ПАРАМЕТРЫ ПЛАЗМЫ И КИНЕТИКА ТРАВЛЕНИЯ SiO₂ В СМЕСИ C₄F₈ + Ar + O₂

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Проведено исследование влияния соотношения Ar/O₂ на параметры плазмы, стационарные концентрации активных частиц и кинетику травления SiO₂ в трехкомпонентной смеси C₄F₈+Ar+O₂ в условиях, характерных для процессов реактивно-ионного травления (индукционный 13,56 МГц ВЧ разряд, общее давление газа 6 мТорр, вкладываемая мощность 700 Вт и мощность смещения 200 Вт). Алгоритм исследования сочетал измерения скоростей травления, диагностику плазмы зондами Лангмюра и мерную (глобальную) модель плазмы для получения данных по стационарным концентрациям и плотностям потоков активных частиц. Было найдено, что полное замещение аргона на кислород при постоянном содержании фторуглеродного компонента (фактически, переход от системы 50% C₄F₈ + 50% Ar к 50% C₄F₈ + 50% O₂): 1) характеризуется слабым немонотонным (с максимумом) изменением скорости травления SiO₂ с близкими абсолютными значениями в точках с нулевыми содержаниями O₂ и Ar; 2) вызывает монотонное снижение плотностей потоков атомов фтора и энергии ионов; 3) способствует снижению толщины фторуглеродной полимерной пленки на обрабатываемой поверхности за счет ее окислительной деструкции. Модельный анализ кинетики травления позволил заключить, что увеличение эффективной вероятности взаимодействия в гетерогенном процессе SiO₂ + F противоречит поведению плотности потока энергии ионов, но качественно согласуется с изменением параметров газовой фазы, отслеживающих толщину фторуглеродной полимерной пленки. Таким образом, рост содержания кислорода в плазмообразующей смеси влияет на эффективную вероятность взаимодействия за счет снижения толщины фторуглеродной полимерной пленки и облегчения доступа атомов F к обрабатываемой поверхности.

Ключевые слова: SiO₂, травление, полимеризация, поток атомов фтора, поток энергии ионов, эффективная вероятность взаимодействия, выход травления

PLASMA PARAMETERS AND SiO₂ ETCHING KINETICS IN C₄F₈ + Ar + O₂ GAS MIXTURE

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The effect of Ar/O$_2$ mixing ratio on plasma parameters, steady-state densities of active species and SiO$_2$ etching kinetics in the three-component C$_3$F$_8$+Ar+O$_2$ gas mixture was studied under typical conditions of reactive ion etching process (inductive 13.56 MHz RF discharge, total gas pressure of 6 mTorr, input power of 700 W and bias power of 200 W). The investigation combined etching rate measurements, plasma diagnostics by Langmuir probes and 0-dimensional (global) plasma modeling in order to determine steady-state densities and fluxes of plasma active species. It was found that the full substitution of Ar for O$_2$ at constant fraction of fluorocarbon gas (in fact, the transition from 50% C$_3$F$_8$ + 50% Ar to 50% C$_3$F$_8$ + 50% O$_2$ gas system): 1) results in weakly non-monotonic (with a maximum) SiO$_2$ etching rate with close values for both O$_2$-free and Ar-free plasmas; 2) causes the monotonic decrease in both F atom flux and ion energy flux; and 3) suppresses the formation of the fluorocarbon polymer film on the etched surface through its oxidative destruction pathway. The model-based analysis of SiO$_2$ etching kinetics allowed one to conclude that an increase in effective probability for SiO$_2$ + F reaction contradicts with the behavior ion energy flux as well as demonstrate the agreement with the change in gas-phase parameters characterizing the fluorocarbon film thickness. Therefore, an increase in O$_2$ content in a feed gas influences the effective reaction probability by decreasing fluorocarbon film thickness and providing better access of F atoms to the etched surface.

