The Physics of Reliability of Future Electronic Devices

IBM Thomas J. Watson Research Center

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THE PHYSICS OF RELIABILITY OF FUTURE ELECTRONIC DEVICES

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## The Physics of Reliability of Future Electronic Devices

### Title:
THE PHYSICS OF RELIABILITY OF FUTURE ELECTRONIC DEVICES

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### Abstract:
Among the major reliability problems in semiconductor devices are semiconductor surface defects, dielectric breakdown, interdiffusion across interfaces, and corrosion. In the area of surface defects, we have used scanned optical techniques to probe defects and recombination centers on silicon surfaces. Crystallographic defects on silicon surfaces, of the type measured by the scanned optical techniques SIP and SSP are known to promote device failure.
The problem of dielectric breakdown in SiO₂, which is particularly important to the reliability of integrated circuits, was investigated both theoretically and experimentally. A simple theory, based on impact ionization and a subsequent field distortion, is able to explain thickness and electrode dependence of the process, as well as the radiation sensitivity of dielectric breakdown.

Interface reactions were found to be particularly important to the reliability of Schottky barriers. Several classifications of metals have been determined and studied with respect to their behavior in the formation of metal silicide Schottky barrier contacts to silicon. On the basis of the classification of reaction properties, the reaction between the metal and either silicon or SiO₂ can be explained and understood.
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I. SUMMARY OF THE RESEARCH PROGRAM

Interfaces between dissimilar materials have become an important part of modern electronic devices, particularly in the area of microelectronics, where materials such as semiconductors, metals and insulators are combined in order to use their unique properties and to achieve the high performance and small dimensions required in modern technology. The devices that will be important in the next decade are classified into several dominant technologies: silicon (MOS, I^2L, and bipolar), magnetic films and sensors, and superconducting films. Reliability problems in microminiature devices, such as those anticipated for the next decade, are frequently due to interface reactions, surface defects in the semiconductor, surface and grain boundary reactions, charging or breakdown in insulating films, and registration errors in lithography. The overall objective of our present work is to investigate the fundamental properties of interfaces between dissimilar materials that may influence the performance of future electronic devices, and to provide a scientific understanding of reliability problems associated with material interfaces in present devices. We concentrated on generic problems, both present and anticipated, for which an adequate scientific basis was lacking; short term problems peculiar to any one specific device were not emphasized. The program, which was funded under ARPA Contract Nos. F19628-73-C-0006 and F19628-74-C-0077 for thirty-six months, was largely concerned with fundamental problems in MOS and bipolar silicon devices, although a few types of bipolar switches found in amorphous materials were investigated.

II. SIGNIFICANT RELIABILITY PROBLEMS

The significant reliability problems found in silicon devices, which will comprise the major technology for the next decade, can be grouped into a few general areas.

A. Local Defects in the Silicon Surface

1. Surface defects in silicon material cause a reduction both in reliability and in device process yield.
   a. Even a small defect density can cause problems in dense, large area chips.
   b. Dislocations, diffusion pipes, and precipitates can cause a long term degradation.

2. Serious defects can be introduced during processing steps.
   a. Heavy local diffusion or ion implantation produces defects.
   b. Thick oxide layers (oxide isolation) or other insulator layers cause surface stresses and defects.
B. Charging and Breakdown in Dielectric Films

1. Dielectric breakdown is a continuing problem, particularly in the thin (~200Å) gate insulator in MOS devices being developed.
   a. Initial failure due to defects in the SiO₂.
   b. Time delayed breakdown, which may be related to interdiffusion of material.

2. A long-term buildup of charge in an insulating layer is caused by the injection of hot carriers into insulator where they are trapped.

C. Packaging and Corrosion

1. Metal film corrosion due to trace amounts of S or H₂O gas is significant.
2. Electromigration in thin metal films will be a continuing problem in fine line LSI logic circuitry.

D. Lithography

1. Mask registration and defects are a cause of both immediate and long term failure of LSI devices.
2. New technologies involving X-ray or electron beam photore sist exposure produce radiation damage.

E. Degradation of Contact Interfaces

1. Most of the Schottky Barriers involved in LSI silicon technology are subject to degradation due to interdiffusion of materials.
2. Small changes in the gate contact barrier of MOS devices causes both immediate and long term problems in newer technologies using thin gate insulators.

III. ACCOMPLISHMENTS OF THE PROGRAM - Several significant reliability problems were studied in order to provide a practical and scientific basis for handling the problems. Our work was successful in developing new methods of characterization and measurement; in formulating models of failure mechanisms; and in better characterizing materials, particularly insulators. The more significant results of the program are outlined.

A. Local Defects on Silicon Surfaces

1. A new type of scanned surface photovoltage measurement has been developed which will non-destructively detect and image surface defects in silicon at various stages of processing.
   a. The photovoltage system has been applied successfully to the evaluation of silicon starting material.
   b. The resolution has been increased to about 0.8 micron, on the basis of a theoretical understanding which we developed.
2. The scanned internal photoemission technique has been developed and used to find and determine the influence of local sodium and phosphorus deposits on the Si-SiO₂ interface.

B. Dielectric Breakdown in Thin Insulating Films

1. A complete model was formulated to describe the process of dielectric breakdown in SiO₂ thin films. This model is the first to describe consistently all of the available data.
   a. The model predicts a significant increase in dielectric strength for films less than 300Å thick.
   b. Dielectric breakdown induced by a small amount of ionizing radiation was found and explained in terms of the theory.

2. The uniformity and dielectric properties of thin SiO₂ films were improved by compaction of the film with a careful, controlled amount of ion irradiation, with a subsequent anneal step.

3. The electronic structures of SiO₂ and Si₃N₄, pieced together on the basis of ESCA, photoemission, and reflectivity measurements, provide a basis for understanding dielectric breakdown and other events in these materials.

C. Fundamentals and Degradation of the MNOS Transistor

1. The energy level responsible for charge or information storage was measured for the first time with the newly developed technique of photocapacitance spectroscopy.

2. Both the Si₃N₄ and the SiO₂ have been characterized by several optical and photoemission techniques.

3. A type of degradation of MNOS structures, observed directly by the scanned surface photovoltage technique, appears to be due to a buildup of nonlocal charge in the double insulator.

D. Schottky Barriers on Silicon

1. A class of metals was found which will decompose a thin layer of SiO₂ on silicon at high temperatures to form a silicon-silicide Schottky barrier.
   a. A new technique of using an ion implanted Xe marker was used to study interdiffusion at the interface.
   b. Metals with a high heat of formation for both the silicide and the oxide phase were found to decompose SiO₂ at low temperatures. Integrated circuits using these metals are expected to exhibit long term failure.

E. Interface Polarization and Degradation

1. The interdiffusion of a metal into SiO₂ was studied by the new technique of field dependent photoemission. The resulting polarization layer can shift the operating point of an MOS device by several tenths of a volt over the life of the device (for particular gate metallizations such as gold, platinum, and others).
2. A small amount of hydrogen was found to produce an atomic polarization layer between SiO₂ and a catalytic metal such as platinum or palladium. This layer can shift the operating point of an MOS transistor by as much as 0.5 volt for certain gate metals. However, aluminum gates were found to be insensitive to hydrogen.

F. Metal - Glass Interaction

1. Phosphorus, a common dopant in glass, was found to crystallize glass at temperatures below those used in device processing. This reaction can cause a serious mode of failure in MOS circuits in which phosphorus is used to dope the SiO₂ gate insulator.

2. The adhesion between certain metals and glass was found to be greatly enhanced by the addition of a small amount of free silicon to the metal. This allows the use of gold on SiO₂, a combination which is normally not used because of poor adhesion.
II. DIELECTRIC INSTABILITY AND BREAKDOWN IN SiO₂ THIN FILMS* 

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ABSTRACT

Dielectric instability and breakdown in SiO₂ have been well characterized by many experimental techniques, including measurements of thickness dependence, contact barrier dependence, and time dependence of breakdown as well as determination of radiation sensitivity and pre-breakdown charge buildup within the insulator. All of the various types of data can be explained consistently by an impact ionization model which predicts a negative resistance type of instability; electrons are injected from the cathode, the electron distribution is heated, hot electrons ionize the lattice, and the residual positive charge distorts the electric field to further increase impact ionization. The model is sensitive to two key parameters, the ionization bandgap \( E_g \) and the electron-phonon scattering length \( \lambda \).

* Work supported in part by the Defense Advance Research Projects Agency and monitored by AFCRL under Contract No. F19628-74-C-0077.
I. INTRODUCTION

Although dielectric breakdown has been studied extensively for several decades\textsuperscript{1-4}, the events which occur during the incipient breakdown process in insulators are not completely understood. In the development of semiconductor technology, dielectric instability and breakdown in insulators such as SiO\textsubscript{2} have become increasingly significant in limiting both performance and reliability. Because of the importance of insulating thin films of SiO\textsubscript{2} in the integrated circuit industry, and because of the new information available on the properties of this material, it is fruitful to reexamine the physical processes which occur at high electric fields in order to gain insight into the possible intrinsic breakdown mechanisms. Reliable data is available on the dependence of breakdown upon insulator thickness, electrode work function,\textsuperscript{5} and ionizing radiation\textsuperscript{6} as well as on several material parameters of SiO\textsubscript{2} including ionization rate,\textsuperscript{7} effective hole mobility,\textsuperscript{8} and recombination cross section.\textsuperscript{9} The problem of sorting out the details of the breakdown process in SiO\textsubscript{2} is considerably easier than in the traditional alkali halides due to the higher development of materials technology as well as the wealth of data on electronic properties which has become available for SiO\textsubscript{2}. The fundamental phenomena which occur during intrinsic dielectric breakdown in thin films of SiO\textsubscript{2} can be explained and understood in a way that is consistent with all available data.

Historically, dielectric breakdown in insulators was first seen as an electronic avalanche process,\textsuperscript{2, 3, 10-13} in which the field is high enough that one initial electron produces a destructively large cascade of ionizing events. The avalanche theories were able to explain the relatively
meager data available at the time, largely on the breakdown strength of alkali halide crystals. However, the significance of the phenomenon of field distortion during breakdown was not appreciated until the proposal of a model by O'Dwyer who included the effect of the hole charge produced during the initial stages of an avalanche. O'Dwyer was able to predict a negative resistance type of instability in which electrons are injected from a cathode electrode by a field enhanced process, a portion of the electrons are accelerated sufficiently to produce impact ionization, and the slow moving holes left behind cause an enhancement of the cathode field which leads to an increased injection current, and so on, ad infinitum. The negative resistance in this model results from the nonlocal feedback produced by the drift of holes to the cathode. The O'Dwyer model resolves several deficiencies of the simple avalanche theories in that it explains why a low rate of avalanche multiplication is measured in insulators near the breakdown field and it predicts a negative resistance type of instability which is impossible in a simple avalanche model. Further refinements in the O'Dwyer model were made to account for the local details in the initial stages of breakdown. The O'Dwyer model characterizes breakdown quite well in semiconductors and in selected insulators, but it cannot be applied directly to insulators such as SiO₂ in which hole mobility is extremely small; holes produced by impact ionization in SiO₂ are removed predominantly by recombination in the insulator leading to dielectric breakdown.

