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1995

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INTRODUCTION

Silicon carbide is the only binary compound of carbon and silicon existing in a solid phase. It crystallizes in either the sphalerite (β-SiC or 3C-SiC) or wurtzite (α-SiC) structure with formation of ion-covalent bonds. A high Si-C bond energy in silicon carbide (~5 eV) provides its resistance to high temperature and radiation, imparts mechanical strength and chemical stability and causes low diffusion coefficients of dopants and host atoms. Among the materials, which impurity conduction of both the n- and p-type can be controlled within wide limits, SiC is distinguished as possessing the widest band gap, which ranges from 2.3 eV for the cubic polytype 3C-SiC up to 3.4 eV for the hexagonal polytype 2H-SiC. Silicon carbide has also a high saturated electron drift velocity which improves performances of devices destined to operate at high speed, high power levels. Its high thermal conductivity permits high-density device integration.

A number of techniques for growth of silicon carbide crystals has been proposed by different researchers. These techniques in general can be sorted over three broad classes. Yet, it seems to be not so easy to carry out this classification according such a scheme. The first technique is known as the method of chemical deposition in reagent gases introduced in a system within which they form the SiC crystals on an appropriate substrate. The second technique consists in crystallization from the silicon melt. The last type of main techniques for growth of SiC crystals is usually referred to as the sublimation method.

1 SECTION
GROWTH AND INVESTIGATION OF SILICON CARBIDE PARAMETERS FABRICATED BY THE SUBLIMATION METHOD

The most successful synthesis of SiC has been performed by the sublimation method. The earliest technological procedure for preparation of SiC is the so-called Acheson process. This method
is used for commercial production of SiC. It consists in that a mixture of silicon and carbide is heated up to 2500 - 2700 °C. The reaction yield is mainly the 6H-modification polycrystalline material. In some hollow cavities the single-crystal plates with the outline size up to 10 to 20 mm are formed. The crystals fabricated by this method cannot be used as substrates for semiconductor devices because the crystals grown are too small. The main shortcoming in the method consists in that one cannot control the process of nucleation, that leads to a large scatter of structural and electrophysical properties resulting in a low yield of suitable crystals.

Another sublimation procedure, which is virtually much better controlled, is the Lely method [1]. This method consists in vaporization of a SiC charge at about 2500 °C into a cooler cavity with subsequent condensation. Initially the charge forms its own cavity, while the more uniform crystals grow when a thin graphite cylinder is placed in the center of the charge. Such a thin cylinder reduces also a number of nucleation centers so that fewer and more perfect crystals grow. The crystals grow as thin hexagonal platelets perpendicular to the growth cavity surface. The 6H modification prevails among these crystals while other modifications such as 15R, 8H and 21R are found as well.

Tairov [2-4] has developed the method of fabrication of the 6H-SiC single-crystal ingots with using seed crystals by employing the modified Lely method in vacuum. When using this method, the crystal ingots with up to 20 mm diameter were obtained by Ziegler et al. [5,6] and the inch-size crystals of 6H-SiC were obtained by K.Koga, Nakata et al. [7-9]. These authors made use of various polytype seed crystals with the (0001) basal planes or the planes deviating by a few degrees from (0001). The crystal growth was carried out at temperatures around 1800 °C under the vacuum 10^-3 to 10^-4 bar with the axial temperature gradient in the growth zone of 30 to 40 °C/cm. Tairov and Tsvetkov [4] stated that using seed crystals with a given polytype is a reliable method for growing ingots of a desired polytype. On the other hand, they
marked problems as to preventing simultaneous growth of several polytype modifications in a single-crystal.

1.1. Growth of SiC single-crystals. The technique and equipment.

The SiC crystals have been grown in the special vertical-type high-temperature furnaces (Реммет-30) made of stainless steel and with water-cooled walls. High temperatures were provided with a heater consisting of a cylindrical graphite tube having a complicated form to provide a required temperature field in the growth zone. The equilibrium conditions in the growth zone to provide a homogeneous deposition take place usually in the closed systems because in such systems a material removal from a container space through a vapor phase is absent. Unfortunately, it is difficult to make an entirely closed system for growing SiC because of absence of a suitable container material. The vacuum-tight graphite container with a thin graphite cylinder placed in the container center has been used for our growth process. The SiC powder as a source material is loaded between the container walls and the graphite cylinder. When heated, the SiC powder evaporates and due to a presence of a temperature gradient it diffuses through pores in the graphite cylinder and then condenses on the walls. One can write the following set of chemical reactions for the Si-C system:

\[
\begin{align*}
\text{SiC}_{\text{sol}} & \leftrightarrow \text{SiC}_{\text{gas}} \\
\text{SiC}_{\text{sol}} & \leftrightarrow \frac{1}{2} \text{Si}_2\text{C}_{\text{gas}} + \frac{1}{2} \text{C}_{\text{sol}} \\
\text{SiC}_{\text{sol}} & \leftrightarrow \frac{1}{2} \text{SiC}_2\text{gas} + \frac{1}{2} \text{Si}_{\text{gas,liq}} \\
\text{SiC}_{\text{sol}} & \leftrightarrow \text{Si}_{\text{gas,liq}} + \text{C}_{\text{sol}}
\end{align*}
\]

We have used the commercial-grade SiC as a source material. Secondly, we have used SiC synthetized at our laboratory. The growth process was carried out within the temperature range 2200 to 2400 °C at the partial pressure of an inert gas (Ar or He) from \(10^{-5}\) to \(10^{-1}\) torr. The temperature was measured using an optical pyrometer in several parts of the growth zone.
The process duration ranged from 5 to 12 h.

The doping level of SiC crystals depends strongly on the impurity composition of an initial material, the inert gas purity and quality of the graphite units as well as on the outgassing cycle duration.

Fig. 1.1 illustrates a cross-sectional view of the sublimation crucible.

1.2. Polycrystalline SiC. The technique and equipment.

To grow polycrystalline SiC we have used the method of synthesis from the Si and C separate sources. This method uses vaporization of Si from a graphite container with a subsequent condensation of SiC on a graphite lid. A large amount of superpure graphite used for a crucible and other units of the high-temperature furnace is indicative of this apparatus. Graphite units in the system represent a reservoir of carbon. The following set of chemical reactions can be written in this case:

\[
\begin{align*}
\text{Si}^{\text{liq, gas}} + \text{C}^{\text{sol}} & \rightleftharpoons \text{SiC}^{\text{gas}} \\
2\text{Si}^{\text{liq, gas}} + \text{C}^{\text{sol}} & \rightleftharpoons \text{Si}_2\text{C}^{\text{gas}} \\
\text{Si}^{\text{liq, gas}} + 2\text{C}^{\text{sol}} & \rightleftharpoons \text{SiC}_2^{\text{gas}}
\end{align*}
\]

The synthesis was carried out at 1800 to 2000°C under the vacuum $10^{-4}$ to $10^{-2}$ torr. The growth rate depends on the temperature of growth and achieves 0.2 to 2 mm/h.

It is possible to obtain layers which are up to 10 mm thick [10]. Formed as a rule at synthesis temperatures $T \leq 1800 ^\circ$C is the 3C-SiC modification. Temperature increasing leads to formation of $\alpha$-SiC. The structure of layers depends on the deposition rate. Dense layers are formed at the maximum deposition rate. A grain growth direction in regard to a substrate is determined by directions of vapor fluxes. The maximum cross-section size of the grains is 250 to 500 µm.

After SiC growth a graphite substrate was removed by burning in air at 750 to 900°C.
Fig. 1.1. Cross-sectional view of the sublimation crucible: 1 - SiC powder; 2 - crucible; 3 - fabricated crystal; 4 - heater; 5 - heat shield; 6 - heat-insulating material; 7 - pyrometer opening.
Fig. 1.2. Scheme of the crucible for the growth of polycrystalline SiC.
Fig. 2 shows the set-up arrangement.