Key words: SiO$_2$, etching, polymerization, fluorine atom flux, ion energy flux, effective reaction probability, etching yield

INTRODUCTION

Silicon dioxide (SiO$_2$) is a well-known dielectric material which found many applications in integrated electronic device structures. The nearest examples are final passivation and protective layers, hard masks and gate dielectrics in field-effect structures [1-4]. As most of mentioned applications require the precise patterning of SiO$_2$ layers, there were many studies reported about the plasma-assisted etching characteristics and mechanisms for SiO$_2$ in fluorocarbon (C$_3$F$_8$) gas plasmas [5-10]. The results of existing works may briefly be summarized as follows:

- The spontaneous chemical between SiO$_2$ and F atoms is thermodynamically prohibited because the Si-O bond of ~ 799 kJ/mol is stronger than the Si-F one of ~ 552 kJ/mol [11]. As such, the SiO$_2$ dry etching process requires the ion bombardment in order to destruct Si-O bonds (in fact, to produce the chemisorption sites for F atoms) as well as to sputter the low volatile fluorinated layer [4-6].

- The ion bombardment energy more than 200 eV is generally enough to provide the reaction-rate-limited etching regime where the SiO$_2$ etching rate is controlled by F atom flux [12, 13]. Particularly, the non-monotonic (with a maximum at 30-40% O$_2$) etching rate for SiO$_2$ in the CF$_4$ + O$_2$ plasma is surely associated with the same non-monotonic behavior of F atom density [12, 13].

- The highest SiO$_2$ etching rates were obtained for low-polymerizing fluorocarbons, such as CF$_4$, while the maximum SiO$_2$/Si etching selectivity was found for high-polymerizing gas systems (C$_3$F$_6$, C$_2$F$_6$, CHF$_3$) [7, 9, 12]. The last phenomenon is connected with the much lower thickness of the fluorocarbon polymer film on the oxygen-containing surface [8-10].

Since the SiO$_2$/Si etching selectivity is the critical issue for many practical cases, the correct interpretation of various etching effects in high-polymerizing fluorocarbon gases with accounting for SiO$_2$ reactive-ion etching mechanism is an important for further process optimization. Particularly, it was found that the variation of gas mixing ratio in both C$_3$F$_8$+Ar and C$_3$F$_8$ + O$_2$ plasmas resulted in non-monotonic (with a maximum at ~ 50% additive gas) SiO$_2$ etching rates [14, 15]. Though the authors attributed these effects to the change in fluorocarbon polymer film thickness, they did not support their suggestions by the analysis of plasma chemistry and/or polymer film deposition kinetics. Moreover, the obtained non-monotonic etching rates formally contradict with monotonic change in measured steady-state polymer film thickness as well as with a maximum on the pol-
mer deposition rate [14]. As such, the SiO$_2$ etching mechanism in C$_2$F$_4$-based gas mixtures is not completely understood yet.

The main goal of this study was to investigate how the O$_2$/Ar mixing ratio in C$_2$F$_4$ + Ar + O$_2$ plasma influences the SiO$_2$ etching rate through the change in gas-phase plasma characteristics (electron temperature, energy of ion bombardment, and densities and fluxes of plasma-active species). In addition, we attempted the model-based analysis of the SiO$_2$ etching mechanisms in order to understand relationships between processing parameters and output process characteristics.

