

Epitaxial lanthanide oxide thin films on Si for high-k gate dielectric application: Growth optimization and defect passivation

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Epitaxial layers of insulating binary lanthanide oxides have been considered as potential alternative to conventional SiO_2 for gate dielectric application in future Si-based MOSFET devices, which was investigated in more detail for epitaxial Gd_2O_3 and Nd_2O_3 as model systems. Additionally, the ability to integrate epitaxial dielectric barrier layers into Si structures can usher also in a variety of novel applications involving oxide/silicon/oxide heterostructures in diverse nanoelectronic and quantum-effect devices. Although epitaxial layers of such ionic oxides with excellent structural quality can be grown using molecular beam epitaxy, they often exhibit poor electrical properties such as high leakage current density, flat band instability, poor reliability etc. owing to the presence of electrically active charge defects, generated either during the oxide layer growth or typical subsequent CMOS process steps. Based on the origin and individual character of these defects, we review various aspects of defect prevention and passivation which lead to a significant improvement in the dielectric properties of the heterostructures.

I. INTRODUCTION

The physical dimension of metal oxide field effect transistors (MOSFETs) which are essential components of semiconductor integrated circuits (ICs) have been subjected to continuous downscaling since the birth of ICs to improve the device performance and reduce the production cost per bit. This shrinking of the semiconductor device dimension follows an empirical trend popularly known as Moore's law, which describes that the number of transistors per IC, or in other words processing capacity and speed doubles, every two years. Moore's prediction was found true for a long while thanks to the rapid advancement of semiconductor device processing technologies such as lithography, selective etching processes etc. On the other hand from the materials aspect, the key material property that enabled the existence of Si-MOSFET devices is the excellent insulating characteristic of SiO₂ thin layers which is used as the gate dielectric to isolate the transistor gate from the Si channel. This is possible due to the large band gap of SiO_2 ($E_g \sim 9$ eV), and the low density of interface traps with Si. For example, in advanced integrated devices SiO_2 gate dielectrics with charge densities of 10^{10} cm⁻², midgap interface state densities of 10^{10} cm⁻², and dielectric strength of

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layer of SiO₂ can be grown by simple thermal oxidation that remains amorphous at typical CMOS process condition which is less prone to any structural defects. However, it was evident that continuing the device scaling as per Moore's law cannot be pushed longer if SiO₂ has to be used further as the gate dielectric. The primary disadvantage of using SiO₂ as the gate dielectric is its relatively low dielectric constant ($k \sim 3.9$). This imposes significant reduction in the thickness of the SiO2 layer as a consequence of downscaling the MOSFET devices to achieve higher gate capacitance necessary to produce the required drive current in submicron MOSFETs. For example, the specific gate capacitance used for todays CMOS technology nodes demands SiO_2 thickness < 1 nm. Unfortunately, such ultrathin layers of SiO₂ cannot act really as an insulator, as direct tunneling through the dielectric barrier dominates the conduction behavior and the leakage current increases exponentially with decreasing gate dielectric layer thickness.⁴ For example, scaling of the gate oxide below 1 nm leads to leakage current density of $\sim 10 \text{ A/cm}^2$ at 1 V which is beyond the acceptable limit for low power device applications. It has also been demonstrated that fabricating CMOS devices with SiO₂ layer thickness in the range of 1.0-1.2 nm does not lead to any further gain in the drive current setting a fundamental limit for gate oxide scaling.⁵ Moreover, in contrast to high performance microprocessors, the leakage current densities of transistors in case of low power consumption devices should be well below 10^{-1} A/cm², making the gate oxide thickness

15 kV/cm have been produced.^{2,3} Further, high quality

issue more stringent. There have been additional limiting factors to SiO₂ scaling, such as boron diffusion from traditional highly doped poly-Si to lightly doped channel layer that results in shift in the transistor threshold voltage and degraded device reliability owing to hot electron injection at the Si-SiO2 interface causing dielectric breakdown of the gate oxide.8 These issues related to material limited device scaling is most prominent in case of MOSFETs than in any other device components and became the showstopper in the historical progress of multibillion semiconductor industry.² In this context, one of the major challenge associated with further scaling of integrated devices was to find suitable replacement gate insulator material that possesses higher dielectric constant than SiO₂ and therefore can be grown with larger thickness which shall allow sufficient FET current drive while maintaining lower leakage current.9

II. CHOICE OF ALTERNATIVE GATE INSULATOR

SiO₂ is the key material that laid the foundation of Si based microelectronics technology and led it to flourish despite the average performance of Si compared to other semiconductors, such as Ge, GaAs, GaN, SiC, etc. which do not form a thermally stable native oxide. Integrating an alternative dielectric to Si would be possible only at the expense of the advantages of SiO2, which makes the choice of alternative gate dielectric a critical issue. The semiconductor industry first found interim solution remaining within the Si-SiO₂ materials framework by selecting silicon oxynitride and oxide/nitride stacks as alternative gate dielectrics which offer slightly higher k value compared to SiO_2 and thus lead to reduced leakage, boron penetration, and reliability issues by using relatively larger gate dielectric layer thickness. 10 However, this was only a temporary solution as incorporation of increasing amount of N gives rise to side effects, such as severe degradation of electron mobility in the transistor channel and reduced band gap of the oxide.⁸ Therefore, the search of really alternative dielectric materials with much higher dielectric constant was on the agenda to keep pace with the historical steep progress of Si technology. The first parameter to be considered for selecting an alternative material is the appropriate value of the dielectric constant that can be useful for transistor application. While leakage current through the dielectric layer in the direct tunnel regime depends exponentially on its thickness, the capacitance varies linearly. Consequently, dielectrics with too high dielectric constant lead to short channel performance degradation, owing to the fringing field from the gate to source/drain region due to large thickness to length aspect ratio of the gate dielectric. 11 Optimum k values within the range of 15 and 50 are predicted to be ideal for continuing scaling of transistors in the future, which shall be applicable for

both low standby power and high performance devices. Besides suitable higher k value, any material selected as alternative to substitute SiO_2 as gate dielectric will also have to satisfy several other criteria to be eligible for gate dielectric application, such as thermodynamic stability in direct contact with Si, thermal stability even at typical CMOS process temperatures of up to $1000 \,^{\circ}$ C, large band gap, low bulk defect density, low interfacial trap densities, CMOS process compatibility and reliability.

A wide variety of insulating metal oxides with much higher k values has been investigated in the past decade as viable alternative to Si-oxynitride and oxide/nitride systems. Many of these metal oxide alternative dielectrics appear to satisfy some of these requirements, but unfortunately only a very few materials are promising with respect to all of the guidelines. Furthermore, the process complexity associated with deposition of the high-k metal oxides in contrast to simple thermal deposition of SiO_2 and oxynitrides pose additional challenges for integrating the alternative dielectrics into mainstream Si technology, such as cost of the deposition tools for large scale production with proper stoichiometry and satisfactory microstructure control, in addition to the requirement of using metal gates, etching and related aspects. 12

Different binary metal oxides have been considered promising because of the relative ease in controlling the stoichiometry in their thin films. But a number of these oxides are not stable in contact with Si at T > 1000 K and gives rise to an interfacial layer formation. 13 Interfacial layer exhibiting lower k value eventually leads to degradation of the effective dielectric constant and limits the scalability of the device. Another critical parameter to consider is the band gap of the metal oxides, which is mostly much smaller than that of SiO₂ (~9 eV), that directly influences their conduction band and valence band offset (VBO) energies with respect to Si. This is related to the ionic character of the metal oxides in contrast to covalent SiO₂. A number of representative dielectric oxides have been found to obey an approximate inverse relation between their static dielectric constant and band gap. This behavior is expected because stronger polarizability implies weaker bonding which leads to smaller separation in bonding and antibonding energies.¹⁴ The band offsets, which determine the electron and hole injection barriers respectively at the oxide/Si interface, should be at least 1 eV (for both conduction and valence bands) to prevent charge carrier injection from Si into the bands of the chosen oxide. 15 Figure 1 illustrates the calculated band offsets of several promising high-k dielectric oxides with respect to Si. It is clearly evident that only a few of the high k oxides are suitable candidates for gate dielectric application, despite superior dielectric constant values of several other oxides.

