<table>
<thead>
<tr>
<th><strong>Title</strong></th>
<th>Origin and passivation of fixed charge in atomic layer deposited aluminum oxide gate insulators on chemically treated InGaAs substrates</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Author(s)</strong></td>
<td>Shin, Byungha; Weber, Justin R.; Long, Rathnait D.; Hurley, Paul K.; Van de Walle, Chris G.; McIntyre, Paul C.</td>
</tr>
<tr>
<td><strong>Publication date</strong></td>
<td>2010</td>
</tr>
<tr>
<td><strong>Type of publication</strong></td>
<td>Article (peer-reviewed)</td>
</tr>
</tbody>
</table>
[http://dx.doi.org/10.1063/1.3399776](http://dx.doi.org/10.1063/1.3399776)  
Access to the full text of the published version may require a subscription. |
| **Item downloaded from** | [http://hdl.handle.net/10468/4343](http://hdl.handle.net/10468/4343) |

Downloaded on 2018-12-21T13:48:59Z
Origin and passivation of fixed charge in atomic layer deposited aluminum oxide gate insulators on chemically treated InGaAs substrates


Citation: Appl. Phys. Lett. 96, 152908 (2010); doi: 10.1063/1.3399776
View online: http://dx.doi.org/10.1063/1.3399776
View Table of Contents: http://aip.scitation.org/toc/apl/96/15
Published by the American Institute of Physics

Articles you may be interested in

Controlling the fixed charge and passivation properties of Si(100)/Al2O3 interfaces using ultrathin SiO2 interlayers synthesized by atomic layer deposition
Journal of Applied Physics 110, 093715 (2011); 10.1063/1.3658246

Presence and origin of interface charges at atomic-layer deposited Al2O3/III-nitride heterojunctions

O3-sourced atomic layer deposition of high quality Al2O3 gate dielectric for normally-off GaN metal-insulator-semiconductor high-electron-mobility transistors

High-κ gate dielectrics: Current status and materials properties considerations
Journal of Applied Physics 89, 5243 (2001); 10.1063/1.1361065

Manipulating the negative fixed charge density at the c-Si/Al2O3 interface

Interface charge engineering at atomic layer deposited dielectric/III-nitride interfaces
Origin and passivation of fixed charge in atomic layer deposited aluminum oxide gate insulators on chemically treated InGaAs substrates

Byungha Shin,1,a Justin R. Weber,2 Rathnait D. Long,1,3 Paul K. Hurley,3 Chris G. Van de Walle,2 and Paul C. McIntyre1

1Department of Materials Science and Engineering, Stanford University, Stanford, California 94305, USA
2Department of Materials, University of California, Santa Barbara, California 93106, USA
3Tyndall National Institute, University College Cork, Lee Maltings, Prospect Row, Cork, Ireland

(Received 23 September 2009; accepted 23 March 2010; published online 15 April 2010)

We report experimental and theoretical studies of defects producing fixed charge within Al2O3 layers grown by atomic layer deposition (ALD) on In0.53Ga0.47As(001) substrates and the effects of hydrogen passivation of these defects. Capacitance-voltage measurements of Pt/ALD-Al2O3/n-In0.53Ga0.47As suggested the presence of positive bulk fixed charge and negative interfacial fixed charge within ALD-Al2O3. We identified oxygen and aluminum dangling bonds (DBs) as the origin of the fixed charge. First-principles calculations predicted possible passivation of both O and Al DBs, which would neutralize fixed charge, and this prediction was confirmed experimentally: postmetallization forming gas anneal removed most of the fixed charge in ALD-Al2O3. © 2010 American Institute of Physics. [doi:10.1063/1.3399776]

In recent years, dimensional scaling of complementary metal-oxide-semiconductor (CMOS) devices has brought about serious interest in introducing high-mobility channel layers composed of III–V materials.1 A longstanding problem for structures that include an oxide/III–V semiconductor interface has been the presence of interfacial defects that can trap charge during device operation. However, defects present in the bulk of a high permittivity (high-k) oxide layer that forms a relatively passive interface with III–V channel materials may be of equal importance because of their potential to increase gate leakage current, and to form fixed charge that will scatter carriers in the channel and alter the threshold voltage of the device.2

Many of the high-k dielectric materials investigated to replace SiO2 in Si-channel MOS devices are grown as thin films with significant areal densities of fixed charge.3 However, the density of fixed charge either near the interface between ALD-Al2O3 and the channel or in the bulk of Al2O3 layers deposited on III–V materials such as InGaAs has not been studied carefully. In this paper, we report experimental and theoretical studies of defects producing fixed charge2 within Al2O3 layers grown on (001) In0.53Ga0.47As channel layers and the effects of hydrogen passivation of these defects.

