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Atomic scale observation of phase separation and formation of silicon clusters in Hf higk- κ silicates

E. Talbot, ^{1,a)} M. Roussel, ¹ C. Genevois, ¹ P. Pareige, ¹ L. Khomenkova, ² X. Portier, ² and F. Gourbilleau² ¹*Groupe de Physique des Matériaux (GPM), Université et INSA de Rouen, UMR CNRS 6634,*

Av. de l'Université, BP 12, 76801 Saint Etienne du Rouvray, France

²Centre de Recherche sur les Ions, les Matériaux et la Photonique (CIMAP), CEA/CNRS/ENSICAEN/UCBN,

6 Bd. Maréchal Juin, 14050 Caen Cedex 4, France

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Hafnium silicate films were fabricated by RF reactive magnetron sputtering technique. Fine microstructural analyses of the films were performed by means of high-resolution transmission electron microscopy and atom probe tomography. A thermal treatment of as-grown homogeneous films leads to a phase separation process. The formation of SiO₂ and HfO₂ phases as well as pure Si one was revealed. This latter was found to be amorphous Si nanoclusters, distributed uniformly in the film volume. Their mean diameter and density were estimated to be about 2.8 nm and $(2.9 \pm 0.4) \times 10^{17}$ Si-ncs/cm³, respectively. The mechanism of the decomposition process was proposed. The obtained results pave the way for future microelectronic and photonic applications of Hf-based high- κ dielectrics with embedded Si nanoclusters. © 2012 American Institute of Physics. [http://dx.doi.org/10.1063/1.4718440]

I. INTRODUCTION

Continuous downscaling of the complementary metaloxide semiconductor technology has evidenced the limitations of the silicon oxide gate dielectrics for thickness lower than 0.7 nm.^{1,2} To overcome this limit, alternative high- κ materials have been proposed. Among them, zirconium and hafnium oxides, as well as their silicates are considered as the best candidates to replace SiO₂.^{3,4} Besides a high dielectric constant, the advantages of these materials are their thermal and chemical stabilities, and good interfacial quality with Si channel.^{3,5–7} Widespread applications of high- κ materials are linked with the floating gate non-volatile memory devices containing semiconductor and/or metallic nanocrystals (NCs) or amorphous nanoclusters (ncs).^{8,9} Their controlled precipitation in high- κ oxide host is a major challenge for such devices.

Among semiconductor NCs (or ncs), silicon and/or germanium ones are the most addressed. Their formation usually results from a two-step process. Initially, silicon (or germanium) is incorporated into high- κ host. For this purpose, different techniques are used, for instance, Si(Ge) ion implantation¹⁰ or deposition of thin Si(Ge)-based layer sandwiched between the high- κ layers.^{11,12} After this step, a specific annealing treatment is carried out to form NCs (or ncs) inside high- κ host.^{10–12} It is worth to note that up to now neither Si-NCs nor Si-ncs formation in Hf-silicate host was demonstrated. To explain this fact, one can refer to a Hf-Si-O phase diagram.^{13–17} This pseudo-binary alloy can be accounted for a mixture of HfO₂ and SiO₂ unit cells such as $(HfO_2)_x(SiO_2)_{1-x}$.¹³ They present a miscibility gap in the range of composition considered. This can lead to a phase separation process either during fabrication or post-fabrication processing. It results, usually, in

the formation of HfO₂ and SiO₂ phases.^{16,17} Meanwhile, to our knowledge, no information was obtained about decomposition process for films as $(HfO_2)_x(SiO_2)_y$ with $x + y \neq 1$. As evoked previously, the formation of Si- or Ge-NCs (or ncs) requires an annealing treatment at 900-1100 °C.^{10,18,25} If these NCs (or ncs) have to be grown in the Hf-silicate matrix, the thermal behaviour of this latter element has to be taking into account.^{5,14,15,20}

In this paper, we present a specific approach developed to fabricate Hf-silicate dielectrics with embedded Si-ncs. The films were grown by reactive RF magnetron sputtering, yielding on a fine control of their chemical composition. An annealing treatment was used to form Si-ncs. A combination of high-resolution transmission electron microscopy (HRTEM) and atom probe tomography (APT) techniques allowed to study the film microstructure at atomic scale level and to obtain information about a phase separation process and Si-ncs formation.

