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Atomic-layer deposited thulium oxide as a passivation layer on germanium

I. Z. Mitrovic,1,a) S. Hall,1 M. Althobaiti,2 D. Hesp,2 V. R. Dhanak,2 A. Santoni,3 A. D. Weerakkody,1 N. Sedghi,1 P. R. Chalker,4 C. Henkel,5,b) E. Dentoni Litta,5 P.-E. Hellsström,5 M. Ostling,5 H. Tan,6 and S. Schamm-Chardon6

1Department of Electrical Engineering and Electronics, University of Liverpool, Brownlow Hill, Liverpool L69 3GJ, United Kingdom
2Department of Physics and Stephenson Institute for Renewable Energy, University of Liverpool, Liverpool L69 7ZF, United Kingdom
3ENEAG, Frascati Research Centre, via E. Fermi 45, 00044 Frascati, Italy
4Department of Engineering, University of Liverpool, Brownlow Hill, Liverpool L69 3GJ, United Kingdom
5School of ICT, KTH Royal Institute of Technology, Isafjordsgatan 22, 164 40 Kista, Sweden
6CEMES-CNRS and Université de Toulouse, nMat group, BP 94347, 31055 Toulouse Cedex 4, France

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A comprehensive study of atomic-layer deposited thulium oxide (Tm2O3) on germanium has been conducted using x-ray photoelectron spectroscopy (XPS), vacuum ultra-violet variable angle spectroscopic ellipsometry, high-resolution transmission electron microscopy (HRTEM), and electron energy-loss spectroscopy. The valence band offset is found to be 3.05 ± 0.2 eV for Tm2O3/p-Ge from the Tm 4d centroid and Ge 3p3/2 charge-corrected XPS core-level spectra taken at different sputtering times of a single bulk thulium oxide sample. A negligible downward band bending of ~0.12 eV is observed during progressive differential charging of Tm 4d peaks. The optical band gap is estimated from the absorption edge and found to be 5.77 eV with an apparent Urbach tail signifying band gap tailing at ~5.3 eV. The latter has been correlated to HRTEM and electron diffraction results corroborating the polycrystalline nature of the Tm2O3 films. The Tm2O3/Ge interface is found to be rather atomically abrupt with sub-nanometer thickness. In addition, the band line-up of reference GeO2/n-Ge stacks obtained by thermal oxidation has been discussed and derived. The observed low reactivity of thulium oxide on germanium as well as the high effective barriers for holes (~3 eV) and electrons (~2 eV) identify Tm2O3 as a strong contender for interfacial layer engineering in future generations of scaled high-k gate stacks on Ge. © 2015 AIP Publishing LLC.

I. INTRODUCTION

Interface engineering plays a pivotal role in new high-k/metal gate technology advancement.1 Germanium has recently gained much interest as a high carrier mobility semiconductor (CMOS) devices together with the use of high-k dielectric materials, such as HfO2.2,3 In contrast to superior interface properties of SiO2/Si, Ge oxides are well-known to be thermally and chemically unstable.4 Therefore, passivation of the Ge surface is a critical step for fabricating high performance Ge-based field effect transistors (FETs). The native oxide GeO2 has attracted renewed research interest5 as a potential passivation layer owing to its excellent interface control on Ge. However, for aggressive oxide scaling with equivalent oxide thickness (EOT) well below 1 nm, the combination of higher-k rare-earth (RE) oxide and ultra-thin GeO2 is required.5 The thermodynamically robust interfacial layer (IL) engineering on Ge using RE Y2O3-doped GeO2 has been shown recently to deliver 0.47 nm EOT on Ge, with superior interface properties and mobilities in Ge nMOSFETs.6 Another approach to achieving sub-nm EOT Ge gate stacks is utilizing Al2O3 barrier IL properties.2,3,7

Rare-earth thulium oxide (Tm2O3) has been considered as the main high-k dielectric8 and as a capping layer for La2O3-based gate stacks,9 but only on Si. A low reactivity of Tm2O3 with the Si substrate has been observed.10 There have been theoretical prediction and some recent estimations of the band gap (~5 eV,11 ~6.5 eV,12 and 5.76 eV13 respectively) on Tm2O3/Si structures. Atomic-layer deposition (ALD) has become one of the preferred methods for thin film deposition in several fields due to the excellent thickness control, uniformity, and conformity. A novel process for atomic-layer deposition of thulium oxide has been recently developed.14 A TmSiO IL layer with EOT of ~0.25 nm has been achieved, which indicates a strong potential for its integration in sub-10 nm technology nodes.15 Conversely, there have been no reports on Tm2O3 as a passivation layer on Ge, apart from our earlier work.16