The starting In0.53Ga0.47As surface was treated with 4 vol % NH4OH(aq) solutions, followed by thermal desorption of residual As from the wet etching step at 380 °C prior to atomic layer deposition (ALD).5 Al2O3 films of different thicknesses, 2.4–16 nm, were prepared by ALD at 270 °C using trimethylaluminum and H2O precursors. Film thicknesses were determined by ellipsometry, the readings of which were calibrated by cross-sectional transmission electron microscopy. For capacitance-voltage (CV) measurements, Pt top electrodes were deposited in an e-beam evaporator through a shadow mask. Postmetallization forming gas anneal (FGA) was carried out under forming gas (95% N2 +5%H2) flowing at a rate of ~2 L/min at 400 °C for 50 min. The dielectric constant of the Al2O3 films was determined—by linear fitting of the inverse of the maximum capacitance at accumulation at 800 kHz versus film thickness—to be 8.2 ± 0.5 and 6.8 ± 0.1 for the as-deposited and post-FGA films, respectively.6

First-principles calculations based on density functional theory were performed for κ-Al2O3; the κ phase was chosen because of the similarity of its density7 to that of amorphous Al2O3. We employed a hybrid-functional scheme8 as a way of overcoming the band-gap problem. We utilized projector-augmented-wave pseudopotentials9 and a plane-wave basis set with a 300 eV plane-wave cutoff as implemented in the VASP code.10 The percentage of Hartree–Fock exchange mixed into the hybrid functional was chosen to be 32% in order to optimally reproduce the experimental gaps of Al2O3 as well as InAs, GaAs, and InGaAs. The calculated Al2O3 gaps are 7.9 eV for the κ phase and 9.1 eV for the α phase, the latter in good agreement with experiment (8.8 eV, Ref. 11). There are no known experimental values for the band gap of κ-Al2O3. The band gap of the amorphous oxide has been measured to be 7.36 eV.12 This is slightly lower than our calculated κ-Al2O3 band gap but this difference does not affect our conclusions.

To investigate the spatial distribution of fixed charge in as-deposited Al2O3 films on In0.53Ga0.47As, CV curves from MOS capacitors with different Al2O3 thicknesses were collected and are shown in Fig. 1(a). The thickness series exhibits a considerable negative shift of the flat-band voltage (VFB) with increasing thickness, indicating the presence of net positive fixed charge in the as-deposited Al2O3. A closer examination of Fig. 1(a), however, reveals that the VFB’s of thinner films (30, 45, 60 TMA/H2O cycles) are more positive than the expected ideal VFB (0.6–1.0 V, depending on the value of Pt work function and the exact substrate dopant concentration assumed in the estimation), indicating that the thinner films contain negative fixed charge. With increasing film thickness, VFB becomes comparable to the ideal VFB (after ~90 cycles of deposition), and then shifts to negative polarity for thicker ALD-Al2O3 layers. The observed...
thickness-dependence of $V_{FB}$ can be rationalized if there is both negative fixed charge near the interface in the In-GaAs channel and positive bulk fixed charge in the Al$_2$O$_3$ films. This nonuniform distribution of fixed charge across the thickness of the Al$_2$O$_3$ films may result from a local variation in aluminum:oxygen stoichiometry. Table I lists the ratio of Al 2p to O 1s X-ray photoelectron spectroscopy (XPS) core level peak intensity for various Al$_2$O$_3$ layer thickness. A take-off angle (with respect to the sample surface) of 90° was used and, for the film thicknesses studied, peaks from the substrate (i.e., In, Ga, and As) were observed, meaning that the Al 2p and O 1s peaks include contributions from the entire thickness of each Al$_2$O$_3$ film. In previous work, we observed an interfacial In oxide from an AlD-Al$_2$O$_3$/In$_{0.53}$Ga$_{0.47}$As sample which was prepared identically to those used in this study. However, any contribution from this interfacial In oxide to the total O 1s peak intensity is negligible because its areal coverage is only ∼12% and it is buried under an Al$_2$O$_3$ layer which attenuates most of O 1s photoelectrons from the interfacial In oxide. A clear trend of increasing Al:O atomic ratio with increasing film thickness is evident, and this suggests that the films are O-rich near the interface and become progressively more Al-rich farther away from the interface. Therefore, it appears that the sign of fixed charge is correlated with the local stoichiometry of the films, such that negative fixed charge exists within O-rich regions near the interface and positive fixed charge within Al-rich regions away from the interface.

