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Author(s)	Kikuchi, Tatsuya; Onoda, Fumiya; Iwai, Mana; Suzuki, Ryosuke O.
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Influence of sub-10 nm anodic alumina nanowire morphology formed by two-step anodizing aluminum on water wettability and slipping behavior

Tatsuya Kikuchi*, Fumiya Onoda, Mana Iwai, Ryosuke O. Suzuki

Faculty of Engineering, Hokkaido University, N13-W8, Kita-ku, Sapporo, Hokkaido, 060-8628, Japan

*Corresponding author: Tatsuya Kikuchi e-mail: kiku@eng.hokudai.ac.jp

Abstract

The influence of the nanomorphology of alumina nanowires (ANWs) fabricated by a two-step anodizing process on the superhydrophobicity was investigated through advancing contact angle (ACA) and receding contact angle (RCA) measurements. Aluminum nanobowl specimens were anodized in pyrophosphoric acid to fabricate an ordered ANW structure with an average diameter of 7.1 nm, and the outermost surface of the ANWs was chemically bonded with fluorinated self-assembled monolayers. The growing ANWs bent immediately their own weight, and pyramidal ANW structures were formed as they joined the surrounding nanowires together. The ACA value increased with the number density of pyramidal ANW structures due to the reducing area fraction of ANWs, and an increased superhydrophobicity with a contact angle of approximately 165° was measured on the low-density pyramidal structure with a density of 8.1×10^{11} m⁻². Additional anodizing led to complete nanowire bending; thus, the advancing contact angle decreased. The pyramidal nanowire structure exhibited a large slipping property with a contact angle hysteresis (CAH) < 10°, whereas the bent nanowire structure exhibited a decreased slipping property with a CAH > 100°. Superhydrophobic surfaces with opposite water slipping properties were demonstrated by a water dropping experiment.

Keywords: Anodizing, Aluminum nanobowls, Alumina nanowires, Contact angle, Superhydrophobicity

1. Introduction

An improvement in the water wettability of industrial aluminum products is one of the key advances needed to improve their self-cleaning properties, heat exchange rate, anti-fog properties, and corrosion protection. Because such wettability greatly depends on the nanoscale and microscale morphologies fabricated on the outermost surface of aluminum products, various surface finishing processes, such as laser irradiation [1-5], coatings [6-9], and plasma electrolytic oxidation (or microarc oxidation) [10,11], have been widely investigated for wettability modification. In particular, an electrochemical passivation process well known as anodizing easily enables the formation of three-dimensional oxide nano- and micron-sized structures [12-20]. Recently, remarkable wettabilities, including superhydrophilicity indicated by a water contact angle lower than 10° and superhydrophobicity indicated by a water contact angle higher than 150°, have been exhibited by micron- and nano-sized structures created on an aluminum surface by anodizing. Because the top surface of the porous oxide film formed by anodizing possesses a unique honeycomb concavo-convex structure, the surface exhibits superhydrophilic and superhydrophobic behaviors [21-23]. In addition, the wettability control on various materials can be achieved via the replication process using the porous oxide film [24-26]. As the porous oxide film is excessively immersed in acidic solutions during or after anodizing, the top surface of the anodic oxide is gradually dissolved into the solution, and alumina nanowires (ANWs), which are different from the typical structures of barrier and porous films, are formed. The aluminum surface covered with these ANWs exhibits higher superhydrophilic and superhydrophobic behaviors [27-33]. These techniques are based on the typical anodizing method using sulfuric, oxalic, and phosphoric acid solutions. However, the nanostructures of ANWs, including their uniformity, length, diameter, and density, could not be accurately controlled via long-term anodizing and chemical etching.

We previously reported the fabrication of superhydrophilic and superhydrophobic aluminum surfaces covered with sub10-nm ANWs by anodizing in pyrophosphoric acid [34-37]. The hydrophilic ANW-covered aluminum surface exhibited a high superhydrophilicity with rapid water evaporation, and we found that the nanostructure consisting of many ANWs strongly affected the superhydrophilic behavior on the surface [38,39]. Conversely, the surface demonstrated a high superhydrophobicity when the ANWs were chemically modified with low surface tension molecules [40,41]. However, the effect of the nanoscale ANWs grown during anodizing on the superhydrophobicity of the aluminum surface is still unsettled in detail; thus, the ANW nanostructures must be precisely controlled during anodizing in pyrophosphoric acid to obtain a deeper understanding of the superhydrophobicity on aluminum surfaces covered with ANWs.

In the current study, we fabricated a superhydrophobic aluminum surface modified precisely with

ANWs via two-step anodizing, and the water wettability and slipping behavior were investigated in detail through advancing and receding contact angle measurements. In contrast to the conventional simple anodizing method, the two-step anodizing process provides higher controllability of the individual ANW structure, such as the length and distance. We revealed the effect of the anodized ANW morphology that consisted of a pyramidal ANW structure on the superhydrophobicity and slipping behavior on the nanostructured surface. Moreover, superhydrophobic surfaces with completely different water sliding properties that were fabricated by precise control of the anodizing time and the corresponding ANW morphology were demonstrated.

