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# Bonding constraints and defect formation at interfaces between crystalline silicon and advanced single layer and composite gate dielectrics

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An increasingly important issue in semiconductor device physics is understanding of how departures from ideal bonding at silicon–dielectric interfaces generate electrically active defects that limit performance and reliability. Building on previously established criteria for formation of low defect density glasses, constraint theory is extended to crystalline silicon–dielectric interfaces that go beyond  $Si-SiO_2$  through development of a model that quantifies average bonding coordination at these interfaces. This extension is validated by application to interfaces between Si and stacked silicon oxide/nitride dielectrics demonstrating that as in bulk glasses and thin films, an average coordination,  $N_{av}$ , greater than three yields increasing defective interfaces. © 1999 American Institute of Physics. [S0003-6951(99)00414-3]

As integrated circuits are aggressively scaled to increase device packing, channel lengths in field effect transistors (FETs) are projected to decrease to  $\sim 50$  nm by 2012 with the oxide-equivalent thickness ( $t_{\rm ox-eq}$ ) for gate dielectrics decreasing proportionally to <1 nm. For  $t_{\rm ox-eq}<2.5$  nm direct tunneling in SiO<sub>2</sub> becomes important in off-state leakage. Since tunneling increases exponentially with decreasing oxide thickness, this necessitates introduction of alternative insulators such as Si<sub>3</sub>N<sub>4</sub>, and Ta<sub>2</sub>O<sub>5</sub> with dielectric constants>SiO<sub>2</sub>. These provide scaled-down values of  $t_{\rm ox-eq}$  required to maintain FET current drive, while reducing tunneling through increases in film thickness.

For oxides <2.5 nm, interfacial defects such as Si dangling bonds, can limit performance and reliability, making it important to establish relationships between interface bonding chemistry and defect properties. Experiments on stacked silicon oxide/nitride dielectrics prepared by remote plasmaenhanced chemical vapor deposition (RPECVD) have provided insights into these issues. RPECVD, followed by rapid thermal annealing (RTA) at 900 °C in a nonoxidizing ambient has yielded device-quality nitrides for n- and p-channel FETs with  $t_{\rm ox-eq}{\sim}2$  nm. 1.2 Minimization of Si and N atom dangling bonds in annealed bulk RPECVD nitrides derives from low levels of bonded H,  ${\sim}15$  at. %.  $^{3,4}$ 

Figure 1 displays current-voltage (I-V) curves for

p-channel FETs for different gate dielectrics with  $t_{ox-eq}$ ~2 nm: (i) a 4 nm RPECVD nitride, (ii) a 0.5 nm plasma oxide with a 2.4 nm RPECVD nitride, and (iii) a 1.5 nm thermal oxide with a 1.0 nm RPECVD nitride. I-V traces for (ii) and (iii) display excellent turn-on behavior and the same current drive, with differences in threshold voltage derived primarily from differences in substrate doping. In contrast, for the FET with the 4 nm nitride: (i) threshold voltage is shifted negative by >1 V, (ii) turn-on is soft, and (iii) channel drive current is reduced by a factor of  $\sim$ 50. Figure 2 displays capacitance-voltage (C-V) characteristics for p-type metal-oxide-semiconductor (PMOS) devices with  $t_{\text{ox-eq}} \sim 4.3 \text{ nm}$ : one with a plasma oxide, and two with stacked dielectrics with RPECVD nitride interface layers of 0.4 and 0.8 nm, respectively. Shifts in threshold (and flat band) voltage relative to Si-SiO2 indicate increased fixed charge for devices with nitride interfaces:  $\Delta q_f = C_{\text{ox}} \Delta V_{\text{th}}$  (or  $\Delta V_{\rm fb}$ ). Qualitatively similar results have been obtained for n-type metal-oxide-semiconductor (NMOS) devices with nitride layer interfaces.4

