling bonds on the missing dimer region. As a result of the first-principles investigation on the S-S bond length of one S dimer, the separations are calculated as 0.2370 nm and 0.3897 nm, with and without one electron transfer, respectively. Experimental result agreed well with the former condition. This indicates that there may be a long-range electron transfer on this (2 × 6) structure. Moreover, the calculated electronic structure of this S dimer with one electron transfer shows a flat unoccupied band which is mainly caused by the antibonding band of the S-S bond. This result also agreed well with conductivity spectrum of this (2 × 6) surface by scanning tunneling spectroscopy, which shows a peak on the conduction band side.

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1. Introduction

The sulfur-terminated (S-terminated) GaAs (0 0 1) surface is an important factor in droplet epitaxy which is a useful technique for fabricating quantum dot structures [1]. In order to establish this technique and to understand the droplet growth mechanism, the details of the S-terminated surface reconstruction needs to be investigated.

The (2 × 1) surface reconstruction is generally understood to be that of the S-terminated GaAs (001) surface as measured by low-energy electron diffraction (LEED) and reflection high-energy electron diffraction (RHEED) studies [2, 3]. However, this 2 × 1 reconstruction is a rather diffused streaky pattern with diffused transmission spots, which is due to the surface roughness caused by the etching

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effect of a sulfide solution such as Na$_2$S [4], (NH$_4$)$_2$S [5, 6], and (NH$_4$)$_2$S$_x$ [6]. Moreover, there are few reports of scanning tunneling microscopy (STM) images of such a surface [7, 8]. Therefore, under these conditions, it is unwise to say that the (2 x 1) surface reconstruction is dominant on the S-terminated GaAs(0 0 1) surface.

Recently, we have developed a novel fabrication method for the S-terminated GaAs(0 0 1) surface, with exposure to S vapor in a molecular beam epitaxy (MBE) system [9, 10]. Using this method, the S-terminated GaAs surface is formed without exposure to air or etching the surface. In this case, a diffused (2 x 1) structure is not observed in RHEED, only the (2 x 6) forms are observed after exposure to S vapor and annealing [9, 10]. Moreover, we have demonstrated that a (2 x 6) is dominant on the S-terminated GaAs(0 0 1) surface by STM [11, 12] and high-temperature STM observations [13]. Even on the (NH$_4$)$_2$S$_x$ treated [12] and H$_2$S irradiated [14] GaAs(0 0 1) surfaces, (2 x 6) structure was observed. Despite all these experimental works, the passivation mechanism of the GaAs surface is still not clearly understood. Knowledge of the actual atomic configurations at the S-terminated surface and the corresponding electronic structures remains insufficient.

Therefore, in this paper, we study this (2 x 6) structure with a scanning tunneling spectroscopy (STS) technique and a first-principles pseudopotential method. The structural and electronic properties of GaAs(0 0 1) surfaces covered with an adsorbed monolayer of S atoms are discussed with the experimental and theoretical results.

2. Termination process

In this study, the two vacuum systems, MBE and STM, are separated by air. The MBE system is a conventional ANELVA-620 with a sample introduction chamber, in which a valved knudsen cell charged with elemental S is installed. The background pressure of the sample introduction chamber is 6 x 10$^{-6}$ Pa.

The Si-doped GaAs(001) 1° off $\langle 1 1 1 \rangle$A ($n = 2 \times 10^{18}$ cm$^{-3}$) sample is prepared by standard solvent cleaning and etching procedures, and then loaded into the MBE chamber. An undoped GaAs buffer layer of 0.2 μm was grown at 590°C, resulting in smooth one-monolayer-stepped terraces. Then, the sample was processed with multi-step S termination method [15]. This method reduces the ununiformity of surface reconstructions, realizing the S-termination on the Ga-stabilized surface with a coverage of 1 [13]. The process of this method is as follows. First, the substrate temperature has decreased to room temperature with c (4 x 4) As-adsorbed surface shown in the Fig. 1a. Next, the sample was immediately transferred into the MBE sample introduction chamber and exposed to the S vapor, which generates a chamber pressure of 4 x 10$^{-4}$ Pa for 3 min (Fig. 1b). Then,

![Fig. 1. Multi-step S termination process.](image-url)
the sample was transferred into the MBE growth chamber and annealed at 370°C for 30 min without the As molecular-beam flux (Fig. 1c). Because of the difference of bond energies of S–Ga and As–Ga, at this temperature, only the As atoms remaining upon the Ga layer were desorbed (Fig. 1d). Then, again, repeating the same process, the sample was cooled down, exposed to the S vapor (Fig. 1e), and annealed at 370°C for 30 min (Fig. 1f). This second step of the S-termination filled the escaped holes of As atoms and realized the highly uniform (2 × 6) surface reconstruction (Fig. 1g). After the sample was cooled down to about 10°C and again transferred into the MBE sample introduction chamber, it was exposed to the S vapor, which generates a chamber pressure of 8 × 10⁻⁷ Pa, at room temperature for 5 min, forming the thick S protective layer [13]. The sample was then transferred from the MBE system through air to the STM apparatus.