II. EXPERIMENTAL

A. Fabrication of Hf-silicate samples

For this study, a 230-nm HfSiO film was deposited on ptype (100) Si substrate cleaned in 10% HF solution to remove native silicon oxide. The RF magnetron cosputtering of a 4-in. HfO₂ target (99.9%) topped by Si chips placed on the electron "race track" was used. The films were fabricated under a RF power density of 0.74 W/cm² and a substrate temperature of 400 °C in mixed argon-hydrogen plasma (with hydrogen flow rate of 20%). The total plasma pressure was kept at 40 μ bar. The hydrogen was used due to its ability to remove the oxygen from plasma and, in consequence, to reach higher Si excess in the films. More details on the reactive fabrication process can be found elsewhere.^{19,20} An annealing treatment of the sample was

^{a)}Electronic mail: etienne.talbot@univ-rouen.fr.

performed in a conventional furnace at 950 °C during 15 min in a nitrogen flow.

B. Methods used for structure analysis

Several techniques were used to study the structural properties of the samples. X-ray diffraction analysis was performed using a Phillips XPERT HPD Pro device with Cu K_{α} radiation ($\lambda = 0.1514$ nm) at a fixed grazing angle incidence of 0.5°. An asymmetric grazing geometry was chosen to increase the volume of material interacting with x-ray beam, as well as to eliminate the contribution from the Si substrate. attenuated total reflectance (ATR)-FTIR spectra were measured in the $600-4000 \text{ cm}^{-1}$ spectral range by means of a 60° Ge Smart Ark accessory inserted in a Nicolet Nexus spectrometer to study the chemical composition and microstructure of the samples. For atomic scale analysis of samples, APT and HRTEM were used. For this latter, the crosssectional specimens were prepared by a standard lift-out TEM lamella procedure. Z-contrast high angle annular dark field (HAADF), scanning transmission electron microscopy (STEM), electron energy loss spectroscopy (EELS), and HRTEM experiments have been performed by means of a JEOL-ARM200F working at 200 kV. APT techniques allow a chemical mapping of materials in three dimensions, in the real space with an atomic resolution.^{21–23} Laser assisted APT has been recently used to study insulating materials.^{24–26} It provides a better understanding of material nanostructure, allowing to quantify a phase separation process and to evidence clustering effects. To investigate Hf-silicate layer by this technique, tip shaped specimens were prepared by means of a dual-beam FIB-SEM (Zeiss NVision 40), using lift out method and an annular milling procedure.^{27,28} To prevent Ga implantation and ion beam amorphization, the samples were beforehand capped with a Pt layer. APT analyses were carried out in a laser assisted wide angle atom probe tomography (LAWATAP) from CAMECA. The experiments were carried out at 80 K, using UV (343 nm) femtosecond laser pulses (50 nJ, 350 fs) with a pulse rate of 100 kHz.

III. RESULTS

A. As-grown HfSiO layers

Structural properties of the sample, examined by x-ray diffraction and ATR-FTIR spectroscopy, showed the homogeneous and amorphous nature of the as-grown sample.^{5,20} These results were confirmed by HRTEM observations (Figure 1). The observed bright contrast at the film/substrate interface evidenced the formation of a thin interfacial HfSiO_x layer (not thicker than 1 nm), as already shown for similar films by Khomenkova *et al.*^{5,20}

B. TEM observation of annealed HfSiO layers

Figure 2 shows a cross-sectional HRTEM image of the HfSiO layers annealed at 950 °C for 15 min in nitrogen flow. The bright and dark contrasts on HRTEM micrograph (see arrows) are associated with the formation of Si-rich and Hfrich phases due to the phase separation process in the HfSiO layer. The Hf-rich phase appears here to be crystallized,



FIG. 1. Cross sectional HRTEM image of the as-grown HfSiO layer. Thin bright region close to substrate is HfSiO_x interfacial layer.

which is consistent with previous study on the amorphouscrystalline transition on thick HfSiO layers.⁵ Thus, the detrimental role of thermal treatment resulting in the phase separation and formation of Hf-rich and Si-rich phases is clearly seen, as predicted by the phase diagram. Similar phase separation was already evidenced in the literature.^{5,16,29}

Besides this phase separation in the film volume, the formation of two thin interfacial layers (IL) was observed. Based on their contrast, we can expect that the first IL (bright contrast, Si-IL, \sim 2-nm) contains a lower amount of Hf than the second one (dark contrast, Hf-IL, \sim 3-nm). To deeply investigate the nature of the phases formed as well as the origin of interface layer between HfSiO film and Si substrate, the analysis of the same sample by APT technique was performed.