The effect of local variations in stoichiometry was investigated using first-principles calculations as described above. In an amorphous oxide, Al deficiency may predominantly manifest itself in the form of oxygen dangling bonds (DBs), and O deficiency in the form of Al DBs. We have therefore performed a study of the atomic and electronic structure of DBs in Al$_2$O$_3$, using geometries previously applied to group-IV semiconductors. The DBs were studied in an 80-atom bulk $κ$-Al$_2$O$_3$ supercell with occupancies of zero, one, and two electrons, as well as with hydrogen passivating them. As shown in Fig. 2, the oxygen DB gives rise to charge-state transition levels at $−0.83$ eV and $0.61$ eV above the valence-band maximum (VBM) of Al$_2$O$_3$, while the Al DB produces levels at $5.12$ and $5.35$ eV above the VBM. Therefore, using a VB offset of $2.8$ eV between In$_{0.53}$Ga$_{0.47}$As and Al$_2$O$_3$, we conclude that O DBs will always be below the Fermi level and thus negatively charged, while Al DBs will be positively charged. This is consistent with the experimental findings that O-rich regions have fixed negative charge, while Al-rich regions have fixed positive charge.

We have also studied hydrogen passivation by comparing calculated total energies of the passivated DBs with un-passivated DBs plus interstitial hydrogen. The resulting binding energies are $1.3$ eV for O–H and $1.4$ eV for Al–H, indicative of stable passivation. Hydrogen passivation neutralizes the DBs, i.e., the O–H and Al–H bonds occur exclusively in the neutral charge state over the relevant range of Fermi levels. We note that complete neutralization may not be possible in the case of vacancies that contain multiple dangling bonds, although addition of hydrogen will still reduce the amount of fixed charge per vacancy.

The prediction that hydrogen can effectively passivate DBs in the oxide can be tested by comparison to experimental data. CV curves from the samples, which received FGA at $400$ °C after electrode deposition, are shown in Fig. 1(b). The flat band voltage, which exhibited a wide variation depending on the Al$_2$O$_3$ layer thickness of the as-deposited films, was shifted in all cases to a value very close to the ideal $V_{FB}$ after the FGA, suggesting that annealing in hydrogen can compensate most of the positive bulk fixed charge and the negative interfacial fixed charge. In addition to the compensation of the fixed charge in Al$_2$O$_3$ by the FGA, we notice significant improvements in the CV characteristics—such as a decrease in frequency dispersion in depletion and accumulation, and a decrease in CV stretch-out, indicative of passivation of interface states by the FGA [compare Figs. 3(a) and 3(b)]. We would like to point out that enhancement of high-$κ$/In$_{0.53}$Ga$_{0.47}$As interface quality by FGA is not limited to an Al$_2$O$_3$/In$_{0.53}$Ga$_{0.47}$As interface; the reduction in interface state density has been observed from an ALD-HfO$_2$/n-In$_{0.53}$Ga$_{0.47}$As interface after postmetallization FGA at $325$ °C.15

![FIG. 1. (Color online) CV curves from Pt/ALD-Al$_2$O$_3$/n-In$_{0.53}$Ga$_{0.47}$As MOS capacitors with various Al$_2$O$_3$ layer thicknesses: (a) before, and (b) after postmetallization FGA at 400 °C. Estimated range of ideal $V_{FB}$ (0.6–1.0 V) is indicated by vertical gray bars in both figures.](image)

![FIG. 2. (Color online) Band alignment between Al$_2$O$_3$ and relevant semiconductors, and position of charge-state transition levels for dangling bonds in the oxide.](image)

### Table I. Ratio of Al 2p to O 1s XPS peak intensity for different Al$_2$O$_3$ film thicknesses.

<table>
<thead>
<tr>
<th>Al$_2$O$_3$ film thickness (nm)</th>
<th>2.4</th>
<th>2.8</th>
<th>3.6</th>
<th>4.8</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ratio: Al 2p peak intensity to O 1s peak intensity</td>
<td>0.141</td>
<td>0.145</td>
<td>0.154</td>
<td>0.157</td>
</tr>
</tbody>
</table>
The authors acknowledge support from SRC (NonClassical CMOS Center Task No. 1437,003 and Grant No. 2009-VJ-1867) and Intel Corp. RDL acknowledges support from Irish Fulbright Commission and Fulbright USA.


4The term, “fixed charge” in this article should be distinguished from conventional interface states whose charging state varies depending on the position of the semiconductor Fermi level.


6There is an apparent decrease in the dielectric constant after the FGA, which we attribute to the incorporation of hydroxyl groups as revealed by the comparison of the O 1s XPS spectra before and after the FGA. A similar phenomenon (an inverse correlation between dielectric constant of ALD-Al2O3 and H content) was observed by other group. See, M. D. Groner, J. W. Elam, F. H. Fahreguette, and S. M. George Thin Solid Films 413, 186 (2002).