The STM (JEOL-JSTM4500VT) is a multichamber UHV system. There are a sample introduction chamber, a treatment chamber with RHEED, and a STM main chamber. The background pressure of both the treatment chamber and the STM main chamber is 1 × 10⁻⁸ Pa. In the STM main chamber, the sample was directly heated by an electric current and the temperature was measured with an optical pyrometer having a detection wavelength of 2 μm. The removal process of the protective layer has been described elsewhere [11, 13]. The STM measurements were performed in the constant current mode and the STM images were taken at a sample bias voltage of −3 V relative to the tip and a constant tunneling current of 0.2 nA. The tips are electrochemically etched tungsten wires.

3. Calculation methods

We performed total-energy calculations by using the local density functional (LDF) approach. The total energy function is minimized with respect to both the plane-wave coefficients of the occupied orbitals and the ionic degrees of freedom by using the conjugate-gradient technique [16]. The pseudo wavefunctions are expanded in terms of a plane-wave basis set corresponding to a kinetic-energy cutoff of 7.29 Ry. Four special k points are employed to sample the primitive surface Brillouin zone. The surface is simulated by a slab geometry shown in Fig. 2. This model uses supercells containing one sulfur layer, two layers of GaAs, and one layer of hypothetical hydrogen used for the surface termination [17]. It is found that the energy cutoff of 7.29 Ry and the supercell containing five atomic layers are sufficient to achieve the convergence of energy differences within 0.1 eV. Here, we only focus on a (2 × 1) unit because a (2 × 6) unit is too large for the calculation. Then, with and without one electron transfer from this (2 × 1) unit, the (2 × 6) and (2 × 1) cases are discussed, respectively.

4. Results and discussion

At first, in order to evaluate the slab geometry for calculation, we compared an S–S bond length from STM results with that from the theoretical calculation. From the STM observation, it was found
that the separation between the S atoms of the S dimer was about 0.23 nm as shown in Fig. 3. According to the electron counting model, in order to form this \( (2 \times 6) \)-S on GaAs(0 0 1) surface, one electron needs to be transferred from each S dimer to the Ga dangling bonds on the missing dimer region [11]. Thus, we investigate the S–S bond length with and without one electron transfer using the first-principles pseudopotential method. As a result, the separations are calculated as 0.2370 and 0.3897 nm, with and without one electron transfer, in the case of \((2 \times 6)\) and \((2 \times 1)\), respectively. Experimental results agreed well with the former condition. This indicates that this slab geometry is only a part of the \((2 \times 6)\) structure but able to describe the whole structure with fulfillment of the electron counting (local-charge neutrality) heuristics.

By the STS measurement on this \((2 \times 6)\) structure, a conductivity spectrum as shown in Fig. 4 was obtained. The spectrum is an average of several spectra taken from a well ordered region of a \((2 \times 6)\) surface. No spatial variation was observed in the spectra. The voltage width of \(dI/dV = 0\) indicates the energy band gap of GaAs (1.45 V) which is reasonable at room temperature, showing that the S layer does not significantly affect the band gap. There is a peak on the conduction band side. The GaAs(0 0 1) \((2 \times 1)\)-Se surface does not show such a peak in STS [18]. In order to understand this spectrum, the electronic structure of this S dimer with and without one electron transfer was calculated using the same slab geometry as in Fig. 2. The effective band gap obtained by this method of calculation is shown in Fig. 5. The discrepancy between this effective band gap and the experimental value (1.45 V) is about 47%. In the case of the one electron transfer, \((2 \times 6)\) case, it exhibits an unoccupied flat band which is mainly caused by the antibonding band of the S–S bond shown in Fig. 5a. This result agreed well with the sharpness and energetic position of the peak in Fig. 4. Therefore, this peak should be related to the S dimers.