EXPERIMENTAL AND MODELING DETAILS

Experiments were performed in the planar inductively coupled plasma (ICP) reactor with a cylindrical chamber ($r = 15$ cm, $l = 12.8$ cm) made from anodized aluminum [10]. Plasma was excited using a 13.56 MHz radio-frequency (RF) power supply connected to a flat 5-turn copper coil on the top side of the chamber. Another RF power supply was connected to the chuck electrode in order to control the ion bombardment energy ($e_i$) through the negative dc bias voltage ($V_b$). The latter was measured by the high-voltage probe AMN-CTR (Youngsin Eng. Korea). The chuck electrode had a built-in water-flow cooling system that allowed one to maintain its temperatures at ~17 °C. Experiments were carried out at constant gas pressure (p = 6 mTorr), total gas flow rate (q = 40 sccm), input power (W = 900 W) and bias power ($V_b$ = 200 W) while the variable parameter was the Ar/O$_2$ mixing ratio. The latter was adjusted by the partial flow rates for corresponding gases at fixed C$_2$F$_4$ flow rate of 20 sccm. Accordingly, the fraction of the fluorocarbon component in a feed gas, $y$(C$_2$F$_4$), was always 50%, and the remaining half was composed of various amounts of $y$(Ar) and $y$(O$_2$). An increase in O$_2$ flow rate from 0-20 sccm corresponded to full substitution of Ar for O$_2$ as well as to the transition from 50% C$_4$F$_8$ + 50% Ar to 50% C$_4$F$_8$ + 50% O$_2$ gas system.

Plasma diagnostic by double Langmuir probe (DLP2000, Plasmart Inc.) provided the data on electron temperature ($T_e$) and ion current density ($J_i$). The latter were extracted from voltage-current curves using the well-known statements of the double Langmuir probe theory [16]. In order to minimize the influence of fluorocarbon polymer film on measured I-V curves, the probe tip was cleaned in 50% Ar + 50% O$_2$ plasma for 1 min before and after each measurement. As a result, the difference between data points recorded under the same experimental conditions within a time period of ~ 5 min did not exceed the standard experimental error.

Etched samples were the fragments of Si wafer with the 500 nm-thick SiO$_2$ layer produced by PECVD method. The sample surface had the size of about 2×2 cm and was partially covered by the photoresist mask (AZ1512, ~1.5 μm) in order to provide the measurement of the etched depth, Δh. For this purpose, we used the surface profiler Alpha-Step 500 (Tencor). The SiO$_2$ etching rates were determined as $R = \Delta h/\tau$, where $\tau = 2$ min is the processing time. In preliminary experiments, it was found that 1) an increase in $\tau$ up to 10 min did not disturb the nearly linear behavior for $\Delta h = f(\tau)$; 2) an increase in amount in simultaneously etched samples up to 10 pieces did not influence the SiO$_2$ etching rate; and 3) the presence of etched sample in the reactor chamber had no effect on results of plasma diagnostics by Langmuir probes. All these suggest the steady-state kinetic regime of the ion-assisted chemical reaction as well as the negligible influence of reaction products on gas-phase plasma parameters.

In order to analyze the influence of Ar/O$_2$ mixing ratio on kinetics and densities of plasma active species, we developed a simplified 0-dimensional (global) kinetic model operated with volume-averaged plasma parameters. Similar to our previous works [17, 18], the model was based on the simultaneous solution of steady-state kinetic equations with experimental data on $T_e$ and $J_i$ as the input parameters. The basic assumptions were that:

- The electron energy distribution function (EEDF) in highly-ionized plasmas ($n_e/N \sim 10^4$, where $n_e$ is the total positive ion density, and N is the gas density) is strongly influenced by electron-electron collisions. This allows one to use the Maxwellian approximation for EEDF when calculating the electron-impact rate coefficients and related plasma parameters [12, 19].

- The measured ion current density is linked with the total density of positive ions as $J_i \approx 0.61 n_e(eT_i/m_i)^{1/2}$ [12], where $m_i$ is the effective ion mass. The latter may be roughly evaluated through known fractions and ionization rate coefficients for dominant neutral species [17].

- The electronegativity of low-pressure C$_4$F$_8$ + Ar + O$_2$ plasma is low enough to assume $n_e \approx n_i$, where $n_i$ is the electron density [19, 20].