1.3. Parameters of the polycrystalline SiC

The resistivity of the SiC material synthetized under different conditions may range within the limits $\rho = 2.5$ to 400 Ohm cm. According to the data of a spectral analysis the percentage content of main impurities is as follows:

<table>
<thead>
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<th>Specimen</th>
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<tbody>
<tr>
<td></td>
<td>No.1</td>
</tr>
<tr>
<td>Ca</td>
<td>$10^{-3}$ to $10^{-5}$</td>
</tr>
<tr>
<td>Cu</td>
<td>-</td>
</tr>
<tr>
<td>Fe</td>
<td>$10^{-3}$ to $10^{-4}$</td>
</tr>
<tr>
<td>Al</td>
<td>$10^{-2}$</td>
</tr>
<tr>
<td>Mg</td>
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</tr>
<tr>
<td>Sb</td>
<td>-</td>
</tr>
<tr>
<td>Ti</td>
<td>$10^{-5}$</td>
</tr>
<tr>
<td>B</td>
<td>-</td>
</tr>
<tr>
<td>P</td>
<td>-</td>
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Fig. 3 presents the results of measuring the Hall effect for the sample with the resistivity $\rho = 270$ Ohm cm. It is seen that the electron concentration increases with temperature almost exponentially over the range 80 to 300 K within the limits $2.9 \times 10^{15}$ to $4 \times 10^{17}$ cm$^{-3}$. The mobility $\mu_n$ within the range 100 to 300 K is nearly constant and equal to 0.15 cm$^2$/V s. Only at temperatures near to 80 K its value somewhat increases, remaining
Fig. 1.3. Temperature dependence of concentration and mobility of polycrystalline SiC.
less than $1.0 \text{ cm}^2/\text{Vs}$. The activation energy calculated from a slope of the $n=f(1/T)$ dependence is 0.04 eV, which corresponds to the energetic position of nitrogen in regard to the c-band of 3C-SiC. Thus, one may assume that the electron concentration in our samples is determined by ionized levels of nitrogen the concentration of which is $(3 \text{ to } 8) \times 10^{17} \text{ cm}^{-3}$.

We suppose that the presence of boundaries between micrograins of 3C-SiC gives insignificant effect on a measured value of the Hall coefficient and an electron concentration determined from it. This is caused by the fact that the micrograin dimensions are by three orders of magnitude larger than the free path of electrons in the case of scattering by impurities at impurity concentrations $\sim 10^{18} \text{ cm}^{-3}$. On the other hand, the presence of intercrystal boundaries significantly decreases the efficient electron mobility in the samples under study. Therefore, the mobility values for polycrystalline samples are by three orders of magnitude less than that for the 3C-SiC single crystal. Invariability of a measured efficient mobility over the range 100 to 300 K is caused by a constant value of energetic barriers at microcrystal boundaries [10].

1.4. Investigation of single-crystal parameters.

1.4.1. Analysis of impurity composition

We have used the X-ray microprobe analyzer being a part of the scanning electron microscope JOEL (Japan), the JAM845 Model, with an auxiliary unit for microprobe analysis (Link-system). This technique is destined for a sample composition analysis with a depth of analysis dependent on the accelerating voltage of an electron microprobe (5 to 50 kV) and atomic numbers of materials under study. The analysis depth is within the limits 0.2 to 1.5 \text{\mu m}. The method is based on the fact that the line intensity of the characteristic X-rays is proportional to percentage of a given element in a sample. The method enables to analyze elements with atomic numbers from 6 to 100 and has the sensitivity of 0.5 \%wt. The characteristic emission is received by the detector with
energetic dispersion, the AN10000 type, through special windows. For light elements with small emission energy one uses a thin window, while for heavy elements the thicker beryllium window is applied. To remove a charge from dielectrics, a thin gold film (~50 Å) is deposited on a surface which because of its small thickness does not manifest itself in a spectral pattern. Three typical samples have been analyzed: No.1 - 15R, No.2,3 - 6H-SiC. As following from the spectral patterns (Fig.1.4-1.9), all the samples have the same element composition in regard to impurity contents. The samples are homogeneous by both their depth and surface area. Present in the samples under study are oxygen and phosphorus at a trace level (no more than 1 %wt). One observes such impurities as Na, S, Cl, K.

1.4.2. Investigation of structural perfection

This investigation has been carried out for the 6H- and 15R-SiC single-crystal plates cut perpendicular to the c axis. The plates were polished with diamond pastes and then etched for 2 min in the KOH-KNO₃ (1:4) melt at 650°C.

The Lang projection method [11] and the transmission hold-frame method [12] have been used as X-ray topographic methods with employing the MoKa₁ emission and the {1120}, {3300}, {1011} reflexes which possess the largest reflectivity. Chosen for studies were the samples with well developed pyramidal {1011} faces and the (0001) face being a base of a pyramid. Such crystals are more perfect.

The most frequently used to obtain projection topographic patterns are the {1120}-like reflexes since they are more informative. The reflexes like {3300} and {1011} were used to determine the Burgers vector directions in separate plane clusters of dislocations from "extinguishing" their reflexes [13].

One should remove a damaged surface layer formed after mechanical processing. For this purpose we used chemical etching. A surface damaged layer consists of individual scratches creating elastic deformations which spread into a large depth. Presented in
Fig 1.4.
Fig 1.5.
Fig 1.6.
X-RAY: 0 - 20 keV
Live: 100s Preset: 100s Remaining: 0s
Real: 111s 10% Dead

6H-SiC N1  SP-9
Be window

Fig 1.7.
X-RAY: 0 - 20 keV
Live: 100s Preset: 100s Remaining: 0s
Real: 111s 10% Dead

6H-SiC N2 SP-6
thin window

Fig 1.8.
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<td>Real: 111s 10% Dead</td>
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<th>SP-10</th>
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<tr>
<td>Be window</td>
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![Graph showing X-ray spectrum with peaks for Si, P, S, C, K, and M, with annotations for 5.040 keV and 10.2 keV, and 6 cts.]

**Fig 1.9.**
Fig. 1.10b is a topographic pattern of a sample along its depth. The regions with a light contrast are damaged regions of the crystal. After etching along the depth down to $dt=5 \mu m$ elastic deformations have decreased and it has become possible to observe a defect structure of the entire volume. It is seen that the sample No.1 (6H-SiC) is a perfect crystal containing in all 8 dislocations which are originated from a polytype inclusion. Moreover, concentrated in the central part of the sample are inclusions of unknown character.

Fig. 1.11 presents the topographic patterns for two samples of 6H-SiC. The sample No.2 is characterized with three plane dislocation clusters lying at different depths. The major centers of dislocation generation are concentrated near to a crystal "leg" which is a place where a crystal adheres to the crystallization bush of the growth furnace.

It is convenient to measure the dislocation density $N_d$ in plane clusters using its number per length unit instead of a number per area unit as in the case of a random distribution. Expediency of such an approach is evident from Fig. 1.11b obtained by the "hold-frame" method for the sample No.2. As seen from this figure, various dislocation clusters lie at different depths in a sample. The dislocation density in such clusters is $N = 10$ to $100 \text{ cm}^{-1}$.

The crystal No.3 is the least perfect. Its majority of dislocations have a loop-like form and they are generated by inclusions of another phase.

The samples No.4 and 5 (15R-SiC) are quite perfect and they have small densities of dislocations whose ends come out on the lateral sides (Fig. 1.12).

Thus, the single crystals of 6H and 15R-SiC prepared by the sublimation method are quite perfect. Most of them have small ($N_d=10$ to $100 \text{ cm}^{-1}$) density of basal dislocations.
Fig. 1.10. Projection topograhpic patterns (a,b) and "hold-frame" transmitting topograhpic patterns (b,c) for sample #1 MoKa1\(\frac{1}{2}120\) - reflex, a,b - after mechanical ground and polishing; c,d - after chemical etching.
Fig. 1.11. Projection (a, c) and "hold-frame" transmit (b) topographic patterns. MoKα₁, \{1120\} - reflex. a, b - crystal № 2, c - crystal № 3.

Fig. 1.12. Projection topographic patterns sample № 4 (a) and sample № 5 (b). MoKα₁, \{1T20\} - reflex.
1.4.3. Identification of SiC-polytypes by the Raman Scattering and X-ray diffractometric methods

The spectral features of optoelectronic devices on SiC depend on the polytype used. The latter one is identified commonly by the spectral position of impurity luminescence bands. But the intensity of the characteristic luminescence at the room temperature is high only at a large enough impurity content. On the other hand, the spectral half width of these luminescence bands is large enough and this fact results in a bad accuracy, especially for specimens that contain several polytypes. Such polytype splices are natural heterostructures and therefore maybe practically important in some optoelectronic devices.