Moreover, in the case of no electron transfer, \((2 \times 1)\) case, there are several surface state bands associated with the surface S atoms in the energy gap region (Fig. 5b). But, in the case of the one electron transfer, there are no surface state bands in that region (Fig. 5a). This indicates that the one electron transfer can noticeably reduce the surface state density in the gap region.

Even though we need to confirm that these electrons from S dimers should transfer to the missing dimer region by a calculation of the complete \((2 \times 6)\) structure as well as the detailed STS mapping of the \((2 \times 6)\) structure. It is suggested by these theoretical and the experimental results that there are long-range electron transfers on this \((2 \times 6)\) structure.
These transfers may change the surface charge density on an atomic scale. We believe that these missing dimer regions will become the first growth sites when Ga atoms are deposited.

5. Conclusions

We have studied the \((2 \times 6)\) structure with the STS technique and the first-principles pseudopotential method. The structural and electronic properties of GaAs\((0 0 1)\) surfaces covered with an adsorbed monolayer of S atoms were discussed with the experimental and theoretical results. By comparing the experimental results with the theoretical calculation, it was found that there was agreement between them, indicating that there are long-range electron transfers on this \((2 \times 6)\) structure and these transfers may change the surface charge density in the atomic scale. Therefore, when Ga atoms are deposited on this \((2 \times 6)\) structure, this changed surface charge density may determine the first growth sites of the Ga atoms.

References


Does scanning tunnelling microscopy provide a realistic picture of the step array of vicinal GaAs(0 0 1) surfaces grown at high temperature?

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Abstract

A 90° double reflection high-energy electron diffraction (RHEED) set-up was used to perform a two-dimensional (2D) study of the morphology of vicinal GaAs(0 0 1) surfaces during molecular beam epitaxy, in particular at high temperatures where the quenching for scanning tunnelling microscopy studies (STM) is a problem. In the transition range between 2D and step flow growth the terrace width fluctuation is found to increase on the B-surface (2° misorientation towards (1 1 1)As), whereas the A-surface (2° misorientation towards (1 1 1)Ga) becomes more uniformly stepped. This is discussed in terms of different barrier heights for downward diffusion of Ga adatoms over A- and B-type steps. For the A-surface different apparent transition temperatures between 2D and step flow growth are found in the (1 1 0) and (1 1 0) azimuths. In accordance with STM results this is explained by an increasing elongation of islands in (1 1 0) direction with increased substrate temperature. Above 580°C kinetically smoothed A and B surfaces transform after growth interruption rapidly to less ordered equilibrium surfaces. On the A-surface in addition the adatom concentration changes rapidly. Both processes occur in less than 1 s which creates problems to properly quench such surfaces for STM studies.

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Keywords: Kinetics of growth; Morphological stability of GaAs(0 0 1); Molecular beam epitaxy

1. Introduction

Step arrays on vicinal GaAs(0 0 1) surfaces have attracted considerable interest in molecular beam epitaxy (MBE) growth studies. The growth on regularly stepped surfaces leads to an improved surface/interface morphology which is reflected in improved device quality [1]. For novel concepts, such as quantum wire growth [2], the requirements on the regularity of the step array are much more stringent and a deeper understanding of the surface processes is needed. The recording of RHEED...
intensity oscillations as a widely used technique to control the transition from 2D to step flow growth [3] and the measurements of intensity profiles for an assessment of terrace width fluctuation and step edge straightness [4] in principle provide 1D information, whereas the morphology of the real surface [5–7] and kinetic processes on it have 2D character. Real space imaging by STM showed quite different step structures for the vicinal surface misoriented towards [1 1 0] (A-surface) and towards [1 1 0] (B-surface), respectively. Ga terminated (A) steps on the A-surface are smoother compared with As terminated (B) steps on the B-surface, but the terrace width is less homogeneous [5]. From RHEED [8] and in situ scanning electron microscopy studies [9] it became evident that the transition between 2D and step flow growth is not abrupt. STM investigations after quenching showed islands elongated in [1 1 0] direction [10–14]. When high substrate temperatures are used, which are of interest for the growth of laterally ordered structures, quenching the actual structure of the growth surface becomes a real problem.

In this work, we use a 90° double RHEED set-up for a 2D real-time study of the GaAs(0 0 1) growth surface. The intensity behaviour during growth and after growth interruptions is discussed in terms of competition between kinetic processes and thermodynamic equilibrium. In particular, we consider an anisotropic barrier for downward diffusion of adatoms over A and B steps. Finally it is pointed out that in situ RHEED and room temperature STM results can differ considerably due to fast changes from local to global thermodynamic equilibrium.

