Ultrathin Zirconium Silicate Films Deposited on Si(100) Using Zr(O-i-Pr)\(_2\)(thd)\(_2\), Si(O-t-Bu)\(_2\)(thd)\(_2\), and Nitric Oxide

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Ultrathin Zr silicate films were deposited using Zr(O-i-Pr)\(_2\)(tetramethylheptanedione,thd)\(_2\), Si(O-t-Bu)\(_2\)(thd)\(_2\), and nitric oxide in a pulse-mode metalorganic chemical-vapor deposition apparatus with a liquid injection source. High resolution transmission electron microscopy, atomic force microscopy, X-ray photoelectron spectroscopy (XPS), and medium energy ion scattering were employed to investigate the structure, surface roughness, chemical state, and composition of the films. The nitric oxide used as oxidizing gas, instead of O\(_2\), not only reduced the thickness of the interfacial layer but also removed the carbon contamination effectively from the bulk of the films. The as-deposited Zr silicate films with a Si/Zr ratio of 1:3.1 were amorphous, with an amorphous interfacial layer 0.3-0.6 nm thick. After a spike anneal in oxygen and a 60 s nitrogen anneal at 850°C, these films remained amorphous throughout without phase separation, but the interfacial layer increased in thickness. No evidence of Zr-C and Zr-Si bonds were found in the films by XPS and carbon concentrations <0.1 atom%. The hysteresis, fixed charge density, and leakage current determined from capacitance-voltage analysis improved significantly after postdeposition anneals at 850°C and the films exhibited promising characteristics for deep submicrometer metal-oxide-semiconductor devices.

For standard SiO\(_2\) or SiON gate insulators with thickness <1.5 nm, leakage currents will become prohibitive in a number of the applications of complementary metal-oxide-semiconductor (CMOS) technology. Thus, to improve reliability and reduce power consumption, there is a worldwide search for a replacement for these gate dielectrics having a higher dielectric constant, \(\kappa\). Recent attention has focused on the silicates of those transition or rare-earth metals that are thermodynamically stable on silicon in the sense that their oxides will not be reduced by silicon to form metal or silicide layers. Zirconium silicate has been shown to be thermodynamically stable on Si while possessing a higher crystallization temperature than the pure metal oxide. The silicate will have a lower dielectric constant than the pure metal oxide but this may be partially offset by a higher bandgap or conduction band offset. Usually experimental films of zirconium oxide or silicate are produced by sputtering or e-beam evaporation, techniques not easily transferred to production. A fully production-compatible chemical vapor deposition (CVD) technique is required, likely atomic layer deposition (ALD) or CVD using metalorganic precursors (MOCVD).

In MOCVD, alkoxide and \(\beta\)-diketonate precursors are commonly used. However, the former are air and moisture sensitive. The \(\beta\)-diketonates are less ambient sensitive, and have higher thermal stability requiring higher deposition temperatures. The films can exhibit severe carbon contamination or, for the \(\beta\)-diketonates with fluorine-containing ligands, fluorine contamination. In order to combine the advantages of alkoxides and \(\beta\)-diketonates, mixed ligand Zr and Si precursors were developed to lower the thermal stability but maintain a higher ambient stability and higher volatility.

In this paper, we report the physical and electrical properties of zirconium silicate films deposited by introducing Zr(O-i-Pr)\(_2\)(tetramethylheptanedione,thd)\(_2\) and Si(O-t-Bu)\(_2\)(thd)\(_2\) by liquid injection and introducing oxidizing gas, nitric oxide (NO), or molecular oxygen, into a CVD system. The zirconium silicate films have been analyzed by high-resolution transmission electron microscopy (HRTEM), atomic force microscopy (AFM), X-ray photoelectron spectroscopy (XPS), and medium energy ion scattering (MEIS). The effects of postdeposition anneals were investigated. The current density vs. voltage (I-V) characteristics and capacitance vs. voltage (C-V) characteristics were analyzed for Al-gated capacitors.