C. APT analysis of HfSiO layers

A key point for high- κ Hf-based microelectronic devices is the nature of film/substrate interface. The results of APT



FIG. 2. Cross sectional HRTEM micrograph in [110] zone axis of the Si substrate. Si-IL denotes thin Si-rich layer at the interface with the silicon substrate, Hf-IL denotes Hf-rich interfacial layer followed Si-rich one. The arrows show the bright and dark regions, corresponding to Si-rich and Hf-rich phases formed in the film volume. The crystalline sequences are seen in the dark regions.

experiments on the interface between HfSiO film and underlying Si substrate are shown in Fig. 3. A 3D reconstruction image of this interface region is displayed by Fig. 3(a). On this image, the hafnium and silicon atoms are presented by blue and red dots, respectively. The oxygen atoms are not shown here for clarity. The region *I* corresponds to the HfSiO film that is separated by the interface layer (IL, region *II*) from the Si substrate (region *III*). At film/substrate interface (region *II*), the higher number of Hf atoms (blue dots) is found close to the HfSiO film (region *I*), whereas an enrichment in Si atoms (red dots) is clearly seen close to Si substrate (region *III*). Thus, this APT experiment confirms the formation of Si-rich IL layer followed by a Hf-rich one in the interface region. These results are consistent with the HRTEM observations described above.

Figure 3(b) represents the concentration profile for Si, O, and Hf (in at. %) to the surface of the film towards the substrate. It has been calculated by counting the atoms of each species in a sampling volume of $25 \times 25 \times 1$ nm³ parallel to the interface. The Si, O, and Hf content were found to be the same for different sampling volumes, chosen in the region *I* (Fig. 3(b), depth before 15 nm). The calculations showed that $X_{\text{Si}} = 30.6 \pm 0.3$ at. %, $X_{\text{O}} = 61.5 \pm 0.3$ at. %, and $X_{\text{Hf}} = 7.9 \pm 0.3$ at. % over all film volume.

The region II (depth between 16 and 21 nm) shows also a mixture of Si-rich and Hf-rich phases. However, it is not



FIG. 3. APT reconstruction of the HfSiO-film/Si-substrate volume $(25 \times 25 \times 50 \text{ nm}^3)$ analysed by LAWATAP. (a) Hafnium atom maps (blue dot) and silicon isoconcentration surface (red zones, Si > 60 at.%). (b) Si, O, and Hf concentration profile (in at.%) along the growth direction.

homogeneous and contains two layers. The first one is enriched in hafnium (depth between 16 and 19 nm) with the concentration of about 13 at. % contrary to 7.9 at. % observed for the film volume. The second layer (depth between 19 and 21 nm) has higher Si content and its composition is close to SiO₂. The thicknesses of Hf-rich and SiO₂ layers were estimated to be \sim 3 nm and \sim 2 nm, respectively, that is in a good agreement with HRTEM results (Fig. 2). The region *III* corresponds to the silicon substrate.

Figure 4 shows more detailed 3D chemical maps of a volume of the HfSiO film (region *I*), reconstructed after APT experiment, in which each dot represents one atom. For clarity, only 10% of Si and O atoms is shown in these maps. The analysis of the distribution of Si and Hf atoms demonstrates the presence of three different phases: (1) enriched in Hf (black arrow on Hf map), (2) depleted by Hf (white arrow on Hf map) as well as (3) enriched in Si (black arrow on Si map). Thus, it is clearly seen a phase separation process in the film volume, caused by high-temperature annealing. This is in agreement with HRTEM observation (Fig. 2).

In order to describe the spatial distributions of these phases, isoconcentration maps of silicon and hafnium atoms were calculated and are presented in Fig. 5. These filtered images show the three-phases system, where each phase is labelled as α (silica), β (pure Si), and γ (hafnia). Local concentrations of Si, O, and Hf have been measured in each region and reported in Table I.

It is worth to note that atom probe measurements of the composition of nanometric clusters or small fluctuation area can be a difficult task due to artefacts generated by the difference in fields of evaporation between Si-ncs, Hafnia, and Silica. Indeed, this effect, already evidenced in similar



FIG. 4. 3D chemical maps of silicon (red), oxygen (green), and hafnium (blue) in a volume of $28 \times 28 \times 80$ nm³. Hf-rich and Si-rich phases are indicated by black arrow and Hf-poor one with white arrow.