2. Experiment

The experimental set-up for the double RHEED experiments (Fig. 1) comprises two RHEED guns installed in such a way that the two 20 keV electron beams intersect on the sample at an angle of 90°. The incidence angle was 0.4°. The intensity of the specular beam was simultaneously monitored in both (110) azimuths parallel and perpendicular to the steps with two CCD cameras connected to two independent image processing systems. Small substrate pieces of 10 × 10 mm² from epiready GaAs(0 0 1) wafers with 2° misorientation towards (1 1 1)Ga or (1 1 1)As were used. The substrate temperature measured with a thermocouple pressed to the backside of the substrate was calibrated to the oxide desorption temperature (580°C). The accuracy of temperature differences for experiments on the same sample is supposed to be better than 5°C, whereas the accuracy of the absolute temperature measurement for different samples has an error margin of up to 20°C. In order to avoid systematic structural changes, measurements at higher and lower substrate temperatures were carried out at random. The growth rate was 0.7 ML s⁻¹ and the AsₓGa beam equivalent pressure ratio was 15 in all experiments to maintain a (2 × 4) reconstruction during growth.

3. Results and discussion

Fig. 2 shows recordings of the specular beam RHEED intensity for GaAs growth on A- and
Fig. 2. Intensity of the specular RHEED beam during and after growth on vicinal GaAs(001)A surface (a, b) and on vicinal GaAs(001)B surface (c, d) for different substrate temperatures, recorded simultaneously in the azimuth parallel (a, c) and perpendicular to the misorientation steps (b, d). The dotted line in the second trace of (a) defines the average intensity.

B-surfaces. Their comparison allows us to check whether the curves are directly correlated with changes in distribution and shape of atomic steps. At high substrate temperatures, where the growth proceeds entirely in the step flow mode and therefore no intensity oscillations are visible, a striking difference is observed between the intensity behavior in the two azimuths. For the A-surface the intensity decreases after growth initiation in the [110] azimuth (Fig. 2a) whereas it increases in the
[1 1 0] azimuth (Fig. 2b). The opposite behaviour is found for the B-surface, where the intensity decreases in [1 1 0] azimuth (Fig. 2c) and increases in [1 1 0] azimuth (Fig. 2d). Both surfaces show, however, a similar intensity behaviour for the direction parallel to the steps (compare Fig. 2a and Fig. 2c) and for the direction perpendicular to the steps (compare Fig. 2b and Fig. 2d). Consequently, the intensity behaviour is dominated by the evolution of the step structure. If it would be governed by changes in surface reconstruction one should expect a similar intensity behaviour in the same azimuths of the different vicinal surfaces. This is not the case. Therefore, it is justified to assume in the later discussion that intensity recordings in the azimuths parallel and perpendicular to the nominal step direction are sensitive to step edge roughness and terrace width fluctuation, respectively. This is based on general belief that the intensity is mainly affected by the step component perpendicular to the direction of the incident beam [12, 15]. In addition, changes in adatom density will affect the intensity in particular in the azimuth parallel to the step edges. The latter explains that under the regime of entire step flow the intensity decay and rise after growth initiation and interruption, respectively, in the azimuth parallel to the step edges is different for B- and B-surfaces (Fig. 2a and Fig. 2c). This is attributed to the build-up of a higher adatom density on the growing A-surface compared to the B-surface due to a higher incorporation rate into B-type steps and/or preferred migration along [1 1 0].

At substrate temperatures of ~580°C for the B-surface and ~600°C for the A-surface, the specular RHEED beam intensity reaches constant values for the non-growing and growing surfaces within less than 1 s. This proves that the kinetic processes are rapid so that the system can approach a local thermodynamic equilibrium on the length scale probed by RHEED.

Below these substrate temperatures the competition between kinetic processes and the system's desire to reach the thermodynamic equilibrium is shifted in favour of the kinetics. A striking difference in the intensity behaviour during growth is observed between A- and B-surfaces. For the A-surface, the average intensity as defined in Fig. 2a, is approximately constant in the azimuth parallel to the steps. In the perpendicular azimuth (Fig. 2b), however, it increases for temperatures > 540°C. In the case of the B-surface the average intensity decreases continuously during growth in both azimuths (Fig. 2c and Fig. 2d). This opposite behaviour of A- and B-surfaces in the azimuth perpendicular to the steps can be understood within models considering the barrier for downward diffusion over steps [16, 17]. If the Schwoebel barrier is large or infinite for downward flow, the preferential adatom attachment to the upper step edge will lead to a terrace width equalisation because the step velocity is directly proportional to the size of the feeding range. In the opposite case atoms arriving on the terrace can approach the step with a higher probability from the upper terrace and hence the terrace width fluctuation will increase. Supposing that the ordered step array leads to a higher specular beam RHEED intensity, the intensity increase observed during growth on the A-surface suggests terrace width equalisation whereas the intensity decrease in case of the B-surface is explained by increased terrace width fluctuation. Therefore, we conclude that the diffusion over the steps is anisotropic, with a higher barrier for A-type steps than for B-type steps.