Experimental

Si(100) substrates, 100 mm diameter, n-type \((\rho = 0.02 - 0.06 \Omega \text{cm})\) were used as starting material. The wafers were given an HF-last RCA clean prior to insertion into a liquid-injection MOCVD system which has been described in detail previously. In order to deposit zirconium silicates by liquid injection MOCVD the precursors for Zr and Si should be chemically compatible if they are to be mixed. Organometallics with similar ligand structures are less likely to exchange ligands so the precursors, Zr(O-i-Pr)\(_2\)(thd)\(_2\) and Si(O-t-Bu)\(_2\)(thd)\(_2\) produced by ATMI Inc., were employed. A general problem with silicon sources for MOCVD is that they have very low deposition rates at the deposition temperatures chosen for this study. However, silicon can be incorporated at higher rates than expected from the deposition rates for pure SiO\(_2\) films due to an interaction between the Si and Zr precursors. The mixed precursors, 45% 0.15 M solution of Zr(O-i-Pr)\(_2\)(thd)\(_2\) and 55% 0.05 M solution of Si(O-t-Bu)\(_2\)(thd)\(_2\) in octane, were pumped at a rate of 0.2 Ml/min because the pump was unreliable at lower pump rates. During the deposition, nitric oxide was used as oxidizing gas to oxidize the silicate films and facilitate the complete desorption of carbon formed by decomposition of the precursors. The components of the vaporizer, the gas ring, and the connecting tube were maintained at a temperature of 190°C with heating tapes and blankets, while the substrate temperature was controlled at 550°C with quartz-halogen lamps and a thermocouple. This deposition temperature was chosen since deposition rates for the Si precursor were too small at lower temperatures.

The base pressure of the MOCVD chamber was \(-10^{-9}\) Torr. Just prior to deposition most of the wafers were heated for 5 min at 580°C in NO to grow an ultrathin oxynitride buffer layer. The deposition of Zr silicate films was done in the “pulse mode” in which the precursor and nitric oxide were introduced separately through different gas distribution rings with intervening pumping periods of 15 s. Nitrogen at a flow rate of 100 standard cubic centimeters per minute (sccm) was introduced into the nitric oxide gas ring and
flowed during the complete deposition cycle. Thus each cycle of deposition consisted of four stages: (i) 100 sccm N₂ for 15 s, (ii) 150 sccm NO + 100 sccm N₂ for 15 s, (iii) 100 sccm N₂ for 15 s, and (iv) 50 sccm Ar + 100 sccm N₂ + precursor for 40 s. Pressures at various stages of the deposition cycle were in the 8-12 mTorr range. Different oxidizing gases (NO, O₂) were used to evaluate the effect on the interfacial layers.

Films were analyzed by XPS using a PHI 5500 system with a monochromatic Al Kα X-ray source in a standard 90° geometry (X-ray source and electron spectrometer 45° off normal) and a pass energy of 58.7 eV. Depth profiling was accomplished with intermit-
tent Ar⁺ sputtering at 1 keV, 50 nA, and 45° incidence. The peak positions were referenced to the Si 2p½ peak at 99.9 eV.

AFM measurements were made using a Digital Instruments Nanoscope III operating in tapping mode with 5 nm diam silicon probes. Cross-sectional samples were made using a low-angle ion-milling technique. The structure of the films and the effect of annealing were observed directly in HRTEM micrographs made on a Philips EM-430T instrument operating at 250 keV.

The thicknesses were performed at the 1.7 MV high current tandetron facility at the University of Western Ontario. Elastically scattered protons were analyzed at 90° via a toroidal electro-
static analyzer having a resolution ΔE/E = 0.003. The measured energy variance at 100 keV was ~150 eV, which corresponds to a depth resolution of 0.5 nm at the surface of a zirconium silicate sample. Scattered particles were recorded via a position-sensitive (channel plate) detector, thereby providing a two-dimensional spect-
trum (energy-angle) over an angular range of ~30°. A double align-
ment configuration with ions incident along the (101) direction and scattered along the (101) direction was employed to reduce the scatter-
ing yield from the crystalline silicon substrate.