As an alternative method to identify SiC polytypes one can use the method based on the Raman Scattering (RS). If it is implemented using a focused laser beam for excitation then it gains advantage over the luminescence and X-ray diffractometric methods. The latter is related with a high space resolution of Raman Scattering and as a result with a possibility to check small inclusions of another polytype in a sample.

It is known that all the SiC polytypes are characterized by the general phonon dispersion in the standard large Brillouin zone, but the frequencies of phonon lines in the Raman spectra are specific for every polytype [14]. From this point of view a certain polytype may be considered as a natural superlattice with a specific period. The most convenient way to use this fact for diagnostics is measuring the Raman spectra in the frequency region of "folded" acoustic phonons, due to a large dispersion of acoustic phonon branches.

With a view to identify a polytype composition of the SiC splices we have carried out investigations of the Raman spectra for a number of samples. Along with it, we have measured the photoluminescence spectra excited by the nitrogen laser
radiation ($\lambda=337.1$ nm). The Raman spectra were recorded with the DFOC-24 spectrometer. An excitation source was the argon laser ($\lambda=488.0;514.5$ nm). A laser beam was directed along the c axis with a view to obtain scattering by phonons of all the polytypes which are present in a splice along the given direction.

Presented for illustration (Fig. 1.13) is the Raman spectrum for the sample which, according to the photoluminescence data from opposite faces, is assigned to a splice of the 6H and 21R polytypes. It is seen that in this Raman spectrum in addition to appropriate phonon lines 148 cm$^{-1}$ (6H polytype) and 130 cm$^{-1}$ (21R polytype) manifested is also the intensive peak 172 cm$^{-1}$, corresponding to the 15R polytype (with a view to compare, Fig. 1.13 presents also the Raman spectra for pure polytypes). Thus, the RS method has enabled to obtain more complete information on the splice polytype composition. One may conclude that the 15R polytype layer is situated inside the sample and the surface layers at both splice sides are formed by the 6H and 21R polytypes. Moreover, since the RS cross-sections for the characteristic phonons under consideration are nearly equal for different polytypes, then the RS line intensity relationship gives an information on the ratio of polytype thicknesses in the splice. The prevailing bulk part of the sample under consideration consists of the "internal" 15R polytype which has not been found in the spectra of photoluminescence from its faces.

Similar investigations of RS have been carried out for a number of splices containing several combinations of the following polytypes: 4H, 6H, 8H, 15R, 21R. In the cases when the samples consist of two polytypes, we observed a correlation with the photoluminescence data. For the splices consisting of three and more polytypes, with a view to check the Raman spectra we have analyzed a polytype composition using the diffractometric method proposed in [15].

Since the samples under study are plates parallel to the (0001) plane, then the analysis of a polytype structure
Fig. 1.13. RS Spectrum fragment for the pure SiC polytypes: 15R(1), 6H(2), 21R(3) and polytype splice 15R+6H+21R(4)

Fig. 1.14. Diffraction patterns for the SiC polytype splices: 15R+6H+21R(a), 6H+15R(b)
is carried out as following from the angle difference \( \omega = \theta - \alpha \) (\( \theta \) is the angle between a crystal plane and the c axis, \( \alpha \) is the Bragg angle). The diffractometric analysis was carried out using the X-ray goniometer and MoK\(_{\alpha}\)-emission. Since for the sample under study one observes the Laue diffraction of X-ray emission, we studied the entire volume.

In the diffraction pattern shown in Fig.1.14a we can see the reflexes (01.2), (01.5), (01.8), (01.11) of the 15R polytype; (10.1), (10.2), (10.3), (10.4), (10.5) of the 6H polytype and (01.5), (01.8), (01.11), (01.14), (01.17) of the 21R polytype. In accordance with RS data the diffraction pattern presented gives evidence that the prevailing part of the sample consists of the 15R polytype and the 6H and 21R polytypes are present with a considerably less amount. Shown in Fig.1.14b is the diffraction pattern for the sample with approximately identical percentage of the 6H and 15R polytypes. The diffractometric analysis data for all the samples studied have entirely confirmed the results of a polytype composition identification for the SiC splices made by the RS method.

Especially convincing advantages of the RS-method manifest themselves with using the MicroRaman analysis. Focusing the exciting laser beam in a 1 \( \mu \text{m} \) size spot enables to check polytype interfaces in a splice with a high space resolution.

In such experiments we have used the triple additive monochromator Dilor (OMARS 89, XY), precision ~ 1 \( \mu \text{m} \), with the Olympus microscope (x10, x50, x100 objectives) and CCD camera as a multichannel detector (1024 pixels 25x2500 \( \mu \text{m}^2 \) intensified by the microchannel tube), the S25 entrance photocathode.

Fig.1.15 presents some RS spectra from a cross-section of the splice which contains three polytype layers (15R+6H+15R). The cross-section of the splice studied is shown schematically in Fig.1.16.

The spectra No.1, No.10 and No.19 in Fig.1.15 correspond to
Fig. 1.15. RS spectra from a cross-section of the splice which contains three polytype layers (15R+6H+15R)

Fig. 1.16. The cross-section of the splice which spectra are shown on Fig. 1.15.
the points No.1, No.10 and No.19 in the first (15R), second (6H) and third (15R) layers of the sample at a distance from polytype interfaces. The rest spectra are obtained with the 0.25 µm step near to the first (No.3, No.5, No.7) and the second (No.13, No.14, No.15, No.16) polytype interfaces. The line 221.04 cm⁻¹ in all the spectra is the Ar⁺ plasma line which was used as a reference point.

1.4.4. Spectrum of nitrogen in 6H and 15R SiC.

ESR spectra of nitrogen are studied for 6H and 15R SiC samples produced by means of Lely method, contained nitrogen in concentrations of $10^{16}$–$10^{19}$ cm⁻³. Measurements were carried out by the EPR spectrometer with operation frequency 140 GHz. The high microwave field frequency and the presence of a static magnetic field up to 7.4 T, generated by a superconducting magnetic system with a high homogeneity (10⁻⁵ per 1 cm⁻³), made it possible to investigate ESR spectra with an extremely high resolution in terms of the g-factor.

The relative values of the g-factor of the unknown spectra were determined to within $2 \times 10^{-5}$ (when the width of an absorption line was 5 G). The adopted procedures made it possible to ensure a high sensitivity of the spectrometer when single crystals of $10^{-4}$ cm³ volume ($5 \times 10^8$ spins/G) were used [16].

The ESR spectrum of 6H and 15R SiC samples with nitrogen concentration from $2 \times 10^{16}$ to $7 \times 10^{16}$ cm⁻³ and different degrees of compensation consisted of two for 6H and three for 15R triplets of the hyperfine structure lines (I₂k₁, I₂k₂, I₂k₃) due to the nitrogen nuclei $^{14}$N, corresponding to two for 6H SiC and to three for 15R SiC in equivalent cubic position of nitrogen, as well as a single line I₁h for 6H SiC and I₁h₁, I₁h₂ for 15R SiC with an anisotropic width corresponding to the hexagonal position of nitrogen (Fig.1.17a, 1.18a) [16].

The appearance of the I₁h for 6H SiC and I₁h₁, I₁h₂ for 15R SiC
ESR spectrum of nitrogen in 6H SiC

6H SiC  T = 4.2 K

f = 140 GHz

Fig. 1.17
ESR spectrum of nitrogen in 15R SiC

15R SiC

T = 4.2 K

f = 140 GHz

b

\( S \)

18 \(-3\)

5 \( \cdot \) 10 cm

a

\( S \)

17 \(-3\)

2 \( \cdot \) 10 cm

Fig. 1.18
line in the ESR spectrum and the intensity of this line depended on the degree of compensation of the investigated samples [17].

An increase in the nitrogen concentration $N_N$ modified the observed spectrum. It clear from Fig.1.17b,1.18b that when $N_N$ was increased up to $5 \times 10^{17}$ cm$^{-3}$, an additional wide S-line appeared against the background of the spectrum of single centers in 6H and 15R crystals. The S-line position did not correspond to the center of gravity of the ESR spectra of the cubic and hexagonal nitrogen positions nor to the center of gravity of these spectra. The S-line has an asymmetric profile and an anisotropic $g$-tensor. Its intensity was much higher than the sum of the intensities of the absorption lines of isolated nitrogen atom.