For the A-surface the transition temperature from 2D to step flow growth is 605°C in the [1 1 0] azimuth parallel to the steps (Fig. 2a) and 585°C in the [1 1 0] azimuth perpendicular to the steps (Fig. 2b), as indicated by the disappearance of the oscillations. This difference between the transition temperatures in the two azimuths can be explained in terms of a changing shape of the growing island when the transition temperature is approached. Following the assumptions that the specular beam RHEED intensity and the step length per area (which for an islanded surface is a better measure than step edge roughness and terrace width fluctuation) are correlated and that the step length per area normal to the electron beam is mainly responsible for the oscillation in this azimuth [12], we must conclude that the step length per area reaches its steady-state value in the [1 1 0] azimuth at a lower substrate temperature than in the [1 1 0] azimuth. In particular at higher temperatures the preferred Ga atom migration along [1 1 0] [18]
(which is disputed by some authors [19]) and/or the higher sticking probability on B-type steps than on A-type steps [19, 20] leads to anisotropic island growth (for an illustration of the surface processes see Fig. 3). At lower substrate temperature the elongated islands preserve their axis ratio more or less during growth leading to periodic changes in step length per area in both the [1 1 0] and [1 1 0] azimuth. With increasing substrate temperature the island elongation is not only influenced by the factors already considered. The tendency to anisotropic growth will even be increased by the downward diffusion over B steps. During island growth and coalescence the length of the A-steps approaches a steady-state value whereas the B-steps are still changing their length per area periodically. Accordingly, intensity oscillations occur in the [1 1 0] azimuth but can no longer be observed in the [1 1 0] azimuth.

For the B-surface (Fig. 2c and Fig. 2d) the oscillations are less pronounced. A strong meandering of the B-steps during growth is evident from the continuously decreasing intensity in the [1 1 0] azimuth. The simultaneous decrease of the specular beam intensity in the [1 1 0] azimuth indicates that the step meandering is coupled with an increase in terrace width fluctuation. Assuming at least two distinguishable intensity maxima to be present, the temperature for transition from 2D to step flow growth is estimated to 560 °C for both the [1 1 0] azimuth and the [1 1 0] azimuth. It is obvious that due to step meandering the B-surface is more isotropic regarding the surface roughness than the A-surface. The surface evolution below the transition to step flow is well explained by the preferential adatom migration along the [1 1 0] direction, the higher sticking probability of Ga atoms at B-type steps and a high probability of an adatom crossing downward B-type steps (Fig. 3b).

Above the transition to step flow (≥ 560 °C), the RHEED intensity behaviour in the [1 1 0] azimuth indicates that after growth initiation the step edge roughness increases but reaches a steady-state value within 1 s (Fig. 2c). Surprisingly, the rapid intensity increase after growth initiation to a constant value in the [1 1 0] azimuth (Fig. 2d) suggests “kinetic smoothing” due to terrace width equalisation. Therefore, in this temperature range another mechanism than adatom crossing downward B-steps must be effective. The fractional-order beams disappear during growth, indicating that the ordering (unbunching) of steps and disordering on terraces are linked and that the energetics of the resulting metastable surface is significantly changed. It is the same mechanism that becomes operative on the vicinal Si(1 1 1) surface in a narrow temperature range beneath the equilibrium
faceting temperature [21]. In this case the kinetic smoothing of the equilibrium faceted surface is accompanied by the disappearance of the \((7 \times 7)\) reconstruction. In contrast to the metastable smooth vicinal Si(1 1 1) surface which is remarkably robust, the kinetically smoothed vicinal GaAs(0 0 1)B surface transforms to the equilibrium bunched surface within less than 1 s.