We have proposed and explored a simple model based on impact ionization and field distortion that is appropriate to describe breakdown in
$\text{SiO}_2$, in which holes are relatively immobile. All of the physically significant processes are shown schematically in Fig. 1. Electrons are injected by field enhanced emission from a cathode, accelerated in the electric field, and scattered by phonon emission. A small fraction of these electrons create hole-electron pairs when they reach the ionization energy at approximately twice the bandgap $E_g = 9.0$ eV$^{23}$, leaving behind a cloud of relatively slow moving positive charge.$^{24}$ Holes are removed largely by recombination, with hole drift apparently playing only a minor role in the breakdown event. The net positive charge cloud leads to a larger current of more energetic electrons. As result, a negative resistance instability develops; in this case a position dependent ionization rate is necessary to provide the nonlocal feedback essential for negative resistance. The two principal differences between the proposed theory and that of O'Dwyer$^{24}$ are the dominance of recombination in removing holes from the charge cloud and the inclusion of a non-local rate of impact ionization. Although physical reality lies somewhere between the two extremes of hole annihilation by recombination and removal by drift, we find that the model based on recombination best describes the breakdown process in $\text{SiO}_2$. The impact ionization-recombination model$^{21,22,25}$, based on the most recent experimentally determined material parameters, is able to explain thickness, electrode, and radiation dependence of dielectric breakdown in thin films of $\text{SiO}_2$. The model is sensitive to only two key parameters, the bandgap of the insulator and the electron-phonon scattering length. Hole mobility is found to be an insignificant factor in the calculated dielectric strength for mobilities less than about $10^{-8}$ cm$^2$/V-sec.
II. IMPACT IONIZATION - RECOMBINATION MODEL

A model calculation based on steady state impact ionization and recombination is used to determine the J-V characteristic of a thin film of SiO₂. Dielectric breakdown occurs at the critical point at the onset of negative resistance, where \( dV/dJ = 0 \). Electron current is injected from the cathode electrode into the SiO₂ by the strongly field dependent process of Fowler-Nordheim tunneling.\(^{26}\) The injected electrons gain energy from the field and lose it to the lattice, primarily by phonon emission. In developing a nonlocal rate of impact ionization, it is assumed that a one dimensional projection of the electron-phonon scattering events is a Poisson process in which the probability of emitting a phonon is \((1/\lambda)\) per unit length in the X direction. The \( \lambda \) in this model is assumed to be energy independent.\(^{27}\) In each phonon emission event, the electron loses a quantum of energy which is assumed to be that of an LO phonon for which \( \hbar \omega = 0.153 \text{ eV} \).\(^{28}\)

Based on the Poisson model, the electron energy distribution is found as a function of distance from the cathode. This distribution is then obtained in terms of a single adjustable parameter \( \lambda \). The probability that a given electron will have energy \( \epsilon = -n\hbar \omega \) as it passes the point \( x \) is \( P \), where

\[
P(\epsilon, x) = \frac{1}{(-\epsilon/\hbar \omega)^{1/2}} \left( \frac{x}{\lambda} \right)^{-\epsilon/\hbar \omega} e^{-x/\lambda}.
\]

By the central limit theorem, the energy distribution is approximately a Gaussian,

\[
D(\epsilon, x) \sim \frac{(1/\hbar \omega)}{[2\pi(x/\lambda)]^{1/2}} \exp - \left\{ \frac{[(\hbar \omega x/\lambda) - \epsilon]^2}{(\hbar \omega)^2 (2x/\lambda)} \right\}.
\]
We allow the electron to ionize the lattice as soon as it attains an energy of $\epsilon_i$ above the conduction band bottom; the rate of impact ionization is then
\[
\frac{\partial \rho_+}{\partial t} \bigg|_i = \tau \frac{\partial}{\partial x} \int_0^\phi(x)+\epsilon_i D(\epsilon,x) d\epsilon,
\]
where $\phi(x)$ is the energy at the bottom of the conduction band. In equilibrium, the rate of generation of holes equals that for hole recombination,
\[
-\frac{\partial \rho_+}{\partial t} \bigg|_i = \frac{J \rho_+}{e} \int_0^\phi(x) \sigma(\epsilon) D(\epsilon,x) d\epsilon,
\]
where $\sigma(\epsilon)$ is the recombination cross section. The equilibrium charge density is independent of the injected current $J$,
\[
\rho_+ = \frac{e}{\langle \sigma \rangle} \frac{\partial}{\partial x} \int_0^\phi(x)+\epsilon_i D(\epsilon,x) d\epsilon,
\]
where $\langle \sigma \rangle$ is the weighted average recombination cross section. The potential $\phi(x)$ is then found by a double numerical integration of Poisson's equation with an initial cathode field (or injected current) as a boundary condition.

The material parameters used in the calculation for SiO$_2$ are the phonon energy $\hbar \omega = .153$eV, $\epsilon = 3.8$, and the bandgap $\epsilon_g = 9$eV. There is considerable range of recombination cross section$^{29-33}$ found in SiO$_2$, the most recent evidence$^{33}$ indicates that a coulomb capture cross section is reasonable for the case of electron recombination with a trapped hole. The model calculations were based on a cross section$^{34}$.
\[ \sigma(e) = \frac{1}{\pi} \left( \frac{3e^2}{16\varepsilon_o E} \right)^2, \]

which is \(2.5 \times 10^{-13}\text{ cm}^2\) at the electron saturation velocity in SiO\(_2\). This \(\sigma(e)\), which is considerably larger than the direct radiative cross section used by the authors in a previous calculation,\(^{22}\) provides a better fit to the available experimental data. A further refinement in the estimate of \(\sigma(e)\), to account for the effect of the electric field upon coulomb capture, may allow an even better fit. Fortunately, the model calculation is quite insensitive to details of the recombination coefficient, so that the predicted breakdown behavior is little influenced by the choice of capture cross section.

The electron-phonon scattering length \(\lambda\) is a critical parameter of the model; the calculated dielectric strength varies inversely with small changes in \(\lambda\).\(^{22}\) The value \(\lambda = 1.74\text{ Å}\) was chosen to fix the calculated breakdown voltage to \(V_b = 92.5\text{ V}\) for a 1000 Å thick film, in agreement with the available experimental data.\(^{35-39}\) Then, the voltage at which dielectric breakdown takes place for a given thickness of SiO\(_2\) is found from the critical point at which \(dV/dJ = 0\).

III. RESULTS

The ionization-recombination model was used to calculate \(J-V\) relationships and dielectric strength for a range of film thickness and contact barriers. The results for \(\lambda = 1.74\text{ Å}\) are compared with the experimental data presently available. Also, the results of an O'Dwyer type of model
calculation are given for a reasonable choice of parameters; the mobility product \(u_+u_-\) was set to fix the breakdown voltage \(V_{10} = 92.5\text{V}\) at 1000 \(\text{A}\).

The thickness dependence of the measured dielectric strength is shown in Fig. 2, along with that calculated on the basis of our ionization-recombination model (solid curve) and O'Dwyer's model (dashed curve). The sharp rise in the experimentally determined breakdown strength \(^{36,38}\) for thickness below \(\approx 400\ \text{Å}\) is well explained by the ionization-recombination model. The reason for this increase is the nonlocal ionization rate necessary in the ionization-recombination model; no ionization is possible for a region within \(x = (\varepsilon_1 + \phi)/E\) of the cathode. Since our model predicts no intrinsic breakdown for voltages less than \((\varepsilon_1 + \phi)\) or \(\approx 12\text{ volts}\), the dielectric strength increases for small thickness as the breakdown voltage asymptotically approaches \(\approx 12\text{ V}\) at an infinitesimally small thickness. On the other hand, the O'Dwyer type of model shows only a logarithmic dependence upon thickness, in part reflecting the use of a local ionization rate.

Presumably, the use of nonlocal ionization would introduce a stronger thickness dependence into the model.

The experimentally determined breakdown strength \(^5\) shows no systematic dependence upon the cathode contact barrier \(\phi\) as seen in Fig. 3. Here, the experimental uncertainty in each point is of the order of \(\pm 0.5 \times 10^6\ \text{V/cm}\). The calculated dielectric strength based on our ionization-recombination model shows a very weak dependence on the barrier \(\phi\), in agreement with the general trend of the experimental findings. In contrast, the calculation based on an O'Dwyer type of model shows a strong dependence on \(\phi\) at the injecting contact. This difference in the barrier dependence of the two
theories results directly from the mechanism of clearing holes from the insulator. Based on the data available, it seems that recombination is the dominant mechanism in SiO₂. More definitive measurements of the barrier height dependence should enable a better determination of the role of hole mobility in the breakdown process.

Although electron-hole recombination appears to be the dominant mechanism of clearing holes from the SiO₂, hole motion may play a part in the breakdown process. In order to determine the influence of hole motion on our ionization-recombination model, the calculation was extended to allow holes to drift toward the cathode before recombining with injected electrons. The nonlocal ionization rate was maintained, with λ = 1.74 Å. The results, shown in Fig. 4, show that dielectric strength is relatively unaffected by hole mobility up to \( \nu 10^{-8} \text{ cm}^2/\text{V-sec} \). For the range of hole mobility reported for SiO₂, the breakdown process and dielectric strength are little influenced by hole motion.

Based on the ionization-recombination model of the breakdown process in SiO₂, we would expect a stable charge cloud to form at fields below that at which breakdown takes place. Recently Shatzkes and Avron have been able to measure this charge Q and its centroid \( \bar{X} \); they used a combination of field emission and C-V measurements, performed at low field, to determine the remnant charge left from a previous high field pulse. The results are shown in Figures 5 and 6, for SiO₂ films of thickness 263 Å and 493 Å. The charge Q and centroid \( \bar{X} \) calculated on the basis of the ionization-recombination model for 1000 Å of SiO₂ are shown as the solid curves in the two figures. The reasonably good overall agreement with experiment indicates
the existence of a stable charge cloud which behaves in a way predicted by the ionization-recombination theory.

IV. DISCUSSION

The impact ionization-recombination model for dielectric breakdown seem to be in reasonably good agreement with available data for SiO₂. From the model, a picture can be formulated of the significant processes which occur during breakdown. As shown in Fig. 7 the electrons are field emitted into the SiO₂ from the cathode electrode. The hottest of the electrons ionize the lattice, leaving behind slow moving holes. A cloud of positive charge formed by ionization and recombination leads to an increased current of hotter electrons. The result is that a negative resistance type of instability develops in which a local charge cloud leads to an enhanced current. The small filament of current drains charge from the capacitor in a destructive breakdown event.