The further growth of the concentration of nitrogen up to $10^{19}$ cm$^{-3}$ results in the shift of all nitrogen ESR lines in 6H and 15R SiC towards the S-line and, at last, one observes the only S-line in ESR spectrum.

Analyse of exchange line in 6H-SiC show that S-line consist of two S1 and S2-lines with different width and intensity (Fig.1.17b). At $T = 4.2$ K $g$-value of S1-line corresponds to the average value of the Ih1 and Ik1 ESR spectra whereas the $g$-factor of S2-line corresponds to the average value of the $g$-factor of the Ih1 and Ik2 ESR spectra. As shown from the temperature dependence of the $g$-value of S1 and S2-lines (Fig.19), $g$-value of S1 and S2-lines approaches the $g$-value of EPR spectra of cubic position Ik1 and Ik1 correspondly at 77 K. Fig.1.20 shows the temperature dependence of the intensity of the S1 and S2 line in the temperature interval 4.2 - 40 K. The intensity increases with lowering temperature according to the Curie low:

$$I(T) \sim C/T \quad (1.1)$$

This data are consistent with the data obtained for 6H SiC in [18].
The temperature dependence g-factor S-lines

\[(g - 20000) \times 10^4\]

Fig. 1.19
The temperature dependence of the intensity of the S-line.

Fig. 1.20
The $S_1$ and $S_2$-line width depends on temperature. Fig.1.21 shows the temperature dependence of the $S_1$, $S_2$-line width in the temperature interval 4.2 - 40 K. When temperature increase from 4.2 to 35 K the $S_1$ and $S_2$-line width decreases. A similar behavior of temperature dependence of the exchange line width was observed for Si heavily doped with phosphorus, described by expression [19]:

$$dH \sim \exp\left(\frac{2R_0}{a_0}\right) \times \tanh\left(D/kT\right) \quad (1.2)$$

Where $R_0$, $a_0$ and $D$ are the inter donor distance; effective Bohr radius and the potential difference between an occupied site and an unoccupied site, respectively.

For 6H SiC $a_0 = 5 - 7$ Å [20]. The temperature dependence for $S_1$ and $S_2$-line are in accordance with equation (1.2) with parameters of $R_0 = 14$ Å, $D = 50$ K for $S_1$- line and $R_0 = 12$ Å, $D = 52$ K for $S_2$- line.

These data suggests that the electrons spend most of the time at the donor sites. The probability of the direct Heisenberg exchange between the paramagnetic nitrogen centers was very low, but the probability that paramagnetic nitrogen atom encountered a nonparamagnetic nitrogen atom in the same of the nearest unit cell of the crystal was high. This elementary jump was on the one hand, an essential part of the hopping conduction process and, on the other hand, it ensured hopping exchange of a donor electron between two paramagnetic systems. As shown in [17] at high donor concentrations and high degree of compensation were many ionized nitrogen centers at the hexagonal positions, which were nonparamagnetic. Therefore the probability that paramagnetic nitrogen atom at the cubic position encountered a nonparamagnetic nitrogen atom in the hexagonal position was high. In this case donor electron could be captured from a cubic by a hexagonal nonparamagnetic nitrogen atom that the latter became paramagnetic. Consequently, the position of the exchange line in the ESR spectrum of nitrogen in
The temperature dependence of the line width.

![Graph showing the temperature dependence of the line width.](image)

Fig. 1.21
the hexagonal polytype of SiC was governed by the nitrogen concentration by the degree of compensation of the sample, by the ratio of the depths of the energy levels of the hexagonal and cubic center in the band gap and by the value of the effective Bohr radius of the donor electron.

It is clearly follows from the temperature dependence of the g-value of S1 and S2-exchange lines. As the temperature increase the ionized nitrogen centers at the cubic position were appeared, probability of the exchange hopping motion between paramagnetic and nonparamagnetic atom in one cubic position was increased and g-value of S1 and S2-line shifts to the center gravity of the ESR spectrum of nitrogen at the cubic position.

This feature suggests that the mechanism of hopping motion for donor electron in 6H SiC heavily doped with nitrogen is working.

The nature of the low-temperature conductivity of 6H SiC was investigated earlier in [18] and it was established that between 4.2 K and 25 K the low governing the electrical resistivity of the sample with the nitrogen concentration $10^{19}$ cm$^{-1}$ was typical of variable-range hopping too.

1.4.5. Surface electron properties of the 6H and 15R modification silicon carbide

The investigation was carried out with the n-type samples of single-crystal silicon carbide, 6H- and 15R-modifications, with the nitrogen impurity concentration of $9 \times 10^{17}$ and $3 \times 10^{18}$ cm$^{-3}$, respectively [20-22]. Their large faces (0001)Si and (0001)C were ground and then polished using diamond pastes. In order to remove a layer with mechanical structural damages, the samples underwent oxidation at 700°C in a humid oxygen flux for a few tens of minutes. Then the oxide layer was etched in the CP-4 etcher (HF:HN$_3$:CH$_3$COOH = 3:5:3). Afterwards, the samples were electrolytically oxidized in the 5% solution of KNO$_3$ in ethyleneglycol for 10min. The samples electrolytically oxidized underwent studies of the surface potential vs temperature
dependencies. The study was repeated after removal of the oxide layer obtained electrolytically by processing in HF. Then after etching the samples in KOH at 650 °C we carried out identification of the (0001)Si and (0001)C surfaces according to their characteristic appearance [23] using an optical microscope.

The surface potential $\phi_s$ (energetic band bending at surface) was determined by the method of measuring the surface photovoltage under the conditions of a large level of electron-hole pair generation by light pulses of the MCI-100 flash lamp (illumination intensity $\sim 10^{24}$ quanta/cm$^2$ s) which straightens bands at a semiconductor surface at the moment of illumination [24]. For this purpose we mounted in a cryostat the measuring capacitor consisting of a mica plate with the SnO$_2$ semitransparent layer deposited on mica. A sample of SiC was attached with one of its surfaces to another side of the micaplate. The photovoltage signal was recorded by a storage oscilloscope that enabled, taking into account the calibration coefficient of the circuit, to determine the $\phi_s$ value.

The temperature dependence $\phi_s(T)$ was recorded under the vacuum $10^{-4}$ Pa with lowering temperature from 300 to 100 K. In the most cases already at the room temperature we have found the surface potential photomemory effect [25]. It consists in the fact that after illumination of a sample by the first lightpulse one observes a capture of minority carriers (holes) at the surface traps which results in variation of $\phi_s$ after action of a light pulse. As a result, using the second or any other subsequent light pulse one measures already a changed value of $\phi_{s2}$ which is less by absolute magnitude as compared with the $\phi_{s1}$ value measured using the first light pulse. The difference $|\phi_{s1} - \phi_{s2}|$ determines the surface potential photomemory. Under the conditions of our experiment it was determined at the light pulse repetition frequency equal to 1 Hz.

It should be noticed that the $\phi_{s2}$ value determined using the second pulse was equal to that for the third or any other subsequent pulse. This gives evidence that traps capturing
nonequilibrium holes are saturated with them already under illumination by the first light pulse. The $\phi_{S1}$ value corresponding to the dark conditions can be determined at a certain temperature only after emptying traps with captured holes which demands to wait a certain time after a previous illumination or to reheat a sample. Below the room temperature the period of storage of the holes captured in traps considerably increases (hours). Therefore, for correct determination of the $\phi_{S1}$ and $\phi_{S2}$ dependencies on temperature every time after measuring at a certain temperature we carried out reheating a sample up to temperatures above 300 K with subsequent cooling in dark down to a temperature for new measurements of $\phi_{S1}$ and $\phi_{S2}$.