RHEED and STM are complementary methods that differ considerably in sample handling. It is, therefore, challenging to relate results of the present experiments with the double RHEED geometry to results from STM investigations. For growth on the A-surface it has been concluded from the intensity oscillation behaviour that the island elongation along [\(\bar{1} 1 0\)] increases in the 2D growth regime with substrate temperature not only due to anisotropy in adatom migration and attachment on A-type and B-type steps but in particular due to downward diffusion over B-type steps. The transition in growth mode from 2D to step flow is completed at 605°C, i.e. in a temperature range for which we expect near equilibrium conditions. To verify this, Fig. 4 summarises data on the axis ratio of islands derived from STM (or atomic force microscopy [7]) studies by several authors for the nominal singular (0 0 1) surface (near A-type steps). If not given by the authors, we have estimated the axis ratios from the smaller islands. For substrate temperatures \(\leqslant 590°C\) the data show a linear increase of the axis ratio with temperature. At a substrate temperature of 590°C the island axis ratio increases from \(\sim 4\) for rapidly cooled samples [11] up to \(\sim 10\) for annealed samples (indicated by arrow) [11, 13]. A shape anisotropy of \(\sim 10\) has been found also for sample growth and 15 min annealing at 610°C suggesting that this represents the equilibrium anisotropy [14]. This agrees well with our conclusion that the island shape anisotropy increases with temperature until a constant value is reached at near-equilibrium conditions. In agreement with our real-time study on vicinal surfaces, the STM analysis of the morphology on the singular surface at near-equilibrium conditions [14] indicated a barrier for transport at least over A steps and a possible anisotropy in the barrier.

The present study shows that at near-equilibrium conditions dramatic changes between the structure of the dynamic and static surface occur in less than 1 s. A change between kinetic smoothing and equilibrium bunching is observed on the A- as well as on the B-surface. On the A-surface we find in addition an abrupt increase and decrease in adatom density after growth initiation and interruption, respectively, indicated by a sudden RHEED intensity change in the [\(\bar{1} 1 0\)] azimuth (Fig. 2a). This on a first glance unexpected high adatom concentration and its rapid reordering with changed surface conditions is evident also from recent STM work [22]. For a 0.15° misoriented surface quenched within 2.0 s from 600°C (faster than in usual MBE/STM work) it has been found that a concentration equivalent to a coverage as high as 0.18 ML was frozen into small islands.

4. Conclusions

Using a 90° double RHEED set-up a 2D analysis of the evolution of the morphology on vicinal GaAs(0 0 1) surfaces during growth and after growth interruption has been performed. On the A-surface the disappearance temperature of the RHEED intensity oscillations is about 20°C higher in [\(\bar{1} 1 0\)] azimuth than in [1 1 0] azimuth due to the formation of elongated islands which change
their axis ratio in the transition range between 2D and step flow growth. During growth under kinetically controlled conditions the A-surface becomes more uniformly stepped whereas on the B-surface the terrace width fluctuation increases. This is explained by an asymmetry in the step attachment with a downward diffusion of Ga adatoms over B-type steps. A kinetic smoothing observed at near equilibrium conditions above ~580°C also on the B-surface is linked with a disordering of the reconstruction and therefore not the result of the step attachment asymmetry but rather a consequence of the energetics. This unbunched surface is metastable and recovers after growth interruption within less than 1 s to the equilibrium bunched surface, which is faster than realizable freezing times in STM studies. On the A-surface in addition a rapid change in the adatom concentration occurs after growth interruption. Our results demonstrate that the 90° double RHEED technique can be advantageous used to study metastable surfaces that are difficult to quench in from high growth temperature for STM studies at room temperature. But even in this case the imaging by STM is valuable for the detailed interpretation of the RHEED results, provided differences due to a transition from local to global thermodynamic equilibrium are taken into account.

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References

Substrate temperature change in III–V molecular beam epitaxy

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Abstract

The surface temperature ($T_s$), as inferred by desorption mass spectrometric determination of the Ga desorption rate, was followed in time after instantaneous changes were made in the setpoint temperature ($T_{set}$) during molecular beam epitaxy (MBE) growth of GaAs. Comparison between the time-evolution of $T_s$ and that of the non-contact thermocouple-indicated temperature ($T_{tc}$) shows that large $T_s$ overshoot is likely to occur under typical MBE growth conditions. Such $T_s$ overshoot can lead to significant compositional grading at nominally abrupt heterojunctions. Details of how $T_s$ evolves in time are found to depend on substrate conductivity, temperature controller settings (proportional band, integral, derivative), and maximum heater current setpoint.