The set of chemical reactions was directly taken from Ref. [17]. Corresponding kinetic scheme has demonstrated an acceptable agreement between model-predicted and measured species densities in C$_4$F$_8$-containing plasmas [17, 19-21].

RESULTS AND DISCUSSION

From Fig. 1, it can be seen that an increase in O$_2$ fraction from 0-50% (i.e. the full substitution of Ar for O$_2$) results in weakly non-monotonic SiO$_2$ etching...
rate that shows a maximum of ~ 125 nm/min at 25% Ar + 25% O₂. This value is about 1.4 times higher than those obtained in CF₄+Ar plasma (~ 97 nm/min) and CF₂+O₂ plasma (~ 92 nm/min). Therefore, the principal questions are 1) why the change in additive gas has the only weak overall effect on SiO₂ etching rate; and 2) what mechanism produces the maximum in SiO₂ etching rate in the three-component gas system.

Fig. 1. Parameters characterizing the SiO₂ etching kinetics in C₄F₄+Ar+O₂ plasma. In Fig. a: 1 – SiO₂ etching rate; 2 – effective reaction probability (γₛ); 3 – effective etching yield (Yₑ). In Fig. b): effective reaction probability (γₛ) as a function of Γpol/ΓF O flux ratio characterizing the fluorocarbon polymer film thickness.

According to Refs. [22, 23], the experimentally obtained rate of the reactive-ion etching process, R, may be represented as a sum of two components, Rphys + Rhem. Here, Rphys is the rate of physical sputtering of SiO₂ etching surface while Rhem is the rate of the ion-assisted chemical reaction between SiO₂ and F atoms. In preliminary experiments, it was found that the SiO₂ etching rate in pure Ar plasma (in fact, Rphys) under the given set of processing condition does not exceed 5 nm/min. As such, one can surely suggest that Rphys << Rhem and R = Rhem = γR ΓF, where ΓF is the flux of fluorine atoms, and γR is the effective reaction probability. From the last expression, one can assume at least two mechanism which may result in the non-monotonic R = f(y(O₂)) curve as well as in the nearly-constant SiO₂ etching rate under the conditions of y(Ar) = 50% and y(O₂) = 50%. These are: 1) the non-monotonic change in ΓF due to the change in formation and/or decay kinetics for F atoms; and 2) the monotonic, but opposite changes for ΓF and γR with increasing y(O₂). Obviously, in given gas system, the parameter γR may be sensitive to many factors influencing the formation of free surface sites (SiO bond breaking) and their accessibility for the adsorption of F atoms (desorption of reaction products, fluorocarbon film thickness). Therefore, the correct interpretation of SiO₂ etching mechanism requires the data on the plasma parameters as well as on the densities and fluxes of F atoms, polymerizing radicals, and positive ions. For this purpose, we performed plasma diagnostics by Langmuir probes and plasma modeling.

From Table 1, it can be seen that the substitution of Ar for O₂ results in decreasing both Tₚ (4.8-3.1 eV for 0-50% O₂) and Jₑ (1.21-0.91 mA/cm² for 0-50% O₂) values. The first effect results from an increase in the electron energy loss due to the low-threshold excitations (vibrational, electronic) for both O₂ and molecular reaction products. The mentioned decreased in Jₑ follows the behavior of nₑ (4.2·10⁷-3.7·10⁸ cm⁻³ for 0-50% O₂). This is because the transition to O₂-rich plasmas 1) suppresses the ionization through the decreasing ionization rate coefficients for all neutral species due to deceasing Tₑ; and 2) accelerates the decay of positive ions in the ion-ion recombination process due to increasing densities of electronegative species (O₂ itself and oxygen-containing reaction products). Accordingly, the similar decreasing tendency was found for ion flux Γ₂ (7.6·10¹⁵-5.7·10¹⁵ cm⁻²·s⁻¹ for 0-50% O₂, see Fig. 2).

