Contamination Particle Behavior of Aerosol Deposited Y$_2$O$_3$ and YF$_3$ Coatings under NF$_3$ Plasma

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Abstract: The internal coatings of chambers exposed to plasma over a long period of time are subject to chemical and physical damage. Contamination particles that are produced by plasma damage to coatings are a major contribution to poor process reliability. In this study, we investigated the behavior of contamination particles produced from plasma damage to Y$_2$O$_3$ and YF$_3$ protective coatings, which were applied by an aerosol deposition method. The coating materials were located at the powered electrode, the grounded electrode, and the grounded wall, which were exposed to a NF$_3$ plasma. The mass loss at the powered electrode, which was exposed to the NF$_3$ plasma etching under an applied bias, showed that the YF$_3$ etching rate was higher than that of Y$_2$O$_3$. Conversely, the mass of coating increased at the grounded electrode and the grounded wall, which were exposed to a NF$_3$ plasma. The mass loss at the powered electrode, which was exposed to the NF$_3$ plasma etching under an applied bias, showed that the YF$_3$ etching rate was higher than that of Y$_2$O$_3$. Conversely, the mass of coating increased at the grounded electrode and the grounded wall, which were exposed to a NF$_3$ plasma etching under zero bias. The mass of the Y$_2$O$_3$ coating increased more than that of the YF$_3$ coating. X-ray photoelectron spectroscopy analysis showed that the Y$_2$O$_3$ coating corroded to YO$_x$F$_y$ in the NF$_3$ plasma, and YF$_3$ existed as YF$_x$. Light scattering sensor analysis showed that the YF$_3$ coating produced fewer contamination particles than did the Y$_2$O$_3$ coating.

Keywords: yttrium oxide (Y$_2$O$_3$); yttrium fluoride (YF$_3$); aerosol deposition (AD); contamination particle; plasma etching; NF$_3$ plasma

1. Introduction

Efforts to minimize the influence of contamination particles, which are a major cause of decreased production yields, have been continuing as the degree of integration of semiconductor circuits has improved. There are various causes of contamination particles, including mechanical, chemical, and environmental factors. In particular, plasma used in processes for manufacturing semiconductors reacts with parts inside the chamber (e.g., electrodes, showerheads, liners, focusing rings, wall shield rings) to generate contamination particles.

Corroded particles grow on the surface of parts of the inner chamber, which are exposed to plasma ion bombardment and radical reactions over a long period of time. The weakly bonded corrosion particles then fall from the surface as the particles grow in size. As the process repeats, the plasma etches the surface and contamination particles fall off, becoming a major cause of process failure [1]. Corrosion and the extent of etching of parts of the surface are determined by the ion density and ion flux of the exposed plasma. A powered electrode which is exposed to a plasma etching under an applied bias is easily corroded and etched owing to collisions with a large energy flux. The grounded electrode and a grounded wall, which are exposed to a plasma etching under zero bias, sustain limited corrosion and etching by collisions with the ions, which have a low energy flux [2,3]. Ceramic coating films are used to prevent plasma corrosion at these surfaces and minimize the plasma corrosion [4–7].
Among ceramic coating materials for preventing plasma corrosion, Y₂O₃ is widely used for inner parts of the chamber because of its low etching rate and low chemical reactivity. Recently, YOF and YF₃ have also drawn attention as ceramic coating materials for parts owing to their ability to suppress chemical reactions with fluoridated gases such as CF₄ and NF₃ [8–13]. Nevertheless, there have been few reports about the behavior of contamination particles by plasma etching [14]. Atmospheric plasma spraying (APS) methods are commonly used to coat parts with ceramic coating materials such as Y₂O₃ and YF₃. The APS method melts a ceramic powder, with particles sizes in the range of several tens of micrometers, in a high-temperature plasma (10,000 K or more), which attaches to the substrate. The APS method enables the coating of thick films in a short period of time, but provides a rough surface and layered structure, such that the adhesion with the surface is poor and the coating peels off easily. Furthermore, such coatings have a low density owing to numerous internal pores [15]. The aerosol deposition (AD) method has been studied as a coating technology to overcome the limitations of APS. The AD method involves spraying a sub-micrometer-sized powder with high-pressure air through a nozzle in a vacuum chamber (>10⁻³ Torr). The AD method coats a high-density film with excellent uniformity of thickness compared with that of the APS method [16–19]. The plasma etching properties of ceramic coatings coated by the APS method have been reported in previous studies [8]; however, there is a lack of studies on the properties of ceramic coatings prepared by AD methods, including plasma etching properties and the generation of contamination particles. The etching properties of CF₄ gas, which is used in the etching process, have been widely studied. However, etching with CF₄ gas is often accompanied by the formation of unwanted fluorocarbon polymer layers. Recently, completely dissociated NF₃ has been used to minimize these unwanted reactions and to achieve high etch rates [20–23].