A nonlocal ionization rate is an essential feature of any model of dielectric breakdown in very thin insulating films. The influence of the nonlocal ionization rate is manifested in the rapid increase of the breakdown field seen in SiO₂ as the film thickness is reduced below about 400 Å. It is hardly surprising that a nonlocal ionization rate is important in films of a total thickness only several times greater than the distance required to accelerate a free electron up to the ionization energy.

During the electronic breakdown process, the experimental evidence indicates that recombination is the most important mode of removing holes produced by impact ionization. Hole drift to the cathode plays a relatively
minor role in determining the conditions leading to dielectric breakdown. One piece of experimental evidence, the dependence of dielectric strength on the cathode contact barrier, shows no systematic variation which would be expected if drift were the dominant mechanism of clearing holes from the insulator. Further, an impact ionization model calculation which includes both hole recombination and hole drift shows no appreciable influence on the dielectric strength for hole mobility $\mu < 10^{-8}$ cm$^2$/V-sec. Although hole mobility does not appear to have a significant effect on dielectric strength, it is thought to play a role in dielectric instability, particularly that induced by ionizing radiation of various sorts.
V. ACKNOWLEDGEMENT

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REFERENCES


24. It can be shown that the O'Dwyer model can be applied to the case of SiO₂ if the model parameters are appropriately chosen. In a previous article, the authors (ref.22) were unable to find a negative resistance for the case of an infinitesimally small ionization rate because of an improper choice of cathode condition.


31. D. J. DiMaria, PhD thesis (Lehigh University, 1973)


40. The parameters used in the O'Dwyer model calculation are $H = 1.8 \times 10^8$ V/cm, $b = 2.2 \times 10^{19}$ sec$^{-1}$ (from P. Solomon, ref. 7); and J(E) for Fowler-Nordheim tunneling from Si into SiO$_2$ (ref. 26). The mobility product $\mu_u \mu_-$ was determined by setting the breakdown voltage to $V_b = 92.5$ V at 1000 Å film thickness.
41. Details of this calculation are to be published elsewhere.
42. M. Shatzkes and M. Av-ron, to be published.
Figure 1. The physically important processes which occur in a thin film of SiO$_2$ at a high electric field. Electrons are injected by a field enhanced process at the cathode electrode. Hot electrons produce holes which induce a distortion of the electric field. Holes are removed predominantly by recombination and by drift to the cathode.
Figure 2. Dielectric strength as a function of thickness of an SiO₂ layer. The solid curve shows the results of our impact ionization-recombination theory for $\lambda = 174 \text{ Å}$. The dashed curve is the calculated dielectric strength based on the O'Dwyer model, for a reasonable set of parameters (ref. 40). The best available experimental data on intrinsic breakdown is represented by the points: ■ Si(p)-SiO₂-Al; • Si(n)-SiO₂-Al (ref. 36); OSi(p)-SiO₂-Al (ref. 36); X K. Hamano (ref. 38).
Figure 3. Dielectric strength as a function of cathode contact barrier $\Phi$.

The points shown for several metals, determined experimentally by Osburn and Weitzman (ref. 5), are only approximate.
Figure 4. Dielectric strength calculated as a function of hole mobility for a model incorporating both recombination and hole mobility as mechanisms of clearing holes from the SiO$_2$. 

$\overline{E}_b (10^6 \text{ V/cm})$

$\mu (\text{cm}^2/\text{V-sec})$

SiO$_2$

1000 Å
Figure 5. The total remnant charge density $Q$ produced in the positive charge cloud in $\text{SiO}_2$ by a cathode field $E_c$. The solid curve is the calculated charge density in the ionization-recombination model with $\lambda = 174 \, \text{Å}$. The experimental points are from Shatzkes and Av-ron (ref. 42).
Figure 6. The centroid $\bar{x}$ of the charge cloud left in SiO$_2$ by a high electric field as a function of the electrode field $E_c$. The solid curve is the centroid calculated on the basis of the ionization-recombination model with $\lambda = 1.74$ Å. The experimental points are from Shatzkes and Av-ron (ref. 42).
Figure 7. Schematic representation of the processes which occur during the critical stages of a dielectric breakdown in SiO$_2$. 
III. PHOTOEMISSION AND PHOTOVOLTAIC IMAGING OF SEMICONDUCTOR SURFACES*

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ABSTRACT: The new electro-optical techniques of scanned internal photoemission and scanned surface photovoltage can be used to obtain images of non-uniformities and defects at semiconductor surfaces or interfaces. The spatial resolution of either method is primarily determined by the wavelength of the laser used as a collimated light source. The images obtained by the photoemission technique are replicas of the local contact barrier at the interface between two materials. Typically, photoemission images show details due to variations of stoichiometry or local concentration of impurities. The photovoltage technique, on the other hand displays recombination centers such as precipitates, dislocation, stacking faults, cracks, and gross damage near a semiconductor surface. Each of these electro-optical imaging techniques provides information about interfaces that was previously inaccessible to observation.

* Work supported in part by the Defence Advance Research Projects Agency and monitored by AFCRL under contract number F 19628-74-C-0077.
Introduction

Focused beams of various types of ionizing radiation have been used in the past to probe semiconductor surfaces and devices, generally in the region near a p-n junction. The signal induced in a junction by a scanned optical beam has been found to provide information on resistivity fluctuations in the material, on gross defects in the device, and on surface irregularities or inversion layers near the junction. Generally, in this type of measurement, a beam of electrons or photons is scanned across an area of a semiconductor to produce hole-electron pairs in the material. An irregularity in the material will produce a fluctuation in a sample current of some sort detected when the beam scans across the defect. In the standard configuration, a p-n junction collects current produced within a diffusion length of the junction. Any change in either the generation or recombination rate near the junction is detected as a fluctuation of the collected current. The beam induced current is displayed on a rastered CRT to produce an image of the defects and inhomogeneities. Imaging techniques of the electron beam induced current (EBIC) or photon induced current type are useful in identifying defects in individual semiconductor devices, but they cannot easily be applied to the study of semiconductor surfaces or interfaces on which no p-n junction exists. The necessity of using a p-n junction to collect the beam induced current makes non-destructive evaluation of material surfaces at various stages of processing virtually impossible by the beam induced current techniques.

Recently, several scanned beam techniques, including scanned internal photemission (SIP), scanned surface photovoltage (SSP), and the
corresponding electron beam analogs \(^{27,28}\) have been developed to probe free semiconductor surfaces. These new techniques provide images of defects near surfaces or interfaces without the necessity of a p-n junction to detect the signal current. As a result, they can be used to examine non-destructively the defects which occur on semiconductors at various stages of material processing. Each of the techniques has characteristic advantages and capabilities. The e-beam techniques provide the best resolution, better than about 1000 Å, \(^{27}\) but they are complicated by beam induced charging of any insulating film on the surface. \(^{28}\) On the other hand, the optical techniques have a resolution capability of on the order of one micron, but they do not charge the surface insulating or passivating film. Hence, the optical techniques can also be used to examine charge non-uniformity in the insulating film on the surface without disturbing the system. Additionally, the optical scanning apparatus can be made to be cleaner than existing e-beam systems because it does not require a complex vacuum system.

Of the optical techniques, SIP determines small inhomogeneities in the contact barrier \(\Phi\) between two materials, one of which is typically an insulator such as SiO\(_2\). The SIP technique is useful in imaging changes in \(\Phi\) which are due to contamination, interface chemical reaction, impurity segregation at the interface, or surface crystallographic defects. In contrast, the SSP technique is sensitive to local recombination centers that are within about one micron of the surface and to charge inhomogeneities in the insulating overlayer. Typical recombination centers seen in SSP images include crystallographic defects, precipitates, and gross surface damage; these defects are usually introduced during growth, cutting,
polishing, and high temperature processing of the semiconductor material. The SSP technique has a unique advantage in that no direct contact to the sample is required if the photovoltage measurement is made by capacitive coupling. Since physical contact is unnecessary, the SSP technique is clean and completely non-destructive.

**Optical Scanning System**

The instrumentation for the two scanned optical techniques is quite similar in that a small spot is rastered on the sample surface while a signal, either photoemission or photovoltage, is detected and displayed on a CRT. A cw laser is used for the light source in order to obtain a measureable signal from a small, nearly diffraction limited light spot. Reasonable signals were obtained with 1.0 mW of light incident on the sample, without any noticeable thermal damage to the sample at the $10^5$ W/cm$^2$ power density at the focused spot. The spot was swept by the sawtooth motion of a galvanometer driven mirror, which was scanned at a 20 Hz rate. In the other direction, the beam was swept slowly in order to complete a 600 line single frame in 30 seconds. The frame rate, which is limited by the rate of the fast scan mirror, could easily be increased an order of magnitude by increasing the fast scan rate to 200 Hz. A further increase, at the expense of a reduced spacial resolution, could be obtained with an acoustic deflection system for the fast scan direction. The total resolution capability of the optical scanning technique is limited by three equally important factors: the divergence of the light output from the laser, the diffraction limit of the objective lens, and the bandwidth of
of the detector. At best, the spacial resolution (of the SIP measurement) is estimated to be about 0.43 microns, with the dominant factors being lens diffraction at $\lambda = 3250$ Å and laser divergence.

The simple optical system shown in Fig. 1 was used to scan a light spot over a square area of the sample surface. Radiation from a laser source was focused onto a 50 micron pinhole in order to eliminate the light that diverged by more than 0.5 mrad. The laser source was chosen to be either an RCA He-Cd laser operating with 3mW at 3250 Å or a Spectra Physics He-Ne laser with 5 mW at 6328 Å. Light emergent from the pinhole was projected a distance of one meter onto a series of two front surface mirrors, which rotate through angles $\theta$ and $\phi$. Light from the two scanned mirrors completely filled the aperture of the objective lens for the full range of the scanned angles. Then, the focused spot of light from the objective lens scanned an area of the sample as the two orthogonal mirrors were rotated. The size of the scanned area, and to some extent, the resolution of the system, were determined by the objective lens. The focal length of the lenses used in the measurements range from 0.54 cm to 20 cm. Reflecting objective lenses were used for the high resolution measurements so that correction of the optics for the wavelength of each laser was unnecessary.

The fast scan mirror was driven at about 20 Hz by a rotary galvanometer, G-306, available from General Scanning, Inc. Care was taken to balance the mirror and rotor in order to minimize parasitic oscillation of the mirror assembly. This parasitic oscillation or vibration causes deflection of the light perpendicular to the direction being scanned by that particular mirror. Although faster galvanometer scanners are available, the relatively
conservative rate of 20 Hz was chosen to minimize the problems caused by
vibration and to optimize resolution.

The problem of positioning and focusing the sample, something which is
particularly difficult in the case of UV light, was solved by detecting
light reflected back into the system from the sample. Initially, light
from the pinhole goes through the scan system which focuses it onto the
sample. When the system is in focus, the light reflected from the sample
is focused by the system back onto the pinhole, independent of the mirror
positions $\theta$ and $\phi$. However, if the system is out of focus, the reflected
light falls outside the pinhole where it strikes a fluorescent screen. The
system is brought into focus by adjusting the sample position along the
optical axis until all of the reflected light is refocused back onto the
pinhole.