Three factors can play a role in promoting interfacial defects: (i) interfacial dipoles due to charge transfer between the Si substrate and gate dielectric,<sup>5</sup> (ii) molar volume differences between the Si substrate and gate dielectric, and (iii) over-constrained bonding due to large values of average coordination in the interfacial region.<sup>6</sup> Charge transfer is smaller at nitride interfaces so that interfacial dipoles cannot

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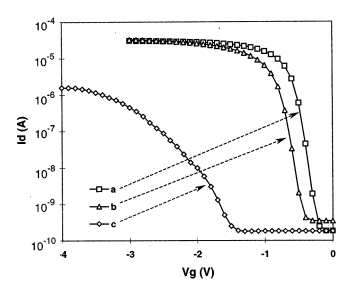


FIG. 1. Drive current-gate voltage  $(I_d - V_g)$  characteristics for PMOSFETs with  $t_{ox-eq} \sim 2$  nm: (a) a 1.5 nm oxide separating a 1.0 nm nitride from the Si substrate, (b) a 0.6 nm oxide separating a 2.4 nm nitride from the Si substrate, and (c) a 4 nm nitride layer. The threshold voltage shift between (a) and (b) is due in part to substrate doping differences (0.16 V) and in part to positive charge at the oxide-nitride interface (0.04 eV).

play the determinant role in increased defect densities.<sup>5</sup> Since the molar volume mismatch between Si<sub>3</sub>N<sub>4</sub> and Si is reduced with respect to that of SiO<sub>2</sub> and Si, residual interface strain is smaller and cannot be the origin of higher defect concentrations at Si-Si<sub>3</sub>N<sub>4</sub> interfaces. The remainder of this letter focuses on interfacial bonding constraints.

The abruptness of Si-SiO<sub>2</sub> interfaces suggests that the defect density of thermally grown oxides and optimally annealed deposited oxides is a characteristic function of their bonding chemistry and structure. Experience with good glass formers such as SiO<sub>2</sub> and As<sub>2</sub>Se<sub>3</sub> has shown that as long as only single bonds are present charge transfer plays a minor part in determining structure. The major factor is the network stress which arises for a given space-filling bonding topology.

Constraint theory has provided a remarkably accurate description of network stress and its consequences.<sup>6,7</sup> The theory is based on the idea that all the bonding forces (stretching, bending, etc.) in the network can be arranged in a hierarchy from strong to weak. The constraining effects of these forces are a linear function of the average coordination number,  $N_{\rm av}$ . If both bond-bending and stretching forces are present, the optimal average coordination number,  $N_{\rm av}^*$ , which matches constraints to degrees of freedom is 2.4 as in  $As_2S(Se)_3$ , however, for  $SiO_2$ ,  $N_{av}^* = 2.67$  is optimal because bond-bending forces at O atoms are too weak to function as significant constraints at growth or annealing temperatures.8 For over-constrained networks such as  $Si_3N_4$  ( $N_{av}=3.43$ ), Si-atom stretching constraints are stronger than bending constraints, so that strain energy accumulates along the bending constraints. The average Si-N-Si bond angle  $\theta_{ij}$  is distorted from the ideal local value  $\theta_{ij}^*$  by an amount

$$\delta\theta \propto \delta N_{\rm av}^* = N_{\rm av} - N_{\rm av}^*. \tag{1}$$

Since total strain energy is proportional to  $(\delta\theta)^2$ , it is then anticipated that defect creation, e.g., dangling Si or N bonds,

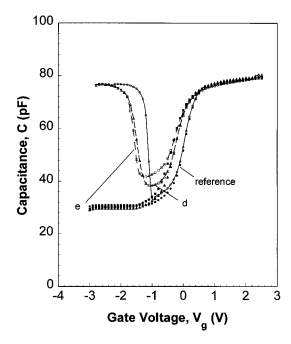


FIG. 2. C-V characteristics demonstrate shift in flat band voltage due to positive charge, and increased separation between high frequency and quasistatic plots due to interface trapping accompanying direct deposition of thin nitride films onto Si. Each of these capacitors has  $t_{\text{ox-eq}} \sim 4.3 \text{ nm}$ : (i) a reference oxide, and (ii) two stacked NO structures with the nitride layer in contact with the Si substrate. The nitride layer thickness is 0.4 nm for (d), and 0.8 nm for (e).