In this study, we discuss the etching properties and contamination particles generation of Y₂O₃ and YF₃ coatings fabricated by the AD method utilizing the NF₃ plasma process. We investigated locations including the powered electrode, grounded electrode, and grounded wall to observe the etching characteristics of the coating material and the dependence on the bias voltage.

2. Materials and Methods

Figure 1 shows a schematic diagram of the capacitively coupled plasma system. The dimensions of the reactor were an inner diameter of 30 cm and a height of 30 cm. The powered electrode and grounded electrode had a radius of 7.6 cm. The discharge gap length from the powered electrode to the grounded electrode was 10 cm. Specimens were placed in the powered electrode, grounded electrode, and grounded wall, as shown in Figure 1.

![Figure 1](image_url)
The NF$_3$ gas was used to generate a plasma supplied through a showerhead with a mass flow controller. A dry pump and turbo pump were used in the vacuum system, and the working pressure of the experiment was 40 Pa. The power was set to be 13.56 MHz (Sizer Generator, Advanced Energy, Fort Collins, CO, USA), and an impedance matching network (Navigator, Advanced Energy, Fort Collins, CO, USA) was used to deliver the maximum power. The RF power applied to the plasma was 400 W. During the plasma etching, the temperatures at the powered electrode, grounded electrode, and grounded wall were about 120, 60, and 50 °C, respectively. The temperature was measured using an IR thermometer (IMPAC Pyrometer IN5, Lumasense Technologies, Santa Clara, CA, USA) and the temperature inside the chamber was measured through a viewport with zinc selenide.

The disc-like substrates were made of Al alloy 6061 and had a diameter of 76 mm and thickness of 1 mm. The substrates were then coated with Y$_2$O$_3$ and YF$_3$ by AD, with the use of an aerosol deposition system (Dandan Corp., Daejeon, Korea), where Y$_2$O$_3$ and YF$_3$ were in a powder form (99.99%, D$_{50}$ = 0.5–5 µm). The sprayed coatings of Y$_2$O$_3$ and YF$_3$ were 8 and 19 µm thick, respectively. The AD coating was performed as follows: the air flow rates were 40–80 SLM and the vacuum pressure was 10 Pa.

Before and after the NF$_3$ plasma etching, the surface morphology and composition of the Y$_2$O$_3$ and YF$_3$ coatings were ex-situ analyzed with a field-emission scanning electron microscope (FE-SEM, S-4800, Hitachi, Tokyo, Japan) and X-ray photoelectron spectroscopy (XPS; Monochromatic Al-Kα, AXIS-NOVA, Manchester, UK), respectively. The mass of the specimen was measured before and after the plasma etching test with a XP205 analytical balance (Mettler Toledo, Greifensee, Switzerland). After the plasma etching was performed for 60 min, the specimen was removed, and the mass loss was measured. The contamination particles produced from the Y$_2$O$_3$ and YF$_3$ coatings were measured in real time according to the NF$_3$ plasma exposure time. An ISPM light-scattering sensor (Stiletto, In Situ Particle Monitor, Inficon, Heidiland, Switzerland) was attached to the exhaust line to measure the amount and size of the contamination particles. This system was capable of measuring contamination particles in real time as they passed through the exhaust pipe. The ISPM was placed in the exhaust line so that it detected particles moving in one direction. The ISPM does not operate properly when exposed to the light of the plasma, because of excess noise. The minimum measurable particle size was ~0.2 µm. The sensor was based on the principles of laser light scattering, and more details of its working principles can be found in previous reports [24,25].

