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Control of atomic layer degradation on Si substrate

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To develop 32 nm node devices, the degradation of atomic layers on the surface of Si substrates must be controlled. During the etching of a $SiO₂$ or $Si₃N₄$ hard mask or sidewall, the surface of Si is attended due to exposure to fluorocarbon plasma. The authors have quantitatively evaluated the relationship between the energy of incident ions and the thickness of the fluorocarbon polymer for a $CH_2F_2/CF_4/Ar/O_2$ plasma in a dual frequency CCP system. At a fixed ion energy the thickness of the damage layer (T_d) basically depended on the thickness of the fluorocarbon polymer (T_{C-F}) . When the T_{C-F} was changed by controlling the O/CF_{*x*} gas ratio, T_d had a minimum thickness under the conditions at balance point: P_b , under which the T_{C-F} was nearly equal to ion penetration depth: D_p . Using molecular dynamics simulation, reaction around the transition from SiO₂ to Si was clarified. The damage was done to the Si before the $SiO₂$ was completely removed, and the largest T_d was observed when the SiO₂ was etched off. After that, T_{C-F} began to increase because there was no longer an outflux of O from $SiO₂$ and the damage decreased as the unstable SiF_x species in the damaged layer desorbed. Once the $T_{\text{C-F}}$ became thicker than the ion penetration depth, the damaged layer got buried and T_d stopped changing. When the ion penetration depth was controlled to be equal to T_{C-F} in a steady state under low ion energy conditions, the T_d was reduced to less than 1 nm. *© 2007 American Vacuum Society.* DOI: 10.1116/1.2713114

I. INTRODUCTION

Fluorocarbon plasmas have been widely used to etch dielectric materials for the fabrication of ultralarge-scale integrated circuit devices. To advance semiconductor device design, 32 nm node devices must be fabricated with an ultrashallow junction.^{1,2} Unfortunately, the surface of Si is degraded by exposure to fluorocarbon plasma during etching of sidewalls. This unstable surface layer can cause defects or fluctuations in transistor properties because the activation of impurities doped into the source/drain region is subtly affected by the surface damage. 3 The minimization of substrate damage is necessary to stabilize transistor properties. Models for controlling surface reactions during etching of Si, $Si₃N₄$, $SiO₂$, and SiOCH films have been extensively discussed.⁴⁻⁶

Steady-state etched surfaces are very sensitive to changes in radical composition as well as to ion energy. In this article, we will elucidate the relationship between the damage to Si substrates and the parameters in fluorocarbon plasma. A method of minimizing Si damage will be demonstrated.

II. EXPERIMENTS

We used a $CH_2F_2/CF_4/Ar/O_2$ plasma in a dual frequency CCP $(60/2 \text{ MHz})$ system⁷ in our experiments. The gap between the two electrodes was fixed at 35 mm. The temperatures of the top electrode, bottom electrode, and sidewall were kept at 60, 20, and 60 °C, respectively. A high-voltage probe was used to measure the peak-to-peak voltage $(V_{\text{p.p.}})$ of the rf-biased bottom electrode; the electron density N_e was measured with a surface wave probe. Radical densities were measured with an optical emission spectroscope. We also used a quadrupole mass spectrometer to determine plasma

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FIG. 1. Damage to Si substrate caused by exposure to $CF_4/CH_2F_2/Ar/O_2$ plasma: (b) as etched and (c) after ashing.

potential *Vp*. Pressure was kept at 30 mTorr in these experiments, while partial pressure of each gas molecule in the $CH_2F_2 / CF_4 / Ar/O_2$ mixture varied. The damaged layer to the Si substrate was oxidized by O_2 plasma after fluorocarbon plasma irradiation. The damage thickness (T_d) was defined by measuring the thickness of $SiO₂$ by using ellipsometry. The thicknesses of both the damage and the C–F polymer were cross-checked using a tunneling electron microscope (TEM). Using molecular dynamics (MD) simulation, δ the time dependence of the surface structure during the early stage of etching was also determined.