shown that  $N_{\rm av} \sim 3$  represents a criterion between low defect  $(\sim 10^{16} \, \text{cm}^{-3}),$ and increasingly density defective materials.10

Extension of constraint theory to Si-dielectric interfaces considers three interfacial contributions to  $N_{\rm av}$ : (i) the Si substrate represented by one-half a Si atom, (ii) an ultrathin oxide or nitride interfacial layer (0.3–0.6 nm), and (iii) the bulk dielectric by one-half a molecular layer. Table I includes calculations of  $N_{\rm av}$  for representative Si-dielectric interfaces. When a demarcation level  $N_{av} \sim 3$  is applied, these calculations are in excellent agreement with experiment (see Figs. 1 and 2). The model confirms that Si-SiO<sub>2</sub> interfaces are expected to display excellent interface properties ( $N_{av}$  $\sim$ 2.8), whereas Si-Si<sub>3</sub>N<sub>4</sub> interfaces are not  $(N_{av}\sim$ 3.5). Equally important, the calculations demonstrate that interposition of ultrathin SiO<sub>2</sub> layers between Si and Si<sub>3</sub>N<sub>4</sub> results in values of  $N_{av} \le 3$ , whereas interposition of ultra thin Si<sub>3</sub>N<sub>4</sub> layers between Si and SiO<sub>2</sub> results in  $N_{av}>3$ . Figure 3 is based on the data of Figs. 1 and 2, and demonstrates that defect scaling for bulk films, Eq. (1), also holds at interfaces.

Nitride layers have been produced by techniques other than RPECVD. For example, films have been prepared by jet vapor deposition (JVD) yielding excellent electrical results.<sup>13</sup> However, nitride films produced in this way can have oxygen concentrations as high as 15-18 at. %, and as such gate stacks incorporating these films are expected to have significant Si-SiO<sub>2</sub> bonding at the Si-dielectric interface. Without more detailed characterizations of interfacial oxide concentrations in the JVD films, it is not possible to make direct comparisons between the performance of these devices and the quantitative aspects of constraint theory.

The model has also been applied to interfaces between will be proportional to  $\{N_{\rm av}-N_{\rm av}^*\}^2$ . Experiments have Si and (i) silicon oxynitride alloys and (ii) alternative high- $N_{\rm av}$  bounloaded 22 Feb 2008 to 152.1.190.114. Redistribution subject to AIP license or copyright; see http://apl.aip.org/apl/copyright.jsp Si and (i) silicon oxynitride alloys and (ii) alternative high-K

TABLE I. Average bonding coordination at Si-dielectric interfaces.

Material system	Average coordination $(N_{av})$	Electrical quality
Si–SiO <sub>2</sub> (1.5 molecular layers)	2.8	excellent, thermal oxides
Si-Si <sub>3</sub> N <sub>4</sub> (1.5 molecular layers)	3.5	very poor [Ref. 4]
$Si-\{SiO_2\}(t)-Si_3N_4$	t = 0.6  nm: 3.0	very good [Fig. 1]
t = oxide layer thickness	t = 1.5  nm: 2.9	excellent [Fig. 1]
$Si-\{Si_3N_4\}(t)-SiO_2$	$t = 0.4 \text{ nm}$ : $3.3^{\text{d}}$	poor [Fig. 2]
t = oxide layer thickness	t = 0.8  nm: 3.4	poor [Fig. 2]
Si-N-SiO <sub>2</sub> {1 monolayer (ML)}	2.8	excellent [Ref. 5]
$Si-(SiO_2)_{0.977} \{Si_3N_4\}_{0.023}$	2.3 at. % N: 2.8	excellent [Ref. 14]
$Si-(SiO_2)_{0.89} \{Si_3N_4\}_{0.11}$	11 at. % N: 3.0	poor [Ref. 14]
Si-TiO <sub>2</sub> } <sup>a</sup> (1.5 molecular layers)	4.0	unreported
$Si-Ta_2O_5$ (1.5 molecular layers)	3.5	unreported
Si-Al <sub>2</sub> O <sub>3</sub> } <sup>c</sup> (1.5 molecular layers)	3.6	unreported