3. Results and Discussion

Figure 2 shows an FE-SEM image of Y$_2$O$_3$ and YF$_3$ powders used as a starting material for coating. The Y$_2$O$_3$ powder (Figure 2a,b) and YF$_3$ powder (Figure 2c,d) were composed of flat particles approximately 1–5 µm in size, and spherical particles approximately 0.5–1 µm in size, respectively. Furthermore, the YF$_3$ powder showed greater aggregation than that of the Y$_2$O$_3$ powder. We expected that the YF$_3$ powder was sprayed as a mixed powder, containing dispersed particles in the size range of 0.5–1 µm together with aggregated particles larger than 1 µm, because aggregated particles are not easily dispersed. When we applied the AD method, the Y$_2$O$_3$ and YF$_3$ particles were sprayed by pressurized gas. The accelerated particles had a kinetic energy, which dissipated on collision with the substrate [17]. In this case, plastic deformation or fracturing of particles might occur owing to the high kinetic energy of large sprayed particles [18]. Particles tend to fracture when they aggregate or have a high kinetic energy owing to their large size, as shown in Figure 3.
we confirmed that the coating surface changed depending on the position in the chamber. The Y particles were coated mainly by plastic deformation; however, fragmentation of larger particles into particles with high kinetic energy, which had a hammering effect. After the NF3 plasma etching, the initial YF3 powder was sprayed as a mixed powder, containing dispersed particles in the size range of 0.5–1 µm together with aggregated particles larger than 1 µm, because aggregated particles are not measurable particle size was ~0.2 µm. The sensor was based on the principles of laser light scattering, measuring contamination particles in real time as they passed through the exhaust pipe. The ISPM is connected to the exhaust line to measure the amount and size of the contamination particles. This system was capable of measuring contamination particles in real time according to the NF3 plasma exposure time. An ISPM light-scattering sensor was placed in the exhaust line so that it detected particles moving in one direction. The ISPM does not measure particles moving in opposite directions. A fast scanning ISPM has the sensitivity to detect individual contamination particles in real time. Values of the contamination mass per unit length were measured using an ISPM light-scattering sensor.

Figure 2. FE-SEM images of Y2O3 and YF3 powders before coating by aerosol deposition: (a,b) Y2O3 powder and (c,d) YF3 powder.

Figure 3. FE-SEM images of the surface of Y2O3 and YF3 coatings before and after exposure to NF3 plasma: (a) Y2O3 and (e) YF3 before etching; (b) Y2O3 and (f) YF3 after etching of the powered electrode; (c) Y2O3 and (g) YF3 after etching of the grounded electrode; (d) Y2O3 and (h) YF3 after etching of the grounded wall.