FIG. 5. Isoconcentration maps of silicon atoms (a); oxygen atoms (b), and hafnium atoms (c) in a selected volume $(5 \times 40 \times 50 \text{ nm}^3)$. Different chemical zones, labelled as α (silica), β (pure Sinc), and γ (hafnia) are detected. Pure Si nanoclusters are highlighted with dashcircle.

materials,^{24,25} can introduce a bias for direct composition measurements. In our case, the concentration of Si-ncs, SiO₂-rich, and HfO₂-rich zones was corrected, applying the procedure developed by Talbot *et al.*²⁴ Based on 3D chemical maps (Fig. 4) and isoconcentration maps (Fig. 5), we identified a thin (<1-2 nm) layer between the different phases (SiO₂, HfO₂, and Si-ncs). It consists of Hf-Si-O demonstrating that the phase separation is not complete.

The diameter of individual Si-ncs was calculated by counting the number of Si atoms in each nanocluster. This latter was supposed to have a spherical shape. A dispersion in the size distribution (from 1 nm to 6.5 nm) is observed with a mean gyration diameter of about 2.8 nm. Taking into account the dimensions of the analysed volume, the Si-ncs density was estimated to be $(2.9 \pm 0.4) \times 10^{17}$ Si-ncs/cm³.

D. HAADF-STEM and EELS study of the annealed samples

Phase separation in a mixture of Hf-rich and Si-rich phases in the Hf-Si-O layer was confirmed by a combined HAADF-STEM and EELS experiments. The corresponding HAADF images (sensitive to the Z-contrast) and EELS spectra (sensitive to local chemical composition) for HfSiO films are presented, respectively, on Fig. 6(a) and Figs. 6(b) and 6(c).

The HAADF-STEM image exhibits high-Z (bright) and low-Z (dark) contrast, identified as HfO₂-rich ($Z_{Hf} = 72$ and $Z_O = 8$) and SiO₂-rich ($Z_{Si} = 14$) phases, respectively (Fig. 6(a)). This result has been confirmed by local EELS spectra performed on each contrasted zones. Both contrasted zones (Fig. 6(a)) contain oxygen (Fig. 6(b)), whereas bright and dark zones are hafnium and silicon rich regions, respectively (Fig. 6(c)). Moreover, O-K edge of HfO₂-rich zones exhibits a low energy-loss shift compared to that of SiO₂-rich ones,

TABLE I. Atomic concentration of Si, O, and Hf atoms in different regions labelled α , β , and γ on Fig. 5.

Label	X _{Si} (at. %)	X _O (at. %)	$X_{\rm Hf}$ (at. %)	Phase
α	32.5 ± 0.2	66.2 ± 0.2	1.3 ± 0.2	SiO ₂
В	98.8 ± 0.2	1.2 ± 0.2	0.0	Si
Г	0.0	68.2 ± 0.2	31.8 ± 0.2	HfO_{2}

confirming the chemical difference between dark and bright zones. Such a result is in good agreement with a recent work of Agustin *et al.*³⁰

IV. DISCUSSION

Let's start the discussion by thermodynamic considerations of the Hf-silicate material. Dealing with thermodynamic equilibrium and Gibbs free energy of bulk HfO₂ and SiO₂ phases (respectively, $G_f^0(HfO_2) = -1238.7 \text{ kJ/mol}$ and $G_f^0(SiO_2) = -981 \text{ kJ/mol}$, HfO₂ phase is expected to be stable on silicon substrate without Hf silicide formation. However, several groups have reported about the formation of interfacial silicon oxide layer, when HfO₂ film is in direct contact with Si substrate. The similar effect is shown in this paper. It was clearly demonstrated in Ref. 6 that the formation of this SiO₂-IL is initiated by different diffusion processes. Among them, the most probable is oxygen out-diffusion from high- κ regions close to the Si substrate towards the Si substrate surface. This can explain the appearance of coupled SiO₂rich and HfO₂-rich layers following each other during an annealing.