III. RESULTS

A. Formation of degraded layer on Si substrate

We used fluorocarbon $\text{CH}_2\text{F}_2/\text{CF}_4/\text{Ar}/\text{O}_2$ plasma to fabricate the sidewall of the transistor gate electrode [see Fig. 1(a)]. As shown by the TEM image, the top surface of the Si substrate was degraded and a fluorocarbon polymer layer was formed on the damaged layer, as shown in Fig. $1(b)$. We were able to remove the polymer layer using O_2 ashing Fig. 1(c)]. The damaged layer, however, remained after ashing and wet treatment. The thicknesses of the damage (T_d) and the C–F polymer $(T_{\text{C-F}})$ depended on the plasma conditions as well as the atomic compositions (especially O concentration) of etched materials.

FIG. 2. Optical emission intensity ratio of CF_2/Ar , O/Ar, and O/CF₂ as a function of $O₂$ flow rate.

B. Effects of radical composition on damage thickness

To clarify the relationship between the plasma conditions and the T_d , we changed the flow rate of O_2 in $CH_2F_2 / CF_4 / Ar/O_2$ plasma in fixed power conditions. The plasma density and $V_{\text{p.p.}}$ were $1 \times 10^{1\hat{1}} \text{ cm}^{-3}$ and 1230 V, respectively. Figure 2 shows the densities of CF_2 and O radicals [optical emission ratios of $I_{CF2}(251.8 \text{ nm})/$ $I_{\text{Ar}}(750.4 \text{ nm})$ and $I_0(777.2 \text{ nm})/I_{\text{Ar}}$ as a function of O₂ flow rate. The ratio of O to CF_2 was observed to increase monotonically with the O_2 flow rate. In Fig. 3, T_d is shown as a function of the O/CF_2 ratio. When we increased the O/CF_2 ratio, T_d first decreased ($O/CF_2 = 0 - 0.07$), then reached the minimum peak (at 0.07; condition B), and then increased again (>0.07) though $T_{\text{C-F}}$ decreased monotonically. Figure 4 shows T_d and T_{C-F} in conditions A, B, and C in Fig. 3 as a function of plasma exposure time. Under condition A, $T_{\text{C-F}}$ increased linearly, indicating a constant fluo-

FIG. 3. Thickness of damaged layer (T_d) and C–F polymer (T_{C-F}) as a function of $O/CF₂$ ratio.

FIG. 4. T_d and T_{C-F} as a function of plasma exposure time. O/CF₂ ratios were 0 (A), 0.07 (B), and 0.3 (C). rf bias power was kept constant at 800 W.

rocarbon polymer deposition rate. Also under this condition, damage was done as plasma exposure began, and T_d did not change as time passed. We observed a slight decrease in T_d and a slight increase in $T_{\text{C-F}}$ under condition B. When the $O/CF₂$ ratio was higher than B, the Si etching rate increased (condition C), $T_{\text{C-F}}$ was thinner than 0.5 nm, and the Si was etched. T_d was relatively thick and was almost constant as time passed.

C. Effect of ion energy on damage thickness

When we changed the substrate bias power, T_d also changed, as shown in Fig. 5. $V_{p,p}$ at bias powers of 1500, 800, and 400 W were 1860, 1230, and 770 V, respectively. The thicknesses of the C–F polymer were not affected by the bias power and almost the same graph (as shown in Fig. 3) was obtained for each bias condition further decrease in ion energy, however, induced excess deposition of C–F polymer). $T_{\text{C-F}}$ depends only on radical composition, and when we fixed the balance between the total amount of incident fluxes of CF_x and O, T_{C-F} could be uniquely determined. The position of minimum peak (condition B in Fig. 3) shifted to a lower O/CF_2 ratio under higher bias power conditions. When the O/CF_2 ratio was lower than this minimum peak, we observed a deposition rate (thickness increased with time) of C–F polymer and no etching of the Si substrate.

FIG. 5. T_d as a function of O/CF₂ ratio. Bias power was varied from 400 to 1500 W.