In order to locate the sample in the beam and to examine the surface
conditions during the measurements, a reflectivity image of the sample can
be obtained and displayed in the coordinate frame of the CRT. A portion
of the light reflected from the sample is directed onto a photocell by a
beamsplitter. The light falling on the photocell comes to focus at a point
which is independent of the same angles $\theta$ and $\phi$. Current from the photo-
cell, when displayed as Z-axis modulation of a rastered CRT, forms an
image of the sample at the wavelength of the laser source and in the same
coordinate frame as the SSP or the SIP image of the same area.

Both the photoemission and the photovoltage signals are detected and
displayed on a CRT screen by the circuitry shown schematically in Fig. 2.
The signal current is detected by a Keithley 18000 picoammeter which has
been modified for a fast response; the modification involves a reduction of the feedback capacitance in each of the two amplifier sections, while maintaining the stability of the overall system. The signal in the SIP measurement is the photocurrent, while the signal in the SSP measurement is the capacitive current induced in the probe electrode by the surface photovoltage. The amplitude and dc level of the signal are adjusted and applied to the Z-axis of the CRT, along with a retrace blanking pulse derived from the fast scanner. The X and Y axes of the CRT are driven by voltages proportional to those directly across the drive motor armatures. Images of the Z-axis modulated display of the SSP or the SIP signal are recorded photographically.

The measurement apparatus, shown in Fig. 3, includes the scan mirrors, the galvanometer motors, the reflecting microscope objective, the sample positioning stage, and the picoammeter. Light, incident on the mirrors from the right, is deflected onto the two orthogonally mounted scanning mirrors by a stationary positioning mirror. From the fast scanned mirror, the beam enters the reflecting objective lens (Beck 36/.5) which focuses the light onto the sample. Because of the curvature of the focal plane of this objective, the overall resolution is noticeably degraded for beam deflections greater than ±0.5°. In order to extend the useable field of view, it is necessary to use a refractive plano-objective which is corrected for flatness of field at the laser wavelength.

The sample is held in position and focused by an X-Y-Z adjustable hot stage. Signals from the sample are detected by the picoammeter located behind the hot stage in Fig. 3. The entire assembly is secured to a two
Scanned Internal Photoemission

Scanned internal photoemission (SIP) is a relatively recent technique for probing interfaces with a beam of light to produce electronically an image or replica of the interface contact barrier. Typical images display a modulation of the internal photocurrent induced by the scanned beam which is due to a lateral inhomogeneity of the contact barrier. The technique is somewhat analogous to scanning electron microscopy (SEM), except that in this case a light beam is used to excite electron emission into a dielectric instead of into vacuum. As in the SEM technique, the emitted current is displayed on the Z-axis of a CRT.

The mechanism of SIP is internal photoemission on a local scale, as shown in Fig. 4. A scanned light beam of energy $\hbar \omega$ penetrates a transparent or semi-transparent electrode and is absorbed in the opposite electrode where it excites electrons into the conduction band. Some of the electrons have sufficient momentum normal to the interface so that they are able to surmount the barrier $\phi$ and reach the opposite electrode. Near threshold, the quantum yield for photoemission is

$$\psi = \gamma (\hbar \omega - \phi)^\alpha,$$

where $\gamma$ is a constant and $\alpha$ is an empirically determined power, typically in the range 2 to 3. A small percentage reduction in $\phi$ leads to a relatively larger increase in yield,

$$d[\ln \psi] = \frac{\alpha \phi}{(\hbar \omega - \phi)} \, d[\ln \phi].$$
The sensitivity of the SIP technique depends upon the rapid increase of photocurrent produced by a small decrease of $\phi$ below the photon energy $\hbar \omega$. For a reasonable image contrast, the laser and any Schottky barrier reduction are chosen so that $\hbar \omega$ is several tenths of an eV above $\phi$.

The Si-SiO$_2$ interface is an interesting candidate for examination by SIP imaging, as can be seen in Figs. 5 and 6. Here, the Si-SiO$_2$ interfaces have been uniformly coated with about $4 \times 10^{12}$ Na/cm$^2$ by electrodeposition in order to reduce the interface barrier$^{31}$ from 4.25 eV down to a level that is accessible to the He-Cd laser operating at 3.81 eV. Actually, the sodium acts as a "staining agent," increasing the local photocurrent in those places where ions accumulate on the interface. Since the sodium coverage is influenced by defects, it was found to enhance the SIP contrast around defects. Fig. 5 shows an interesting pattern of dark spots seen in the emission from an Si-Si$_2$ interface grown on $\sim 10^{20}$ P/cm$^3$ doped (100) Si. These spots, which are about 10 microns in diameter, have been identified as phosphorous that has segregated onto the Si-SiO$_2$ interface during high temperature growth$^{32,33}$ of the 1000 Å thick SiO$_2$. Apparently, sodium that does reach the Si-SiO$_2$ interface is rendered ineffective by chemical combination with the phosphorous lying within a few atomic layers of the interface. These phosphorus islands, observed for the first time by the SIP imaging technique, were found to be correlated with the phosphorous doping density in the silicon substrate. Another type of defect on the Si-SiO$_2$ interface is seen in Fig. 7 as a line, about 30 microns long, with bright spots at either end. The defect, which lies along the $<110>$ crystallographic direction, is thought to be a stacking fault that has been decorated
by sodium ions. The contrast of the stacking fault is enhanced by the sodium decoration.

The SIP technique was also used to study metal-insulator interfaces such as Nb₂O₅–Bi, which is of interest because of the fast switching found in this structure. The SIP image in Fig. 7 represents photocurrent emitted from a 200 Å Bi electrode into Nb₂O₅. The light source was a He-Ne laser at 1.96 eV, which is close to threshold where the technique is most sensitive. In this system, the contrast mechanism is not simple because of the large number of traps in the Nb₂O₅.

**Scanned Surface Photovoltage**

Photovoltage images show defects that cause minority carrier recombination on a semiconductor surface. The SSP images have also been found to show any non-uniform charge in the passivating layer on a semiconductor. An SSP image is formed by displaying the first or second derivative of the photovoltage induced by a scanned beam of light on a semiconducting surface. The travelling light spot produces a comet shaped cloud of minority carriers which are captured on the slightly depleted surface, as represented in Fig. 8. Carriers from the moving cloud cover the surface to produce a photovoltage. When the beam passes over a defect, some of the charge cloud is lost to recombination, leading to a dip in the photovoltage. This dip is detected by a capacitively coupled electrode near the surface, and displayed on a CRT. Since the charge cloud is quite large, on the order of the diffusion length, the resolution of images produced directly from the photovoltage is rather poor. The high resolution of the SSP technique is due...
to the use of the first or second derivative of the photovoltage to form
the image. The differentiation accentuates the signal in the region around
the light spot, where the charge cloud density has a logarithmic singularity.
In practice, the first derivative is obtained directly by measuring the
capacitive current induced by fluctuations of the surface photovoltage.

By using conduction of minority carriers along the slightly depleted
surface, it is possible to obtain SSP images at some distance from the
electrode. The local photovoltage is coupled to the entire surface by the
slightly conducting surface. In some cases, it is necessary to enhance the
surface conductivity by generating extra minority carriers by a dc blanket
illumination. The resolution of remotely detected images was found to be
reduced if the surface conductivity is too low. Also, in order to develop
a photovoltage signal, the surface must be depleted, either by a blanket
electrode (the electrode can be an aqueous solution, as was shown by Lile
and Davis\textsuperscript{26}), or by a surface treatment such as that produced by dry oxida-
tion of p-type silicon.

The signal $S'$ is proportional to a convolution of the excess local
recombination rate $R$ with an experimental sampling function, which is
approximately equal to the $x$-derivative of the charge cloud density $n$.
Mathematically,

\[
S'(x_0,y_0) \sim \frac{Gy_c \delta}{\epsilon_b \tau_0 t} \int \int dx dy \frac{dn(x-x_0,y-y_0)}{dx} R(x,y),
\]

where $G$ is the total generation rate, $\delta$ is the effective depletion width,
$\epsilon_b$ is for the dielectric, and $\tau_0$ is the surface minority carrier life-
time. Ideally, $dn/dx$ would be a singly differentiated Dirac delta function.
and the signal $S'$ would then be proportional to $dR/dx$. Because of the finite size of the light spot, however, the resolution is reduced. In a coordinate system centered on the moving spot, a modified diffusion equation\(^{25}\) is solved to obtain the charge density $n$ in the region around the scanned spot. The charge density is

$$n = A \exp \left\{ - \frac{Vx}{2D} \right\} K_0(\kappa r),$$

where

$$\kappa = \frac{1}{v \sqrt{D \tau_0}} \left\{ 1 + \frac{2 \tau_0}{4D} \right\}^{1/2},$$

and where $K_0$ is a modified Bessel function of the second kind and $D$ is the surface diffusion coefficient. The sampling function $dn/dx$ is plotted in Fig. 9 for several scan velocities, with the reasonable parameters $D = 15 \text{ cm}^2/\text{sec}$ and $\tau_0 = 0.01 \text{ sec}$. The form of $dn/dx$ deviates little from $1/r$ near the light center at $r = 0$. Based on this simple theory, the resolution at $1/e$ is approximately 2.7 times the light spot diameter for singly differentiated SSP and nearly 1.0 for the doubly differentiated technique.

Several examples of SSP images, obtained from (100) silicon surfaces, display the resolution, sensitivity, and comparative advantages of this relatively simple technique. An SSP image of several clusters of emergent dislocations which penetrate the (100) surface of a sample are shown in Fig. 10. The dislocations were produced by mechanically damaging the reverse side of a silicon wafer, and then annealing it in dry He at 800°C for 30 min. It is known that such damage will cause dislocations to propagate through to the front surface during high temperature processing.
Surface defects were also found in SSP images of nominally defect free silicon, an example of which is shown in Fig. 11. Here, a semitransparent blanket electrode was used in order to maximize resolution. The image shows scratches, some as long as 1000 microns, which are not parallel to a low order crystallographic direction. Presumably, these scratches were introduced during the polishing process. Another type of defect, commonly seen in device grade silicon, occurs throughout the area of Fig. 12. These small spots are thought to be crystallographic defects associated with precipitates of oxide or other foreign material. The density of these defects was found to be correlated with oxygen concentration, suggesting that the defect may be associated with an oxide precipitate. The density was found to be as high as $10^5$/cm$^2$ (within several microns of the surface). A rather uncommon surface defect, displayed in Fig. 13, was found on a device-grade silicon wafer. It appears to be similar to the surface defects, identified as microsplits by Schwuttke, which are small cracks along the $<110>$ direction in the silicon surface with stacking faults at either end. Schwuttke has found that microsplits like that in Fig. 13 are introduced during the saw cutting of the silicon.

