Effects of ion irradiation in the thermal oxidation of SiC

C. Radtke and I. J. R. Baumvol
Instituto de Física, UFRGS, Av. Bento Gonçalves 9500, Porto Alegre, RS, Brazil 91509-900

F. C. Stedile*
Instituto de Química, UFRGS, Av. Bento Gonçalves 9500, Porto Alegre, RS, Brazil 91509-900
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We investigated the thermal growth of SiO$_2$ films on 6H-SiC (0001) samples irradiated with 170 keV Ar$^+$ ions. Electronic, compositional, and structural features arisen from the irradiation process were accessed. All results are compared to those of nonirradiated substrates, revealing major differences due to ion bombardment such as the enhanced rate of oxidation, the variation of roughness of both oxide surface and SiO$_2$/SiC interface, and of the stress in the formed oxide layer. Modifications on the electronic structure and crystallographic characteristics of the substrate underneath the oxide were also observed in samples in which the ion irradiation damaged layer was partially consumed by the oxidation step. The present work evidences that a careful choice of sample preparation conditions is necessary to avoid undesired characteristics in the final SiO$_2$/SiC structure and points to a way of preventing irradiation related degradation of electrical characteristics while maintaining the high oxidation rate.

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I. INTRODUCTION

Silicon carbide (SiC) is the material of choice for high-power, high-temperature, high-speed, and/or high frequency electronic devices.\textsuperscript{1,2} This is due to certain properties of this material including wide band gap, high breakdown electric field, high thermal conductivity and high-saturated electron drift velocity.\textsuperscript{3,4} However, the feature that really sets SiC apart from other wide band-gap semiconductors is the possibility of growing a well-known dielectric film on this material by thermal oxidation, namely, SiO$_2$. This is highly desirable since a substantial part of the technology already developed for Si can be adapted to the fabrication of SiC devices. Nevertheless, some barriers must still be overcome.

The electrical passivation characteristics of SiO$_2$ thermally grown on SiC are worse than those of SiO$_2$ on Si.\textsuperscript{1} Various explanations were already proposed to this bad performance, most of them attributing it to carbon present in the oxide, in particular at the SiO$_2$/SiC interface.\textsuperscript{5–8} Another processing problem of SiC is the low oxidation rate of this material: high temperatures and/or long oxidation times have to be used to achieve the desired oxide thickness.\textsuperscript{3}

Thus, the increase of the oxide growth rate and the improvement of the electrical quality of the near-interface region of SiO$_2$ on SiC are critical issues for device fabrication. In an attempt to enhance the oxidation rate of SiC, a couple of techniques were already investigated. Chudoba et al.\textsuperscript{9} studied the oxidation of β-SiC (100) surfaces modified by deposition of a Rb monolayer. The oxygen uptake was enhanced by four orders of magnitude in comparison with clean β-SiC (100) surfaces. In addition to the oxidation rate enhancement, the Rb overlayer was completely removed from the surface by thermal treatment below 780 °C. Despite the excellent results obtained, this technique is only suitable to grow thin oxides while thicker films (~30 nm) are needed in actual SiC devices. In another approach, ion irradiation of SiC single-crystal samples was used.\textsuperscript{10–12} Oxidation kinetics studies of irradiated SiC (Ref. 10) evidenced that the damaged layer oxidizes much faster than crystalline SiC, which allows oxidation at substantially lower temperatures (typically 1100 instead of 1300 °C). Despite the larger rate of oxidation, oxide films thermally grown on ion irradiated SiC presented so far an electrical quality inferior\textsuperscript{13} to that of oxides grown on nonirradiated SiC. We tentatively attribute this result to the fact that the damaged layer was not totally oxidized. In order to test this hypothesis, an atomic scale investigation of the reasons for these characteristics was undertaken. The aim is to modify conveniently the irradiation and oxide growth processes such that they lead to better quality oxides while maintaining the high oxidation rate.