In order to deduce a physical length scale for ion scattering measure-
ment, a density value must be assumed. In the MEIS analysis, values of the density for silicon dioxide and silicon nitride of 2.30 and 2.95 g/cm³, respectively, have been used. These values were deduced from previous nuclear reaction analysis measurements on 100 nm thick films whose thicknesses were determined by ellipsometry.11 A value of 5.06 g/cm³ was used for ZrO₂ and 2.32 g/cm² was used for Si.12

A simulation program, QUARKMEIS, has been used to deter-
mine the composition of the films using a number of layers of varying composition.13 In the simulations, the elastic scattering yield was calculated for each layer, further divided into identical sublayers sufficiently thin, and the composition was varied using a Levenberg-
Marquardt fitting procedure until the agreement with the experimen-
tal spectrum is optimized. The line shapes are determined by con-
volving calculated yields with the response function of the MEIS spectrometer, which is nearly Gaussian but exhibits a low energy tail similar to response functions observed for semiconductor detectors. The response function was determined using the energy spectrum produced by 100 keV protons incident on a thin Au layer (~3 Å) sputtered onto a diamond-like carbon (DLC) substrate. In order to account for the fraction of charged protons transmitted through the MEIS electrostatic analyzer system to the detector, the results of Mao et al.14 were used.

Samples were annealed in a Heatpulse 610 (Steag RTP systems) rapid thermal processor. Keeping the temperature of the nitrogen anneals below 900°C avoids the possibility of phase separation and crystallization.15 For the “spike” anneals in O₂, the temperature was ramped to 1250°C to 850°C and the process was set to pause at this temperature for 1 s before the quartz lamps were turned off. The nitrogen anneals were done at 850°C for 60 s. Forming gas anneals (FGAs) were done in a 4% H₂ in N₂ mixture at 380°C for 20 min.

By comparing the film thickness determined from HRTEM measure-
ments with the accumulation capacitance determined at a fre-
quency of 100 kHz, the equivalent oxide thickness (EOT) and di-
electric constant of a film can be determined. Measurements were made on Al-gated capacitors formed by evaporating aluminum through a shadow mask. No postmetallization annealing was performed to avoid reaction of the Al dots with the Zr silicate films and back contacts were made with In-Ga eutectic. The area of the cap-
acitors was measured to an accuracy of ±2% with a calibrated digitizing camera. Electrical measurements were made by probing the Al gates in a probe station attached to two instruments, a multi-
frequency LCR meter (HP model 4275A) for C-V characteristics, and a picocommernic/duvoltage source (HP model 4140B) for current-
voltage characteristics. High frequency C-V measurements were made by stepping with 0.1 V steps each second, first from −2.5 to +2.5 V and then in the reverse direction to look for hysteresis. The EOT was obtained from the 100 kHz C-V characteristics by using the NCSU C-V fitting routine,16 which includes quantum effects in the channel.

Results and Discussion

In Fig. 1 the HRTEM images of Zr silicate films with a Si/Zr ratio of 1.3/1 are shown. These samples were all grown using 12 cycles of pulse mode deposition at a substrate temperature of 550°C. For these HRTEM images, the spacing of the Si(111) planes (0.313 nm) provided a built-in length reference. In Fig. 1a, the film was deposited with nitric oxide after the growth of the oxynitride layer in nitric oxide at 580°C for 5 min. In Fig. 1b, the film was deposited with oxygen after the growth of the oxynitride layer in nitric oxide at 580°C for 5 min. In Fig. 1c, the film was deposited with oxygen after the growth of the oxide layer in oxygen at 580°C for 5 min. In all cases, the Zr silicate layers and the interfacial layers are amor-
phous. The interfacial layer in Fig. 1a is 0.6 nm thick, which is thinner than those in Fig. 1b and 1c. This is an indication that the oxynitride layer retards oxygen diffusion to the Si interface.

Fig. 1d shows the sample of Fig. 1a after a spike anneal in O₂ at 850°C, followed by a nitrogen anneal at 850°C for 60 s and a FGA at 380°C for 20 min. The Zr silicate layer and the interfacial layer are still amorphous without any phase separation or crystallization. The thickness of the interfacial layer increases to 1.4 nm, while the thickness of the Zr silicate layer remains unchanged. The increase of the interfacial layer is likely due to oxygen or residual water vapor in the thermal processor diffusing to the Si interface during the spike anneal in O₂.

The deposition conditions of the sample in Fig. 1a were used for the films on which the following physical and electrical analysis was performed. In Fig. 2, the root mean square (rms) surface roughness was measured from the AFM image of the Zr silicate film. This 0.25 nm, which is somewhat larger than the value ~0.1 nm obtained on the Si substrates.