An additional roadblock for integrating high-k metal oxides into Si technology is their lower crystallization

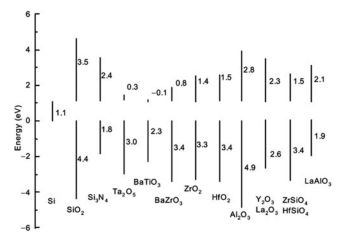


FIG. 1. Calculated conduction band and valence band offsets of various oxides on Si. (Reprinted with permission from Ref. 15. Copyright [2000], American Vacuum Society).

temperatures. Following the trend with SiO₂, the replacement gate dielectric is expected to involve amorphous metal oxides to avoid surface roughness and additional leakage due to the formation of grain boundaries as expected in crystalline material. The higher atomic coordination numbers and ionic character of the metal oxides render them poor glass formers.¹⁶ Further, with increasing dielectric constant the crystallization temperature of the oxides also reduces due to increased ionicity, and even the recently preferred binary oxides (e.g., HfO₂, ZrO₂) do not withstand crystallization at typical CMOS processing temperatures. Alloying with Si or Al (i.e., silicates or aluminates) results in superior thermal stability and higher crystallization temperature, 12,17-19 however, again at the expense of the highest achievable dielectric constant.

Based on the dielectric performance and process condition optimization, the first generation of high-k dielectrics was expected to consist of Hf-based alloys. Interface engineering schemes have been developed to form oxynitride and oxide/nitride reaction barriers between these high-k oxides and Si to minimize the reaction at the interface.²⁰ The amorphous barrier layers used for Si surface passivation was not only promising for minimizing interfacial reaction but to maintain also high channel carrier mobility, which could otherwise be affected by the interfacial defects and the soft phonons of the high-k oxides in direct contact with Si. In 2007 Intel successfully demonstrated for the first time a 45 nm logic technology using Hf based high-k metal gate stack transistors in a high volume manufacturing process, where an equivalent oxide thickness (EOT) of \sim 1 nm was realized.²¹ This has gained wide adoption in the industry.6 While the integration of Hf based gate dielectric to the CMOS device process represented a remarkable success, it is obvious that the lowest possible EOT achievable using alternative

dielectric material cannot be realized in presence of the interfacial layer due to their lower k values.

Therefore, clearly further downscaling of CMOS devices must involve usage of high-k oxide materials which should be integrated to Si without any interfacial layer to realize EOT <0.8 nm. Therefore, it is certainly not exaggeration to state that the quality of the interface between the high-k oxide and Si will be the most important parameter to impact the device performance. A good interface quality can be expected either if the oxide is amorphous so that the number of Si dangling bonds at the interface can be minimized by local bonding adjustment, or if it is lattice matched to Si and grown epitaxial with atomically sharp interface. Epitaxial grown high-k oxides could be a smart choice in this regard since this approach has the advantage of defined interface engineering and hence offers more control over the device properties despite larger process complexity related to the epitaxial growth process.

III. EPITAXIAL GATE INSULATORS: CHOICE OF ALTERNATIVE METAL OXIDES

In the recent past various epitaxial complex oxides, especially insulating binary rare earth oxides (REOs, mainly lanthanide oxides, LnOs), have drawn significant interest as prospective 2nd generation of alternative gate dielectrics.²² Insulating LnOs form a very interesting group of materials for integration on Si⁴ as they offer several advantages, such as large dielectric constant, large band gap and band offsets to Si, superior thermal and thermodynamic stabilities, which make them suitable for Si-based technologies. 13,22 However, the choice of appropriate LnOs for gate dielectric application depends strongly on their stoichiometry. For example, some rare-earth metals exhibit multiple oxidation states (+2, +3,and +4) which lead to oxides with different stoichiometry (LnO, Ln₂O₃, LnO₂),²³ where not all the compositions are insulating. Moreover, the position of the charge neutrality level of ionic oxides depends strongly on their stoichiometry.²⁴ Therefore, their band alignment to silicon and, finally, the leakage behavior becomes strongly dependent on the oxygen content. Therefore, LnOs which exhibit more than one stoichiometry (e.g., Ce-, Eu-oxides) or those which exhibits stable mixed valence structure (e.g., Pr₆O₁₁) are also not the best choices as alternative high-k materials in Si-based devices. Consequently, for practical application it is wise to focus on those insulating LnOs in which the Ln metal possesses only a single valence state. Based on these criteria we focused mainly on LnOs containing Ln³⁺ cations (exhibiting the composition Ln₂O₃).

Additionally, Ln₂O₃ oxides can occur in different crystal structural phases (polymorphs) depending on temperature, e.g., cubic bixbyite structure (Mn₂O₃ type),

hexagonal lanthanum oxide structure (La_2O_3 type), and monoclinic structure (Sm_2O_3 type). The different structures exhibit different dielectric properties. Since temperature-dependent crystal phase transitions are not reversible or not completely reversible, and accompanied by interfacial layer formation on Si, the controlled tuning of dielectric properties is difficult for Ln_2O_3 s which can undergo structural phase transformation already within a temperature range typical for CMOS processing²⁵ which make them not suited for gate dielectric applications.

Based on these criteria we focus mainly on Gd₂O₃ (and to some extent on Nd2O3) for its epitaxial integration on Si as 2nd generation alternative high-k dielectric material. Bulk Gd₂O₃ occurs in the bixbyite cubic type of structure with single stoichiometry and structural transformation occurs only at $T > 1400 \text{ K.}^{26}$ In addition, this oxide has a large band gap of about 6 eV and nearly symmetrical band offsets to Si, ²⁷ a dielectric constant of 14 as well as a low lattice mismatch ($\sim 0.5\%$) with Si. The bixbyite structure of Gd₂O₃ has lattice symmetry suitable for epitaxial growth on Si(100) and Si(111). Although the lattice mismatch between Gd₂O₃ and Si are identical for (100)//(100), (110)//(110), and (111)//(111) epitaxial arrangements based on the symmetry matching, it is to be noted that in practice Gd₂O₃ grows with (111) orientation on Si(111), whereas (110) oriented growth of Gd₂O₃ is observed on Si(100) surface under usual epitaxial growth conditions. A detailed discussion of the impact of symmetry matching and surface energy criteria on the epitaxial relation for Gd₂O₃ growth on Si(100) can be found elsewhere. 22,28

A. Epitaxial growth

The epitaxial layers of Gd₂O₃ on different Si surfaces were grown by evaporating commercially available stoichiometric granular Gd₂O₃ using electron gun in a cluster solid source cluster molecular beam epitaxy system that includes a growth, an annealing, and an analytic chamber connected by an ultra-high vacuum (UHV) transfer system. The layers were grown on silicon substrates with different orientations. Substrates were cleaned ex situ using the last step diluted HF etch followed by a dilution rinse, and then were immediately inserted into the vacuum system. Growth temperatures were in the range of 800–1000 K. Typical growth rates were 0.005-0.01 nm/s. For some growth experiments, additional molecular oxygen was inserted in the chamber without any activation to yield an oxygen partial pressure $(P_{\rm O_2})$ in the range of 10^{-7} mbar. For the fabrication of metal electrodes, the wafers were transferred immediately after the growth in vacuo into another chamber, where metal dots were evaporated through a shadow mask using electron beam heating. The surface and layer structure was evaluated during the growth by

reflection high-energy electron diffraction (RHEED). The chemical natures of the oxides and the band alignment with Si have been investigated using in situ X-ray photoelectron spectroscopy (XPS) in a separate chamber without exposing the layers to atmosphere. Ex situ, the layers were characterized by X-ray diffraction (XRD) ($\theta/2\theta$, ω and Φ -scans), by X-ray reflectivity for layer thickness determination (XRR) and by transmission electron microscopy (TEM). Figure 2 shows the X-ray (θ –2 θ scan) diffraction patterns of Gd₂O₃ thin films grown on different Si substrates. The scan for Gd₂O₃ layer on Si(001) surface exhibits a distinct peak at $2\theta = 46.4^{\circ}$ corresponding to the d(440) inter planar spacing of cubic Gd₂O₃ along [110] orientation. The layers on the Si(111) substrate also exhibit single orientation without any indication of disoriented crystallites. The peaks at 28.5° and 59.4° corresponding to d(222) and d(444) interplanar spacing are hidden under the appropriate Si peaks. The scan for Gd₂O₃ thin films grown on Si(110) displays a peak at 28.5° confirming the preferential growth along (111) direction.