In the present work a set of samples was prepared in order to portrait different stages of thermal oxide growth on SiC samples amorphized by irradiation with energetic Ar$^+$ ions. Chemical composition was probed as a function of depth in these SiO$_2$ films thermally grown on Ar-bombarded SiC substrates. For this purpose, x-ray photoelectron spectroscopy (XPS) analyses in conjunction with step-by-step etching of the oxides were accomplished. The same procedure was carried out with a nonirradiated sample for comparison. Rutherford backscattering spectrometry in channeling geometry (c-RBS) was used to probe the crystallinity of the substrate underneath the oxide and to determine the oxide thickness. Atomic force microscopy (AFM) scans were performed to analyze the surface morphology of the oxide and of the substrate after oxide removal by chemical etching. In this way, differences arisen from the irradiation procedure and possible changes of the SiO$_2$/SiC structure in the course of oxidation could be studied.

II. EXPERIMENTAL DETAILS

Si-faced nitrogen doped 6H-SiC samples cut from a wafer with resistivity 0.075Ω cm supplied by Cree, Inc. were used
in this experiment. After a standard RCA cleaning, samples were dipped for 30 s in a 5% HF solution in ethanol, followed by a 30 s ethanol rinsing aiming at removing the native oxide. Immediately after this washing procedure the samples were introduced in the ion implantation chamber which was pumped down to $5 \times 10^{-7}$ mbar. Irradiation was performed with $\text{Ar}^+$ at an energy of 170 keV and fluence of $1.1 \times 10^{15}$ cm$^{-2}$ in a 500 kV HVEE ion implanter, yielding a superficial amorphized layer observed by Rutherford backscattering spectrometry in channeling geometry.

Four different types of samples were investigated. Samples designated by $N360$ were nonirradiated SiC substrates which underwent wet oxidation for 360 min at 1100 °C, using $\text{N}_2$ bubbled through deionized water kept at 95 °C. Samples $I60$, $I150$, and $I720$ were irradiated substrates oxidized in the same temperature and pressure conditions as samples $N360$ for 60, 150, and 720 min, respectively (see Table I).

RBS in channeling geometry ($c$-RBS) analyses (detection angle was 170°) were performed in a 3 MV HV Tandetron ion accelerator with 2 MeV $\text{He}^+$ ions aligned in the $\langle 0001 \rangle$ crystal axis in order to determine the oxide thickness in different samples and to infer about the residual damage in the SiC substrate. The areal density of O in the films was determined from the areas of the corresponding O signals in the RBS spectra, calibrated against a SiO$_2$ film reference ($5.70 \times 10^{15}$ O cm$^{-2}$). Film thicknesses were obtained using the equivalent relationship $10^{15}$ O cm$^{-2}$ → 0.226 nm of SiO$_2$, obtained assuming an oxide density of 2.21 g cm$^{-3}$, valid for SiO$_2$ thermally grown on Si. Since oxides thermally grown on Si and on SiC are very similar, this assumption should not lead to appreciable errors. The values of oxide thicknesses obtained in this way are shown in Table I. After this procedure, the oxide surface morphology of the different samples was probed using AFM, performed on a Digital Instruments Nanoscope IIIa in tapping mode. After etching away the oxide with a 8:2 $\text{H}_2\text{O}:/\text{HF}(50\%)$ solution, a new set of channelled $c$-RBS measurements and AFM scans were carried out in all samples.

The chemical environment of Si in the oxide was investigated as a function of depth in samples $N360$, $I150$, and $I720$. The oxide was progressively thinned by immersion of each sample in a 50:1 $\text{H}_2\text{O}:/\text{HF}(50\%)$ solution with a well-characterized removal rate (1 Å/s). After each etching step, stopped by water rinsing, XPS spectra were recorded using Mg $K\alpha$ excitation ($h\nu = 1253.6$ eV) in an OMICRON UHV station based on an EA125 hemispherical analyzer. Detection angle $\theta$ of the photoelectrons with respect to the normal to the sample surface (take-off angle) was fixed at 45°. The step-by-step etching procedure was carried out until no change in the Si $2p$ region spectrum was observed. Au $4f_{7/2}$ photoelectrons from a foil covering part of the sample were used as the binding energy (BE) reference (83.8 eV).