The Y2O3 powder consisted of particles approximately 1–5 µm in size (Figure 2), and the Y2O3 particles were coated mainly by plastic deformation; however, fragmentation of larger particles into smaller particles was caused by their high kinetic energies at the moment of collision with the substrate. In this case, a film failed to form or only a residue formed (Figure 3a). The average particle size was 0.5–1 µm, but most aggregated YF3 particles were separated when they collided with the substrate, and formed a porous film. For this reason, before the NF3 plasma etching, the initial YF3 coating film (Figure 3e) and the Y2O3 coating film (Figure 3a) showed different surface properties. However, the final coating was denser than the initial coating film, because the initial coating was subjected to particles with high kinetic energy, which had a hammering effect. After the NF3 plasma etching, we confirmed that the coating surface changed depending on the position in the chamber. The Y2O3 (Figure 3b) and YF3 (Figure 3f) coatings on the powered electrode were rapidly corroded and etched. The NF3 plasma etching under an applied bias removed the initial surface and exposed a uniform surface, which was subjected to the hammering effect. Conversely, the Y2O3 (Figure 3c,d) and YF3 (Figure 3g,h) coatings at the grounded electrode and the grounded wall appeared to mix with the surface and took on a rough morphology similar to that of the powered electrode surface.
We attribute these surface properties to two factors. First, the surface was hardly corroded or etched by the NF₃ plasma etching under zero bias. Rather, some by-products (contamination particles) were deposited, such as YOₓFₚ or YF₃, which were generated from etching of the powered electrode by NF₃. As evidence for this change, we considered the mass change after exposure to the NF₃ plasma (Figure 4). After exposure to NF₃, the masses of the Y₂O₃ and YF₃ coatings located on the powered electrode decreased (Figure 4a). The masses of the Y₂O₃ and YF₃ coatings located on the grounded electrode and the grounded wall increased (Figure 4b,c).

Next, the surfaces of the coating located on the grounded electrode and the grounded wall were maintained in the same state as before exposure to the NF₃ plasma. The plasma density was highest at the center of the chamber and lower at the edges [26]. The grounded electrode and grounded wall were exposed to NF₃ plasma etching under zero bias. Thus, the etching reaction occurred very slowly, and even after exposure to NF₃ plasma, the Y₂O₃ coating on the grounded electrode and grounded wall contained very small particles generated by the fracturing of larger particles (Figure 3c,d), and the YF₃ coating had a porous structure composed of fractured and aggregated particles (Figure 3g,h).

Finally, the coating of the grounded electrode showed a greater mass change than that of the grounded wall coating. If sputtering of by-products occurs, the grounded electrode will be more affected than the grounded wall because it is located directly below the powered electrode. Figure 4b,c shows that both the Y₂O₃ and YF₃ coatings had a considerably greater mass increase for the grounded electrode compared with that for the grounded wall coating. In addition, the initial surface of the grounded wall (Figure 3d,h) showed better resistance than that of the grounded electrode (Figure 3c,g) for both the Y₂O₃ and YF₃ coatings.

Compared with Y₂O₃, the YF₃ coatings were more rapidly etched and showed greater mass loss under the NF₃ plasma (Figure 4a). The standard boiling temperatures defined by IUPAC of Y₂O₃ and YF₃ are 4570 and 2500 K, respectively, and the sublimation enthalpy of Y₂O₃ is also greater than that of YF₃. Thus, Y₂O₃ forms stronger bonds than does YF₃ [8]. Therefore, for an etching reaction occurring under the same conditions, YF₃ is etched more rapidly than is Y₂O₃. We estimated the etching rates of the Y₂O₃ and YF₃ coatings on the powered electrode from mass loss to be 48 and 186 nm/min/m², respectively.

Notably, although the mass of the YF₃ coating located on the powered electrode decreased considerably compared with that of the Y₂O₃ coating, the mass increases of the YF₃ coatings located on the grounded electrode and the grounded electrode were less than those of the Y₂O₃ coating. This result is difficult to explain if simply considering the sputtering of by-products. We posit that fluorine deposition occurs in addition to the sputtering of by-products. Schaepkens et al. reported that fluorine gas takes precedence over etching where a self-bias voltage is present, and deposition occurred where the self-bias voltage was absent [27]. On this basis, we speculate that fluorine deposition occurred on the grounded electrode and grounded wall where the self-bias voltage was absent. In particular, the Y₂O₃ coating was more greatly affected by fluorine deposition than the YF₃ coating because of the
many oxygen atoms and vacancies that can react with fluorine. As a result, by-product sputtering and fluorine deposition occurred at the same time in the coating of the grounded wall and grounded electrode during exposure to NF$_3$ plasma. The Y$_2$O$_3$ coating mass greatly increased because the Y$_2$O$_3$ coating was affected more by the fluorine deposition reaction than was the YF$_3$ coating. On the other hand, the YF$_3$ coating was affected more by the sputtering of by-product reaction than the fluorine deposition reaction.