IV. DEFECTS IN EPITAXIAL HIGH-*k* RARE EARTH OXIDES: SELECTING OPTIMUM GROWTH CONDITION

The existence of various defects inside the layer and/or at the oxide/Si interface often degrades the device performance. Therefore, it is of paramount importance to understand the nature of the defects inside the metaloxide stack and at the oxide/Si interface and address their effective passivation for successful integration of epitaxial LnOs to typical CMOS technology.

Defects in the epitaxial LnOs can be broadly classified into two categories, e.g., structural defects and charge defects. The structural defects include domain boundary, misfit dislocation, stacking faults etc., whereas the charge defect encompasses oxide fixed charge, mobile oxide charges, and interface traps. In the present article we shall

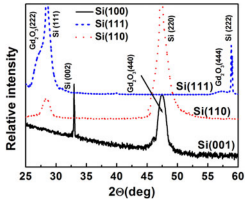


FIG. 2. X-ray diffraction patterns of Gd_2O_3 thin films grown on Si substrates with different orientation.

focus mainly on the charge defects normally encountered in the epitaxial LnOs, their effect on the electrical properties of these heterostructures, and their effective passivation. It should be noted that to understand the role of the charge defects in the LnO thin films it is necessary to realize good structural quality of the epitaxial layers and atomically sharp interfaces with Si. This is because in addition to the charge defects, the quality of interface between the oxide layer and Si surface also significantly influences the dielectric properties of the heterostructures. As a case study to understand the interface formation between the lanthanide oxides and Si, Fissel et al. reported a detailed investigation of the interface formation between Nd₂O₃ and Si(100) during epitaxial growth.²⁹ The RHEED investigation during MBE growth under UHV condition ($P_{O_2} < 5 \times 10^{-8} \text{ mbar}$) revealed that during the initial stage of oxide layer deposition the surface reconstruction of Si(100) changes from usual (2 \times 1) of a clean surface to an unreconstructed (1×1) . Further deposition leads to additional fractional order RHEED spots appearing at the 1/3 positions between the main (0,1) spots of the zero order Laue zone corresponding to a 3-fold surface periodicity at the surface [Fig. 3(a)]. Similar observation was already reported for the epitaxial growth of Pr₂O₃ on Si(001).³⁰ During subsequent deposition, the surface changes back to the (1×1) structure before bulk spots appear related to diffraction from oxide islands [Fig. 3(b)]. The double spots in Fig. 3(b) correspond to two in-plane lattice constants nearly equal to the lattice constant of silicon and $\sqrt{2}$ times of that constant, respectively. This indicates that similar to other LnOs crystalline Nd₂O₃ grows in (110)-oriented domains, with orthogonal in-plane orientations. This is also evident from the X-ray diffraction pattern in the θ –2 θ geometry, where only the (440) diffraction peak from the oxide layer is visible for the growth on Si(100) (Fig. 4).

The Nd₂O₃/Si interface formation was investigated in situ by XPS at different stages of the oxide layer growth. For layers with a thickness in the range up to 1 nm, where the 3-fold periodicity in RHEED was found, the Si 2p core level binding energy mainly corresponds to elemental Si (Si⁰ state) (Fig. 5). However, additionally a significant broadening was observed also toward the low-energy side of the Si⁰ peak. Deconvolution of the spectra by Gaussian fit indicated that for layers with thickness \leq 1 nm the broadening can be explained by the additional occurrence of a Si¹⁻ bonding state due to the formation of Si-Nd bonds in the initial stage of growth. The relatively small shift in the peak position indicates the well-known NdSi₂ phase formation. ³² This finding agrees to an earlier report by Norton et al., where the formation of silicide like surface structure was also associated with the appearance of additional 1/3-order reflections on a (2×1) reconstructed surface during Sr deposition on Si(100).³³ With increase in the Nd₂O₃ layer thickness of above 1 nm the spectra exhibit additional peaks at higher binding energies which could be attributed to the different kinds of silicate like bonds.³⁴ The corresponding

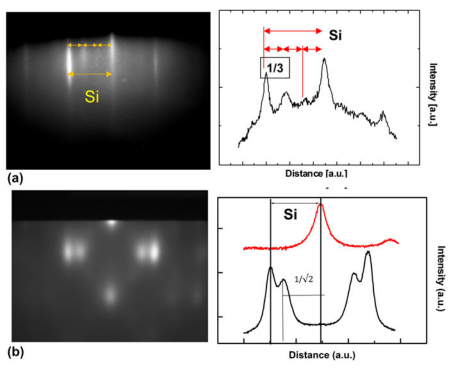


FIG. 3. RHEED pattern corresponding to Nd_2O_3 layer on a Si(100) wafer (a) 0.5 nm Nd_2O_3 and (b) 2 nm Nd_2O_3 layer thickness. The right hand side displays the intensity profiles along the marked line across the RHEED patterns.

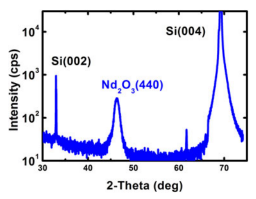


FIG. 4. X-ray diffraction pattern in the θ – 2θ geometry of a 12 nm Nd_2O_3 layer grown on a Si (001) substrate.

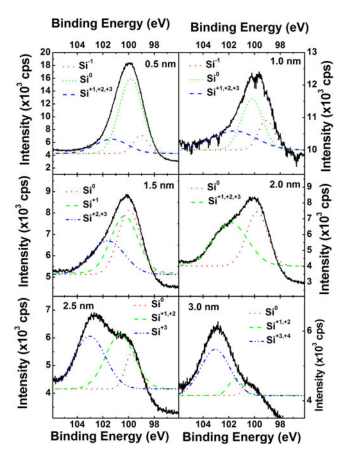


FIG. 5. Si 2p spectra together with the appropriate multi peak fitting results of a Si wafer at different stages of Nd_2O_3 layer growth.

neodymium $3d_{5/2}$ core-level spectra also supports the silicide-like bond formation at the interface during initial stage of the oxide layer growth (Fig. 6). For the second layer, a shift toward lower binding energies is observed which can be expected if the bonds become metal-like. That behavior together with the observed shifts in the Si spectra can only be explained by an exchange of oxygen bound to silicon by neodymium or by a movement of

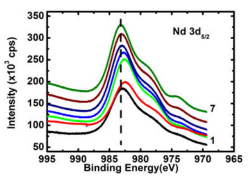


FIG. 6. Nd $3d_{5/2}$ core-level XPS spectra for Nd₂O₃ layers of various thicknesses ranging between 0.5 nm (Spectra 1) to 3 nm (spectra 7). The dashed line represents the expected peak position for bulk Nd₂O₃.

oxygen atoms from a Si-O-Nd interface site into the layer during the initial stage of growth. TEM investigations also support this assumption. Figure 7 shows TEM micrographs of an Nd₂O₃/Si(001) heterostructure grown under UHV conditions. The plan-view micrograph exhibits typical Moire fringes originating from interfacial inclusions [Fig. 7(a)]. The crystalline inclusion with the interface extending some nanometers into the Si substrates is clearly seen in the cross-sectional micrographs [Fig. 7(b)]. Additionally, voids were also found in such layers. Based on a detailed analysis of the TEM results, the crystallographic structure of these inclusions could be identified to be identical to the known structure of tetragonal NdSi₂.35 Other LnOs deposited on Si under identical growth condition exhibits similar results. For LnOs growth, the formation and stability of the silicidelike phase depend on the oxygen chemical potential.³⁶ Considering the low P_{O_2} under UHV conditions (used in standard MBE growth), the chemical potential of oxygen can also become negative which can facilitate silicide formation over oxide formation. The silicide growth can continue as long as the oxygen chemical potential remains strongly negative due to low oxygen content. This effect predominates at a surface or interface region where the thermodynamic equilibrium is perturbed by different factors, such as stress.

However, such silicide inclusion at the Ln₂O₃/Si interface can potentially degrade their dielectric properties, especially in case of thin layers. Therefore, for high-*k* dielectric application the interfacial silicide inclusion has to be avoided. One possible means to circumvent the silicide inclusions is annealing the deposited layers in an oxygen-rich atmosphere. However, such post deposition annealing normally gives rise to an amorphous interfacial layer. Fissel et al. reported that such interfacial layer consists of two regions (Fig. 8), one with lighter and one with darker contrast which was attributed to two different silicates.²⁹

These results indicate that to achieve a smooth interface between the epitaxial Ln₂O₃ and Si, further growth

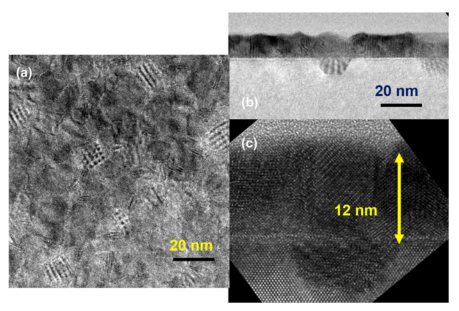


FIG. 7. TEM images of a Nd_2O_3 layer grown on Si(001), (a) plane-view micrograph, exhibiting pronounced Moiré fringes and (b, c) cross-sectional micrographs with different magnifications exhibiting the formation of crystalline interfacial inclusions. The two $Nd_2O_3(110)$ domains are indicated by white lines parallel to the atomic planes.