A reliable measurement method to determine the band offsets is essential for modelling the carrier transport properties. The offsets reported at the GeO2/Ge interface show large scattering in the range of about 1 eV for data obtained by x-ray photoelectron spectroscopy (XPS) and internal photoemission (IPE) (see Ref.17 and references therein). Detailed mechanism responsible for such discrepancy is not clear. There are assumptions made that this is due to different GeO2 growth methods.5,18–23 Furthermore, it has come recently to focus that the XPS requires careful attention to
charging effects as a result of electron emission from the insulator, while the IPE data demand careful interpretation. A clear understanding of the physical phenomena behind the charge accumulation and neutralization in dielectric/semiconductor heterojunction during XPS measurements seems still to be elusive. A recent XPS study on HfO₂/Ge heterostructures suggests that the role of germanium is not negligible in the neutralization mechanisms beyond the differential charging effect. Charging can occur in an XPS experiment when the holes that are created by the ejection of photoelectrons accumulate in a sample. This build-up of charge results in an increase in the binding energy (BE) of spectral features. Bersch et al. have shown that not correcting for charging effects in overestimation of valence band offset (VBO) by ~0.5 eV on average. It is common practice for the VBO to be determined from XPS measurements by Kraut’s method using the valence band (VB) and core-level (CL) photoemission from bulk-like samples of the two constituent materials and a thin interfacial sample forming the interface of interest. The overlayer of this heterojunction sample must be sufficiently thin (usually <5 nm) to allow XPS core-levels from the underlying material to be probed due to the finite escape depth of the photoelectrons. The binding energy values are referenced to the valence band maximum (VBM) of each sample, determined by extrapolating a linear fit of the leading edge of the VB photoemission to the baseline in order to account for broadening of the photoemission spectra. Then, the VBO for oxide/semiconductor substrate sample can be determined as

\[ \text{VBO} = \delta_{\text{SUB}} + \delta_{\text{INT}} - \delta_{\text{OXIDE}}, \]  

where \( \delta_{\text{SUB}} \) and \( \delta_{\text{OXIDE}} \) are the energy differences between chosen reference core-levels in substrate and bulk oxide samples and their respective VBMs, while \( \delta_{\text{INT}} \) refers to the BE difference for the former two core-levels for the interfacial sample.

This paper conveys three important findings: (i) the valence band offset for Tm₂O₃/Ge of 3.05 ± 0.2 eV, determined by Kraut’s method using a single sample consequently sputtered with core-level spectra taken at different sputtering times, shows consistency within experimental error with the offset result obtained using three distinctive samples (bulk, interfacial, and substrate); (ii) the VBO for thermal GeO₂/Ge is in agreement with the most recent report from Toriumi’s group; (iii) Tm₂O₃ shows even lower reactivity on Ge than on Si, with rather atomically sharp interface indicating possible barrier properties.