In Fig. 3, the XPS C 1s, Si 2p, and Zr 3d spectra of the films are shown. The C 1s spectrum of Fig. 3a reveals three peaks at 285.79, 286.73, and 287.99 eV, The peak at 285.79 eV is ascribed to C-C or di-
atic oxidation state. The peak at 103.20 eV is 3.3 eV from the Si 2p½ and Si 2p 1/2 lines. A spin-orbit splitting con-
stant of 0.61 eV and the intensity ratio Si 2p 3/2 :Si 2p 1/2 o f2w a s
1.37 used to fit these peaks. The peak at 101.26 eV is due to an interme-
planes
0.313

In Fig. 3c, the Zr 3d spectrum consists of one doublet with peaks at 183.59 and 185.99 eV, and the expected Zr 3d\textsuperscript{5/2}:Zr 3d\textsuperscript{3/2} intensity ratio of 3/2. This doublet can be ascribed to Si-O-Zr bonds. There is no evidence of Zr-C or Zr-Si bonds in the Zr 3d spectrum.\textsuperscript{18,20}

In Fig. 3d, the O 1s spectrum reveals two peaks at 532.45 and 533.15 eV. The peak at 532.45 eV can be ascribed to the Si-O-Zr bonds. The smaller peak at 533.15 eV is likely associated with the oxynitride layer underneath the silicate film. The binding energy difference $E_{\text{b}}(\text{O 1s}) - E_{\text{b}}(\text{Zr 3d\textsuperscript{5/2}}) = 348.86 \text{ eV}$ and $E_{\text{b}}(\text{O 1s}) - E_{\text{b}}(\text{Si 2p}) = 429.25 \text{ eV}$ are close to the values of 348.35 eV and 429.5 eV, respectively, reported in Ref. 21.

Figure 4 shows the MEIS spectrum for the as-deposited film of Fig. 1a. The solid line is a simulation of the data using the Quarkmeis code with a four-layer model including the substrate whose surface gives rise to a contribution to the Si peak. The first layer had the composition CSi\textsubscript{0.24}O\textsubscript{0.36}Zr\textsubscript{0.07} with a total carbon areal density which amounts to approximately 3 monolayers of carbon on the surface of the film. Assuming this layer had the same density as silicon dioxide, it is 0.5 nm thick. The Si, Zr, and O in the top layer are probably due to unreacted or partially reacted precursor molecules. Since the film was maintained in flowing NO at the deposition temperature for several minutes after deposition, the amount of undepleted precursor was not expected. Its existence implies that subsequent precursor molecules are involved in the completion of the decomposition reactions of the molecules underneath.

The second layer has the composition Zr\textsubscript{0.33}Si\textsubscript{0.44}O\textsubscript{2} with a Si/Zr ratio of 1.3. A density for this layer can be estimated by assuming that the molecular volume is the sum of molecular volumes of the appropriate amounts of Si, SiO\textsubscript{2}, and ZrO\textsubscript{2}. This gives a density of 2.91 gm/cm\textsuperscript{3} and a layer thickness of 1.5 nm.

A layer at the interface was best modeled as $\text{ZrSi}_{1.3}\text{N}_{0.6}\text{O}_{0.7}$ with a total carbon areal density which amounts to approximately 3 monolayers of carbon on the surface of the film. Assuming this layer had the same density as silicon dioxide, it is 0.5 nm thick. The Si, Zr, and O in the top layer are probably due to unreacted or partially reacted precursor molecules. Since the film was maintained in flowing NO at the deposition temperature for several minutes after deposition, the amount of undepleted precursor was not expected. Its existence implies that subsequent precursor molecules are involved in the completion of the decomposition reactions of the molecules underneath.

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agreement considering the errors in determining thickness from HRTEM micrographs.

The C, N, and O peaks for pieces of the same wafer before and after a spike anneal in O$_2$ at 850°C is shown in Fig. 5. The top layer had the composition $\text{CSi}_0.24 \text{O}_0.13 \text{Zr}_0.15$, and was 0.64 nm thick, assuming it had the density of silicon dioxide. The areal density of carbon decreased, but after annealing, the carbon may have been just adventitious. The Si and Zr in the top layer evidently oxidized. This is corroborated by the oxygen peak in Fig. 5 where the leading edge had shifted to higher energies after oxygen anneal, indicative of some oxide above the Zr silicate layer. The most important change in the MEIS spectra is observed in the broadening of the O peak on the low energy side. The simulation indicates that the interfacial layer increased to a thickness of 0.9 nm and the best fit was obtained assuming that it had the composition $\text{SiN}_0.20 \text{O}_1.73$. Attempts to add a zirconium component to this layer did not improve the fit. The thickness was again somewhat smaller than the 1.4 nm observed by HRTEM as shown in Fig. 1d. The fact that the silicon dioxide density resulted in an underestimate of the interfacial layer thickness suggests that the interfacial layer was underdense. Given that the com-