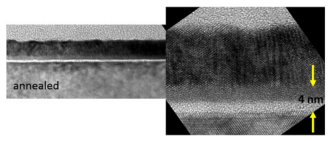


FIG. 8. Cross-sectional TEM micrograph of the Nd_2O_3 layer shown in Fig. 7, after annealing in nitrogen at a pressure of 2×10^{-2} mbar, and temperature if 600 °C for 10 min. In addition to the disappearance of the silicide inclusions, formation of an amorphous interfacial layer is visible.

engineering is necessary. In this view, a simple approach is to vary systematically the oxygen chemical potential by variation of the $P_{\rm O_2}$ during oxide layer. This has been demonstrated by Czernohorsky et al., where a modified MBE growth process was developed in which the chemical potential of oxygen was optimized with respect to the electrical properties by introducing additional nonactivated molecular oxygen during growth to keep the partial pressure in the range of 10^{-7} mbar.³⁷ This was found to result in a sharp interface between Gd₂O₃ and Si (Fig. 9), free from any silicide inclusions as well as silicate or silicon oxide. Therefore, such layers are ideal candidates for investigating the manifestation of charge defects and their influence on the dielectric properties of epitaxial Ln₂O₃s.

Preliminary investigation of electrical properties of epitaxial layers grown under optimized conditions exhibited very promising characteristics, such as low

capacitive equivalent thickness and low leakage current density (CET < 1 nm, $J \approx 10^{-1}$ A/cm²). Gottlob et al. reported CET ~0.86 nm and a leakage current density of ~ 0.5 A. cm⁻² at 1 V gate bias for MOS stacks consisting of ultra-thin Gd₂O₃ layers with CMOS-compatible fully silicided nickel-silicide (FUSI-NiSi) metal gate electrodes yields. 38 The CET trend observed for the MOS structures with Gd₂O₃ gate dielectric and FUSI-NiSi electrode combination met all ITRS targets for oxide thickness as well as leakage current values for high performance (HP), low operating power (LOP), and low standby power (LSTP) applications (Fig. 10). The reported CET values (without quantum-mechanical corrections) represent an upper limit of the achieved effective oxide thickness (EOT). Assuming a quantum mechanical correction of 0.3 nm to obtain EOT values shifts the trend line to an even more desirable region of specifications. These results indicated that NiSi/Gd₂O₃ gate stacks with optimized oxide/Si interface properties could be quite promising candidates for scaling efforts to the limit of CMOS. However, despite the outstanding structural qualities, compatibility with CMOS process techniques including metal gate incorporation and encouraging electrical properties the epitaxial Gd₂O₃ layers (or for that matter other insulating Ln₂O₃s) suffer from several performance related issues which are required to be addressed prior to real device integration. For example, the MOS structure exhibiting EOT \sim 0.86 nm shows strong hysteresis (~120 mV) in the capacitancevoltage (C-V) profile when the gate bias between forward and backward sweeps at 500 kHz. The hysteretic feature of the C-V profile indicates presence of charge defects inside

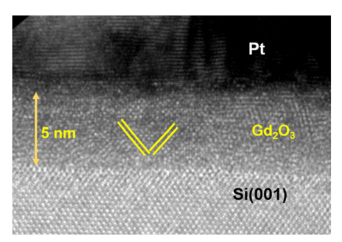


FIG. 9. High-resolution cross-sectional TEM micrograph of a $Pt/Gd_2O_3/Si(001)$ stack. The oxide layer is grown at under a molecular oxygen partial pressure of 5 \times 10⁻⁷ mbar. Sharp interface without any amorphous interfacial layer is visible.

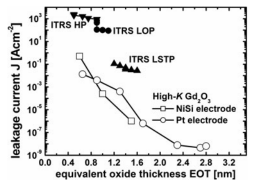


FIG. 10. Leakage currents versus EOT for ITRS (edition 2005) HP, LOP, LSTP target values, and data epitaxial Gd₂O₃ layers with different electrodes.

the oxide layer and/or oxide/Si interface. Gottlob et al. attributed this to oxide traps ($Q_{\rm ot}=3\times10^{12}~{\rm cm}^{-2}$). Such hysteresis represents flat band voltage ($V_{\rm FB}$) instability of the dielectric layer which leads to unstable threshold voltage in transistors. This clearly establishes that excellent structural quality of the alternative high-k dielectric does not guarantee satisfactory device performance due to the presence of electrically active charge defects.

A. Charge defects in epitaxial lanthanide oxides

In MOS structures there can be different types of charge defects that can degrade their electrical performance, such as oxide fixed charge (Q_f) , interface traps (Q_{it}) , and mobile oxide charges (Q_m) . In addition to unstable threshold voltage and instability of transistor operating characteristics, charge defects scatters carriers in the channel and lowers the carrier mobility. Further,

defects cause unreliability; which is the starting point for electrical failure and breakdown of the oxide. SiO₂, which is an almost ideal insulating oxide, has very low defect density. The low co-ordination number and high heat of formation of covalent SiO₂, that allow it to remain amorphous at typical CMOS processing temperature, helps to minimize the defect density via localized bonding rearrangement. Further, the defects at the Si/SiO₂ interface related to Si dangling bonds can be removed by rearranging the bonding network at the interface. Compared to SiO₂, high-k metal oxides normally contain more defects. The bonding in high-k metal oxides is highly ionic with higher coordination number which make them poor glass formers.¹⁶ Therefore, it is difficult to rearrange the bonds in the metal oxides for defect passivation. Since the charge defects in high-k dielectrics are undesirable for device applications for obvious reasons, much of the present-day engineering of these oxides consists of pragmatic strategies of trying to reduce defect densities by processing control and annealing. In this section we discuss about the nature of different charge defects existing in epitaxial LnOs and at the Ln₂O₃/Si interface and their effective passivation to improve the dielectric performance.

Let us start with the issues related to the dielectric properties of as-grown epitaxial Gd₂O₃ layers. The as-grown single crystalline Gd₂O₃ thin films on Si usually suffer from strong flatband voltage (V_{FB}) instability and are not process robust. This could be due to incomplete passivation of intrinsic dangling bonds induced by the existing binding mismatch at the Gd₂O₃/Si interface. When a Gd₂O₃ layer is deposited on clean Si surface; inevitably, the layer always consists of some defects pertaining to the Si dangling bonds at the interface and also due to the very different chemical nature of Si substrate (covalent) and oxide (highly ionic). Despite the oxide being grown epitaxially on Si substrate, not all Si surface atoms would have an opportunity to bond with overgrown epi-Gd₂O₃ layer's atoms. Such defects remain unsaturated due to the difference in the lattice parameters and space groups of the oxide and Si substrate.34

For example, MOS structures consisting of epitaxial Gd_2O_3 grown on clean Si substrates with atomically sharp interface, exhibit a large shift in the flat band voltage toward negative gate bias direction during multiple C-V sweeping (Fig. 11), which infers to a large number of intrinsic/extrinsic defects present at the interface. Such instability occurs due to the accumulation of mostly positive fixed charges at the Gd_2O_3/Si interface created during C-V sweeping, 39,40 which could significantly degrade the electrical performance of any device. 41,42 Laha et al. reported $Q_f \sim 9.5 \times 10^{12}$ cm $^{-2}$ for epitaxial Gd_2O_3 on Si which is only two orders of magnitude lower than the atomic density of dangling bonds on

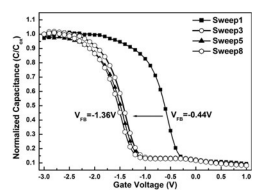


FIG. 11. C–V characteristics of single crystalline ${\rm Gd_2O_3}$ on ${\rm Si}(100)$ substrate.

a nonreconstructed Si(001) surface. This implies that approximately 1% of Si surface bonds remain unsaturated even after oxide growth onto it.⁴⁰ Each Si surface atom possessing a dangling bond usually seeks an additional electron to become electrically neutral. They tend to capture the required electrons from the oxide layer near the interface, thereby creating large number electron vacancies, i.e., positive charges, which results in a large shift of the flat band voltage.