1. Introduction

The ability to control substrate temperature during molecular beam epitaxy (MBE) is critical for realizing high quality semiconductor heterostructures. The most common substrate temperature sensor is the non-contact thermocouple, which is inexpensive and simple to implement. However, its use is prone to large systematic errors, both under nominally constant temperature conditions and when implementing intentional temperature changes. Intentional temperature changes are often implemented at a heterointerface (e.g., growth of GaAs at 600°C followed by growth of AlGaAs at 700°C). In the present work the relationship between the actual surface temperature and the thermocouple-indicated temperature is determined when implementing a stepwise increase in the setpoint temperature. The goal is to examine how closely the thermocouple temperature follows that of the wafer surface; i.e., to determine time-lags and overshoot phenomena. Effects of substrate conductivity and substrate temperature-controller settings are investigated.

2. Experimental procedure

The experiments were performed on a Gen II equipped with an apertured line-of-sight mass spectrometer on the center port of the source flange. The commercial substrate temperature sensing
thermocouple floats in vacuum between the heater filaments and the substrate and is radiatively shielded from the heater filaments. Further details in the experimental arrangement can be found in Ref. [1].

The relationship between the thermocouple-indicated temperature ($T_{TC}$) and the GaAs surface temperature ($T_s$) was followed in time after instantaneous changes were made in the controller setpoint temperature ($T_{set}$). The GaAs surface temperature was inferred from the Ga desorption rate ($F_d(Ga)$), which was measured by desorption mass spectrometry (DMS). Fig. 1 shows how $F_d(Ga)$ and $T_{TC}$ respond to a stepwise increase in $T_{set}$ (dashed line) from 615 to 727°C at $t = 100$ s during growth of GaAs on a semi-insulating (SI) wafer. The Ga flux and As$_2$ beam equivalent pressure were 0.75 ML/s and $5 \times 10^{-5}$ torr, respectively. The behavior of $T_{TC}$ (right) relative to $T_{set}$ is characterized by an initial ~70 s lag, with little if any overshoot. In contrast, $F_d(Ga)$ (left) overshoots to a value which is about 50% of its eventual steady-state value at the new setpoint. The overshoot in $F_d(Ga)$ corresponds to an overshoot in $T_s$ of approximately 35°C [2] and the peak magnitude of $F_d(Ga)$ is equal to 75% of the magnitude of the incident Ga flux. Additionally, the maximum in $F_d(Ga)$ occurs roughly at a time at which $T_{TC}$ settles to the new $T_{set}$.

Similar data was taken for a variety of temperature setpoints, substrate conductivity, and temperature controller (PID and maximum current) settings.

3. Results and discussion

The effect of maximum heater current was examined by repeating the experiment for maximum heater current settings of 7 and 8 A; the results are shown in Fig. 2. (All other conditions were held constant.) The higher maximum current setting, which corresponds to the data displayed in Fig. 1, is seen to give rise to approximately a threefold larger overshoot in $F_d(Ga)$ and a significantly more rapid settling time than the smaller maximum current setting. It is concluded that large stepwise increases in heater current provide large and rapid overshoots in $T_s$. It is also clear that low maximum current settings are consistent with minimization of the overshoot magnitude but not necessarily the time to settle to the new setpoint. Note that the thermocouple response time is most sluggish for the lower current setting, as one would expect. Additionally, the maximum in $F_d(Ga)$ occurs roughly at a time at which $T_{TC}$ settles to the new $T_{set}$, regardless of the maximum current setting.

It is common practice for the MBE grower to alter or tune the proportional band, integral, and
derivative settings of the temperature controller. The data displayed in Fig. 3 is a comparison of two different integral times of 3 and 50 s, with all other conditions held constant. The lower integral setting leads to a larger magnitude of overshoot but gives rise to a similar rise time and eventual settling time for $T_{TC}$ to reach the new $T_{set}$. Also, the response of $F_d(Ga)$ for the lower integral setting shows significantly under-damped behavior, which also is observed in the thermocouple output, although not as pronounced.

The parameter space of temperature controller settings for experiments such as those under investigation is rather large. However, it is thought that careful examination of various combinations of proportional band, integral, derivative, and maximum heater settings would likely result in the determination of conditions giving rise to reasonably rapid response times with minimal $T_s$ overshoot. However, such an investigation would be very time-consuming and the information obtained would not be easily transferred to another set of growth conditions and temperature setpoint changes.