Figure 3. XPS spectra with background subtracted showing (●●●) data and (-----) fitted peaks for the as-deposited Zr silicate film with 12 cycles at 550°C: (a) C 1s spectrum with two peaks at 285.79, 286.73 and 287.99 eV, (b) Si 2p spectrum with four peaks at 99.8, 100.51, 101.26, and 103.20 eV, (c) Zr 3d spectrum with two peaks at 183.59 and 185.99 eV, and (d) O 1s spectrum with two peaks at 532.45, and 533.15 eV. The solid lines are the sums of the fitted peaks.

Figure 4. MEIS spectrum of 2.7 nm thick as-deposited Zr silicate film with a Si:Zr ratio of 1.3:1 (●●●) data, (-----) simulation.
position is close to that of stoichiometric silicon oxynitride, the magnitude of the discrepancy seems to be much larger than expected. However, errors in determining the thickness of the layers from the HRTEM measurements can be as high as 0.4 nm. The areal density of nitrogen in the interface layer remained constant, to within experimental error, after the spike anneal in O₂.

The 100 kHz C-V characteristics scanned from inversion to accumulation of these films are shown in Fig. 6 for different postdeposition annealing (PDA) treatments. Each sample was spike annealed in O₂ at 850°C first, followed by subsequent anneals. In Fig. 7, the effect of PDA on the flatband voltage shift (ΔVFB), the fixed charge density (Nf), and the effective density of trapped charge at the interface (ΔNt) are shown for the samples in Fig. 6. For the film spike annealed in O₂ at 850°C, there is a significant hump in the C-V curve. Considering the ideal flatband voltage (VFB = Wms − 0.2 V, where Wms is the work function difference), the flatband voltage VFB = 2.01 V corresponds to a positive fixed charge density (Nf = 7.3 × 10¹²/cm²). A counterclockwise hysteresis, with ΔVFB reduced to −26.2 mV, corresponding to ΔNf = 2.4 × 10¹¹/cm². Thus the FGA reduced the charge trapping/detrapping and passivated the P₃ centers effectively.

For the sample spike annealed in O₂, then nitrogen annealed at 850°C for 60 s and annealed in forming gas at 380°C for 20 min, the flatband voltage shifted to −0.43 V (Nf = 1.8 × 10¹²/cm²). The value of ΔVFB reduced to −22.5 mV (ΔNf = 1.9 × 10¹¹/cm²). The N₂ anneal reduced not only the charge trapping/detrapping but also the leakage current density (shown later). For the same sample, negligible hysteresis was observed in the inset of Fig. 7a. In Fig. 7, it is seen that ΔVFB, Nf, and ΔNt all decreased after the additional nitrogen anneal. The reduction in Nf might have been due to the
introduction of negative fixed charge and/or the annealing of the positive fixed charge.

Figure 8 shows the 10 kHz, 100 kHz, and 1 MHz C-V curves of the Zr silicate film spike annealed at 850°C in O₂, then nitrogen annealed at 850°C for 60 s and annealed in forming gas at 380°C for 20 min. The area of the capacitor is 2.8 × 10⁻⁴ cm². From the 100 kHz C-V curve, the EOT is 2.32 nm with the quantum correction. Compared with the frequency dispersion of the sample spike annealed in O₂ at 850°C and then given a FGA at 380°C for 20 min (shown in the inset of Fig. 8), there is only a small frequency dispersion in the C-V curves due to interface states, mainly below the midgap. This indicates that most of the interface states, not passivated by the FGA, were eliminated by the nitrogen anneal at 850°C for 60 s. The slight dispersion due to the remaining interface states might be further reduced with a postmetallization anneal because the reaction between the metal and residual water on the surface can generate atomic hydrogen which is more effective in passivating interface states than molecular hydrogen.