In addition to the shift in the $V_{\rm FB}$ under applied bias, MOS structures involving epitaxial Gd₂O₃ layers on Si also suffer from significant C-V hysteresis when the gate bias is cycled from accumulation to the inversion region and back. This hysteretic nature of the C-V characteristic is related to some sort of recharging process inside the dielectric. The origin of the observed hysteresis may not be unambiguously associated with some definite process. Nazarov et al. suggested that the C-V hysteresis in epitaxial Gd₂O₃/Si system could be attributed to both trapping of the holes inside the dielectric and recharging the electrons near-surface traps. 43 One of the major sources of charge trapping in the metal oxides is the omnipresent oxygen vacancies (Vos) inside the layer, which give rise to defect states inside the band gap of the metal oxides. Vos have been identified as critical trapping centers in diverse high-k metal oxides, such as HfO₂, ZrO₂, LaAlO₃ etc. 44,45 Further, in ionic metal oxides V_{OS} redistribution can also occur under sufficiently high electric field, 46,47 which can act as potential source of $Q_{\rm m}$, thereby contributing at least partially to the observed C-V hysteresis. Czernohorsky et al. demonstrated that the hysteretic C-V characteristics of Gd_2O_3/Si heterostructures varies with P_{O_2} up to a certain limit, which clearly establishes the definitive role of V_{OS} on the hysteretic V_{FB} instability of epitaxial Gd₂O₃ layers.³⁷ It is obvious that such charge defects, which degrade the dielectric properties of epitaxial Ln₂O₃s, need to be minimized to improve their electrical performance prior to their integration into microelectronic devices.

V. PASSIVATION OF THE CHARGE DEFECTS IN EPITAXIAL LANTHANIDE OXIDES

In semiconductor industry one of the most commonly practiced method for passivation of charge defects in MOS structures is a thermal anneal in forming gas ambience. As a first measure for passivation of interface defect states in epitaxial Ln₂O₃s/Si heterostructures Sun et al. investigated forming gas anneal for the Gd₂O₃/Si system and found an influence of metal electrode.³⁹ For example, a post MOS fabrication anneal was found to be effective in case of tungsten electrode, whereas annealing prior to the electrode deposition was found necessary for Pt electrodes. Annealing epitaxial Gd₂O₃/Si system in forming gas environment at around 450 °C has been reported to reduce the $Q_{\rm f}$ s which stabilizes the $V_{\rm FB}$. However, this approach is not only very specific to the electrode type, but is not effective for complete elimination of O_f s in the epitaxial Ln₂O₃s. A negative shift in the flat band voltage (\sim -0.44 V) for Pt/Gd₂O₃/Si stack has been reported even after the forming gas anneal of the MOS structures fabricated on p-type Si. The negative flat band voltage results from $Q_f \approx 2.3 \times 10^{12} / \text{cm}^2$, which is estimated using the relation: $Q_{\rm f} = C_{\rm ox}(V_{\rm FB} - \Phi_{\rm ms}),$ where $C_{\rm ox}$ is the oxide layer capacitance and $\Phi_{\rm ms}$ is the metal work function. Additionally, Pt/Gd₂O₃/Si stacks possess a large density of trapped interface charges $(D_{it} \sim 7.63 - 8.8 \times 10^{11} \text{ cm}^{-2})^{39}$ These findings clearly indicate that forming gas anneal is an inadequate method to provide acceptable electrical properties for epitaxial Ln₂O₃s. Moreover, post deposition anneal also involves the risk of creating uncontrolled interfacial layer, exhibiting smaller dielectric constant as well as results in loss of epitaxy. As an alternative to forming gas anneal, Laha et al. demonstrated that the electrical properties of Gd₂O₃/Si structures can be improved significantly by interface engineering that involves atomic control growth of a silicate-like interfacial structure prior to the Gd₂O₃ layer.⁴⁸ However, controlling the interface composition is quite tedious and it involves multiple steps for substrate's surface preparation which renders the method less preferred for device fabrication. Another viable alternative surface preparation approach could be the passivation of Si dangling bond states by suitable chemisorbed species prior to the oxide layer growth. For examples, it has been widely reported that an ultrathin Si layer is very effective to passivate Ge surfaces for deep sub-micron p- and n-field effect transistor (FET) application, ^{49–51} whereas similar investigation for the effect of Ge on Si based devices is not commonplace. But it can be expected that deposition of Ge onto Si surface has also significant impact on the surface charge states of Si. It is known that a submonolayer (ML) Ge deposited onto a Si(111) surface leads to a charge transfer from the center Si adatoms

near the faulted half unit cell (FHUC) of a Si(111) (7×7) unit cell to the nearby Ge atoms, which makes the dangling bond states of Si adatoms empty.⁵² In principle this effect might have significant impact on subsequent properties of devices built on such Si surfaces. Laha et al. investigated the influence of chemisorbed Ge layer on the growth and dielectric properties of epitaxial Gd₂O₃ layers on Si.⁴⁰ In this approach different amount of Ge (0.15-3 ML) was deposited by e-beam evaporation on clean Si surfaces prior to the Gd₂O₃ layer growth. The in situ RHEED patterns of the Gd₂O₃ layers grown on such Ge-covered Si surfaces (Fig. 12) are very similar to what has been observed for the layers grown pristine Si surfaces.⁵³ This demonstrates that despite the large lattice mismatch between Si and Ge, few MLs of Ge pre-coverage (Φ_{Ge}) does not have any impact on the growth and structure of the Gd_2O_3 layer for Φ_{Ge} at least up to 2 ML. Figure 13 compares the C-V characteristics of Gd₂O₃ layers grown on pristine Si surface and on a Ge pre-covered Si surface. A strong $V_{\rm FB}$ instability corresponding to $Q_{\rm f}$ $\sim 9.5 \times 10^{12} \, \text{cm}^{-2}$ was observed in case of Gd₂O₃ grown on Si surface without Ge passivation, whereas for Si passivated with 1.5 ML Ge a much stable and ideal-like C-V characteristics was obtained. A systematic increase in $\Phi_{\rm Ge}$ leads to a monotonic increase in $V_{\rm FB}$ up to a maximum value of 0.3 V for 1.5 ML of Ge coverage [Fig. 14 (a)]. Further increase of Ge coverage was reported to result in nonuniformity in C-V measurements across the wafer. 40 The estimated $V_{\rm FB}$ in this case (+0.3 V) indicates negligible amount Q_f present at the interface. This improvement in the dielectric properties of the Gd₂O₃ layers was found to be independent of the Si surface orientation. It was also found that Ge pre-coverage on Si surface improves the density of interface traps significantly [Fig. 14 (b)], which also led to significant reduction in the hysteresis of the *C–V* characteristic. ⁴⁰ These results clearly indicate that Ge deposition prior to the oxide layer growth significantly influence the electronic properties of Si

surfaces and, consequently, the oxide/Si interface. To understand the influence of Ge on the surface electronic properties, Laha et al. also investigated the Si(111) surface states with different Φ_{Ge} using UPS. Figure 15 displays the photoemission spectra of Si(111) surfaces with different $\Phi_{\rm Ge}$. For pristine Si(111)-(7 × 7) surfaces ($\Phi_{\rm Ge} \sim 0$), the well-known signals related to the adatom state (S_1) , the rest atom state (S_2) , and the back-bond state (S_3) , are visible at energies close to the Fermi level (E_F) . ^{54,55} In contrast, the spectra from the Ge covered Si(111) surface exhibits only two surface states measured under the same condition, one (B_1) about 0.38 eV and the other about 1.70 eV (B₃) below the Fermi level. The first peak corresponds to the adatom state of clean Si(111)-(7 \times 7) surface, whereas the energetic position of the second peak (B₃) is located in between the S₂ and S₃ states of the clean Si surface. These results are in accordance to the earlier report by Carlisle et al., 56 where it was found that with increasing Φ_{Ge} (>0.3 ML), rest-atom and back-bond surface states merge into one broad feature located approximately halfway between them.⁵⁷ Such changes in the surface states indicate significant change in the atomic arrangement of the Si surface due to Ge deposition. For example, in case of Si(111) the chemisorbed Ge atoms randomly occupy the Si adatom positions at the very early stage (Φ_{Ge} < 0.1 ML). However, with increasing coverage they preferentially occupy the corner adatoms' position in the FHUC of Si $(111)(7 \times 7)$ surface structure. 58,59 At higher coverage $(\Phi_{Ge} > 0.5 \text{ ML})$, Ge atoms form nanoparticles on FHUC side consisting of approximately eight atoms. 52,60 In the presence of Ge nanoparticles, the Si center adatoms transfer charges to the nearby Ge atoms and thereby dangling bond states of such adatoms become partly empty. 52 Such charge transfer from Si to Ge occurs due to the difference in electronegativity and the compressive strain in Ge layer. Consequently, dipoles form at the Si/Ge interface with an excess of 0.1 electrons on the Ge side. 61 Similar behavior has also been reported