II. EXPERIMENTAL

The 10 nm (nominal) thick Tm₂O₃ samples were prepared by ALD on 35 nm p-Ge epitaxial layer/Si(100) and on Si(100). The reference samples of GeO₂ (5 and 10 nm nominal thicknesses) were grown on 35 nm n-Ge epi/Si(100) by thermal oxidation at 525°C under 1 atm O₂. Prior to the gate oxide deposition, epi Ge/Si(100) samples were cleaned in a HF 0.5%/Isopropanol 1% /H₂O mixture to remove (minimize) the native Ge oxide layer. The Tm₂O₃ layers were deposited using Tris(cyclopentadienyl)thulium, heated to 140°C, and water vapor as precursor gases. An ultra-high vacuum (UHV) system operating at 4 × 10⁻¹⁰ Pa base pressure and equipped with a VG Al Kα monochromatized x-ray source and a CLAM2 hemispherical analyzer was used for XPS data acquisition at normal emission. The electron analyzer was set at constant 20 eV pass energy mode and calibrated. The total energy resolution is found to be <0.7 eV from the fitting of the Fermi edge of a clean Au sample. The binding energy is referred to the position of the Fermi level measured on a clean Ta strip in good electrical contact with the sample. In order to reach the Tm₂O₃/Ge interface, the samples were mildly sputtered with 0.5 keV Ar⁺ ion energy (0.25 nm/min). The main XPS core-levels and VB edge in the bulk Tm₂O₃ have been monitored as a function of sputtering time in order to reveal possible sputtering-induced effects. No change in the VB edge positions and in the Tm 4d and O 1s CL lineshapes, relative intensities, and BE positions could be detected by XPS after sputtering cycles, indicating that Ar⁺ bombardment did not induce preferential sputtering and any observable surface modifications. The XPS spectra for GeO₂/n-Ge samples were recorded on a separate UHV system consisting of an Al Kα x-ray (hν = 1486.6 eV) source and a P5 Vacuum Technology electron energy analyzer. This spectrometer was calibrated so the Ag 3d₅/₂ photoelectron line had a BE of 368.35 eV, a full width at half maximum (FWHM) of 0.8 eV being the spectral resolution for this study, and a 10 eV pass energy. Charge compensation was achieved using a VG Scienta FG300 low energy electron flood gun with the gun settings adjusted for optimal spectral resolution. The electron BEs were then corrected by setting the C 1s peak in the spectra (due to stray carbon impurities) at 284.6 eV for all samples. The probe area during the XPS measurements was 1 mm². The error bar (±0.2 eV) we defined in this paper is due to VBM determination through the linear interpolation method. The core-level binding energy determination by fitting a Voigt curve to a measured peak introduces typically much smaller (±0.05 eV) error. A Shirley-type background is used during the fitting of all spectra. The vacuum ultra-violet variable angle spectroscopic ellipsometry (VUV-VASE) measurements were performed using a spectral range from 0.5 to 8 eV, and the angles of incidence of 55°−75°, by 10° as a step, to maximize the accuracy. The atomic structure and elemental analysis were investigated with high-resolution transmission electron microscopy (HRTEM) and electron energy-loss spectroscopy (EELS) performed on a field emission image-corrected FEI Tecnai™ F20 microscope operating at 200 kV. For local EELS studies, the microscope was also equipped with a scanning stage (STEM), allowing a focused one nanometer-sized probe to be scanned over the sample area of interest (in our case, a line crossing the Tm₂O₃/Ge interface), and an imaging filter (Gatan GIF TRIDIEM) used as a spectrometer.