Fig.1.22 and 1.23 present the temperature dependencies of the surface potential: $\phi_{S1}$ (curves 1, 2) and $\phi_{S2}$ (curves 1', 2'), respectively for the SiC-6H and SiC-15R surfaces. The dependencies for the (0001)Si surfaces are given in Fig.1.22a and 1.23a. Those for (0001)C are given in Fig.1.22b and 1.23b. The curves 1, 1' are obtained for the surfaces etched electrolytically, the curves 2, 2' - for the surfaces processed after that in HF. The $\phi_{S1}$ and $\phi_{S2}$ values are always negative. As following from calculations, this means that with all kinds of processing then-SiC-6H and 15R surfaces a surface layer depleted with electrons is formed at these surfaces. The depletion decreases after illumination of a sample by the first light pulse since the $\phi_{S2}$ value by absolute magnitude are always less than corresponding values of $\phi_{S1}$.

The behavior of the surface potential vs temperature dependencies $\phi_{S1}(T)$ is different. For the (0001)Si surfaces of SiC-6H and 15R they have an N-like form. For the (0001)C surfaces they are V-like. At the same time, the N- and V-like forms for various polytypes and surfaces are different. The increase of $|\phi_{S1}|$ with lowering temperature, which is always observed near to the room temperature, is related to filling-in the fast surface electron states (SES) with electrons in consequence of shifting the Fermi level in a semiconductor to the c band. The decrease of
Fig. 1.22. Temperature dependencies of the surface potential $\Phi_S$ on the (0001) Si (a) and (0001) C (b) surfaces of SiC-GH:
Surface processing: $1, 1' -$ oxidized electrolytically; $2, 2' -$ processed in HF. $1, 2 - \Phi_{S1}; 1', 2' - \Phi_{S2}$. 
Fig. 1.23. Temperature dependencies of the surface potential $\phi_s$ on the (0001) Si (a) and (0001) C (b) surfaces of SiC-15R. Surface processing: 1, 1' - oxidized electrolytically; 2, 2' - processed in HF: 1, 2 - $\phi_{s1}$; 1', 2' - $\phi_{s2}$.
observed at lower temperatures is related to decreasing the negative charge in SES. It may be explained only assuming that with lowering temperature the fast SES system is reconstructed itself. We observed a similar phenomenon earlier at certain physical and chemical conditions of the germanium [26,27], silicon [28,29], and gallium arsenide [30] surfaces. It was related to reversible structural changes occurring with lowering temperature at a semiconductor interface with a covering oxide film. Reversibility and reproducibility of the ϕ_{S1}(T) dependencies were confirmed also by checking measurements for all the surfaces of SiC-6H and 15R.

A different behavior (N-like or V-like) of the ϕ_{S1}(T) dependencies at the (0001)Si and (0001)C surfaces of SiC-6H and 15R is related to features of oxide coverage on them. It is known that the (0001)Si and (0001)C planes formed respectively only by the Si and C atoms are covered by the SiO_2 oxide films which are formed under natural air conditions or during special oxidizing processing [23]. For the (0001)C surface an oxide coverage is usually uniform, while for the (0001)Si surface it is non-uniform and changing by thickness [23,31]. It should be reasonable to expect that structural changes at the SiC-SiO_2 interface would be different with lowering temperature for different characters of oxide coverage. From our point of view, an N-like behavior of the ϕ_{S1}(T) dependencies on the (0001)Si surfaces is caused first of all by the SiO_2 oxide coverage non-uniformity. We observed earlier a similar effect on the silicon surface when the oxide coverage uniformity was damaged specially by deposition of gold with a micro island structure [28]. Processing in HF etches electrolytically the SiO_2 layer formed on the SiC surface, but instead of it under the air conditions thin layers (~10 Å) of a natural SiO_2 oxide are formed which because of their non-uniformity on the (0001)Si surface provide for it an N-like behavior of ϕ_{S1}(T) as well, although strongly changed. A different character is also observed for V-like characteristics of the (0001)C surfaces, oxidized electrolytically and natural (after processing...
in HF, which are covered by more uniform oxide films of SiO₂, having however different thickness before and after processing in HF.

As for absolute values of the surface potential \( \phi_{s1} \) on the (0001)Si and (0001)C surfaces of SiC-6H and 15R oxidized and processed in HF, then the situation at first is unexpected. One would think that etching an oxide should lead for all the surfaces to increasing \( |\phi_{s1}| \), as it is observed for silicon [28], due to removal of a positive charge built in an oxide. However, such a situation is implemented only on the (0001)C surface of SiC-6H (Fig. 1.23b) and on the (0001)Si surface of SiC-15R (Fig. 1.23a). On the contrary, the values of \( |\phi_{s1}| \) on the surfaces (0001)Si of SiC-6H (Fig. 1.22a) and (0001)C of SiC-15R (Fig. 1.23b) decrease after processing in HF. It seems that values of a positive charge built in an oxide are different for the (0001)Si and (0001)C surfaces of various SiC polytypes. Moreover, processing in HF leads to formation of different SES systems with predominance of their acceptor or donor character.

2. In order to calculate charge variation in SES with lowering temperature, a magnitude of a hole charge captured by surface traps as well as to determine the Fermi level position on surfaces at various temperatures, one should know the temperature dependence of the electron bulk concentration in the semiconductors under study. Investigations of the Hall effect have shown that with lowering temperature in the region 300 to 100 K the electron concentration in the conduction band of SiC-6H and SiC-15R changes, respectively, within the limits \((4.2 \text{ to } 0.2)\times 10^{17} \text{ cm}^{-3}\) and \((1.7 \text{ to } 0.2)\times 10^{18} \text{ cm}^{-3}\).

Equal to the charge in SES of a semiconductor, which determine a value of the surface potential \( \phi_s \), is a charge \( Q_{sv} \) with an opposite sign situated in a semiconductor region adjacent to surface. A variation of \( Q_{sv} \) with temperature (with an appropriate variation of \( \phi_s \) with temperature) has to be equal to variation with temperature of \( Q_s \) in the fast SES of a semiconductor. In our case to compute values of \( Q_{sv} \) one has to use the theory taking
into account a variation of electron charge in the conduction band and donor states with changing $\phi_{s1}$[32]. In this case, $Q_{sv}$ at the surface potentials depleting the region adjacent to surface with electrons is equal to

$$Q_{sv} = q \left( \frac{\varepsilon_k T}{2\pi q^2} \right)^{1/2} \left\{ n \left( \exp \left( \frac{q\phi_S}{kT} \right) - 1 \right) + N_d \ln \left( 1 + \exp \left( \frac{E_d - E_F}{kT} \right) \right) \right\}^{1/2}$$

Here, $q$ is the electron charge, $\varepsilon$ is the dielectric permittivity of SiC, $n$ is the bulk electron concentration in the c band of SiC at the temperature $T$, $N_d$ is the nitrogen donor center concentration, $E_d$ and $E_F$ are the energetic positions of donor centers and the Fermi level, respectively, in the bulk SiC at the temperature $T$ in regard to the middle of the energy gap $E_1$.

In order to compute $Q_{sv}$ using Eq.(1.3), except of measurement of the $\phi_{s1}$ value, one has to know for a certain $T$ values of $n$ and $E_d - E_F$. A value of $n$ was determined from the Hall effect and that of $E_d - E_F$ was determined from the neutrality condition:

$$n = \frac{N_d}{\exp(-\frac{E_d - E_F}{kT}) + 1} \quad (1.4)$$

The calculated values of $Q_{q}/q = -Q_{sv}/q$ for the (0001)Si and (0001)C surfaces of SiC-6H oxidized and processed in HF are presented in Fig.1.24 and those for SiC-15R are shown in Fig.1.25. Note again that the dependencies $Q_s(T)/q$ give variation...
Fig. 1.24. Temperature dependencies of the $Q_S$ charge in SES of SiC-6H. 1, 2 - (0001) Si surface; 3, 4 - (0001) C surface; 1, 3 - oxidized surface; 2, 4 - surface processed in HF.
Fig. 1.25. Temperature dependencies of the $Q_s$ charge in SES of SiC-15R. 1,2 - (0001) Si surface; 3,4 - (0001) C surface; 1,3 - oxidized surface; 2,4 - surface processed in HF.
of a charge in the fast SES as a function of temperature within the accuracy of a constant charge built in an oxide. As seen from Fig.1.24 and 1.25, the $Q_s(T)/q$ behavior is identical to that of $\phi_\text{S1}(T)$ for the (0001)Si surfaces it has an N-like form, and that for (0001)C is a V-like. For all the $Q_s(T)$ dependencies an increase of a negative charge in SES is characteristic when lowering temperature from 300 down to 240-220 K, which is related, as indicated above, to filling SES with electrons along with the Fermi level shifting to the c band.