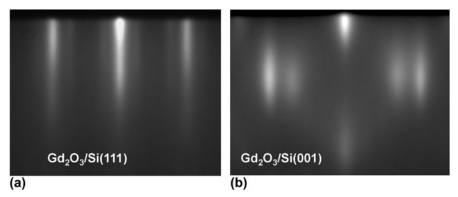


FIG. 12. RHEED patterns of epitaxial Gd_2O_3 on (a) Si(111) and (b) Si(001) substrates. 2 ML Ge was deposited on Si surface prior to the oxide layer growth. (Reprinted from Ref. 40 with permission of AIP publishing).

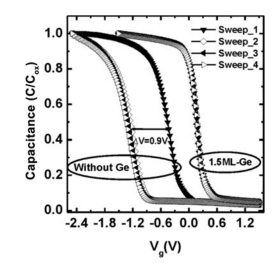


FIG. 13. *C–V* characteristics of Pt/Gd₂O₃/Si(001) MOS capacitors depicting the effect of 1.5 ML Ge coverage on Si(001) substrates. The capacitors fabricated on Ge passivated Si surface exhibit improved *C–V* behavior. (Reprinted from Ref. 40 with permission of AIP publishing).

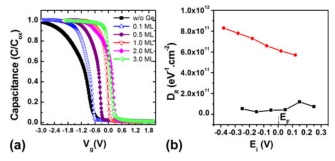


FIG. 14. (a) The C-V characteristics of Pt/Gd₂O₃/Si(111) MOS capacitors as a function of Ge coverage. The flat band voltage increases monotonically with increasing Ge coverage up to \sim 1.5 ML. (b) Density of interface traps for Pt/Gd₂O₃/Si capacitors fabricated on clean and Ge passivated Si(111) wafers, exhibiting significant improvement on $D_{\rm it}$ values.

by various other groups for the Si(001) surfaces, ^{62,63} despite their different microstructure. ⁶⁴

Therefore, the observed reduction in fixed oxide charge density in the Gd_2O_3/Si interface can mainly be attributed to the charge transfer from Si dangling bonds toward Ge. In case of Gd_2O_3 growth on Si without predeposited Ge the remaining dangling bonds in the oxide/Si interface attempt to capture additional electrons from oxide regions close to the interface to become electrically neutral. Thereby, large number electron vacancies will be created, i.e., positive charges, which results in a large shift of $V_{\rm FB}$. In contrast, when epitaxial Gd_2O_3 is grown on a Ge passivated surface, the contribution of the adatom dangling bond states significantly decreases. Moreover, due to the higher electron density at the top Ge layer owing to dipole formation, it is very unlikely that the electrons from the oxide interface

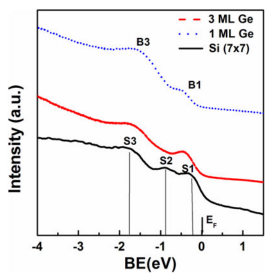


FIG. 15. UPS spectra from clean Si(111) (7 \times 7) surfaces with different Ge coverage, obtained at a photon energy of 21.2 eV. The spectra were recorded along $\langle 11-2 \rangle$ azimuth at an emission angle of 5° with respect to surface normal.

would be captured by the underlying substrates, which leads to much more stable interface.

A reduction in fixed oxide charge density in the Gd_2O_3/Si interface and resultant improvement in the V_{FB} stability of epitaxial Gd_2O_3/Si structures was observed also for oxide layers grown on Si covered by submonolayer carbon, ⁶⁵ indicating a similar charge transfer effect. Moreover, it was established that V_{FB} can be tailored by tuning the concentration of the chemisorbed species which can provide additional flexibility for device operation. ^{40,65,66} These observations establish the proof of principle about defect minimization at the oxide/Si interface by Si dangling bond passivation via charge transfer from Si adatoms to isovalent chemisorbed species with higher electronegativity.

Although surface passivation of Si by isoelectronic elements (Ge or C) prior to the oxide layer growth is a simple and effective way to reduce the fixed oxide charge and trap densities at the epitaxial Gd_2O_3/Si interface, this cannot completely eliminate the hysteretic C–V characteristics of the MOS structures. As discussed in the previous section, in addition to charging effect, the hysteresis in C–V could be partially related to the mobile charge defects inside the oxide layer owing to V_Os . Earlier we observed that with an increase in oxygen chemical potential during the oxide layer growth by an increase in P_{O_2} from 1×10^7 mbar to 5×10^7 mbar, the hysteresis also reduces significantly (Fig. 16).³⁷ However, a monotonic increase in P_{O_2} does not ensure complete elimination of the V_Os in the oxide layers.³⁷ Moreover, the magnitude of the flatband voltage change from voltage cycling $(\Delta V_{FB} > 40 \text{ mV})$ was still beyond the acceptable value $(\Delta V_{FB} < 20 \text{ mV})$ for stable transistor operation.¹²

Therefore, to minimize the Vos related defects various post growth processing techniques have also been attempted to compensate those defects in different ionic oxides. However, as already mentioned above with regard to the thermal anneal in forming gas ambience, post growth processing which often involves annealing at high temperature sometimes degrades the overall electrical properties due to the formation of interfacial layers with inferior dielectric constants. 67,68 Alternatively, incorporation of dopants has been reported to be a viable route to improve the electrical properties of different oxides by reducing the influence of Vos. For example, first principle calculations suggested that nitrogen (N) incorporation in Hf based dielectrics can reduce the leakage current conduction by effectively deactivating V_Os related gap states.⁶⁹ Substantial improvement of electrical properties has also been reported for HfO₂ thin films doped with fluorine, nitrogen, or lanthanum. 70–72

To explore the possible influence of dopants on the electrical properties of epitaxial Ln₂O₃s, we investigated the influence of nitrogen incorporation on the dielectric properties of epitaxial Gd₂O₃ layers. Nitridation of the layers was achieved by using nonactivated molecular N₂O as process gas instead of O₂ during growth. The incorporation of nitrogen into the oxide layers was checked by secondary ion mass spectroscopy (SIMS). The formation of Gd-N bonds in the Gd₂O₃:N layers was confirmed by in situ XPS measurements from the chemical shift of the Gd 4d peaks toward lower binding energy compared to pure Gd_2O_3 layers.⁷³ However, a quantitative estimation of the N incorporated into the Gd₂O₃ layers by this approach is yet to be achieved. Layers with different amount of N were grown by maintaining different partial pressure of N_2O (P_{N_2O})

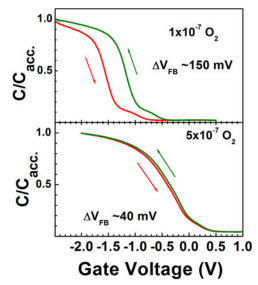


FIG. 16. *C–V* characteristics of two epitaxial Gd₂O₃ layers on Si(001) substrates grown at different oxygen partial pressures.

during deposition. All Gd_2O_3 :N samples grown with different P_{N_2O} exhibited similar uniform distribution of N concentration across the oxide layer. Figure 17 displays the SIMS profile of a representative Gd_2O_3 :N sample which confirms the presence of N inside the oxide layer.

Further, to understand the impact of N incorporation on the growth of Gd_2O_3 and the electrical properties of MOS stacks with epitaxial Gd_2O_3 :N on Si, these layers were compared with control Gd_2O_3 layers grown with similar partial pressure of molecular oxygen. The streaky RHEED patterns [Fig. 18(a)] exhibited by the Gd_2O_3 :N layer surface indicate the same two-dimensional growth on Si(111) as observed for undoped oxide. Figure 18(b) compares the X-ray diffraction patterns of two Gd_2O_3 :N layers grown at different N_2O partial pressures with a control Gd_2O_3 layer in the θ -20 geometry. The shoulder observed at $2\theta \sim 28.9^\circ$ beside the Si(111) peak corresponds to the (111) oriented epitaxial Gd_2O_3 with cubic bixbyite type of structure.