III. RESULTS AND DISCUSSION

A. Estimation of VBO for Tm₂O₃/Ge gate stack

The VBO determination of the Tm₂O₃/Ge system is addressed first. Fig. 1 shows high-resolution Tm 4d, valence
band and Ge 3p XPS spectra taken at three different sputtering times referring to bulk Tm$_2$O$_3$ (Figs. 1(a) and 1(b)), interface (Figs. 1(c) and 1(d)) and Ge substrate (Figs. 1(e) and 1(f)). Since for Ge 3d, a strong presence of neighboring O 2s and Tm 5p doublet complicates the interpretation of the spectra, in this work, Ge 3p$^{3/2}$ and the centroid value of Tm 4d core-levels were used for VBO estimation. Figure 2(a) shows the peak areas of two components of O 1s core-level: from the main Tm$_2$O$_3$ (circle symbol) and from IL (triangle symbol). Each symbol point on the graphs refers to a single sputtering time, when the spectrum was taken. At first, the signal is dominated by the thulium contribution (until ~1100 s); however, as the sputtering through the film continues, the IL becomes more prominent until only interfacial layer species are left (at ~2000 s). These changes are further reflected in the plot of the Auger parameter, also shown in Fig. 2(a). The Auger parameter was calculated using the centroid values of the O 1s peak and from the O KLL Auger peak. Note that in interfacial Tm$_2$O$_3$/Ge heterostructures, the Tm 4d core-levels exhibit a monotonically decreasing shift towards lower BEs of ~0.12 eV when sputtering Tm$_2$O$_3$ film (Figs. 1(c) and 2(b)), thus providing clear fingerprints of charging phenomenon. On the contrary, a very small variation (~0.05 eV) of the Ge 3p BEs was observed (Figs. 1(d) and 2(c)). To account for the effect of differential charging, the positions of Tm 4d and Ge 3p peaks were estimated by extrapolating the measured BEs to zero Tm$_2$O$_3$ thickness (i.e., to the highest value of sputtering time in our experiment, see Fig. 2(a)), and hence ideally to zero charge. The difference of Tm 4d and O 1s peaks was found to be 354.29 ± 0.03 eV, being indicative of the same stoichiometry of the films sputtered <2000 s. The value Tm 4d – Ge 3p$^{3/2}$ = $\delta_{\text{INT}}$ = 54.89 eV was extracted from the extrapolated values in Figs. 2(b) and 2(c). Comparing the Ge 3p$^{3/2}$ peak of the Ge substrate and the same peak with Tm$_2$O$_3$ on top, an energy shift towards higher BEs of 0.06 eV is observed. This is a signature of a small downward band bending, which agrees with the presence of p-type Ge. The result suggests negligible bending of Ge core-levels despite the charging of the Tm$_2$O$_3$ film during x-ray exposure; a converse scenario has been observed for HfO$_2$/n-Ge. The BE differences between Tm 4d centroid and VBM for bulk Tm$_2$O$_3$ ($\delta_{\text{OXIDE}}$), and Ge 3p$^{3/2}$ and VBM for the Ge substrate ($\delta_{\text{SUB}}$) measured from Figs. 1(a) and 1(b) and Figs. 1(e) and 1(f), respectively, are summarized in Table I. By inserting $\delta_{\text{OXIDE}}$, $\delta_{\text{INT}}$, $\delta_{\text{SUB}}$ values in Kraut’s equation (1), the VBO = $\delta_{\text{SUB}}$ + $\delta_{\text{INT}}$ – $\delta_{\text{OXIDE}}$ = 3.05 ± 0.2 eV is calculated for Tm$_2$O$_3$/Ge. The result is in agreement with our
B. Estimation of VBO for GeO\(_2\)/Ge gate stack

We now look to the estimation of VBO for the reference GeO\(_2\)/Ge system. Figs. 3(a) and 3(b) show high-resolution Ge 3d core-levels taken for bulk and interfacial GeO\(_2\)/Ge after prolonged (at least an hour) x-ray exposure, until the point they reached constant BEs.\(^1\) The GeO\(_2\) film shows two main peaks, fitted to doublets of Voigt functions with spin orbit splitting of 0.6 eV and branching ratio \(1/2\). The difference between Ge 3d\(_{5/2}\) of the substrate (28.95 eV) and GeO\(_2\) (32.34 eV) for the bulk sample is 3.4 eV, showing a stoichiometric GeO\(_2\) and negligible differential charging.\(^1\)

Comparing the Ge 3d\(_{5/2}\) peak of the bare Ge (not shown) and the same peak from Ge with GeO\(_2\) on top (Fig. 3(a)), an energy shift of 0.05 eV towards lower BEs is observed. The shift is consistent with n-Ge\(^2\) and implies a 0.05 eV upward band bending at the GeO\(_2\)/n-Ge interface, in agreement with the formation of a superficial p-inversion layer in the n-type Ge substrates.\(^3\) Note in Fig. 3(a) that the peak at \(\sim\)23 eV refers to O 2s CL, while three minor peaks in the energy range from 5 to 15 eV are part of VB and relate to O 2p, Ge 4s, and Ge 4p species in agreement with the literature.\(^1\) The measured BE differences, \(\delta\)\(_{\text{OXIDE}}\), \(\delta\)\(_{\text{INT}}\), and \(\delta\)\(_{\text{SUB}}\), for GeO\(_2\)/Ge are listed in Table I and the literature values\(^1\) are also inserted for comparison. Applying Kraut’s equation (1), yields a VBO = \(\delta\)\(_{\text{SUB}}\) + \(\delta\)\(_{\text{INT}}\) - \(\delta\)\(_{\text{OXIDE}}\) = 3.55 \(\pm\) 0.2 eV, consistent with a value of 3.6 \(\pm\) 0.2 eV reported by XPS\(^1\) and by synchrotron radiation photoemission spectroscopy.\(^5\)