<sup>&</sup>lt;sup>a</sup>Average coordination [Ti]=6, [O]=3.0 [rutile/anatase bonding].

dielectrics. For oxynitride alloys, Vogel et al., 14 reported only a small reduction in electron channel mobilities in an alloy with about 2.3% nitrogen corresponding to  $N_{\rm av} \sim 2.8$ . While the criterion developed by the application of constraint theory predicts that the interfacial defect density should not become appreciable until the alloy concentration exceeds approximately 15-20 at. % N, interfacial defects in n-channel FETs degrade channel mobilities by more than a factor of two at a nitrogen concentration of only 11 at. % corresponding to  $N_{\rm av} \sim 3.0$ . This demonstrates that factors other than interfacial bonding constraints can contribute in a significant way to the formation of interfacial defects. However, it is important to note that for the oxynitride alloys of Ref. 14, the experimentally determined defect levels are higher than what is predicted by the application of constraint theory, so that constraint theory may still provide a guideline for estimating the minimum defect densities. The model calculations for Ta<sub>2</sub>O<sub>5</sub>, TiO<sub>2</sub>, and Al<sub>2</sub>O<sub>3</sub> explain the necessity for ultrathin SiO2 layers between the Si substrate and these high-K oxides. As such the model suggests important limitations for gate dielectric interfaces other than Si-SiO<sub>2</sub>. Specifically: (i) Si<sub>3</sub>N<sub>4</sub> cannot be directly substituted for SiO<sub>2</sub> at

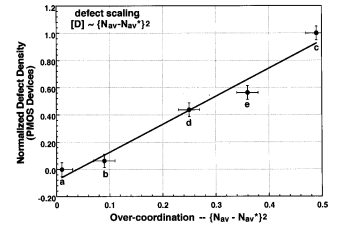


FIG. 3. Plot of normalized defect density as a function of  $\{N_{av} - N_{av}^*\}^2$ . Data points a, b, and c are from Fig. 1, and d and e from Fig. 2.

Si substrates; and (ii) substitution of more highly coordinated high-K dielectrics such as  $Ta_2O_5$ , etc., will generally require  $SiO_2$ , or monolayer nitrided  $SiO_2$  interfaces, thus establishing a limitation on the extent to which  $t_{\rm ox-eq}$  can be reduced below 1 nm. Finally, the bonding constraint model should be taken as a guideline for anticipating differences in interface quality that arise solely from differences in average bonding coordination. As such it focuses on one important aspect of interfacial bonding and structure. It is anticipated that as for the oxynitride dielectrics of Ref. 12, additional aspects of interface bonding and structure will also generate significant concentrations of electrically active defects at other Sidielectric interfaces.

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<sup>&</sup>lt;sup>b</sup>Average coordination: [Ta]=6, [O]=2.4 [Ref. 11].

<sup>&</sup>lt;sup>c</sup>Average coordination: Al=[4.5], [O]=3.0 [3:1 ratio of tetrahedral to octahedral sites, see Ref. 12].

<sup>&</sup>lt;sup>d</sup>Sample calculation of  $N_{av}$  for Si-{Si<sub>3</sub>N<sub>4</sub>}(t)-SiO<sub>2</sub>: t=0.4. Substrate: 1/2 atomic layer: 0.5 atoms, 2 bonds. Interface layer: 1 molecular layer: 7 atoms (Si<sub>3</sub>N<sub>4</sub>), 24 bonds. Dielectric film: 1/2 molecular layer: 1.5 atoms (SiO<sub>2</sub>), 4 bonds. 30 bonds/9 atoms= $N_{av}$ =3.3 bonds/atom.

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