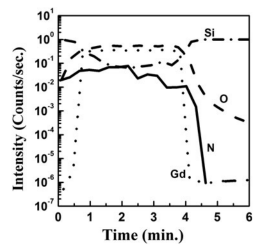


FIG. 17. SIMS depth profile of Gd, O, N and Si elements of a representative Gd_2O_3 :N sample as function of etching time.

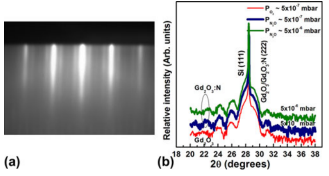


FIG. 18. (a) RHEED pattern along the $\langle 110 \rangle$ azimuth of a representative Gd₂O₃:N layer ($P_{\rm N_2O} \sim 5 \times 10^{-7}$ mbar) and (b) X-ray diffraction patterns in the θ -2 θ geometry of a control Gd₂O₃ layer and two representative Gd₂O₃:N layers.

These results indicate that usage of N₂O as a process gas incorporates N into the Gd₂O₃ layers only at a doping level and it does not affect the growth and structure of the layers.

However, a dramatic change in the dielectric properties of the epitaxial Gd₂O₃ layers was observed due to N incorporation. Figure 19 compares the C-V characteristics of a Gd₂O₃ and a Gd₂O₃:N sample grown under lower partial pressure of the process gases ($\sim 2 \times 10^{-7}$ mbar). The control sample prepared under reduced $P_{\rm O}$ [Fig. 19(a)] exhibits a strong hysteresis in the C–V behavior inferring that there exist a large number of mobile oxide charges inside the oxide layer. A density of $Q_{\rm m} \sim 2 \times$ 10¹² cm⁻² was estimated from the hysteretic shift of the $V_{\rm FB}$ (~140 mV) assuming only $Q_{\rm m}$ contribution to the C-V hysteresis. In contrary, in case of N doping [Fig. 19(b)] hysteresis in C-V was nearly vanished with $\Delta V_{\rm FB} < 10 \text{ mV}$ corresponding to a $Q_{\rm m} < 1 \times 10^{11} \text{ cm}^{-2}$. This clearly demonstrates that nitridation of Gd₂O₃ even at a low level (lower P_{N_2O}) already results in a nearly ideal dielectric behavior, likely due to an effective compensation of mobile oxide charges.

The influence of doping by other species has also been investigated by introducing carbon into Gd_2O_3 . C incorporation in different high-k oxides was reported to improve their dielectric properties. In our investigation, incorporation of C into the epitaxial Gd_2O_3 layers was found to result also in a significant improvement in the dielectric properties of MOS structures (i.e., minimize C-V hysteresis and lower leakage current density) for C concentration below 0.5%. However, at higher concentration, C segregates to the oxide/Si interface during growth. That leads to polycrystallinity and degradation of the electrical properties due to the formation of additional defects, 47,66 which makes it less preferred choice for doping Gd_2O_3 compared to N.

In addition to their impact on the dielectric properties, the dopants can also have significant influence on the electronic structure of high-k oxides. For example, density functional calculations revealed that incorporation of N into high-k oxides leads to a reduction of

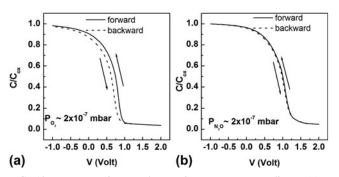


FIG. 19. Representative C-V hysteresis curves corresponding to (a) a control epitaxial Gd_2O_3 samples and (b) an epitaxial Gd_2O_3 :N sample.

their band gap and the VBO with Si which could result in degradation of the MOS electrical properties. Therefore, to gain a holistic understanding, the impact of N incorporation into epitaxial Gd₂O₃ layers on its band structure and dc leakage properties were also investigated. The electronic structure of the oxide layers were probed using *in situ* XPS measurement. Valence band photoelectron spectra of the layers were used to estimate the impact of N doping on the VBO of Gd₂O₃ with Si, which was determined from the relation:

$$\Delta E_{\rm V} = (E_{\rm CL}^{\rm Ox} - E_{\rm V}^{\rm Ox}) - (E_{\rm CL}^{\rm Si} - E_{\rm V}^{\rm Si}) - \Delta E_{\rm CL}$$
, (1)

where $\left(E_{\rm CL}^{\rm Ox}-E_{\rm V}^{\rm Ox}\right)$ and $\left(E_{\rm CL}^{\rm Si}-E_{\rm V}^{\rm Si}\right)$ are the core level to valence band maximum (VBM) energy difference for bulk Gd_2O_3 or Gd_2O_3 :N and Si, respectively, and ΔE_{CL} is the separation between a certain core level of the oxide (O 1s) and Si (Si 2p). The parameters for Si were measured from an identical bare Si wafer. The Si 2p peak position, measured under the presence of a thin oxide layers (\sim 2 nm) at the Si surface, was found to be independent on the nature and the partial pressure of the processing gas (O2 or N2O) applied during the epitaxial layer growth, and, moreover, no notable difference in the oxide peak position was observed for samples with different thicknesses. Since the VBM for Si was nearly zero, the VBO between Gd₂O₃:N or Gd₂O₃ and Si could be directly determined from the position of the VBM of the oxide layers on Si. Figure 20 presents the VB photoelectron spectra of a control Gd₂O₃ sample and Gd_2O_3 :N samples grown on Si at different P_{N_2O} . A value of 2.4 ± 0.2 eV was obtained for VBO for all control Gd_2O_3 samples grown at different P_{O_2} , which is in good agreement to an earlier report. 48 Doping of Gd₂O₃ with N leads to a reduction of the VBO with Si. Thereby, the VBO value of Gd₂O₃N was found to reduce progressively with increasing $P_{\rm N,O}$ from 2.2 \pm 0.2 eV

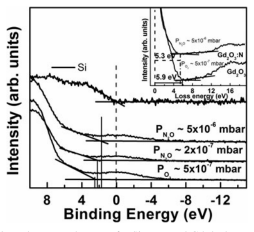


FIG. 20. Valence Band spectra for Si, a control Gd_2O_3 layer and two Gd_2O_3 :N layers, (inset) O 1s energy loss spectra of a control Gd_2O_3 layer and a Gd_2O_3 :N layer.

for $P_{\rm N_2O} \sim 2 \times 10^{-7}$ mbar up to a value of 1.7 \pm 0.2 eV at $P_{\rm N_2O} \sim 2 \times 10^{-6}$ mbar. Further increase of $P_{\rm N_2O}$ did not influence the VBO under present experimental conditions, inferring saturation in the concentration of N incorporated into the Gd₂O₃ layer.

In further investigations, the band gaps of the Gd_2O_3 :N and control Gd_2O_3 layers were estimated in XPS from the threshold energy of the plasmon energy loss spectrum observed at higher binding energy of the O 1s core level peak, as already demonstrated for other metal oxides. As shown in the inset of Fig. 20, the extrapolation yields a band gap energy of 5.9 ± 0.2 eV for the control sample, which agrees very well to the previously reported values for epitaxial Gd_2O_3 layers on Si. In contrast, the band gap of the Gd_2O_3 :N layers was found to decrease from 5.7 ± 0.2 eV for a layer grown at $P_{N_2O} \sim 2 \times 10^{-7}$ mbar to 5.2 ± 0.2 eV for a layer grown at $P_{N_2O} \sim 2 \times 10^{-6}$ mbar.