2. Experimental methods

Substrates comprising 4N aluminum plates were cleaned ultrasonically in 99.5 wt% ethanol for 10 min and then polished by an electrochemical technique at a constant voltage of 28 V in 22 vol% perchloric acid (70 wt%)/78 vol% acetic acid at 280 K for 1 min. The mirror-polished aluminum specimens were anodized in a 0.3 M oxalic acid solution at 290 K at a constant voltage of 40 V for 120 min using a direct power supply (Kikusui Electronics, PWR400M) to form a porous oxide structure on the aluminum surface (first anodizing). This procedure induced the rearrangement of the bottom porous oxide distribution during the anodizing process, and an ordered honeycomb configuration was fabricated [42]. A platinum plate was employed as the cathode, and the current density was monitored with a digital multimeter during the anodizing process (IWATSU, VOAC7602).

After the first anodizing process, the porous oxide structure was chemically dissolved in a 0.2 M chromic acid/0.51 M phosphoric acid solution to fabricate an ordered nanobowl structure on the aluminum surface. The aluminum nanobowl specimens were anodized in a 74 wt% pyrophosphoric acid solution at 283 K at a constant voltage of 40 V for 240 min to form ordered ANWs on the surface (second anodizing process). After the second anodizing process, the specimens were immersed in a 0.5 mM perfluorooctylphosphonic acid (PFOPA)/ethanol solution at 313 K for 24 h, and the outermost surface of the ANWs was chemically bonded with fluorinated PFOPA self-assembled monolayers (SAMs).

The nanomorphologies of the anodic oxide formed via the first and second anodizing processes were examined by field-emission scanning electron microscopy (SEM, JEOL, JSM6500F). The SEM observations were carried out after the specimens were coated with a thin electroconductive platinum layer. High-resolution images of the ANW structures were obtained by transmission electron microscopy (TEM) and scanning transmission electron microscopy (STEM, FEI, Titan Cubed G2 60-300). Atomic force microscopy (AFM, Hitachi High-Technologies, Nanocute) was also employed to characterize the height and the number density of ANW structures. The wettability was evaluated through advancing

contact angle (ACA) and receding contact angle (RCA) measurements with an ultrapure water droplet using an optical contact angle meter (Kyowa Interface Science, DM-501). The sliding behaviors of the water droplet, which was dropped with a needle and splayed with a micropipette, on the 5°- and 90°-tiled superhydrophobic surfaces, were recorded using a digital video camera.

3. Results and Discussion

Fig. 1a shows the current density-time curves during a) the first anodizing process in oxalic acid and b) the second anodizing process in pyrophosphoric acid at the same constant voltage of 40 V. Typical current density-time curve was obtained by the first anodizing process in oxalic acid. This porous oxide formed by the first anodizing process for 120 min was chemically removed, and then the corresponding aluminum nanobowl surface was observed by SEM (Fig. 1b). A honeycomb structure consisting of numerous nanobowls with an average size of approximately 100 nm was formed on the aluminum surface. After chemical dissolution, the aluminum nanobowl specimen was anodized in pyrophosphoric acid at 40 V and 283 K. We previously reported that the ANWs grew stably during anodizing under this operating condition [35], and the anodizing process provides higher controllability of the ANW morphologies by controlling the anodizing time. The current density gradually decreased with the anodizing time for up to 240 min after the early rapid increase and decrease (Fig. 1a). Compared with the current density-time curve in oxalic acid, the current density obtained in pyrophosphoric acid is considerably smaller than that in oxalic acid. Such low current density is typically measured during anodizing in pyrophosphoric acid under various applied voltages [35], and the growth rate of the anodic oxide is slow. This is may be due to the high viscosity and the low electroconductivity of the concentrated pyrophosphoric acid solution.

The surface of the specimens anodized in pyrophosphoric acid was modified with PFOPA to provide a water-repellent finish. Fig. 2a represents the changes in the advancing contact angle (ACA, red) and the receding contact angle (RCA, blue) with the anodizing time using error bars. Here, the measurements of the ACA and RCA values were performed at five different positions on each aluminum specimen, and the values, then, were averaged without the maximum and minimum values. The ACA value for the aluminum nanobowl surface was 105.6°, which indicated that it possessed slight hydrophobicity. The ACA value rapidly increased to approximately 140° after a short anodizing process for 5 min and gradually increased when the anodizing time increased 20 min. The ACA value then rapidly increased once again when the anodizing time increased from 20-25 min, and superhydrophobicity of the anodized surface was indicted by an ACA that was higher than 150°. The ACA value increased gradually and reached a maximum value of 164.5° after the surface was anodized for 120 min, and then decreased to approximately 155° when the anodizing time increased to 240 min. In summary, the SAM-modified