It is interesting to compare the SSP technique with the useful but destructive technique of selective etching. A sample was prepared for this purpose by polishing a piece of saw cut silicon at a 2° bevel off normal to the (100) surface. The sample geometry is represented in Fig. 14. A Siritl etch was applied to half of the surface to reveal the saw damage. The sample was then oxidized and the series of SSP images in Figs. 14 A' - D' were measured. For comparison, Figs. 14 A - D are micrographs of the
etched surface at points of polish depth equal to that of the equivalent SSP image. Both the etch pattern and the non-destructive SSP technique appear to provide the same type of information. However, it is thought that the SSP measurement is sensitive to sub-surface structure such as precipitates which may not appear in an etch pattern.

Conclusion

Each of the scanned optical techniques, scanned internal photoemission and scanned surface photovoltage, has unique capabilities for examining semiconductor surfaces and interfaces. Photoemission imaging is useful in determining interface reaction or contamination. However, it is not amenable to wide use in process control or material evaluation because of the difficulty of the measurement and the requirement of a contacting counterelectrode. On the other hand, photovoltage imaging is a simple, fast, non-contacting, non-destructive technique that offers reasonable resolution, about half a micron for a double differentiated measurement at a wavelength of 3250Å. Since scan speed does not degrade resolution, there is no inherent physical limit on the frame repetition rate. The photovoltage technique has been used routinely to detect stacking faults, dislocations, precipitates, gross surface damage, and local insulator charging.

The photovoltage imaging technique SSP has great potential for use in semiconductor material evaluation and in process control in the production of semiconductor devices. By observing the defects introduced during the various stages of production, it is possible to identify and control the
factors that introduce defects. In the particular case of silicon inte-
grated circuits, both the production and the reliability of the devices
is critically dependent upon crystalline perfection of the material which
can be monitored by SSP imaging. The use of photovoltage imaging is some-
what less useful as a research tool than the electron beam analog, as
developed by Bottoms 28, because the spacial resolution is considerably
lower.
REFERENCES


[36] The bevel polished sample was prepared by J. W. Philbrick, IBM Hopewell Junction, NY.
Figure 1  Optical system and measurement apparatus for the scanned optical measurements. Laser radiation is focused through a 50 μm pinhole and then deflected by two scanning mirrors rotating about perpendicular axes. The deflected light fills the aperture of a microscope objective lens for all deflection angles $\theta$ and $\phi$. The system is brought to focus by adjusting the position of the sample along the optical axis to minimize the size of the light spot reflected onto the fluorescent screen on the back of the 50-μm pinhole. A portion of the reflected beam is diverted and brought to a stationary focus on a photocell in order to detect a reflectivity image of the scanned area.
Figure 2  A schematic representation of the scanned optical system showing the instrumentation required to obtain SIP and SSP images. The x and y inputs to the CRT are driven directly by the voltage applied to the mirror galvanometer motors.
Figure 3 The optical scanning apparatus showing the scan mirrors, galvanometer drivers, microscope reflecting objective, sample positioning stage, and the preamplifier head.
Figure 4  A representation of the processes involved in scanned internal photoemission. A small reduction in the effective interface barrier allows an enhanced photocurrent to flow.
Figure 5  Scanning internal photoemission images of three samples of 
Si-SiO$_2$ interfaces covered with $4 \times 10^{12}$ sodium atoms per 
square centimeter, and measured at a photon energy of 3250 Å. 
The light areas indicate a high photoyield produced by the 
presence of sodium on the interface. The dark areas are 
thought to be due to regions of phosphorus rich layers near 
the Si-SiO$_2$ interface. Phosphorus segregates at that inter-
face during the formation of the SiO$_2$ layer, produced by the 
oxidation of $10^{20}$ P/cm$^3$ doped silicon.
Figure 6  Scanned internal photoemission measurements of an Si-SiO$_2$
interface which has been uniformly coated with $4 \times 10^{12}$ Na/cm$^2$.
The Na is found to decorate a 30 μm, long defect which lies in
the <100> direction on the silicon surface.
Figure 7  Scanned internal photemission image of a Bi-Nb$_2$O$_5$ interface, measured at $\lambda = 6328$ Å. The vertical striae are due to a dc level shift of leakage currents.
Figure 8

The rudiments of the scanned surface photovoltage measurement. A spot of light is swept across a semiconductor surface, generating a comet shaped cloud of minority carriers in a surface depletion region. The surface potential, capacitively detected by a remote electrode, fluctuates when the spot traverses a surface recombination center.
Figure 9  The spatial derivative of the surface minority carrier density, along a line in the direction of travel of a 1 μm spot of light. The derivative is relatively insensitive to the scan velocity \( v \) for the values shown in the figure. The quantity \(|dn/dx|\) is the sampling function in the singly differentiated scanned surface photovoltage measurement.
Figure 10  Scanned surface photovoltage image of several clusters of emergent dislocations through the (100) surface of a silicon wafer. The dislocations were generated by sandblasting a spot on the reverse side. During subsequent annealing and oxidation steps, the dislocation loops generated at the damage sites grow through the wafer, eventually penetrating the front surface.
Figure 11  Scanned surface photovoltage image of surface damage produced during the cutting and polishing of a silicon wafer. The damage line does not follow the $<100>$ direction on the (100) surface.
Figure 12  Scanned surface photovoltage image of a (100) silicon surface showing small, randomly distributed defects.
Figure 13  Scanned surface photovoltage image of a defect, apparently a microsplit, which was introduced during saw cutting of the (100) surface. The micropain is thought to be flanked by stacking fault arrays in the <100> direction on the surface.
Figure 14  Scanned surface photovoltage images (A', B', C', D') of saw damage in a silicon wafer which has been polished at a 2° angle. For comparison, micrographs (A, B, C, D) of an adjacent area of the surface were made after etching. Both sets of images, taken on a highly polished area of the beveled surface, show the transition from the damaged region to the undamaged region occurring at about the same depth, as is observed on an etched portion of the bevel.
IV. FIELD DEPENDENT INTERNAL PHOTOCHESSION PROBE OF THE
ELECTRONIC STRUCTURE OF THE Si-SiO₂ INTERFACE*

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ABSTRACT

Details of the SiO₂ conduction band near the Si-SiO₂ interface were examined by the technique of determining the field dependence of the threshold φ(E) for internal photoemission for fields up to 7 x 10⁶ V/cm. From a measurement of φ(E), the conduction band bottom φ(x) in the SiO₂ was determined to within approximately two lattice units of the interface. In calculating the effective potential φ(x), corrections were made for the effects of the photon-induced tunneling and field penetration into the silicon. The results of conduction threshold measurements show only a small deviation from a flat band, approximately +0.05 eV, up to a point 4Å from the silicon surface. There is no indication of a large ionic charge or of a greatly reduced SiO₂ bandgap to within about two lattice units of the interface.

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INTRODUCTION

Although the Si-SiO₂ interface has been studied far more intensively than any other because of its technological importance, the details of the electronic and physical structure of the interface are little understood. The lack of information about the region near the Si-SiO₂ interface is due to the difficulty of probing only the first few atomic layers around the interface without otherwise disturbing the system. Certainly, one would not expect that the interface between a given crystallographic surface of Si and an amorphous SiO₂ layer is precisely atomically smooth. In fact, it is not unreasonable that the first few molecular layers of SiO₂ and Si might be irregular. Evidence¹ based on He backscattering data suggests that SiO₂ which has been thermally grown on silicon contains excess silicon near the Si-SiO₂ interface. Recent He backscattering measurements, which employed atomic channeling to minimize background, show approximately $6 \times 10^{15}$ Si-SiO/cm² in excess of stoichiometry near a thermally grown Si-SiO₂ interface. However, ion backscattering measurements are somewhat insensitive and difficult to interpret because of the background from the substrate, even if channeling techniques are employed³.

Several sputter-etch profiling techniques have been used to examine the Si-SiO₂ interface, including Auger⁴,⁵ and ESCA⁶ measurements. Data from the sputter-etch measurements show some irregularities near the Si-SiO₂ interface which have been variously interpreted as excess Si in the SiO₂, a graded bandgap interface Si-SiOₓ-SiO₂, or as free silicon inclusions in the SiO₂. However, the data are difficult to interpret unequivocally due to non-uniformities in the microscopic sputter-etch rate in SiO₂ which causes the
interface to appear distorted because some portions of the silicon surface and exposed by the sputter-etch process regions before others. Additionally, these profiling techniques are complicated by details of the electron escape mechanism and by the physical damage induced by the sputtering process itself. Thus, the sputter profiling techniques are particularly difficult for the Si-SiO₂ interface, in which the physical interface is to extend over a region which is on the same order as the uncertainties in the measurement.

In this paper, we provide new information on the local microscopic structure of the Si-SiO₂ interface, obtained by the field dependent internal photoemission technique. In this technique, the electric field dependence of the threshold for internal photoemission is measured and used to determine the conduction band energy in the insulator as a function of distance away from the interface. In this way, the SiO₂ conduction band was probed from 4Å to 12Å from the Si-SiO₂ interface, and was found to bend only 0.05 eV to 0.03 eV, up to a point in the SiO₂ about 4Å from the surface. In view of the 9.0 eV bandgap of SiO₂, the 0.05 eV band bending is small. We see no substantial evidence for an SiOₓ graded bandgap, or for significant excess charge extending farther than 4Å from the silicon surface.
FIELD DEPENDENT INTERNAL PHOTOEMISSION

The Si-SiO₂ interface has been extensively studied\textsuperscript{8-12} by internal photoemission measurements\textsuperscript{13}. The interface barrier is determined by finding the threshold for photoemission over the barrier from spectral photoresponse measurements. Monochromatic light is passed through the SiO₂ to the Si, where photons excite electrons over the Si-SiO₂ interface barrier. The threshold is determined\textsuperscript{14} by extrapolating a power law fit to the photoresponse, where the best fit is obtained with a power\textsuperscript{11} of 2 or 3. Typically the threshold determined from a third power fit is 0.15-0.2 eV lower than that obtained from a second power fit. Thresholds determined by a third power fit, after correction for Schottky barrier reduction, are 4.35 eV\textsuperscript{11} and 4.30 eV\textsuperscript{12}. In the present work we find an interface barrier of 4.44 eV by using a parabolic fit to the photoyield.

The Si-SiO₂ interface may not be an ideal, abrupt surface, but rather, it may extend over a finite distance. Several possibilities for the influence of any excess Si on the microscopic local structure of the interface are represented schematically in Fig. 1. The solid lines show the valence and conduction bands in SiO₂ near an ideal, abrupt surface, while dashed line (b) illustrates the bands in a graded bandgap junction and dashed line (a) shows the effect of positive charge in the SiO₂ bands near the interface. For the graded bandgap model, the transition from Si through SiOₓ to SiO₂ is the continuous, with the bandgap of SiO₂\textsuperscript{15} increasing from 1.1 eV to 9.0 eV, as shown by curve (b). The graded bandgap model implies that the conduction band in the SiO₂ deviates below the ideal straight line near the interface. The other possibility, shown as the dashed curve (a), involves a band
curvature due to charged Si near the interface. Either of the two models for the influence of possible excess silicon near the interface implies a large curvature of the SiO$_2$ conduction band. This curvature can be examined by the field dependent photoemission technique.