C. Band gap evaluation and nature of Tm\(_2\)O\(_3\)/Ge interface

The band gaps of Tm\(_2\)O\(_3\) and GeO\(_2\) were determined by VUV-VASE. This was accomplished by first determining the thickness in the non-absorbing (transparent) region of the spectra. The dielectric function of the Ge film (31.9 nm) with the native oxide as Cauchy layer (1.4 nm) was modeled first. Then, another Cauchy layer is added and fitted for the thickness of Tm\(_2\)O\(_3\) film (10.4 nm) or GeO\(_2\) (4.6 nm). Subsequently, the optical constants (real and imaginary part of dielectric function) were extracted. The Tm\(_2\)O\(_3\) (GeO\(_2\)) film was modelled with Cauchy layer at long wavelengths and extended into VUV with the B-spline and then converted to a general oscillator layer. The dielectric function converts to refractive index and extinction coefficient (k) using Kramers-Kronig relations. The absorption coefficient (\(a\)) is calculated from the extinction coefficient as \(a = 4\pi k/\lambda\), where \(\lambda\) is wavelength. The absorption coefficient vs photon energy plots for GeO\(_2\)/Ge and Tm\(_2\)O\(_3\)/Ge stacks are shown in Figs. 3(c) and 3(d), respectively. The band gap can be estimated by linear extrapolation of the segments on the curves in the non-absorbing regions, and is found to be 5.95 eV for GeO\(_2\) and 5.77 eV for Tm\(_2\)O\(_3\). The schematics of derived band line-ups for GeO\(_2\)/Ge and Tm\(_2\)O\(_3\)/Ge are depicted in Figs. 3(e) and 3(f).

Note that both band gap values are slightly higher than those reported using the Tauc-Lorentz...
method, in agreement with the finding of Di et al. The band gap value for GeO2 compares to Lange et al., where the optical band gap has been measured from an increase of the absorption edge and found to vary from 5.21 eV to 5.95 eV, depending on O2 flow rate during reactive DC magnetron sputtering deposition. The band gap of 5.95 eV refers to highest O2 flow and polycrystalline films of GeO2. The band gap of GeO2 of 6.0 eV has been reported from SE measurements from absorption edge. The band gap value of Tm2O3 compares to 5.76 eV reported from optical reflectance on Tm2O3/Si stack. It is worth noting a pronounced absorption (at 5.3 eV) below the band edge for the Tm2O3/Ge, and an Urbach tail (see inset of Fig. 3(d)) as a signature of the poly-crystalline nature of the thulium oxide film.

The polycrystalline nature of the Tm2O3 deposited on Ge is directly seen from the HRTEM image and the electron diffraction pattern of Figs. 4(a) and 4(b) from which the cubic Tm2O3 structure has been identified. What is noticeable from the HRTEM image is the direct and sharp interface between the projected atomic structures of Ge and the Tm2O3 film (see white arrows in Fig. 4(b)), which is not the case for Tm2O3 deposited on Si, where a thin amorphous interfacial layer is observed (not shown). This feature is common to RE oxide or RE oxide-based films. Some roughness is observed at this interface. From the chemical point of view, there is a transition region between the Ge
substrate and the Tm$_2$O$_3$ film, where the three elements Tm, O, and Ge are present, as can be observed from calculated EELS elemental profiles in Fig. 4(c). Due to the 1 nm probe used for the EELS analysis, the transition region may point out to the roughness of this interface observed at the nanometer level and possibly to a chemically modified interface, at the sub-nanometer level, germanate in nature (Tm-O-Ge).

The latter is further substantiated by the presence of a negligible IL peak (<3% area) from microscopic XPS measurements of Ge 3p CL shown in Fig. 1(d).

IV. SUMMARY

In summary, a consistent valence band offset value of ~3 eV has been obtained for atomic-layer deposited Tm$_2$O$_3$/Ge from core-level and valence band XPS spectra measured at different sputtering times from a single bulk oxide layer. This method allows for more authentic probing of the interface at different sputtering times from a single bulk oxide layer.

Furthermore, this study points unambiguously to both Tm$_2$O$_3$/Ge and GeO$_2$/Ge exhibiting sufficient conduction band offsets (>1.5 eV) to adequately suppress leakage current in real applications.

The barrier role of Tm$_2$O$_3$ interlayer could suppress the growth of unstable GeO$_2$ and bring effective passivation route in future Ge-based scaled CMOS devices.

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