Fig. 2 illustrates the influence of O₂ content in the C₄F₄+Ar+O₂ gas mixture on densities and fluxes of neutral species. It can be seen that the main gas-phase components in the 50% C₄F₄ + 50% Ar gas system are CF₂ (x = 1-3) radicals, C₄F₄, and C₄F₂ [17, 19, 20]. Among these, CF₂ and C₂F₄ represent the first-step dissociation products of original C₄F₄ molecules (R1: C₄F₄ → C₂F₄ + CF₂ + e and R2: C₂F₄ + e → 2CF₂ + e) while the condition [CF₂] >> [C₂F₄] is provided by R3: C₄F₄ + e → 2CF₂ + e (k₃ = 1.3·10¹⁸ cm³/s), R4: C₂F₄ + e → C₂F₃ + F + e (k₄ = 3.0·10¹⁰ cm³/s), and R5: C₂F₃ + F → CF₂ + CF₃ (k₅ = 4.0·10⁻¹¹ cm³/s). The
high density of CF radicals is provided by their fast formation in R6: \( \text{CF}_2 + e \rightarrow \text{CF} + \text{F} + e \) and R7: \( \text{C}_2\text{F}_3^+ + e \rightarrow \text{CF}_2 + \text{CF} + e. \) Accordingly, the high density of CF is supported by R5 and a group of heterogeneous processes R8: \( \text{CF}_x + F \rightarrow \text{CF}_{x+1} \) with \( x = 2. \) The main source of F atoms is represented by R9: \( \text{CF}_x + e \rightarrow \rightarrow \text{CF}_{x+1} + \text{F} + e \) with \( x = 1-3 \) while the decay of these species is mainly due to R8 and R5. Since the last process provides an effective conversion \( F \rightarrow \text{CF}_x \) with \( x = 2 \) and 3, the condition \( [F] \ll [\text{CF}_x] \) always takes place.

The substitution of Ar for O\(_2\) at constant \( y(\text{C}_4\text{F}_8) \) noticeably reduces the rates of R9 (due to the simultaneous decrease in \( T_e \) and \( n_e \)) as well as introduces additional pathways for the decomposition of CF radicals through R10: \( \text{CF}_x + O \rightarrow \text{CF}_{x+1}O + F \), R11: \( \text{CF}_x + O(\text{D}) \rightarrow \text{CF}_{x+1}O + F \) and R12: \( \text{CF}_x + O + e \rightarrow \rightarrow \text{CF}_{x+1}O + F + e \). This provides the monotonic decrease in \([\text{CF}_x]\) \((4.7 \times 10^{12}-1.2 \times 10^{13} \text{ cm}^{-3}) \) for \( x = 2 \) and \( 2.9 \times 10^{12}-1.3 \times 10^{12} \text{ cm}^{-3} \) for \( x = 1 \) at 0.50% \( O_2 \), see Fig. 2(a) toward \( O_2\)-rich plasmas. At the same time, the effective loss of \( O_2 \) molecules through R13: \( \text{CF} + O_2 \rightarrow \text{CF}O + O \) \((k_{13} \sim 3.2 \times 10^{-11} \text{ cm}^3/s) \) and R14: \( C + O_2 \rightarrow \text{CO} + O \) \((k_{14} \sim 2.0 \times 10^{-11} \text{ cm}^3/s) \) limits formation rates for O and \( O(\text{D}) \) species in R16: \( O_2 + e \rightarrow 2O + e \), R17: \( O_2 + e \rightarrow O + O(\text{D}) + e \) and R18: \( O + e \rightarrow O(\text{D}) + e. \) The lack of oxygen atoms reduces the effect of R10–R12 on the F atom formation kinetics (in fact, keeps the condition \( R_9 > R_{10} + R_{11} + R_{12} \) for 0-50% \( O_2 \)) and thus, leads to monotonically decreasing F atom density \( ([F] = 8.0 \times 10^{12}-8.6 \times 10^{11} \text{ cm}^3) \) for 0-50% \( O_2 \) and flux \( (\Gamma_F = 1.6 \times 10^{17}-1.8 \times 10^{16} \text{ cm}^2/s) \) for 0-50% \( O_2 \).