From the field dependence of internal photoemission from Si into SiO$_2$, the variation with distance from the interface of the SiO$_2$ conduction band was determined. The field dependence of the threshold of photoemission from various materials into SiO$_2$ has been found approximately to follow the simple Schottky barrier reduction that one would expect from an ideal abrupt junction. However, any curvature of the SiO$_2$ conduction band near the interface would result in a deviation from this simple Schottky barrier reduction, as is illustrated in Fig. 2.

The experiment consists in illuminating the Si with photons of energy $\hbar \omega$, which excite electrons $e^-$ in the Si. Electrons with sufficient energy can surmount the interface barrier formed by the maximum of the sum of the conduction band potential $\phi(x)$, the image potential $(-e/16\pi\varepsilon_0 x)$, and the applied potential $-x E$. As can be seen in the series of curves for fields $E_1$, $E_2$, and $E_3$, an increase in the applied electric field moves the position of the barrier maximum closer to the interface and lowers the total barrier height. An additional effective reduction in the measured photoelectric threshold is introduced by photon assisted tunneling. Electrons with energy nearly sufficient to surmount the interface barrier have a finite probability of tunneling through the effective barrier for high applied electric fields for which the barrier is relatively narrow. The apparent decrease $\Delta \phi$ in the measured threshold, shown in Fig. 3 for the parameters of SiO$_2$, is relatively
small, even at the highest electric fields used in the present work. The correction for photon-assisted tunneling is included in the results, although the effect introduced a deviation of only 0.025 eV in the resulting determined \( \phi(x) \). From a measurement of the photoelectric threshold \( \phi(E) \) as a function of electric field, the conduction band potential \( \phi(x) \) can be determined. By this field dependent photoemission technique, the SiO\(_2\) band curvature was measured as a function of distance away from the silicon surface.

The measurements were performed on samples formed by the oxidation of 10 \( \Omega \)-cm n-type Si (100) in dry \( O_2 \) at 1050 C. The thickness of the SiO\(_2\), determined by ellipsometric measurements, was 1090 \( Å \). A semitransparent electrode of 160\( Å \) of aluminum, 0.3 cm in diameter, on top of the SiO\(_2\) was used to apply the electric field and to collect the photocurrent. Photoemission measurements were made up to the highest electric field at which the sample was stable. Three independently fabricated samples were measured, with virtually identical results.

The photoemission measurements were performed with light chopped at 6 Hz in order to minimize 1/f noise and a small residual surface photovoltage, as well as to reject the Fowler-Nordheim dark tunnel currents at the higher electric fields. The spectral photoresponse was corrected for the reflectivity of the sample. From a plot of the half-power of the photoyield at a given applied field, the threshold was determined by extrapolation. At the higher electric fields, the photoyield near threshold was complicated by a component due to the photoemission of electrons from the conduction band of the inverted silicon surface. This additional component was similar to the photoemission spectra found by Goodman\(^{16}\) for emission from the conduction band of heavily n-type silicon.
RESULTS

The photoelectric threshold $\phi(E)$ for emission from Si into SiO$_2$ is shown as a function of electric field in Fig. 4. The experimental points were determined from a fit of the spectral photoyield to a parabolic dependence on photon energy. Extrapolated to zero electric field, the threshold indicates an interface barrier of 4.44 eV. A fit of the spectra to a third power dependence on photon energy was found to shift the points for $\phi(E)$ lower in energy by about 0.15 eV, without changing the apparent electric field dependence of the points. A fit to the second power rather than the third power was chosen because it facilitates a separation of the photoyield due to emission from the Si conduction band from the emission for the valence band. In a fit to the third power of the photon energy, the excess emission from the conduction band introduces some uncertainty in the determination of the threshold. The information on the interface region derived from either the second or third power fit is substantially the same, since the field dependences of $\phi(E)$ determined by both methods are virtually identical.

The experimental points for $\phi(E)$ at low field $E$ (or equivalently, large distances of the effective barrier from the interface) fit well to the solid line, which is the simple theoretical Schottky barrier reduction, using the value of $\epsilon = 2.15$ for SiO$_2$. The dashed line is the Schottky theory including a correction for photon-assisted tunneling. The fit of the Schottky lowering theory to the experimental results for Si-SiO$_2$ is reasonably good, much closer than that for Pt - SiO$_2$ or Au - SiO$_2$.

From the electric field dependence of $\phi(E)$ we determined $\phi(x)$, the potential of the conduction band bottom in the SiO$_2$, by a procedure outlined
previously\(^7\). In case of Si-SiO\(_2\), few iterations are necessary because the measured \(\phi(E)\) deviates little from the Schottky theory. For very small deviations from the ideal, \(\phi(x)\) is approximately,

\[
\phi(x) = \phi_o + \Delta\phi(E)
\]

where \(\phi_o\) is the threshold extrapolated to \(E = 0\) and \(\Delta\phi\) is the difference between the measured threshold \(\phi(E)\) and the theoretical Schottky barrier reduction at the field \(E\). The distance \(x_o\) is the position of the maximum of the total effective barrier at the field \(E\). Neglecting the correction for photon-assisted tunneling, this distance \(x_o\) is,

\[
x_o = \left\{ \frac{e}{16\epsilon_o \epsilon_o E} \right\}^{1/2}
\]

where \(\epsilon_o = 2.15\) is the high frequency dielectric constant of SiO\(_2\). The SiO\(_2\) conduction band potential \(\phi(x)\) determined in this way is shown as a function of distance \(x\) from the interface in Fig. 5. The solid curve is the potential found without correcting for photon-assisted tunneling, while the dashed curve includes this correction.

The band bending in SiO\(_2\) near the Si-SiO\(_2\) interface, as shown in Fig. 5, is relatively small. At \(4\angstrom\) from the silicon surface, the deviation from the ideal flat band is only 0.95 eV, or only 0.55\% of the bandgap of SiO\(_2\). The small distortion of the conduction band in the SiO\(_2\) may be due in part to a small amount of positive charge in the SiO\(_2\). From Fig. 5, an upper bound on the maximum amount of positive charge beyond \(4\angstrom\) is \(2.5 \times 10^{12} \text{e/cm}^2\).
SUMMARY

The technique of field dependent internal photoemission has been used to determine the conduction band potential in SiO$_2$ near the Si-SiO$_2$ interface. Only a small deviation from an ideal abrupt junction was found. The total band bending up to a point 4Å from the silicon surface is only 0.05 eV, at most with an estimated experimental error of ±0.03 eV.

The lack of any significant bending of the conduction band rules out the possibility that a graded bandgap junction extends beyond 4Å from the silicon. Further, the small observed band bending is in the opposite direction to what one would expect from a graded bandgap type of model. Also, no significant charge (<2.5 x 10$^{12}$ e/cm$^2$) exists in the SiO$_2$ beyond 4Å from the silicon surface.
REFERENCES


Figure 1  Schematic representation of the electronic bands near the Si-SiO$_2$ interface. The solid line shows the ideal valence and conduction bands of SiO$_2$ near the interface, while dashed curves a. and b. represent deviations from the ideal interface due to excess positive charge and a graded bandgap junction, respectively.
A schematic representation of a Schottky barrier for three different applied electric fields, $E_1$, $E_2$, $E_3$, for the case of an arbitrary initial potential in the insulator. At one particular value of field, the position $x_1$ of the maximum of the barrier and the potential $\phi(x)$ can be obtained from the dependence of the photoemission threshold $\phi(E)$ upon electric field.
Figure 3  The correction to the photoemission threshold due to photon-assisted tunneling of electrons through the interface barrier. The tunneling parameters are for SiO₂.
The threshold $\phi$ for internal photoemission from Si into SiO$_2$ as a function of the electric field in the SiO$_2$. The threshold was determined by a parabolic fit to the photoemissive yield.

The solid line is the ideal Schottky dependence of the threshold $\phi$ on the electric field, while the dashed curve is the ideal threshold after correction for photon-assisted tunneling.
Figure 5  The potential $\phi(x)$ at the bottom of the conduction band in SiO$_2$ as a function of distance $x$ from the Si - SiO$_2$ interface. The solid curve is the band bending calculated without the photon-assisted tunneling correction, and the dashed curve is that calculated by including the correction.
V. INTERFACIAL REACTIONS IN SOLIDS

This program is devoted to the study of reaction at metal/silicon and metal/glass interfaces. These two kinds of interfaces are essential elements in MOS devices where the basic structure is multilayered thin films. Multilayered thin films resemble diffusion couplers, hence diffusion and interface reaction have been recognized as the prime cause of many reliability problems. Due to the small film thickness, the reaction study demands experimental tools that are capable of detecting structural and compositional variation in a distance of the order of 100Å. Reaction kinetics within such a small distance often is distinctly different from its counterpart in a bulk sample. We have combined a glancing angle Deeman-Bohlin x-ray diffractometer and a MeV $^4\text{He}^+$ ion backscattering technique to analyze interfacial reactions. The diffractometer is sensitive enough to obtain the structural information from a metallic film of thickness of 200Å, and the backscattering is capable of profiling the composition variation in a film with a depth resolution of 200Å. These two techniques have been used to measure the growth kinetics of silicide formation. In addition, we have applied implanted noble gas atoms as diffusion markers to determine the dominant diffusing species during the formation.

We now recognize that contrast reaction between Si and near-noble transition metals can occur at a much lower temperature than that between Si and other transition metals. The latter requires a reaction temperature above 500°C yet the former can proceed below 200°C. The large discrepancy in reaction temperature indicates atomic mobilities in these two kinds of metal/Si interfaces are different. The silicide tends to
be metal-rich in the low temperature reaction, and to be Si-rich in the high temperature reaction. The use of silicide as a barrier layer on Si has resulted in a wide range of study on properties of silicides, in particular, its formation and growth kinetics. One aspect of silicide formation which has been studied in detail in this program is the identity of the diffusing species.

In conventional bulk couples the faster of the two diffusing species is identified by the direction of displacement of embedded wire markers. To apply this concept to thin film systems requires markers much smaller in dimension than the film thickness. This can be achieved by ion implanting rare gas species which have a depth distribution smaller than the film thickness. Backscattering techniques with MeV $^4\text{He}^+$ ions can be used to detect the marker species and their location in depth within the sample.

Implanted noble gas atoms of Ar and Xe have been used as diffusion markers in growth studies of six silicides, $\text{Ni}_2\text{Si}$, $\text{Mg}_2\text{Si}$, $\text{FeSe}$, $\text{VSi}_2$, $\text{TiSi}_2$, and $\text{Pd}_2\text{Si}$, formed by reacting metal films with silicon substrates. MeV $^4\text{He}$ ion backscattering has been used to determine the displacement of the markers. The displacement relative to the surface determines the identity of the diffusing species. We found that Si atoms predominate the diffusion in $\text{VSi}_2$, $\text{TiSi}_2$ and FeSi, while Ni atoms are the moving species in $\text{Ni}_2\text{Si}$ and Mg in $\text{Mg}_2\text{Si}$. In $\text{Pd}_2\text{Si}$ both Pd and Si are diffusing species with Si faster of the two. Details of kinetics of silicide formation has been reported in the previous semi-annual reports.