We had previously confirmed that the Y$_2$O$_3$ coating reacts with NF$_3$ plasma to form YO$_x$F$_y$, which is etched by ion attack, and the YO$_x$F$_y$ coating is etched in the form of YO$_x$F$_y$ without a principal chemical reaction [8]. On the basis of previous studies, the YF$_3$ coating is also expected to be etched in the form of YF$_x$ without notable chemical corrosion reactions (Figure 5). We measured the amounts of Y, O, and F in the Y$_2$O$_3$ and YF$_3$ coatings by XPS analysis after exposure to NF$_3$ plasma to confirm the etching mechanism (Table 1). The YF$_3$ coating showed no notable component changes before and after exposure to NF$_3$ plasma. However, the Y$_2$O$_3$ coating had a lower O 1s content and a greater F 1s content.

![Figure 5. Schematic diagram of yttrium oxyfluoride film deposition and etching behavior of (a) Y$_2$O$_3$ and (b) YF$_3$ coatings.](image)

**Figure 5.** Schematic diagram of yttrium oxyfluoride film deposition and etching behavior of (a) Y$_2$O$_3$ and (b) YF$_3$ coatings.

**Table 1.** X-ray photoelectron spectroscopy (XPS) analysis results of Y$_2$O$_3$ and YF$_3$ coatings before and after exposure to NF$_3$ plasma: (a) Y$_2$O$_3$ and (e) YF$_3$ before etching; (b) Y$_2$O$_3$ and (f) YF$_3$ after etching of the powered electrode; (c) Y$_2$O$_3$ and (g) YF$_3$ after etching of the grounded electrode; (d) Y$_2$O$_3$ and (h) YF$_3$ after etching of the grounded wall.

| Compound Content (at.%) | Y$_2$O$_3$ Coating | YF$_3$ Coating |
|------------------------|---------------------|----------------|
|                        | (a) | (b) | (c) | (d) | (e) | (f) | (g) | (h) |
| Y 3d                   | 31.8 | 29.2 | 27.3 | 25.4 | 33.7 | 30.1 | 28.7 | 29.0 |
| O 1s                   | 65.1 | 13.0 | 11.6 | 14.2 | 5.7  | 3.1  | 2.9  | 3.0  |
| F 1s                   | 3.1  | 57.8 | 61.1 | 60.4 | 60.6 | 66.8 | 68.4 | 68.0 |

Figure 6 shows the XPS spectra of yttrium in Y$_2$O$_3$ and YF$_3$ before and after the NF$_3$ plasma treatment for more precise analysis. We designated the XPS peaks for the cations of Y 3d$_{5/2}$ and Y 3d$_{3/2}$ as blue, green, and orange lines, respectively. The binding energy difference of the two peaks was 2.05 eV, which represents an intensity ratio of 3:2. The Y 3d$_{5/2}$ and Y 3d$_{3/2}$ spectra were fitted by a ratio of 10:90 (Gaussian/Lorentzian). In the case of Y$_2$O$_3$ before the reaction with NF$_3$ plasma, the Y 3d$_{5/2}$ peak positions (green line) were 157.8 and 156.2 eV, and the Y 3d$_{3/2}$ peak positions (orange line) were 160.1 and 158.3 eV. However, after the reaction with the NF$_3$ plasma, the binding energy of Y 3d$_{5/2}$ (blue line) and Y 3d$_{3/2}$ shifted to 159 and 161.5 eV, respectively; hence, Y–F bonds formed.
were reasonable. When the Y–O bond was substituted with the X–F bond [7]. After the reaction with the NF3 plasma. Because the F ion has a very high electron affinity, the binding length with the cation X was shortened, the electron density was decreased, and the binding energy increased when the X–O bond was substituted with the X–F bond [7]. After the reaction with the NF3 plasma, the Y–O bonds were substituted with Y–F bonds, and the binding energy increased owing to the difference in electronegativity between the two bonds. This result confirms that the Y2O3 coating of the powered electrode featured coexisting Y–O and Y–F bonds because the surface of Y2O3 coating was etched. At the grounded electrode and the grounded wall, the Y–O bonds on the surface were completely substituted by Y–F owing to the high reactivity of fluorine radicals; hence, the surface had an apparent binding energy peak similar to that of YF3. Conversely, as shown in Figure 6e–h, the YF3 coating exhibited less change in its composition before and after NF3 plasma treatment. As a result, we confirmed that the previously expected mechanisms for the reaction of the NF3 plasma with Y2O3 and YF3 coatings were reasonable.