Since non-contact-mounted substrates are heated largely through radiative coupling, the substrate bandgap is an important issue in determining the fraction of heater radiation which is absorbed. One might expect to observe different responses between conductive and semi-insulating substrates since the effective bandgap of the former is smaller. In Fig. 4 we compare the responses of a SI substrate with that of a conductive $n^+$ substrate. The different stepwise changes in $T_{set}$ effect for the two different substrates, as shown by the dashed lines, were chosen to yield the same magnitude of $F_d(Ga)$ after full settling, in order to show the difference in setpoints required to yield the same steady-state surface temperature. Consistent with the effective bandgap argument are the following observations: (1) $F_d(Ga)$ rises more rapidly for the $n^+$ substrate, (2) the time it takes for $F_d(Ga)$ to peak is shorter; and (3) the setpoint required to reach the same settling value of $F_d(Ga)$ is lower for the $n^+$ substrate than for the SI substrate. One conclusion from these observations is that one must be careful when setting growth parameters for a given substrate conductivity if they were determined or optimized while growing on a substrate of different conductivity.

Thus, heater controller settings and substrate conductivity all affect the relationship between the surface temperature and the thermocouple output when effecting stepwise temperature changes during MBE growth. These observations are the result of the relatively loose coupling between $T_s$ and $T_{TC}$. What is needed is a way of controlling $T_s$ directly. One way of achieving that is by decoupling the

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Fig. 3. Variation of thermocouple output temperature (right axis) and Ga desorption rate (left axis) as a function of time as a stepwise increase in the setpoint temperature (dashed line) is implemented. Results are shown for two different cases corresponding to temperature controller integral settings of 3 and 50 s.

Fig. 4. Variation of thermocouple output temperature (right axis) and Ga desorption rate (left axis) as a function of time as a stepwise increase in the setpoint temperature (dashed line) is implemented. Results are shown for two different substrates: one is conductive ($n^+$) and one is semi-insulating (SI).
thermocouple output from the temperature controller and, instead, directly coupling the signal corresponding to \( F_d(Ga) \) [1]. In Fig. 5 we compare the responses of \( T_{TC} \) and \( F_d(Ga) \) to stepwise changes in their respective setpoints, during normal thermocouple control (TC control) and desorption rate control (DMS control), respectively. The temperature controller settings used were identical for the two data sets; the desorption signal was appropriately conditioned to fall into the appropriate input voltage range for the temperature controller for the case of DMS control. The data show that DMS control provides a rapid time response with very minimal overshoot of \( T_s \), in marked contrast to the significant overshoot and long settling time associated with TC control.

Note that, in the case of DMS control (Fig. 5), the thermocouple output rises in a nearly two-step linear-grade fashion. Thus, another potentially useful approach to improving the time response and overshoot characteristics of the time evolution of \( T_s \), during normal TC control would be to ramp up the setpoint temperature instead of increasing it in a stepwise fashion. This may be especially useful if the luxury of implementing DMS control is not available.

It is important to note that related difficulties in controlling surface temperature can also affect heterostructure growth when purposely reducing the substrate temperature. Previous measurements [3] on MBE growth of (Al,Ga)As/InGaAs structures have shown that significant In desorption can occur even when the thermocouple output temperature indicates that it should not. This comes about when dropping the substrate temperature from that used for AlGaAs growth to that used for InGaAs growth. Since one typically tries to avoid desorptive loss of In, the drop in temperature is implemented during growth of the last portion of AlGaAs. We have found that under typical growth conditions for our system, the thermocouple temperature settles to its new (lower) setpoint more rapidly than the actual surface temperature, and thus it is propitious to allow extra settling time.

4. Conclusions

The relatively loose relationship between the surface temperature and the non-contact thermocouple-indicated temperature gives rise to significant overshoot in surface temperature when effecting temperature setpoint increases under normal thermocouple control. The response time and the magnitude of overshoot associated with the surface temperature, when effecting stepwise changes in setpoint temperature, are dependent on the substrate conductivity and on the temperature controller settings, including the maximum current setting. By decoupling the thermocouple output from the temperature controller and, instead, coupling to it the suitably conditioned output of the desorption sensor, direct feedback control of the substrate surface temperature is effected, giving rise to a rapid response of the surface temperature with very little temperature overshoot.

Since the optimum growth temperature is generally different for adjacent epitaxial layers in a semiconductor heterostructure, stepwise changes in substrate temperature are routinely implemented during MBE growth. The results of the present study indicate that significant variation of growth rate and layer composition at heterointerfaces are likely to result if the temperature ramping is done in an uncontrolled fashion.
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