IV. DISCUSSION

A. Balance point

As shown in Figs. 3 and 5, there is a specific point where T_d has a minimum in each bias condition. We called this point the "balance point," P_b . Since the position of P_b seemed to depend on both the ion energy and polymer thickness $T_{\text{C-F}}$, we quantitatively analyzed the relationship between T_d and the ion penetration depth into the C–F polymer. Figure 6(b) shows the ion energy distribution function (IEDF) calculated using the Monte Carlo simulation.⁹ Highenergy IEDF peaks at bias powers of 400, 800, and 1500 W were observed at about 780, 1230, and 1840 eV, respectively. The C–F polymer on the etched surface decelerated the ion energy during etching, showing that the amount of energy loss, ΔV , depends on $T_{\text{C-F}}$. The ion penetration depth D_p was roughly estimated using data from Tatsumi *et al.*,¹⁰ as shown in Fig. 6(a). For example, ions can penetrate through a 7.8-nm-thick C–F polymer at a bias power of 800 W. $(D_p$ was experimentally obtained for ions with the averaged ion mass of all of incident ions. Therefore, D_p also depends on ion composition.) By comparing D_p in Fig. 6(a) to $T_{\text{C-F}}$ in Fig. 7(a) (or Fig. 3), we found that P_b can be related to the point where $T_{\text{C-F}}$ is equal to D_p . This indicated that, to minimize the damage, it is necessary to control the balance between the high-energy IEDF peak and the C–F polymer thickness on a steady-state etched surface. Next, we classify the plasma conditions into regions where $T_{\text{C-F}}$ was thinner than D_p (Sec. IV B) and thicker than D_p (Sec. IV C) and discuss why T_d changed as radical composition and ion energy in both regions varied.

B. Damage under "thin polymer" conditions

 $T_{\text{C-F}}$ decreased monotonically as the O/CF₂ ratio increased. When the O/CF_2 ratio was larger than P_b , the Si substrate was etched and T_{C-F} was kept constant in a steady state. The ion energy deposited at the interface between the

FIG. 6. (a) Penetration depth of ions (data from Ref. 10 were used) and (b) ion energy distribution.

C–F polymer and the Si substrate can be determined by both the incident ion energy (high-energy IEDF peak) and the energy consumption by the polymer (which depends on ion mass and $T_{\text{C-F}}$). T_d basically depends on ion penetration depth, which is thought to depend on the net ion energy supplied below the C–F polymer, as shown in Fig. 7. T_d became thicker when incident ion energy increased or when $T_{\text{C-F}}$ became thinner, under relatively large O/CF₂ conditions. To decrease T_d (and to decrease the etching rate of Si), it is necessary to decrease the O/CF_2 ratio under the "thin polymer" condition.

C. Damage under "thick polymer" condition

When we used conditions where O/CF_2 ratio was lower than P_b , ions could not penetrate into the Si substrate in a steady state because $T_{\text{C-F}}$ was larger than D_p . However, the degradation of the Si surface was also observed under these conditions. Furthermore, T_d became thicker when we decreased the O/CF_2 ratio, even though the deposition rate of the C–F polymer was relatively high. As shown in Fig. 4, this damage was formed as plasma exposure to the Si wafer began. To determine the mechanism of the formation of the degradation layer under "thick polymer" conditions, we calculated the reaction on the etched surface during the early stages of polymer formation on the Si surface using MD simulation (see Fig. 8). First, we fabricated the SiO_2/Si substrate and irradiated CF_3 (250 eV) to form a steady-state C-F polymer during $SiO₂$ etching, as shown in Fig. 8(a).

FIG. 7. (a) $T_{\text{C-F}}$ and (b) T_d as a function of O/CF₂ ratio. T_d has a minimum at balance point (P_b) where T_{C-F} was equal to ion penetration depth.