### III. RESULTS AND DISCUSSIONS

RBS spectra obtained in channeling geometry of all samples after etching away the oxides are presented in Fig. 1. RBS spectra from a 6H-SiC virgin crystal and from a sample just after Ar$^+$ irradiation are also plotted for comparison. The residual damage after oxidation of samples $I60$ and $I150$ can be clearly observed by the higher yield next to the Si energy leading edge in comparison to the channelled spectrum from the virgin crystal. Different thicknesses of the residual damaged layer are due to different oxidation times. Comparing the estimated amounts of Si present in the oxide with the amount of Si in the irradiated layer of the as-irradiated sample, one can observe that the oxidation times used for samples $I60$, $I150$, and $I720$ consumed 15, 45, and 100% (within an error of 10%) of the amorphized layers.

![FIG. 1. Rutherford backscattering spectrometry spectra in random (shaded curve) and channeling (nonshaded curves) geometries of samples $N360$, $I60$, $I150$, and $I720$, after removal of the oxide layer by HF etching. Spectra of a virgin SiC sample and of the same sample just after ion irradiation are also plotted for comparison. The vertical arrow indicates the energy position of Si at the sample surface.](image-url)
respectively. Performing similar calculations with data from the sample of Ref. 13 we concluded that, in that work, approximately half of the damaged layer was consumed, resembling sample 1150 of the present work.

Another process taking place during oxidation of irradiated SiC can be observed when simulating the c-RBS spectra of the irradiated substrate and of sample 1150 after oxide etching with the RUMP code, as shown in Fig. 2. In the first case, the simulation agrees with experimental data of the amorphized layer, but in the spectrum from sample 1150 the disagreement between experimental and simulated curves evidences a higher degree of ion channeling near the interface between the amorphized layer and the single crystal, resulting from partial crystallization of the amorphized layer. In a previous work, crystallization of the implanted region during oxidation was also observed: selected-area electron diffraction (SAED) measurements evidenced an epitaxial structure of 6H then another one of 3C-SiC polytype close to the interface of the implanted layer with the 6H-SiC substrate, and then hexagonal and cubic SiC grains in the uppermost part of the layer. They also observed by cross-sectional transmission electron microscopy (XTEM) that the top and bottom interfaces of the oxide grown on this crystallized layer were rough.

Considering these findings, we determined the roughness of both oxide and SiC substrate (after chemical removal of the oxide layer) surfaces. Figure 3 shows images of all samples with their respective calculated root-mean-square (RMS) roughness values. Sample N360, the oxidized virgin crystal, has the smoothest surfaces, with roughnesses almost equal to the virgin crystal prior to oxidation (0.66 nm, not shown). Samples I60 and I150 evidence a large increase in roughness values as compared to sample N360. The origin of the observed morphology changes is attributed to the crystallization of the amorphous layer underneath the oxide, leading to a nonuniform oxidation front of the growing oxide. The longer the oxidation time undergone by the sample, the higher the degree of recrystallization of the remaining damaged layer. As a result of this process, roughness of sample 1150 is greater than that of sample I60. Morphology is changed in the case of sample I720, whose surfaces are much smoother, deviating from the previous tendency of roughness increase. In this case the whole bombarded layer was consumed during oxidation, such that the previously existing grains of the polycrystal were totally oxidized, resulting in the observed smoothening of both surfaces. Thus, further oxidation should not lead to increasing roughness since a single crystal is being consumed for oxide growth.