### Table 1

**Electro-physical plasma parameters in \( \text{C}_4\text{F}_8+\text{Ar}+\text{O}_2 \) gas mixture**

<table>
<thead>
<tr>
<th>( y(\text{O}_2) )</th>
<th>( T_e )</th>
<th>( J_e )</th>
<th>( n_e \times 10^{10} )</th>
<th>-U&lt;sub&gt;dc&lt;/sub&gt;</th>
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<td>3.12</td>
<td>0.91</td>
<td>3.66</td>
<td>177</td>
</tr>
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From the comparison of Figs. 1(a) and 1(b), one can conclude that the non-monotonic \( \text{SiO}_2 \) etching rate contradicts with the behavior of \( \Gamma_F. \) Such situation corresponds to monotonically increasing effective reaction probability in the range of 0.02-0.2 for 0-50% \( O_2 \) (Fig. 1(a)). In non- or low-polymerizing
plasma, the non-constant $\gamma_R$ at constant surface temperature is normally associated with the change in the rate of ion-stimulated desorption for reaction products and thus, with the ion bombardment intensity. According to previously published works [17, 18], the intensity of ion bombardment may be traced by the parameter $(M_i e_i)^{1/2} \Gamma_+$, where $M_i$ is the effective ion molar mass, $e_i = e(U_f - U_{ek})$ is the ion bombardment energy, and $U_f$ is the floating potential. It was found that, under the given set of processing condition, a weak increase in both $U_{ek}$ and $e_i$ toward O$_2$-rich plasmas (Tabs. 1 and 2) did not compensate the deeper fall of $\Gamma_+$. As a result, the parameter $(M_i e_i)^{1/2} \Gamma_+$ decreases monotonically in the range of $7.3 \cdot 10^{17}$–$5.4 \cdot 10^{17}$ eV$^{1/2}$cm$^{-2}$s$^{-1}$ for 0-50% O$_2$. These data clearly show that the almost tenfold increase in $\gamma_R$ cannot be explained by the simple ion-stimulated desorption mechanism.

Another reasonable assumption is that $\gamma_R$ may be influenced by the fluorocarbon polymer deposition/decomposition kinetics through the fluorocarbon film thickness, $h_{pol}$. Since the film retards the access of F atoms to the etched surface, the decreasing tendency for $\gamma_R = f(h_{pol})$ is quite expected. According to Refs. [5-7, 9], basic approaches for the analysis of the fluorocarbon film deposition/decomposition kinetics are as follows:

- The formation of the fluorocarbon polymer film is provided by CF$_x$ radicals with $x = 1, 2$ as well as appears to be faster in fluorine-poor plasmas. The last effect is because the polymer surface contains more “open” bonds and thus, easier joins CF$_x$ species from a gas phase. As such, the polymer deposition rate may be traced by the $\Gamma_{pol}/\Gamma_F$ ratio, where $\Gamma_{pol}$ is the total flux of polymerizing radicals.

- The decomposition of the fluorocarbon polymer film is provided by both physical (fragmentation by ion bombardment) and chemical (interaction with oxygen atoms) pathways. Therefore, the change in fluorocarbon polymer film thickness due to these factors may be traced by the parameters $\Gamma_{pol}/(M_i e_i)^{1/2} \Gamma_+ \Gamma_F$ and $\Gamma_{pol}/\Gamma_0 \Gamma_F$, respectively.