In this connection we computed the Fermi level position on a semiconductor surface $E_F$ and its variation with lowering temperature for various conditions of the (0001)Si and (0001)C surfaces of SiC-6H and SiC-15R. It turns out that at 300 K the Fermi level for the (0001)Si surface of SiC-6H, oxidized and processed in HF, is situated by 0.29 and 0.2 eV below the c band. For the same surfaces of SiC-15R it is by 0.1 and 0.21 eV below the c band. The $E_F$ position on (0001)C of SiC-6H is by 0.13 and 0.29 eV, respectively, below the c band and that on the (0001)C surfaces of SiC-15R is by 0.33 and 0.32 eV below the c band. With lowering temperature down to 240-220 K the $E_F$ level position on the surfaces indicated above changes slightly because of a high density of SES. Estimates made for a number of surfaces of SiC-6H and SiC-15R show that the SES density in the vicinity of the Fermi level on these surfaces is within the limits (2 to 10)$\times 10^{12}$ cm$^{-2}$ eV$^{-1}$. Below 220-180 K, when one observes in all the cases a decrease of a negative charge $Q_s$ in SES (Fig.1.24 and 1.25), the $E_F$ level for most of the surface conditions of SiC-6H and 15R approaches considerably to the c band. At minimum values of $|Q_s|$ its position, for example, for the SiC-15R surfaces characterized by curves 1-4 (Fig.1.25), is, respectively, by 0.07, 0.15, 0.24, 0.15 eV below the c band.

The decrease of a negative charge in SES with the Fermi level $E_F$ approaching simultaneously to the c band gives evidence that, as already mentioned above, there is a reconstruction of the SES
system occurring with lowering temperature. With such a reconstruction, the SES density below the $E_{FS}$ level decreases or the density of positively charged SES above $E_{FS}$ increases. The SES system reconstruction seems to be caused by structural changes at the interface between the silicon carbide crystal and the $\text{SiO}_2$ oxide film covering its surfaces. Reversibility and reproducibility of the $\phi_{S1}(T)$ and $Q_s(T)$ dependencies are indicative of the fact that the structural changes are reversible as well. They may be caused by the elastic stress at the interface SiC-$\text{SiO}_2$ arising from lowering temperature. This can lead, for example, to changing angles between the silicon-oxygen tetrahedrons and in the tetrahedrons themselves situated at the interface, that changes reversibly the fast SES system [33]. The SES system changing, which begins at a certain temperature, gives a V-like behavior of $\phi_{S1}(T)$ and $Q_s(T)$ on the (0001)C surfaces. An N-like behavior of these temperature dependencies on the (0001)Si surfaces covered by non-uniform oxide films is caused by competition of two mechanisms of changing charge: reconstruction of the SES system, when a negative charge in SES decreases, and filling SES with electrons, when it increases with lowering temperature.

3. Now, let us consider the surface potential photomemory effect related to capture of non-equilibrium holes in the surface traps under illumination of the SiC surface. According to that described above, the number of holes captured at a certain temperature, which gives a variation of the potential from $\phi_{S1}$ to $\phi_{S2}$, is equal to:

$$P = \frac{Q_{sv}(\phi_{S1}) - Q_{sv}(\phi_{S2})}{q} \quad (1.5)$$

The dependencies calculated using (1.5) for two kinds of conditions for the (0001)Si and (0001)C surfaces of the 6H and 15R modification silicon carbide are presented in Fig.1.26 and 1.27, respectively. It is seen that with lowering temperature from 300 down to 220 K in all cases the concentration of holes
Fig. 1.26. Temperature dependencies of the hole concentrations captured in traps of SiC-6H. 1, 2 - (0001) Si surface; 3, 4 - (0001) C surface; 1, 3 - oxidized surface; 2, 4 - surface processed in HF.
Fig. 1.27. Temperature dependencies of the hole concentrations captured in traps of SiC-15R. 1, 2 - (0001) Si surface; 3, 4 - (0001) C surface; 1, 3 - oxidized surface; 2, 4 - surface processed in HF.
captured in traps increases. Since during the first light pulse action the traps are saturated with holes, then one may state that the P(T) dependence reflects the temperature dependence of the trap concentration for the traps capturing holes. The increasing number of traps, which capture holes with lowering temperature, is related to the fact that there is a set of traps distinguishing by the hole lifetime in them. At higher temperatures the ones manifested as photomemory centers are the traps situated farther from the v-band of SiC. With lowering temperature, because of increasing the time of excitation of holes into the v-band, the traps, which are situated nearer to it, are added to the first ones. Investigations of the photomemory effect relaxation at various temperatures confirm this statement and show that there are several time constants of the relaxation. Energetic positions of traps estimated from the temperature dependencies of the relaxation time for the charge captured in them overlap the 0.7 eV-wide energetic interval above the silicon carbide valence band.

As seen from Fig. 1.26 and 1.27, the number of holes captured and, consequently, the number of traps are maximum (approximately $8 \times 10^{11}$ cm$^{-2}$) at 220 K on the (0001)C surfaces of SiC-15R and they are minimum ones ($\sim 2 \times 10^{11}$ cm$^{-2}$) on the (0001)Si surface of SiC-6H processed in HF. At temperatures below 220-180 K on the (0001)C surface of SiC-6H one observes intervals of slowing increase of the trap concentration with lowering temperature (curves 3 and 4, Fig. 1.26) or even some intervals of decreasing this concentration on the (0001)Si surface of SiC-6H (curves 1 and 2, Fig. 1.26) and on all the SiC-15R surfaces (Fig. 1.27). In this case the P(T)-dependencies for the (0001)Si surfaces of SiC-6H and 15R have an N-like behavior. As following from the $\phi_{s1}(T)$ dependencies, this is the very temperature region where the reconstruction of the SES system occurs for the states situated in the upper half of the SiC energy gap. Therefore, one may conclude that the traps capturing holes have a nature of origin which is common with that of SES and they differ from the latter only by the larger cross-section of
hole capturing. The structural reconstruction at the interface between a semiconductor and an oxide film occurring with lowering temperature changes the SES parameters including those states, which manifest themselves as traps for non-equilibrium holes.
II Section

AVALANCHE LIGHT EMITTING DIODES

Avalanche light emitting diodes (LEDs) are semiconductor devices using optical emission which arises in the p-n-junction under the avalanche breakdown. Such LEDs are characterized by a wide emission spectrum from 200 to 2500 nm, high temperature and temporal stability of emission, and their speed is limited by technical potentialities of modern electric pulse generators of the subnanosecond range. In connection with availability of these useful properties the avalanche LEDs may be used as reference emission sources to check stability of photoreceiving tracts, as sources of subnanosecond light pulses, as sources of UV emission as well as to imitate emission of space objects in astronomy and astronautics.

2.1. Investigation of parameters and technology

Under the conditions of reverse bias an electric field in the p-n-junction easily achieves values of $10^5$ to $10^6$ V cm$^{-1}$ in such fields, as a rule, the electric breakdown of the p-n-junction takes place, which is accompanied by sharp increase of a current and emission of photons. The emission spectrum of SiC diodes overlaps a wide range of photon energy and for this reason it differs from those corresponding to other electroluminescence mechanisms (Fig.2.1).

In the case, when the effects of selfabsorption can be made negligible, the emission is in fact independent of the concentration and given impurity composition in a crystal, and it is also slightly sensitive to variation of temperature [34].

Investigations show [34] that a wide-band structureless emission is caused by the intraband non-direct transitions of high-energy carriers. High-energy carriers are capable to excite also the recombination luminescence bands. However, these bands
Fig. 2.1. Emission spectrum of LEDs based on SiC.
1 - LED with "open" design; 2, 3 - LEDs with "closed" design and uncompensated donor concentration:
2 - $2 \times 10^{17}$ cm$^{-3}$; 3 - $3 \times 10^{18}$ cm$^{-3}$.
Y-axis: $N_\nu$ arb. units.
are superimposed on the continuous spectrum and may manifest themselves as some structure on the background of this spectrum.