XPS results aiming at probing the chemical environment of Si are shown in Figs. 4–7. Owing to the complexity of photoelectron signals, a fitting procedure routine for the Si 2p region was adopted, which consisted of assuming two components, one related to SiO$_2$ (Si$^{ox}$) (FWHM = 1.5 eV, 80% Gaussian + 20% Lorentzian) and the other to SiC (Si$^{Sc}$) (FWHM = 1.7 eV, 80% Gaussian + 20% Lorentzian). The width and shape of each component were fixed using values obtained from fitted regions of a SiO$_2$ thick film on SiC and a virgin SiC sample, respectively.
Two binding energy (BE) constrains were primarily adopted: (i) these two Si 2p components are separated by 2.7 eV and (ii) Si 2p and O 1s components related to SiO$_2$ are separated by 429.5 eV. These initial approaches were used in fitting spectra from samples N360, I150, and I720 submitted to step-by-step etching. In the case of samples I720 and N360, no additional components were needed in the fitting procedure but the proposed binding energy separation between the Si$^{\text{ox}}$ and Si$^{\text{SiC}}$ components had to be changed and depended on oxide thickness. In the case of sample I150, however, the modifications were more pronounced. Figure 4 shows the Si 2p and O 1s regions of the spectrum of sample I150 after a total accumulated etching time of 1700 s, which exposed the SiO$_2$/SiC interfacial region. In addition to the two components used in the first fitting trial, another component with an intermediate BE had to be added. This new component (Si$^{\text{DC}}$) was attributed to the formation of Si oxyxides ($\text{Si}_x\text{C}_y\text{O}_z$), compounds where Si is bonded to a variable number of O and C atoms. This kind of compound was already observed in the interfacial region of thermal oxides grown on crystalline SiC. Concerning the O 1s region of the spectra, the predominant contribution to it is a component related to oxygen in SiO$_2$ ($\text{O}^{\text{ox}}$), centered at 534.5 eV in the case of Fig. 4. In order to obtain good fits, the BE differences between Si$^{\text{SiC}}$ and Si$^{\text{ox}}$ components had to be changed as a result of BE shifts in both components, which will be analyzed further in Figs. 5 and 6. However, it is worth to mention that the binding energy constrain of 429.5 eV between Si$^{\text{SiC}}$ and O$^{\text{ox}}$ components was kept in fittings of all samples and of all thicknesses.

Figure 5 shows Si 2p regions spectra of samples I150, I720, and N360 after being etched in a 8:2 H$_2$O:HF(50\%) solution for 1 min. After this step, the oxide layer should have been completely removed in all samples and the Si 2p signals should be due solely to SiC. However, reoxidation of the samples due to their exposure to air between the etching procedure and their introduction in the UHV chamber was observed by the appearance of another component around 104 eV. In the case of sample I150, its Si 2p region spectrum evidenced a large contribution of Si$^{\text{DC}}$ in addition to the other two components. This behavior was attributed to the SiO$_2$/SiC interface roughness, observed by AFM, and will be further discussed with Fig. 7. The Si 2p regions from the other two samples presented different contents of Si$^{\text{ox}}$ due to different exposure times to air. Comparing the BE’s of the Si$^{\text{SiC}}$ component, one observes different values. Other authors also observed distinct Si$^{\text{SiC}}$ BE’s recorded from SiC surfaces that underwent different sacrificial oxidation preparation procedures. They attributed this change to the modification of both the density of surface states and their position within the band gap. In view of that, the observed change in BE may be related to electronic changes of the substrate underneath the oxide. Thus, ion irradiation yields (in addition to morphological differences already observed by AFM and c-RBS) an electronic modification of the semiconductor region close to the SiO$_2$/SiC interface. Moreover, the observed energy shift depends on how much of the amorphous layer was oxidized: as the oxidation front approaches the single crystal, the closer to the nonirradiated sample is the BE value of the Si$^{\text{SiC}}$ component.