Finally, we measured the contamination particles generated in real time in the NF3 plasma environment for 60 min in order to confirm the formation behavior of contaminating particles from the Y2O3 and YF3 coatings (Figure 7). Figure 7 shows the concentration and the cumulative concentration of contamination particles generated in real time. During the initial 10 min, contaminating particles were rarely generated at the Y2O3 and YF3 coating films. This is thought to be due to the growth mechanism of the contamination particles and the surface temperature change of the powered electrode by plasma applied bias. The contamination particles initially grow on the coating surface, and the grown contamination particles are dropped off at the weakened bond sites. Also, the surface temperature of the powered electrode was increased to 120 °C while being heated by plasma applied bias, and then maintained at 120 °C by cooling systems. When the coating material reached 120 °C, the coating films became stressed because of the difference in the thermal expansion coefficient between the aluminum used as the base material and the coating material, and grown contamination particles more easily fell off from the surface. Hence, during the initial 10 min, contaminating particles were rarely generated because it takes time to grow contamination particles and raise the temperature of the powered electrode. The results show that more contamination particles were generated in the Y2O3 coating. This finding contradicts the results of mass loss measurements by etching of the YF3 and Y2O3 coatings, as discussed previously. As mentioned above, Y2O3 reacted with the NF3 plasma to produce YOF, a volatile gas, and NOx. The YF3 produced YF2 and NF3 gas without any notable chemical changes (Figure 5). Hence, the chemical corrosion of the Y2O3 coating was faster when subjected to
NF₃ plasma than was the YF₃ coating, which means that the contamination particles grew faster on the Y₂O₃ coating surface than on the YF₃ coating. However, Y₂O₃ has a higher binding energy than that of YF₃; hence, the growth of contamination particles from the Y₂O₃ coating was slower than for the YF₃ coating.

![Graph showing accumulated contamination particle concentration over 0.2 µm size](image)

**Figure 7.** Real-time detection of accumulated contamination particle concentration, over 0.2 µm size, generated from Y₂O₃ and YF₃ during the 60 min NF₃ plasma treatment.

As a result, YF₃ is stable to chemical corrosion when subject to the NF₃ plasma; however, the weak physical binding force resulted in rapid etching and produced fewer contamination particles of a critical size, which could act as defects in the actual plasma process. The Y₂O₃ was slowly etched because of the strong physical binding forces, but was susceptible to chemical corrosion by the NF₃ plasma, which caused large contamination particles, which act as defects in the actual plasma process [18,22].

4. Conclusions

We prepared Y₂O₃ and YF₃ coatings by AD and subjected these to NF₃ plasma. Owing to the characteristics of the starting materials, the initial surfaces had different properties. However, the Y₂O₃ and YF₃ coatings placed at the powered electrode were rapidly etched by the NF₃ plasma treatment. Less etching occurred at the grounded electrode and the grounded walls, and the mass at these surfaces increased owing to sputtering of by-products and fluorine deposition. Additionally, the Y₂O₃ coating showed more rapid chemical corrosion than did the YF₃ coating, and many contamination particles were generated, which acted as defects. This study demonstrates that the YF₃ coating has the potential to be used as a candidate material to reduce contaminating particles in the semiconductor etching process.

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