The technology of avalanche LEDs has a number of peculiarities. Used for their fabrication are the 6H- and 15R-SiC crystals of n-type with the uncompensated donor concentration of \((0.5 \text{ to } 6) \times 10^{18} \text{ cm}^{-3}\). In the case of fabrication of p-n-junctions using customary methods (diffusion, epitaxy, MIS-technology) the breakdown voltage varies within the limits \((0.3 \text{ to } 2) \times 10^2 \text{ V}\). Application of the alloying technique enables to decrease the breakdown voltage down to \((15 \text{ to } 30) \text{ V}\) and to fabricate fast LEDs with a small area \([35,36]\). However, use of such junctions has a number of shortcomings: small number of microplasmas (MP) and large spread by the breakdown voltage of individual MPs in the same p-n-junction. As a rule, these microplasmas are situated in the peripheral and widely spaced parts of p-n-junctions.

Authors of \([37-40]\) have developed an improved alloying technique for fabrication of LEDs with a small area \(S < 10^{-4} \text{ mm}^2\). It consists in a preliminary treatment of crystal surface with a view to obtain p-n-junctions with a large quantity of microplasma regions with nearly identical breakdown voltages.

Fig.2.2 and 2.3 present typical reverse-bias I-V characteristics of the diodes and their dependence of the differential resistance on the reciprocal current \((1/I)\). Analysis of the characteristics enables to distinguish the following four parts:

1. In the region of small currents \(10^{-7} \text{ A} \ (10^{-3} \text{ A/cm}^2) < I < 10^{-5} \text{ A} \ (10^{-1} \text{ A/cm}^2)\) one observes, as a rule, a linear dependence of the current on the voltage.

2. In the region of \(10^{-5} \text{ A} \ (10^{-1} \text{ A/cm}^2) < I < 10^{-3} \text{ A} \ (10 \text{ A/cm}^2)\) the tunneling breakdown is predominant. This part manifests itself in the most LEDs and it is characterized by a linear dependence of the differential resistance on the reciprocal current \((1/I)\).

3. Intermediate region: \(10^{-3} \text{ A} \ (10 \text{ A/cm}^2) < I < 10^{-2} \text{ A} \ (100 \text{ A/cm}^2)\). Current flowing in this region is caused by simultaneous
Fig. 2.2. Current voltage characteristics of the diodes with breakdown voltage: 1 - 21.6 V; 2 - 21.2 V; 3 - 20.02 V; 4 - 19.05 V; 5 - 17.99 V; 6 - 17.12 V.
Fig. 2.3. Dependence of the differential resistance of the diodes $R_d$ vs current ($1/I$). Breakdown voltage: 1 - 21.9 V; 2 - 19.2 V; 3 - 15.4 V; 4 - 16.2 V.
action of the tunneling effect and impact ionization.

4. High current density region: \( I > 10^{-2} \text{ A} \) \((100 \text{ A/cm}^2)\). In this region the I-V characteristics are described by the expression indicative of the avalanche breakdown:

\[
I = \frac{(U - U_{br})}{R_d}
\]

The differential resistance value \( R_d \) as a function of the breakdown voltage 16 to 30 V varies within the limits from 8 to 30 Ohm. As following from these investigations the LED equivalent circuit may be presented as shown in Fig.2.4.

Fig.2.5 presents the dependence of the emission intensity on the current. One may distinguish two typical parts. At high current densities under the conditions of avalanche breakdown we have \( P \sim I \), i.e. in this regime the quantum yield of emission is independent of the current. In this case difference in the external quantum yield of LEDs with different breakdown voltages is mainly caused by absorption of emission during its leading-out through a crystal. LEDs with high breakdown voltages (material with a lower uncompensated donor concentration) have a higher external quantum yield.

In the intermediate current region \( (P \sim I^n, \text{ where } 1<n<2) \) the quantum yield is 2 to 3 times as low than that in the avalanche breakdown region and it increases with increasing current.

The LED characteristic of importance is their long-term stability of parameters, especially of emission intensity. We have developed the special automated stand [6] to measure this important parameter. The main idea of the method consists in comparison of the emission intensity of the LEDs under study and that of reference sources. At the same time it is assumed that the emission intensity variation for reference sources (long-term instability) is small and does not exceed some known value. Such a concept of measurements enables to eliminate instability of a photodetector transfer function. A set of the radioluminescent
Fig. 2.4. Equivalent circuit of LED for: a - direct current; b - high frequency supply. Here $I_T, R_{ST}$ - tunnel components of spilling current and resistance; $I_A, R_{SA}$ - avalanche components; $R_L, R_J$ and $R_{SA}$ - leakage resistance, junction resistance and series avalanche resistance respectively of the avalanche breakdown; $L_C, C_e$ - reactances of a can.
Fig. 2.5. Dependence of the emission intensity vs current for LEDs with breakdown voltage:
1 - 27.5 V; 2 - 16.9 V; 3 - 19.2 V; 4 - 15.4 V.
emitters (RIE) based on a mixture of a crystallophosphor with a radioactive substance containing $^{14}C$ is used as reference sources. The temporal instability of the reference sources was no more than $5.7 \times 10^{-5}$ h$^{-1}$. Measurements were carried out using a photomultiplier in the photon-counting regime. The measurements carried out for a group of three RIE and six avalanche LEDs have shown that both kinds of sources with a high accuracy are described by the Poisson distribution and LEDs after preliminary training have a high long-term stability of emission, which is no more than 3 % per 1000 h of operation.

Measurements of the avalanche LED speed were carried out in the Institute of Physics of Solids at Latvian State University. We made use of the single-photon statistical method with the time resolution of 0.25 ns and the subnanosecond generator shown in Fig.2.6.

Fig.2.6 shows a light pulse from the avalanche LED. The pulse duration at half height is no more than 0.5 ns at the 0.2 level of amplitude – no more than 0.7 ns. The pulse decay is exponential with the time constant of 0.15 ns within the limits of than 3 decimal orders. The dynamical range of measurements is 4 orders of magnitude and it is limited a photomultiplier interference.

The physical limit of speed for our LEDs cannot be determined at present because of insufficiently high level of the subnanosecond equipment. Further decreasing the light pulse duration may be possible after improving the pulse generator design.
2.2. Technical parameters

- spectral emission range (×) 200 to 2500 nm
- flash intensity at the 1 ns duration (photon number, no less) 5000
- temperature coefficient of emitted intensity (no more) 0.1 %/degree
- time constant (no more) (××) 0.2 ns
- emission zone diameter 30 to 150 μm
- operation temperature range (×××) -60 to +80 °C
- operation voltage range: dc 13 to 30 V
  pulse current up to 100 V
- operation current range: dc up to 150 mA
  pulse current up to 10 A (××××)
- operation life (no less) 100000 h

(×) the spectral range is indicated for the "open" design;
(××) resolution limits of the measuring set-up;
(×××) determined by a capsule design; fabrication of LEDs with a wider range is possible;
(××××) permissible limits of the pulse current are determined by the pulse duration; for short pulses T<3 ns I_{max}=10 A, at T<10 ns I_{max}=3 A.

Figures below present the LED design (proposed to a customer). There is a possibility to apply other design versions coordinated with a customer.

By their electrical characteristics and operation principle the avalanche LEDs are similar to the semiconductor voltage-reference diodes. Therefore they work under a reverse bias. Note: in the case of direct current supply the LED device must be mounted on a radiator with an effective area corresponding to the power dissipated by LED (~3 W). For the design versions presented here the polarity of power supply corresponds to positive voltage on a can. In the pulse operation regime LEDs can be used without radiators in case the mean dissipated power is no more than 100 mW. With the pulse duration...
T>10 ns the maximum pulse current should be no more than 200 mA. In the case of short pulses T<3 ns the current may reach 10 A.