The BE of the Si$^{\text{ox}}$ components from samples N360, I150, and I720 as a function of the oxide thicknesses (etched step-by-step in the HF solution mentioned in Sec. II) is shown in Fig. 6. Curves were aligned by the point corresponding approximately to the oxide/SiC interface. A depth scale was built based on the etching time scale. In this ap-
FIG. 6. Binding energy of the Si\(^{2+}\) component of the Si 2p regions as a function of the remaining oxide thickness after successive HF etching steps of samples N360 (solid circles), I720 (open squares), and I150 (open circles). The depth scale was obtained assuming a fixed etching rate of the HF solution.

approach, the etching rate was assumed constant through the whole oxide length. This is an approximation, since this rate changes, particularly in the SiO\(_2\)/SiC interfacial region. However, this effect will not change the essence of our conclusions. The BE variation of this component can not be attributed to the detection of different compounds, since a synchronization of BE shifts of SiO\(_2\) components from Si 2p and O 1s regions is always observed. All curves evidence a similar trend, namely, a lowering of the BE as the interfacial region is approached. In addition, the BE values corresponding to sample I720 are always higher than those of sample I150 for the same oxide thickness. Next to its oxide surface region, sample I720 evidences BE values comparable to those observed in the analogous region of sample N360.

Differences between oxides thermally grown on Si and on SiC were investigated\(^{20}\) previously, and the authors claimed that the observed BE difference between Si\(^{2+}\) components of oxides grown on Si and on SiC were due to structural differences associated with the formation of the dielectric film: SiO\(_2\) films thermally grown on SiC (with smaller BE) appear to be under less stress than on Si (100) (with larger BE). The present results can be understood using the same reasoning developed in Ref. 20, namely, correlating BE values with the amount of stress. Thus, the oxide formed on sample I150 is the least stressed one, since the BE of its Si\(^{2+}\) component is the lowest. This is in agreement with what is expected from the oxidation of a more disordered material, since the bombarded SiC layer should induce a lower strain in the oxide as compared to the single crystal. The observed differences in stress manifest the distinct degrees of crystallinity reached by the damaged layers of samples I150 and I720. The substrate underneath the oxide in sample I150 recrystallized in a smaller extent than that of sample I720. This last sample, due to the longer oxidation, had more time to recrystallize and consumed all bombarded layer by oxidation yielding an oxide on a nearly single crystalline substrate. Thus, consistently with Ref. 20, the Si\(^{2+}\) BE at the SiO\(_2\)/SiC interface of sample I720 has values similar to those from sample N360, the one with the oxide layer grown on a c-SiC substrate. This assumption is also reinforced by the results obtained by c-RBS and AFM.

Si 2p regions of XPS spectra of samples N360, I150, and I720 recorded after each etching step were fitted by the above mentioned procedure. The area of each component was normalized by the total area of the respective Si 2p regions. These values were plotted as a function of the accumulated etching time in Fig. 7, yielding composition profiles throughout the oxide films. Data from sample I720 are not shown since they are very similar to those of sample N360. An approximate depth scale can be obtained assuming a constant etching rate as already described. The composition profile for sample N360 evidences expected facts: a SiO\(_2\) signal as the only component until a certain etching time, which decreases to a lower areal value due to progressive contribution from the substrate underneath the oxide. The lowest areal value of the SiO\(_2\) signal corresponds to the native oxide layer formed during sample exposure to air between successive analyses. The width of the transition region shown in Fig. 7 top (\(~10\) nm), defined by the range of etching times limited by near constant behavior of the components, is wider than the real SiO\(_2\)/SiC interface due to variations in the etching rate and to the finite probing depth of XPS (\(~1.5\) nm). Composition profile of sample I150 evidences a different interfacial region in comparison with sample N360. Besides the additional curve related to SiC\(_x\)O\(_y\) compounds, a wider transition region is observed (\(~18\) nm). The actual value is certainly lower, but it is anyway much larger than the one obtained from sample N360. We propose that the roughness of the SiO\(_2\)/SiC interface (observed by AFM) is responsible for this artifact. Due to roughness, the photoelectrons detected are in fact probing regions at differ-
ent distances relative to the interface and the resulting spectrum would be an average of regions more or less close to the SiC substrate. In addition to the wider transition region, the observed interfacial roughness gave rise to another artifact: the amplification of the signal from the Si$^{2+}$C component. SiC$_{2}$O$_{2}$ compounds were observed in much thinner oxides analyzed by angle-resolved XPS. In that work, clear signals of these compounds were only obtained with an experimental setup privileging the analysis of the sample near-surface region, using high take-off angles values ($>60^\circ$). In the present study, the amplified sensitivity was provided by the increased roughness of the SiO$_2$/SiC interface, which leads to an increased areal density of this interfacial probed region.