A strong connection exists between silicide formation and glass/metal
reaction. Many metals are known to show detrimental effects on glass; phase separation, devitrification and decomposition. When a metal decomposes SiO_2, it may form oxide and silicide. From the point of view of metal/glass reaction, a metal can be classified as whether or not it forms oxides and/or silicide based on the following reaction equation:

\[ M + SiO_2 \rightarrow \text{silicides} + \text{oxides} \]

where \( M \) represents a metallic element. In Table I, we show the four kinds of reactions that can be expected.

<table>
<thead>
<tr>
<th>Metallic Elements</th>
<th>Products of the Reactions</th>
</tr>
</thead>
<tbody>
<tr>
<td>(example)</td>
<td>Silicide</td>
</tr>
<tr>
<td>Au, Ag</td>
<td></td>
</tr>
<tr>
<td>Al, Sn</td>
<td></td>
</tr>
<tr>
<td>Pt, Pd</td>
<td>( x )</td>
</tr>
<tr>
<td>Cr, V</td>
<td>( x )</td>
</tr>
</tbody>
</table>

We have used the Table as a guide for our study and have investigated and reported the reaction with metals in the first group (e.g. Au) and the 4th group (e.g. V). We note that even though Au itself does not react with SiO_2, its eutectic alloy with Si wets and reacts with SiO_2. A case in which such reaction has been found is the MOS structure of Au/SiO_2/Si. A wetting experiment with a Au-Si eutectic melt on surfaces of fused quartz has been carried out in He atmosphere. The wetting shows that the melt reacts with quartz. Kinetics of the reaction was studied by annealing an MOS structure of Au/SiO_2/Si at 200 to 800°C in a dynamic vacuum of \( 2 \times 10^{-6} \) Torr. Details of the study is reported in the following section.

So far, glass-metal reactions have been found to show detrimental...
effects on the integrity of glass. Thus, it is desirable to strengthen glass to show a better resistance to attack by metallic atoms. Structurally, the network configuration of a glass is very loose and could be packed into a denser form. A denser form of glass is ideally better whenever a very thin glass layer (< 1000Å) is required because thin glasses are especially vulnerable to pin holes. We have developed a new technique which potentially may be applied to densify a glass layer and to reduce its pin hole density. The technique is to expose the glass to a low dosage, high energy ion beam. The amount of compaction has been measured by Talystep and confirmed by the changes of density and reflective index obtained by ellipsometry. The densified glass layer has also been found to show an improved dielectric breakdown strength. Details of the study is reported here.
VI. SURFACE REACTIONS ON MOS STRUCTURES

by

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ABSTRACT: A wetting experiment with a Au-Si eutectic melt on surfaces of fused and single crystal quartz was carried out in an atmosphere of He. The wetting showed that the melt reacted with quartz and the wetting angle was about 10 degrees. Kinetics of the reaction was studied by annealing an MOS structure of Au/SiO$_2$/Si at 700 to 800°C in a dynamic vacuum of 2x10^{-6} Torr. In the initial stages, reduction rate of the oxide layer by Au-Si eutectic followed an approximately linear dependence on time and its temperature dependence was characterized by an activation energy of about 26 kcal/mole. The reduction ceased when depletion of Au took place.
INTRODUCTION

We have reported a low temperature surface reaction in MOS structures which results in the degradation of the oxide layer.\(^1\) The MOS structure was made by depositing arrays of dots of Au film on 800Å thick thermal oxide layers on Si wafers. When this structure was annealed at temperatures ranging from 500 to 800°C in a dynamic vacuum of 2x10\(^{-6}\) Torr, the Au was observed to diffuse through the oxide and to form a liquid Au-Si eutectic with the underlying Si. The eutectic dissolved the oxide and then spread laterally from edges of the Au dots along the SiO\(_2\)/Si interface into areas that were not covered by the Au. A detailed description of the morphology and reaction products has been given,\(^1\) and the most significant finding was that in the reacted areas there was no oxygen, indicating the destruction of SiO\(_2\). We have proposed that the destruction of SiO\(_2\) is due to its reaction with liquid Au-Si eutectic, since no such reactions have been observed with pure Au on fused quartz or on bare Si wafers.\(^1\)

If our explanation is correct, it would suggest that the liquid Au-Si eutectic wets a SiO\(_2\) surface, otherwise there is no reaction between the two. This paper is a study of the wetting of Au-Si eutectic on fused and single crystal quartz surfaces. The results show that the eutectic does indeed react with quartz surfaces. A preliminary study of the kinetics of the eutectic-glass reaction was performed using the Au/SiO\(_2\)/Si structure, and the rate and activation energy were given in a earlier report.\(^2\) These and additional kinetic results will now be analyzed further to give a more detailed picture of the reaction mechanisms.
EXPERIMENTAL

(a). Wetting of Au-Si Eutectic on Fused and Single Crystal Quartz Surfaces

A eutectic Au-Si alloy was made by mixing pellets of Au of six-9 purity and pieces of Si from a Si wafer, and annealing the mixture in a sealed quartz tube at 500°C for 24 hours. The mixture had a composition of 69 at % Au and 31 at % Si, in agreement with the eutectic composition in Hansen’s book. The alloy was broken into pieces for subsequent wetting experiments and each piece weighed about several grams. The wetting experiments were carried out on a hot stage maintained at 450°C in a titanium-gettered, He atmosphere. A sketch of the hot stage is shown in Fig. 1. The temperature was measured by means of a thermocouple inserted in the A block which held the quartz wafer. The fused quartz substrates were 1" diameter wafers 1/16" thick. The single crystal quartz had the C-axis normal to its surface and had dimensions of 1"x1"x 1/16 ".

To determine the wetting angle, we spread the liquid alloy in the form of a stripe on quartz surfaces, kept them at 450°C for 15 minutes to allow local equilibrium to be reached at the edges and then solidified the stripe. The angle was measured on the cross-section of the stripe by cutting across it.

(b). Reaction Kinetics in Au/SiO₂/Si Structure

The layered structure was made by depositing arrays of circular dots of Au film 30 mil in diameter and 500Å in thickness on 80 mil centers onto Si wafers oxidized with 800Å of SiO₂. An optical micrograph of the top view of a Au film dot reacted at 700°C for 4 hours is shown in Fig. 2a. The oxide under and in the vicinity of the Au film was consumed, forming
the circular opening, due to the eutectic-oxide reaction. In Fig. 2b, a scanning electron micrograph of a circular opening is shown. The channels in the opening are parallel to [110] directions of the Si substrate. At the end of each channel where it meets the edge of the oxide we have found a bead of Au-Si. A replica electron micrograph of the channels, the beads, and the interface between the reacted and unreacted areas is shown in Fig. 2c. We have measured the migration rate of the interface between the reacted and unreacted areas, and the spacing between channels as a function of time and temperature. The measurement of the rate was not straightforward since the reaction often does not proceed uniformly over the entire circumference of an Au dot. We chose to measure its average rate over a wide reaction front. To determine its time dependence, we have followed the reaction at a specific area over a selected sequence of annealing times. The spacing between the central line of two channels was determined by averaging measurements over 10 to 20 spacings.

RESULTS AND DISCUSSION

When we broke the sealed quartz tube to take out the Au-Si eutectic, part of the eutectic alloy was found to stick to the wall of the tube. The tube fractured as it was forced apart from the eutectic and pieces of the fractured quartz still adhered to the eutectic. This is a clear indication that the eutectic has wetted the quartz surface.

When the eutectic pieces were placed on the hot stage, they melted at roughly 390°C, which indicates that the pieces were richer in Si than the alloy at the eutectic point. At first, no wetting on the fused quartz surface could be detected, this was probably due to the presence of a surface oxide coating on the melt which existed before the melting and had prevented
the melt from making direct contact with the quartz. When we broke the coating with a pointed Au wire, the shiny, silver-colored liquid wetted the quartz and stuck to the quartz surface wherever we spread it. It also maintained its shiny surface at 450°C throughout the experiment ( ≈ 30 minutes).

The wetting angle as measured from cross-sections of eutectic stripes is about 10 degrees. In Fig. 3 an optical micrograph of such a cross-section is shown. Since there is a volume shrinkage at solidification, the measured angle may differ slightly from the true wetting angle. The measurement on the quartz crystal was performed in the same manner and the wetting angle is about the same. There is also a wetting between the bead and the oxide in Fig. 2. However, the wetting angle is not well defined since the oxide was thin and the underlying Si might have altered the angle.

To investigate the eutectic-oxide reaction kinetics, an MOS structure of Au/SiO₂/Si was used instead of samples of Au-Si eutectic on quartz. This is because the progressive enlargement of the oxide-free circular opening is easily monitored from optical micrographs such as the one shown in Fig. 2a. Taking radial measurements of the circular opening at selected intervals of time allows one to plot growth curves as shown in Fig. 4. The basic features of these curves are the linearly rising portion at early times and the leveling-off at long times. The leveling-off can logically be attributed to the depletion of Au.

Before we analyze these reaction curves, the driving force of the reaction should be discussed. It is most likely the lowering of free energy of liquid Au-Si alloy by dissolving oxygen. In the dynamic vacuum condition,
if we assume that the Au rich alloy can release oxygen to the vacuum rather quickly, the alloy is always below its solubility limit and therefore, under a driving force to dissolve more SiO$_2$. In dissolving the oxide, the alloy gains some silicon in addition to oxygen. We believe, however, the solution of Si to the alloy is not the driving force since the alloy can obtain sufficient Si from the underlying substrate. It appears that the pumping out of oxygen must also be accompanied by the pumping out of some of the Si. This is because each of the channels as shown in Fig. 2c is, on average, 5 microns in width and 0.4 microns in depth; the Si that was in the channel has disappeared. Presumably the partial vapor pressure of Si of liquid Au-Si alloy at 700 to 800°C is high enough so Si can be pumped out easily. The morphology of the channel with a bead of Au-Si at its end appears similar to the reverse process of vapor-liquid-solid (VLS) mechanism of crystal growth of Si whiskers.$^{5,6}$ In our case, the reason that the bead stays at the interface rather than drills a hole into the Si substrate as in the VLS case might be its wetting adhesion to the oxide.