For the as-deposited and spike annealed samples of Fig. 8, electron-energy-loss spectroscopy (EELS) spectra confirmed the conclusion from the MEIS spectra that the Zr content in the interfacial layer was below the detection limit. Assuming a dielectric constant appropriate for SiO₂ of 3.9 for the interfacial layer, a dielectric constant of 8.5 was determined for the Zr silicate layer.

Figure 9 shows the corresponding leakage current densities as a function of gate voltage for the samples under substrate injection (positive gate voltage). The leakage current of the film is reduced by orders of magnitude after the spike anneal in O₂. Some of the reduction of leakage current is due to the increase of the EOT associated with the increase in interfacial layer thickness noted in the Fig. 9 and the rest may be due to the decrease of traps which contribute to trap-assisted tunnelling current. The small change observed after the FGA may be due to variations in film thickness over the wafer. A further significant current decrease in leakage current is noted after nitrogen annealing, probably related to a further decrease in trap-assisted tunnelling. For the annealed samples, the J-V curves are similar to those associated with soft breakdown in SiO₂ films and are characteristic of the trap-assisted tunnelling in the Zr silicate films.

In Fig. 10, the leakage current densities as a function of gate bias under substrate and gate injection are shown, indicating the polarity dependence. The leakage current density at V_G-V_FB = 1 V is 1.4 × 10⁻⁴ A/cm², which is about 20 times lower than that of SiO₂ with the same EOT (3 × 10⁻³ A/cm²). The leakage was higher under substrate injection than under gate injection. The tunnelling probability is inversely proportional to the tunnelling area defined by the electron tunnelling distance and the barrier height. Under gate injection, the tunnelling area was larger, i.e., the electron tunnelling probability is lower as described in Ref. 30, thus the leakage current is lower for gate injection.

Conclusions

In this study, ultrathin Zr silicate films were deposited on Si(100) using Zr(O-i-Pr)₂(thd)₂ and Si(O-i-Bu)₂(thd)₂ dissolved in octane. Either O₂ or NO was used as the oxidizing gas. Nitric oxide forms
an oxynitride buffer layer which retards oxygen diffusion to the Si interface. Both NO and O₂ reduce the carbon contamination effectively. No evidence of C-Si, Zr-C and Zr-Si bonds were found in the XPS spectra. HRTEM images showed that the as-deposited Zr silicate films were amorphous. After postdeposition anneals at 850°C, Zr silicate films with the composition Si:Zr ratio of 1:3:1 were still amorphous and no phase separation was observed.

The HRTEM images and MEIS analysis showed that a substantial interface layer, 0.9-1.4 nm thick, was produced by a spike anneal in O₂ at 850°C. The Zr concentration in the interfacial layer was below the sensitivity of the MEIS technique (~5 atom %) after annealing and this has been confirmed by EELS analysis. Limiting the growth of the interface layer will require a reduction in the thermal budget during any O₂ anneal, and this needs to be further investigated.

One of the functions of the O₂ anneal was to eliminate the C from precursor evidently left on the surface of the film. The fact that this remaining precursor was not oxidized during the final oxidation stage in the deposition chamber suggests that the decomposition of the precursor is a rather complicated process requiring both a separate oxidant and species produced during the succeeding precursor pulse.

The ultrathin Zr silicate films described in this work exhibit promising C-V and J-V characteristics. The P₉ centers at the interface were effectively passivated by FGA, consistent with the hypothesis that the interfacial layer was silicon oxide. The hysteresis, fixed charge density, and the effective density of trapped charge at the interface are also improved significantly by anneals at 850°C. Most of the frequency-dependent interface states, not passivated by the low temperature FGA, were eliminated after the 60 s nitrogen anneal at 850°C. The slight remnant frequency dispersion below midgap may be reduced if a proper FGA were performed after, rather than before, gate electrode definition. Higher temperature annealing might also reduce the remnant dispersion but the films may crystallize at temperatures above 900°C. After a spike anneal in O₂ at 850°C and an anneal for 60 s in N₂ at 850°C, a Zr silicate film with an EOT of 2.3 nm exhibited a leakage current about 20 times lower than that of a SiO₂ film with the same EOT. These electrical properties demonstrate that a thin Zr silicate film formed by ZrO₂-Pr₃(THD)₂·SiO₂-Bu₃(THD)₂ and nitrile oxidemight be a promising gate dielectric for deep submicron CMOS devices, if ways can be found to decrease the interface layer thickness.

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