Locquet *et al.*³¹ have demonstrated the stability of such a layer introducing Gibbs free energy of an interfacial silicon oxide phase in the thermodynamic equations. Nevertheless, formation and stability of the SiO₂-IL do not explain the origin of the observed continuous Hf-rich layer parallel to the substrate. This leads to an assumption of another decomposition mechanism. The thermodynamic analysis of thermal behaviour of bulk high- κ silicates was performed by Kim and McIntyre.²⁹ The silicates were considered as a supercooled liquid for typical transistor processing temperatures (600-1000 °C). The trend for the spinodal decomposition at lower temperatures is caused by negative Gibbs free energy of the silicates. Thus, a driving force for the decomposition process exists for initially homogeneous silicates and results in the formation of SiO₂-rich phase (that contains more than 98 mol. % of SiO₂) and high-*k*-rich (consists of more than 20 mol. % of high- κ oxide phase) upon annealing. This phenomenon was found to be the main decomposition mechanism for the silicates with 40-90 mol. % of SiO₂ content. The consideration of the kinetics of this process revealed the presence of longitudinal composition waves in the bulk



FIG. 6. Cross sectional HAADF-STEM image showing Hf-rich zones (bright contrast) and Hf-poor zones (dark contrast). EELS spectra obtained on bright and dark zones of the HAADF image for (b) O-K edge and (c) Hf-M_{4.5} and Si-K edges.

silicate. Meanwhile, for the film approach, this wave forms in the growth direction due to an effect of Si-subtrate/film interface. Consequently, formation of a layer structure, where SiO₂-rich phase is alternated by high- κ -rich phase parallel to Si subtrate, occurs. It was shown that the silicate composition controls the number of these alternated layers (at other constant annealing parameters) and for the case of more than 50 mol. % of SiO₂ content in high- κ silicate films, only one SiO₂ layer appeared at Si substrate surface followed by one high- κ -oxide-rich layer. As one can see from Fig. 2, these coupled layers are observed in our case. This is an evidence that interface-directed spinodal decomposition plays the main role in the structure evolution of samples investigated in this work. The similar formation of coupled SiO₂-rich and HfO₂-rich continuous layers due to spinodal decomposition mechanism of the film upon an annealing treatment was also observed in Refs. 5, 14, and 15. Meanwhile, any formation of Si-NCs (or ncs) was not evidenced in those works. In the present work, Si-ncs are imaged for the first time, despite that, this was not predicted by the model proposed by Kim and McIntyre.²⁹

As one can see from Fig. 2, the annealing step at 950 °C during 15 min leads to the formation of a dual mixture of SiO₂ and HfO₂ phases as well as Si-ncs in the volume of silicate films, which is confirmed by the analysis of APT results. The formation of Si-ncs in the films investigated can be explained in the next way. Upon HfSiO film annealing, the formation of HfO₂ phase is preferable due to ionic nature of Hf-O bonds. As-grown films have a composition as Hf_{0.07}Si_{0.33}O_{0.60}. Assuming that upon an annealing all Hf atoms are "catched" by HfO₂ phase, then non-stoichiometric SiO_x phase (with O/Si ratio of about 1.4) is formed instead of stoichiometric SiO₂. Consequently, the formation of Si-ncs can occur due to decomposition of the SiO_x phase, similar to that observed for Si-rich-SiO₂ materials.^{9,19} It is worth to note that for HfSiO materials

with about 20 at. % of Hf content, the formation of HfO₂-rich and SiO₂-rich phases occurred only upon post-deposition processing.^{5,15} No evidence of Si-ncs was reported in these works,^{5,15} confirming the effect of the Hf content in the film on the Si-ncs formation.

In the present work no crystalline plane sequences, which could correspond to Si-NCs, have been observed by HRTEM experiments. Moreover, either EFTEM or STEM-EELS did not reveal their presence. This could be due to low density, small size as well as amorphous nature of these Sincs. To go further in the conclusion and limits of EFTEM techniques to detect such clusters, additional experiments in the low energy-loss domain should be performed. On the contrary to TEM technique, APT showed that our sample contains about $(2.9 \pm 0.4) \times 10^{17}$ Si-ncs/cm³. Thus, one can conclude that Si nanoclusters are mainly amorphous, which could be expected due to the annealing conditions.

V. CONCLUSION

In summary, new deposition approach has been proposed for the fabrication of HfSiO layers with specific composition. They have been characterized at the atomic scale by atom probe tomography and transmission electron microscopy. The information about decomposition of $(HfO_2)_x(SiO_2)_y$ $(x + y \neq 1)$ was obtained. A phase separation process in these materials occurs through the formation of HfO_2 , SiO_2 , and Si phases. The appearance of the last one depends strongly on the film composition and is possible for Hf-silicates with lower Hf content. The formation of Si-ncs in such films evidences the capability of magnetron sputtering technique for the production of future Si-rich-high- κ materials, paving way for nanomemory applications. The results also demonstrate that APT and TEM are complementary techniques for atomic scale investigation of clustering and structural characterisation in higk- κ materials.

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