It was found that, as the O$_2$ content in a feed gas increases, the parameter $\Gamma_{pol}/(M_i e_i)^{1/2} \Gamma_+ \Gamma_F$ keeps the nearly constant value in the range of $9.3 \cdot 10^{18}$–$1.4 \cdot 10^{17}$ eV$^{1/2}$cm’s for 0-50% O$_2$ (Table 2). In fact, this means that there are no principal changes in physical etching pathway of the fluorocarbon polymer film. Oppositely, the parameter $\Gamma_{pol}/\Gamma_0 \Gamma_F$ exhibit the sufficient decrease toward O$_2$-rich plasmas ($3.0 \cdot 10^{14}$–$2.4 \cdot 10^{15}$ cm’s for 5-50% O$_2$, see Table 2) that probably corresponds to a nearly proportional decrease in $h_{pol}$.

As such, the mentioned change in $\gamma_R$ has an opposite trend but is very close in relative scale to that expected for $h_{pol}$. In our opinion, this finding may be accepted as the indirect proof that and increase in $\gamma_R$ is really connected with decreasing $h_{pol}$ and increasing F atom flux on the polymer film/etched surface interface. In addition, Fig. 1(b) represents the correlation between $\gamma_R$ and $\Gamma_{pol}/\Gamma_0 \Gamma_F$ ratio. Both shape of the curve and the tendency itself are very close to those obtained for reaction probabilities and etching yields as functions of $h_{pol}$ in several experimental works related to the etching of Si and SiO$_2$ in fluorocarbon gas plasmas [5, 6, 24]. Therefore, one can surely assume that, under the given set of processing conditions 1) a decrease in $h_{pol}$ toward O$_2$-rich plasmas provides an increase in $\gamma_R$; and 2) an increase in $\gamma_R$ produces the non-monotonic SiO$_2$ etching rate with monotonically decreasing F atom flux.

<table>
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<tr>
<th>$\gamma$(O$_2$)</th>
<th>$\varepsilon_i$</th>
<th>$\Gamma_{pol}/\Gamma_F$</th>
<th>$\Gamma_{pol}/(M_i e_i)^{1/2} \Gamma_+ \Gamma_F$</th>
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<td>9.12</td>
<td>$1.40 \cdot 10^{-17}$</td>
<td>$2.4 \cdot 10^{-15}$</td>
</tr>
</tbody>
</table>

Another approach which is sometimes used to analyze the reactive ion etching kinetics is to represents the etching process as the chemically-enhanced sputtering. In this case, the etching rate is $Y_R \Gamma_+$, where $Y_R$ is the ion-type-averaged etching yield [12]. In reactive gas plasmas, the latter depends on both ion bombardment energy and heterogeneous reaction kinetics through the nature and composition of sputtered material [23]. The data of Fig. 1(a) show that the variation of Ar/O$_2$ mixing ratio causes the non-monotonic $Y_R$ with the overall increasing tendency toward O$_2$-rich plasmas. The fact that the behavior of $Y_R$ repeats one for etching rate is connected with the lower sputtering threshold for reaction products compared with SiO$_2$ itself. As such, the higher etching rate formally corresponds to higher amount of easier sputtered material on the etched surface. Accordingly, an increase in $Y_R$ toward O$_2$-rich plasmas correlates with the behavior of ion bombardment energy.
CONCLUSIONS

In this work, we investigated the effect of Ar/O₂ mixing ratio on plasma parameters, steady-state densities of active species and SiO₂ etching kinetics in the three-component C₂F₆ + Ar + O₂ gas mixture. It was found that the substitution of Ar for O₂ at constant fraction of fluorocarbon gas 1) causes the monotonic decrease in both F atom flux and ion energy flux; and 2) results in non-monotonic (with a maximum) SiO₂ etching rate with close values for both O₂-free and Ar-free plasmas that corresponds to monotonically increasing effective reaction probability. From the model-based analysis of SiO₂ etching kinetics, it was concluded that an increase in O₂ content in a feed gas influences the effective reaction probability through decreasing thickness of fluorocarbon polymer film (due to the oxidative destruction) and providing better access of F atoms to the etched surface.

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