 $T_{\text{C-F}}$ is relatively thin during SiO₂ etching. As CF₃ dosage on the same substrate increased, $SiO₂$ became thinner than the penetration depth of CF_3 and some of the CF_3 started to reach the underlying Si [Fig. 8(b)]. When $SiO₂$ was just etched off, the C-F polymer was still thin [Fig. $8(c)$] and the degradation layer was formed on Si. When the deposition rate was relatively high, the C–F polymer started to grow thicker because C–F polymer cannot be removed by O supplied from an etched surface (Fig. 3 and A in Fig. 8). The damaged layer was covered by a thickly deposited C–F layer. On the other hand, when the polymer deposition and etching were balanced in a steady state, the damaged layer decreased slightly as dosage increased (increasing etch time), as shown in B in Fig. 8. Figure 9 shows the atomic composition of mixed layer as a function of CF_3 dosage. The surface composition was basically consisting of C, F, (O), and Si: and the ratios of F/C and Si/C were gradually increased along with

FIG. 8. Image of change in surface structure as a function of plasma exposure time. A: Thick polymer condition $(T_{C-F} > D_p)$. Damage [formed similarly to B (a)–(c)] was buried under polymer. B: Balance point $(T_{C-F} = D_p)$. Surface structure was calculated using molecular dynamics simulation. CF₃ beam of 250 eV was introduced to SiO₂/Si structure. (a) During SiO₂ etching. Thin polymer was formed on surface. (b) Remaining thickness of SiO₂ became thinner than D_p . (c) SiO₂ was etched off and damage was formed. T_{C-F} started to grow thicker. (d) Unstable C–F or Si–F species desorbed. (e) Steady-state C–F polymer was formed on a minimum amount of damage. C: Thin polymer condition $(T_{C-F} < D_p)$. Thinner T_{C-F} induces thicker T_d . Etching of Si also occurred.

the depth from the top surface. The interface between mixing layer and Si substrate has a F-rich composition. By increasing the dosage of CF_3 injection, Si atom in mixed layer was decreased. MD calculation also suggested SiF*^x* was released from surface during over etching [Figs. $8(c) - 8(e)$]. The top part of mixing layer where Si content was low could be removed during exposure to O_2 plasma [see Fig. 1(c)]. Consequently, the thinner damage was observed after longer plasma exposure under condition B. The relaxation of the degraded layer (release of unstable SiF_x) is the reason why T_d has a minimum around P_b . As mentioned in Sec. IV B,

when steady-state $T_{\text{C}-\text{F}}$ was thinner than D_p , the degraded layer was kept at a constant thickness, as shown in C in Fig. 8. A thinner polymer layer induces a thicker damage layer under these conditions.

Thus we have identified the mechanism of damage formation on Si. To minimize T_d , it is necessary to lower the highenergy ion peak by controlling the power and the frequency of bias power. It is also necessary to adjust the steady state of the C–F polymer so that it is the same thickness as ion penetration depth. There are many kinds of ions $(H^+, CH_2F^+,$ $CHF₂⁺, CF₃⁺, and Ar⁺ were observed) and ion energy also has$

FIG. 9. Atomic composition of mixed layer (MD). CF_3 dosages were (a) 1.908×10^{-16} , (b) 2.544×10^{-16} , and (c) 3.392×10^{-16} cm⁻², respectively.

a certain dispersion, as shown in Fig. 6. The reaction is, therefore, more complex in the actual etched surface, and $T_{\text{C-F}}$ and T_d vary depending on the etching system and gas composition used in the actual manufacturing. Quantitative monitoring and control of plasma parameters and quantitative modeling of the surface reaction will be required to minimize the fluctuation in surface damage.

V. CONCLUSION

The mechanism of damage formation on a Si substrate during etching by fluorocarbon plasma was identified. Damage thickness T_d depends on the penetration depth of ions whose energy was decreased by a steady-state surface polymer. When the ion penetration depth was shallower than the thickness of the surface polymer, the damage was formed in the early stage of plasma exposure, when Si started to be exposed to plasma, but the polymer has not grown thicker yet. This damage was covered by the deposition of a thick C–F polymer and did not change as time exposed to plasma increased. There is an optimum point (balance point P_b) where ion penetration depth is nearly equal to the thickness of the steady-state polymer. Under this condition, unstable bonding was released from the mixing layer of the fluorocarbon species and Si, and the thickness of the damage layer decreased. To minimize the damage, we need to quantitatively control the radical composition (thickness of C–F polymer) and ion energy (high-energy IEDF peak) to find the P_b .

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