The band gap narrowing and reduction in VBO of epi-Gd₂O₃:N with Si can be understood by considering the nature of the VB structure of the binary Ln sesquioxides. In case of the Ln sesquioxides the valence band maximum is formed mainly by the O 2p states.⁷⁹ In case of substitutional N incorporation in different lanthanide and transition metal oxides (e.g., La₂O₃, HfO₂), their upper VB levels are shared between both O 2p and N 2p states.⁷⁸ The energy level of the N 2p states lie above the O 2p states, which leads to a shift of the VBM toward higher energy, thereby reducing the VB offset of the N doped oxide layer with Si. The same is valid for Gd₂O₃, the increase of N content in the Gd₂O₃ layers causes the VBM to possess increasing N 2p character, which leads to the gradual reduction of the band gap and VBO with Si, until saturation. The conduction band offsets (CBOs) estimated for the Gd₂O₃/Si structures considering the differences in the VBO, the energy gap of the Si substrate (~1.12 eV) and the measured band gap of the oxide were found to be nearly identical (2.4 \pm 0.2 eV) for all the Gd₂O₃ and Gd₂O₃:N samples; indicating that the CB of the oxide layers remains nearly unaffected by N doping, that agrees well with ab initio calculations reported for related oxides.⁷⁸

The reduction in the VBO with Si due to N incorporation can be expected to degrade the insulating properties of the Gd_2O_3 :N layer owing to lower hole barrier. However, as demonstrated for a representative set of samples in Fig. 21(a), the room temperature dc leakage measurements revealed that at applied electric fields below 4 MV/cm all the Gd_2O_3 :N samples grown under different P_{N_2O} exhibit lower leakage current densities compared to the control Gd_2O_3 samples grown under similar P_{O_2} indicating their superior insulating properties. For example, at 1 V ($V_{\rm FB}$ –1 V), the leakage current densities of the Gd_2O_3 :N layers were found to be about

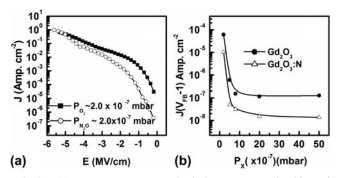


FIG. 21. (a) Room temperature dc leakage current densities of a representative Gd_2O_3 and a Gd_2O_3 :N layer. (b) dc leakage current densities measured at 1 V for different Gd_2O_3 and Gd_2O_3 :N samples as functions of ambient gas partial pressure (P_X) during growth. X denotes O_2 in case Gd_2O_3 and N_2O in case of Gd_2O_3 :N layers. The lines are to guide the eyes.

one order of magnitude lower compared to the control samples [Fig. 21 (b)].

The superior insulating properties of the Gd₂O₃:N layers can be explained by considering the influence of N on the electrically active defects in Gd₂O₃, especially on Vos, which are the most dominating electrically active defect in complex oxides grown at higher temperatures.^{80,81} Experimental investigations and *ab initio* calculations revealed that Vos induce electron conduction paths in complex oxides which give rise to enhanced leakage current densities. 82–85 The same can be suggested for epitaxial Ln₂O₃s. In case of ionic oxides, this occurs via Vos induced energy states within the band gap, which are occupied by two electrons for the neutral vacancy $(V^0_{\rm O})$ and are strongly localized on the orbitals of the adjacent metal ions. 66,80,83 Moreover, migration of Vos under sufficiently large electric field can also give rise to an increase in the leakage current densities in complex oxides. 86-88 Lowering of the leakage current densities by incorporation of N into Gd₂O₃ layers can be understood in much the same way as the elimination of the hysteresis effect in C-V measurements of nitrogen-doped Gd₂O₃ by considering the specific role of N in compensating the adverse effects the Vos related defects. In analogy to the effect of N incorporation in various other ionic oxides, this can only happen either due to the formation of N-V_Os complexes and/or substitution of the V_Os by N atoms in the Gd₂O₃:N. First principle calculations of N incorporation in different oxides suggested that in a minimum free energy configuration two N atoms occupy the nearest neighbor O sites to Vos. ^{69,89} In this arrangement each N atom extracts one electron from the neutral V_O sites to form a closed shell electronic configuration. Consequently, neutral Vos transform into positively charged V_O²⁺ forming electric dipoles, which effectively elevates the energy of the V_O related gap states above the conduction band of these oxides. In this way the dc leakage conduction is reduced. ^{69,80}

In analogy, the observed improvement in the dielectric properties of the Gd₂O₃:N layers could be attributed to a similar mechanism when N atoms combine favourably to the V_O sites in Gd₂O₃ and modify their charge states. Moreover, N atoms can also immobilize the Vos which can in turn reduce the Vos motion related leakage current conduction under an applied electric field.⁶⁹ The effect gets more pronounced with increasing N content in the Gd_2O_3 :N layers grown at higher P_{N_2O} . Further, the electric field corresponding to dielectric breakdown was found to be in the range of 4.5-5 MV/cm for all the samples, which agrees to the thermochemical description of dielectric breakdown in high-k materials. 90,91 However, in case of the Gd₂O₃:N layers the breakdown field was found to reduce with increasing N content. Such reduction in the dielectric breakdown field could be attributed to a collective effect of the reduced VB offset and an enhanced local electric field in the Gd₂O₃ layer due to N incorporation. 90,91

VI. SUMMARY AND OUTLOOK

We discussed different ways of growth optimization for improvement of dielectric properties of insulating Ln₂O₃ epitaxial grown on Si using molecular beam epitaxy. The growth optimization and discussion involves the density of charge defects inside the oxide layer as well as the oxide/Si interface, their impacts on the dielectric properties of oxide/Si heterostructures and different methods to passivate the charge defects. As a standard material we have considered epitaxial Gd₂O₃ layers to discuss the proof of principles. We demonstrated that the best vacuum condition achieved in the MBE chamber during oxide layer growth ($\sim 10^{-8}$ mbar) leads to silicide formation at the interface, which degrades the dielectric properties of the oxide layers. A controlled supply of molecular oxygen during the growth improves the interface quality significantly, however too high oxygen partial pressure leads to formation of lower permittivity interfacial layer and finally limits the achievable minimum effective oxide thickness. Partial pressure of oxygen in the range 10^{-7} mbar was found to be optimal to achieve epitaxial layers having atomic sharp interface to Si. Although such layers exhibit good dielectric properties in terms of the effective oxide thickness, as grown epitaxial Ln₂O₃s exhibit strong shift of the flatband voltage toward negative gate bias direction and also suffer from hysteretic flat band instability which render them unsuitable for transistor applications. In view of understanding the origin of flat band instability, we discussed the nature of fixed oxide charges in the epitaxial layers and their impact on the dielectric properties. The unsaturated dangling bond states in the oxide/Si interface were found to be responsible for the creation of positive charges inside the oxide layer, since they captures electrons form the oxide layer near to the interface. In addition to the fixed oxide charges, interface traps and mobile oxide charges originating from omnipresent oxygen vacancy related defect states also degrade the dielectric properties of the oxides. Consequently, these defects are required to be minimized prior to integrating these oxides into the Si technology for device application.

We have discussed different possible methods of defect passivation, such as annealing, doping and surface passivation by foreign atoms. Annealing the Ln₂O₃ layers in forming gas ambience, which is one of the most commonly used method for defect passivation in microelectronic materials, stands inadequate due to its sensitivity to the type of the metal gate used. As a viable alternative to this approach we have demonstrated that few monolayers of chemisorbed Ge or C on Si surface significantly improves the electrical properties of MOS structures with epitaxial Gd₂O₃ layers by passivation the Si dangling bond states. This happens due to the charge transfer from the dangling bonds of Si to the Ge or C atoms owing to the difference in their electronegativity. The passivation of the fixed oxide charges and interface traps stabilizes the flat band voltage and reduces hysteresis. Moreover, the flat band voltage can be tuned systematically by precisely controlling the amount of the chemisorbed species.

Incorporation of nitrogen into the oxide layers in a doping level was found to be a suitable way for passivation mobile oxide charges inside the layers, which results in vanishing flat band voltage hysteresis. Incorporation of N into the epitaxial Gd₂O₃ layer results in a band gap narrowing, which is related to the shift of valence band edge in Gd₂O₃:N layers toward higher energy due to their presumably increasing N 2p character in the valence band edge. Despite the declining VBOs with Si, the dc leakage current densities of epitaxial Gd₂O₃ layers is reduced significantly by N doping. These observations suggest that N incorporation in a doping level can improve the overall electrical properties of epitaxial Ln₂O₃s by passivation the detrimental effects of the electrically active Vos related defect states. Therefore, it appears that suitably doped epitaxial Ln₂O₃ layers deposited on surface Si surfaces, which are passivized with chemisorbed Ge, can be ideal candidate materials for next generation high-k dielectric applications. However, it should be noted that the extent to which the band gap of Gd₂O₃:N and its band offsets with Si can be tuned by increasing the N doping concentration still remains unclear. Furthermore, the impact of oxide doping and oxide/Si modification by Ge predeposition in active devices, such as CMOS based FET, is not yet tested in practice.

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