**IV. CONCLUSIONS**

In summary, we investigated structural, compositional, and electronic characteristics of thermally grown oxide layers on ion irradiated SiC substrates. Several differences were observed in comparison with a nonirradiated sample. The most striking effect observed is the enhanced rate of oxidation: using the same oxidation conditions, much thicker oxides were obtained on the irradiated sample. In addition to the enhanced oxidation rate, the atomic scale processes arisen from ion bombardment, which can affect the electrical characteristics of the SiO$_2$/SiC structure, were investigated. The roughnesses of both oxide surface and SiO$_2$/SiC interface were greater for samples submitted to ion irradiation. XPS analyses of different samples, in which oxide films were step-by-step thinned by chemical etching, evidenced electronic and structural differences induced by the irradiation process. Indeed the BE shifts of the Si$^{2+}$C component near the SiO$_2$/SiC interface are different from those in the same region of oxidized virgin SiC wafers. Irradiation also gave rise to less stressed oxides, as evidenced by the analysis of the BE values of Si$^{2+}$ components. The different structure of the semiconductor underneath the oxide and the irradiation-induced roughnesses may be responsible for the bad electrical characteristics of devices obtained following this route.

In addition to the differences attributed to irradiation itself, remarkable changes became evident when comparing irradiated substrates oxidized for different times. Distinct degrees of recrystallization were reached by the remaining damaged layer, which is progressively consumed in the course of oxidation. This recrystallization process gives rise to an increasing roughness of the oxide surface and SiO$_2$/SiC interface. When the whole damaged region is consumed, roughness decreases since further material consumption for oxide growth is now single crystalline. Hence, the electrical characteristics of the final device should depend on the thickness of this remaining damaged layer, which is a function of the oxidation time. A previous work concerning electrical characterization of metal-oxide-semiconductor structures built on irradiated SiC considered this route not suitable for the growth of gate oxides. However, in that work, only part of the irradiated layer (50%) was oxidized, which certainly influenced the observed electrical characterization results. Thus, the present findings do not exclude ion irradiation as a processing tool while pointing to the need of a total consumption of the damaged layer as an essential requirement for device quality oxides thermally grown on irradiated SiC. Following this reasoning, thick oxides grown with high oxidation rates and presenting the required electrical characteristics are expected.

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9 Electronic address: fernanda@iq.ufrgs.br


21 The amount of Si present in each oxide film was obtained from the amount of O determined as described in the experimental details section. This last quantity was divided by two (assuming a stoichiometric layer of SiO$_2$) yielding the areal density of Si. To determine the amount of Si in the irradiated layer before oxidation, we simulated the channeled RBS spectrum of the as-irradiated sample as shown in Fig. 2. Simulation using the RUMP
(Ref. 15) code consists of assuming a certain areal density of a material with a defined stoichiometry and stopping power; analyzed under certain experimental conditions (beam energy, beam ion, incident charge, detector geometry, resolution, and solid angle). Then, the program generates a theoretical spectrum that should be compared to experimental data. In this way we determined the areal density of Si in the amorphized layer, since in the single crystal the alignment of atoms respective to the incident beam makes the detected ion yield to decrease.