Parameters of fabricated LEDs

<table>
<thead>
<tr>
<th>No.</th>
<th>Design</th>
<th>$V_{br}, V$</th>
<th>$R_{dip}(I=100 \text{ mA})$ Ohm</th>
<th>P(450–600 nm) mcWt</th>
</tr>
</thead>
<tbody>
<tr>
<td>505</td>
<td>&quot;Closed&quot;</td>
<td>20.52</td>
<td>18.44</td>
<td>0.129</td>
</tr>
<tr>
<td>532</td>
<td>&quot;</td>
<td>21.06</td>
<td>16.01</td>
<td>0.135</td>
</tr>
<tr>
<td>610</td>
<td>&quot;</td>
<td>20.75</td>
<td>11.65</td>
<td>0.160</td>
</tr>
<tr>
<td>630</td>
<td>&quot;</td>
<td>21.25</td>
<td>16.02</td>
<td>0.152</td>
</tr>
<tr>
<td>641</td>
<td>&quot;</td>
<td>20.86</td>
<td>15.05</td>
<td>0.141</td>
</tr>
<tr>
<td>649</td>
<td>&quot;</td>
<td>23.23</td>
<td>16.51</td>
<td>0.167</td>
</tr>
<tr>
<td>673</td>
<td>&quot;</td>
<td>22.42</td>
<td>16.96</td>
<td>0.157</td>
</tr>
<tr>
<td>674</td>
<td>&quot;</td>
<td>22.40</td>
<td>15.00</td>
<td>0.149</td>
</tr>
<tr>
<td>684</td>
<td>&quot;</td>
<td>27.32</td>
<td>25.63</td>
<td>0.158</td>
</tr>
<tr>
<td>1</td>
<td>&quot;Open&quot;</td>
<td>20.50</td>
<td>10.00</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>&quot;</td>
<td>19.80</td>
<td>15.00</td>
<td></td>
</tr>
</tbody>
</table>

The light emitting diodes of given design have been tested and are applied in the Institute of Physics of Solids in the Latvian State University, Institute of Nuclear Physics (SDAC, Novosibirsk), Physical Institute (PIAN, Moscow), Institute of Theoretical and Experimental Physics (ITEP, Moscow).
Section III

HIGH-TEMPERATURE THERMAL RESISTORS BASED ON SILICON CARBIDE

The modern industry needs in temperature sensors which are capable to operate under the conditions of high temperature and aggressive media. Such demand, in particular, exist for semiconductor thermal resistors. They gain interest in them first of all due to high sensitivity as compared to the wire-type thermal resistors and thermocouples. Yet, devices based on customary semiconductor materials cannot work under extreme conditions. The most attractive material for this purpose is silicon carbide. It is caused first of all by its thermal and chemical durability and a wide band gap. Moreover, the silicon carbide devices are characterized by a high performance stability. Many laboratories in various countries exert every effort to develop technology of thermal sensors based SiC. It is customary to use for this purpose either single crystals doped with boron or thin films on the insulating substrata prepared by the RF plasma sputtering. Such thermal resistors distinguish by their high stability and relatively low extreme operation temperature [41-44].

In this connection we exert an effort to develop technology to fabricate cheap thermal resistors for a wide temperature range.

3.1. Thermal resistors based on the polycrystalline SiC

The method of synthesis from the separate Si and C sources had been used for fabrication of the polycrystalline SiC. The synthesis proceeded on surface of a graphite substrate. At the synthesis temperature (1800 to 2000 °C) the cubic modification SiC (3C-SiC) was formed, as a rule. To provide uniformity of the SiC layer along the structure depth, we have developed the rig with special design, which provides desired distribution of temperature over the operation space. The SiC layer deposition rate was up to
0.2 to 2.0 mm/h. It was possible to obtain layers up to 10 mm thick. The method productivity is determined by the graphite rig dimensions and, consequently, furnace dimensions. For example, the improved set-up "Редмет-30" during a technological cycle provides fabrication of material for 50 to 100 thousand thermal resistors. The material has the n-type conduction. Investigations of the material structure has shown that the SiC layers directly adjacent to the graphite substrate are, as a rule, fine-grain and in fact textureless. The grain growth direction in regard to the substrate is determined by the flux direction of vapors: Si, SiC, SiC$_2$ and Si$_2$C, and, consequently, it depends on the furnace rig design. With thickening the film a pillar-like shape of grains is manifests explicitly itself. The maximum cross-section size of grains is 250 to 500 μm.

Having grown SiC we removed the graphite substrate by burning it in air. Using the diamond disk saw we cut flat plates with a desired shape. The plate surface was processed using diamond pastes.

To obtain high-temperature thermal resistors with high sensitivity one needs the material conduction to be determined by an impurity with a high energy of thermal activation. For SiC such an impurity is boron ($E_i=0.39$ eV). The diffusion coefficients of impurities in SiC are small. Therefore, in order to dope a single crystal over the entire volume, high temperatures and rather long time are needed. Application of the polycrystalline SiC gives new potentialities for fabrication of thermal resistors. When doping this material, boron easily penetrates in interstitials between microcrystals and diffusion proceeds in fact simultaneously from the entire surface. As a result, the p-n-p structure is formed in every microcrystal. Thus, the thermal resistor consists of a series chain of microcrystals and p-n-p, p-n-p,...,p-n-p structures. Such an arrangement provides high resistivity of material and its high thermal sensitivity. The diffusion procedure was carried out by known technique at the temperature 1800 °C in the argon medium, 100 to 200 tor. A diffusant was the boron oxide.
Fig. 3.1 presents typical I-V characteristics of the thermal resistors. It is seen that they are linear in the small current region. Current increasing leads to increasing the power dissipated by a thermal resistor. Already at \( P > 2 \times 10^{-6} \) W the I-V characteristic deviates from its linear law. It seems that this is caused by self-heating of thermal resistors. In order to investigate what causes this deviation, we measured the I-V characteristics in various media: in oil, in air and in vacuum. As seen from comparison of curves 3 and 4 in Fig. 3.1, sample self-heating takes place indeed. Fig. 3.2 illustrates variation of resistance for a typical sample as a function of the air pressure under the conditions of constant current through a resistor.

\[
\delta R = \frac{R_0 - R}{R_0} \times 100\% \quad (3.1)
\]

\( R_0 \) is the sample resistance at \( I = I_{\text{meas}} \) under the standard conditions. As seen from this Fig. 3.2, the thermal resistors based on SiC may be applied as sensitive sensors of vapor/gas pressure.

Fig. 3.3 presents typical temperature dependencies of the thermal resistor resistance. As it is seen, they are well described by

\[
R = R_0 \times \exp\left( \frac{B}{T} \right) \quad \text{or} \quad R \sim \exp\left( \frac{E_i}{kT} \right) \quad (3.2)
\]

where \( B = 4500 \) and \( E_i = 0.39 \) eV, which corresponds the thermal activation energy of boron.

3.2. Thermal resistors based on bulk crystals of SiC

The thermal resistors have been fabricated by the radiation method of doping the 6H-SiC single-crystals with subsequent processing [45, 46]. Such technology enables to obtain material with high resistivity \((10^{12} \text{ to } 10^{14} \text{ Ohm cm})\), with a high thermal activation energy \((E_i > 0.9 \text{ eV})\), and high sensitivity. Its
Fig. 3.1. Current voltage characteristics of thermal resistors. 1, 2 - samples No 211 and 12 respectively measured in air under the standard conditions; 3 - sample No 284 measured in vacuum oil; 4 - sample No 284 measured in vacuum \( \times 10^{-2} \) torr.
Fig. 3.2. Dependence $\delta R$ vs air pressure at constant current through the thermal resistor for three samples.
Fig. 3.2. Dependence of resistance vs temperature of thermal resistor: 1 - sample No 121; 2 - sample No 213.
The temperature dependence of resistivity is described by expression (3.2) with $B=10000$.

**Main parameters**

**Main Technical Characteristics**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Polycrystalline</th>
<th>Single-Crystalline</th>
</tr>
</thead>
<tbody>
<tr>
<td>Available range of $R_0$ at temperatures, $\degree C$</td>
<td>100 kOhm to 10 MOhm</td>
<td>100 kOhm to 100 MOhm</td>
</tr>
<tr>
<td>Operation temperature range, $\degree C$</td>
<td>20 to 600</td>
<td>200 to 800</td>
</tr>
<tr>
<td>Val. of $B$ in Eqn. (3.1)</td>
<td>4500</td>
<td>10000</td>
</tr>
</tbody>
</table>
Fig. 3.4. a) Arrangement of thermoresistors:
1 - SiC crystal, 2 - Ni contacts.
b) Resistance vs temperature dependencies:
1 - polycrystalline sample; 2 - single-crystalline sample.
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