In analyzing the reaction kinetics, two pieces of information seemed significant. We know from the wetting experiments that there is an interfacial reaction between the eutectic and the oxide and from the leveling-off of reaction rates at longer times we know that the supply of Au to the reaction interface is crucial. These suggest that the kinetics might be controlled by one or a combination of the following steps; the interfacial reaction, the diffusion of Au to the reaction interface, and the diffusion of Au along the reaction interface. We have assumed here that the formation of the eutectic alloy at the reaction temperatures is so fast that it is kinetically irrelevant.
Also, the time needed for penetrating the oxide layer under the Au dot is treated as an incubation time of the lateral reaction. The constant reaction rates show clearly that at the early stage they do not depend on diffusion of Au from the circumference of the dot to the reaction interface, rather they may depend either on interfacial reaction or on diffusion of Au along the reaction interface as both can give rise to a constant reaction rate. In the latter case, its morphology is similar to cellular reaction, e.g. precipitation or dissolution of Sn from Pb-rich Pb-Sn alloys. In such reactions, the rate takes the following form,

$$\frac{dr}{dt} = \frac{D_b \delta}{S^2}$$  \hspace{1cm} (1)

where $D_b$ is the diffusion coefficient along the reaction interface, $\delta$ is the effective interface width and $S$ is the interlamellar spacing. Assuming this is the rate controlling step, we can estimate the term $D_b \delta$ in the present case. Values of measured $\frac{dr}{dt}$ and $S$ in the temperature range of 700 to 800°C together with the estimated $D_b \delta$ are listed in Table 1.

In the Table, there are two features that are worth considering. First, the spacing seems to be independent of temperature in the temperature range measured. This is not what one would expect in a diffusion controlled reaction. Secondly, the effective interface width is hard to guess but is conceivably larger than 5Å, the value accepted for most abrupt solid-solid interfaces. This is because the interface considered here is between solid SiO$_2$ and liquid Au-Si alloy, so the flux of Au will not be limited in the interface but rather throughout an extended region in the liquid. This is supported by microprobe scanning x-ray micrographs which have shown the presence of Au along and behind the interface. Then if we
take $\delta$ to be much wider than $5\lambda$, we find $D_h$ to be much smaller than $10^{-5}$ cm$^2$/sec, the value which is accepted as a lower bound for diffusion in liquid metals and alloys. So we conclude that the observed reaction was not controlled by diffusion along the interface.

We have, in the above, eliminated the possibilities of rate controlling by diffusion to and along the reaction interface, so the remaining one that may control the rate is the interfacial reaction, the dissolution of SiO$_2$ by the eutectic. Assuming there is always an ample supply of Au-Si eutectic at the edge of the oxide, the volume rate of dissolution of oxide will depend on the area of oxide exposed to the eutectic. That is,

$$\frac{dV}{dt} = 2\pi rh K(T,P(O_2)).$$

(2)

Where $V$ is the volume of oxide film consumed, $r$ is the radius of an assumed circular opening, $h$ is the oxide thickness and $K(T,P(O_2))$ is the reaction rate constant which will depend on temperature and the partial pressure of O$_2$. The above expression may be reduced to

$$\frac{dr}{dt} = K(T,P(O_2)).$$

(3)

This result indicates that an initial loss of oxide should occur at a constant rate for any given $T$ and $P(O_2)$. Since there was no provision in our temperature controlled vacuum furnace for measuring O$_2$ partial pressure, we assumed that a constant total gas pressure was indicative of a steady state condition in which the pressures of ambient gases such as O$_2$ remained sensibly constant. Therefore, the slope of the linearly rising portion of the growth curve was employed to determined $dr/dt$. Furthermore, the slope of a plot of ln $dr/dt$ vs $1/T$ (Fig. 5) yielded an activa-
tion energy in the customary manner for Arrhenius plots. Proceeding in this manner the activation energy for oxide reduction under a dynamic vacuum of \(2 \times 10^{-6}\) Torr was approximately 26 kcal/mole.

This result is necessarily very approximate as any detailed model would certainly have to include the diffusion of Au through \(\text{SiO}_2\) and to the reaction interface. We mentioned that the diffusion of Au through \(\text{SiO}_2\) and the destruction of the oxide under the Au have been treated here as steps included in the incubation period of the reaction. Since our experimental approach was geared to observing the lateral reactions, it was not possible to draw any direct inference about the vertical or through-the-thickness reaction of Au on \(\text{SiO}_2\). However, calculated values of diffusivity using the data of Badaiev and Shuman\(^8\) are sufficiently large, \(7 \times 10^{-16}\) and \(4 \times 10^{-14}\) cm\(^2\)/sec at 700 and 800°C, respectively, to account for several hundred Å penetration of Au during typical incubation times of \(\sim 10^3-10^4\) sec. Ordinary diffusion processes do not seem adequate to explain the extensive chemical changes which must accompany or precede the destruction of the underlying oxide. Perhaps the Au penetration process is more complex involving a Au-\(\text{SiO}_2\) reaction analogous to what Madam, Morgan and Howes\(^9\) reported in their back-scattering study of Au migration through \(\text{SiO}_2\) films. Clearly, the nature of the diffusion process is not well understood and could warrant further study. It can be stated, however, that the energy we measure is considerably less than the dissociation energy of \(\text{SiO}_2\) by thermal energy alone (\(\approx 9\) eV) and demonstrates the efficiency of the Au-Si eutectic melt for oxide reduction.

Finally, at longer reaction times, the longer diffusion distance might become the rate limiting factor and should affect the constant
reaction rate. But more critical at the end of the reaction is the depletion of Au at the edge of the oxide. This is due to the initial finite amount of Au which has to be distributed to an increasingly bigger area and the loss of Au to the substrate. As the depletion takes place, the rate will decrease and eventually the reaction stops.

CONCLUSIONS

(a) A eutectic melt of Au-Si has been found to wet surfaces of fused and single crystal quartz. The wetting shows that the eutectic reacts with quartz and the wetting angle is about 10 degrees. The eutectic, after solidification, was found to bond rather strongly to quartz surfaces.

(b) An MOS structure of Au/\(\text{SiO}_2\)/Si has been used to study the kinetics of reaction between Au-Si eutectic and thermally grown \(\text{SiO}_2\) at temperatures between 700 to 800°C in a dynamic vacuum of 2x10^{-6} Torr. The reaction proceeds at a constant rate with an activation energy of 26 kcal/mole. The reaction will slow down and stop if depletion of Au occurs.

Acknowledgements

The authors are grateful to K. Asai, W. Kateley and J. Kucza for sample preparation, J. Kuptsis for the SEM picture, C. Aliota for the replica micrograph, and J. C. McGroddy for helpful comments.
References


4. Private communication with Prof. D. Turnbull and Dr. C. C. Tsuei.


Table 1

Measurements of Reaction Rate and Spacing

<table>
<thead>
<tr>
<th>Temp. °C</th>
<th>Reaction rate, ( \frac{dr}{dt} ) cm/sec</th>
<th>Spacing, cm</th>
<th>( \frac{D_b}{3} ) cm³/sec</th>
</tr>
</thead>
<tbody>
<tr>
<td>700</td>
<td>7.8x10⁻⁶</td>
<td>1x10⁻³</td>
<td>7.8x10⁻¹²</td>
</tr>
<tr>
<td>750</td>
<td>16.2x10⁻⁶</td>
<td>1x10⁻³</td>
<td>16.2x10⁻¹²</td>
</tr>
<tr>
<td>800</td>
<td>22.2x10⁻⁶</td>
<td>1x10⁻³</td>
<td>22.2x10⁻¹²</td>
</tr>
</tbody>
</table>
Fig. 1. A sketch of the hot stage used for wetting measurement.
Fig. 2. (a) Optical micrograph of a Au/SiO$_2$/Si structure after reaction at 700°C for 4 hours. The circular open areas are areas where reaction has taken place and no oxygen can be detected.

(b) Scanning electron micrograph showing the channels of a circular open. The channels have been found to be parallel to [110] directions of the Si substrate.
(c) Replica electron micrograph of a reaction front showing the channels, Au-Si eutectic beads and the interface. The sphere with a white shadow is a latex ball of 0.5 microns in diameter.
Fig. 3. A wetting edge of Au-Si alloy (near eutectic composition) on fused quartz surface. The angle was measured after the solidification of the Au-Si stripe.
Fig. 4. Reaction rate of liquid Au-Si eutectic with thermally-grown SiO$_2$ using MOS structure of Au/SiO$_2$/Si annealed at 700, 750 and 800°C in a dynamic vacuum of $2 \times 10^{-6}$ Torr. The thickness of Au and SiO$_2$ were 500Å and 800Å, respectively.
Fig. 5. Plot of $\log \frac{dr}{dt}$ vs $\frac{1}{T}$ for the reaction of liquid Au-Si eutectic with thermally-grown SiO₂.

$Q = 25.6 \text{ Kcal/mole} = 1.1 \text{ eV}$
VII. Publications under F19628-73-C-0006 and F19628-74-C-0077


Oral Presentations under F19628-73-C-0006
and F19628-74-C-0077

1. T. H. DiStefano, "Impurity Induced Dielectric Breakdown in SiO$_2$,"

2. D. R. Campbell, E. I. Alessandrini, K. N. Tu and J. E. Lewis
   "Phosphorus Diffusion and Catalyzed Crystallization in Amorphous Films

3. K. N. Tu, Chairman of the Session on Advances in Amorphous to
   Crystalline Phase Transformation, 1972 Materials Engineering Congress,

   Engineering, Caltech, Pasadena, California, Nov. 13, 1972.

5. T. H. DiStefano, "Electrical Interfaces," (Invited), Princeton Univ.,
   Nov. 20, 1972.

6. T. H. DiStefano hosted an AVS regional symposium on "Electrical
   Properties of Material Interfaces," at IBM Yorktown Heights, December
   7, 1972. His paper entitled, "Alteration of Electronic Barriers," was
   presented.

7. T. H. DiStefano, "Electronic Interfaces," (Invited) ECS, Ottawa,

8. T. H. DiStefano, "Electronic Interfaces," (Invited), Yale Univ.


11. E. I. Alessandrini, D. R. Campbell, "Catalyzed Crystallization and
    Transformation in SiO$_2$, Thin Films," ITL Meeting on Thin Film and

12. R. B. Laibowitz, P. Chaudhari, S. Herd, and S. Mader, "Direct
    Observation of Structural Changes in Switchable Materials,

13. T. H. DiStefano, "Thermionic Emission into Insulating SiO$_2$," APS


34. T. H. DiStefano, Panel Review of ARPA funded research on Surface Science, ARPA National Symposium on Surface Science, December 4-6, 1974, University of Pennsylvania.


37. D. E. Eastman and J. L. Freecouf (Invited) "Photoemission Measurements of Filled and Empty Surface States on Semiconductors and Their Reaction to Metal-Semiconductor Barrier Energies," 2nd Annual Conf. on the Physics of Compound Semiconductor Interfaces, UCLA, LA, California, February 4-6, 1975.


43. T. H. DiStefano served on the Program Committee for the 2nd Annual Conference on the Physics of Compound Semiconductors.


50. K. N. Tu organized a two-day symposium on "Interfacial Reactions in Solids," held at Cincinnati, Ohio, November 11-12, 1975.

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