aluminum surface anodized in pyrophosphoric acid possessed a highly hydrophobic nature, and the hydrophobic behavior can be divided into the following three stages: the initial hydrophobic stage with an ACA of 140-145°, the middle superhydrophobic stage with an ACA of 160-165°, and the latter superhydrophobic stage with an ACA of 155°. The RCA values were substantially changed by increasing the anodizing time. The RCA value measured on the aluminum nanobowl surface was 25.5° and then decreased significantly to 0° when the surface was anodized for 5-10 min. However, the RCA value rapidly increased to approximately 150° when the surface was anodized for 15-25 min and gradually increased to 155° as the anodizing time increased. Finally, the RCA value suddenly dropped to 0° when the surface was anodized for 180 min. The contact angle hysteresis (CAH) is an important parameter of the slipping behavior of a water droplet formed on a surface. The CAH value is defined as the difference between the ACA and PCA values, and the variation of the CAH value as a function of the anodizing time is summarized in Fig. 2b. The early and late stages exhibited a sticky state with high CAH values, whereas the middle stage displayed a slippery state with low CAH values. Compared with the previously reported superhydrophobic surfaces fabricated by the typical anodizing method (ACA values: approximately 150-170°, CAH values: less than 10-70°), similar high superhydrophobicity measuring 150-165° in the ACA value could be achieved via anodizing in pyrophosphoric acid. Moreover, extremely wide CAH values measuring from less than 10° to more than 150° could be obtained by controlling of the anodizing time, thus the slipping and sticking properties of the water droplet on the aluminum surface can be easily controlled by choosing an appropriate anodizing time.

The SEM images of the hydrophobic aluminum specimens formed by the second anodizing process in pyrophosphoric acid are summarized in Fig. 3. When the aluminum nanobowl specimen was anodized for 5 min, a honeycomb porous oxide possessing a periodic nanoscale pore structure was obtained on the surface. Although similar honeycomb oxide structures were also observed on the specimen after anodizing for 10 min, 15 min, and 20 min, the diameter of each pore gradually increased with the anodizing time due to the chemical etching of the aluminum oxide in the pyrophosphoric acid solution. The complete dissolution of the honeycomb oxide structure started after it was anodized for 21 min, and clear convex parts were formed at the triple points of the honeycomb structure. The continued anodizing process enabled the formation and growth of ANWs at the triple points, and the pyramidal ANW structures consisted of several ANWs that appeared on the surface (22 min). The number of ANWs contained in each pyramidal structure gradually increased with anodizing time from 23 min to 30 min due to the growth of the ANWs. Therefore, the size and height of the pyramidal ANW structures increased with anodizing time, and as a result, the number density of the pyramidal structures decreased. Finally, the aluminum surface was covered completely with numerous long ANWs for anodizing times longer than 180 min.

Based on Figs. 2 and 3, a slippery superhydrophobic surface with higher ACA values and lower CHA values was formed on the aluminum surface covered with pyramidal ANW structures. During advancing and receding contact angle measurements on these surfaces, the water droplet was supported by the top of each hydrophobic pyramidal structure consisting of water-repellent ANWs. To deeply understand the effect of the nanomorphologies on the wettability and slipping behavior, the pyramidal ANW structures were examined by AFM. Fig. 4a shows AFM images of the pyramidal structures formed by the second anodizing process for 22 min, 25 min, and 60 min. Many narrow pyramidal structures were distributed on the whole aluminum surface after 22 min. These pyramidal structures increased in size as the anodizing time increased due to the growth of the ANWs and the joining of surrounding ANWs, and "high ANW mountains" with a low number density were observed on the surface formed after anodizing for 60 min. Figs. 4b and 4c show the changes in the height and the number density of the pyramidal ANW structures with the anodizing time. Here, the average height was measured from the bottom honeycomb structure to the top of the pyramidal structure. The height of the pyramidal structure increased with anodizing time for up to 60 min (from 201 µm for 22 min to 591 µm for 60 min) due to the growth and tangling of the ANWs. However, the height started to decrease after anodizing for more than 80 min, and clear pyramidal structures disappeared from the surface due to the bending of soft amorphous ANWs. Accordingly, the number density of the pyramidal ANW structures drastically decreased with anodizing time (from 3.0×10¹³ m⁻² for 22 min to 8.1×10¹¹ m⁻² for 100 min).

Fig. 5a represents the effect of the number density of the pyramidal structures, ρ , on the advancing contact angle, and the ACA value increased as the number density decreased. Consequently, an increased superhydrophobic contact angle of approximately 165° was caused by the low-density pyramidal structures. This superhydrophobicity was due to the formation of a composite structure consisting of water/ANW and water/air interfaces, and in this state, the water droplet did not penetrate into the concave parts of the composite structure (Cassie-Baxter state) [43]. Cassie and Baxter developed an equation for the steady-state contact angle on the composite structure as follows [44]:

$$\cos \theta_{\rm CA} = f_1 \cos \theta_1 + f_2 \cos \theta_2 \qquad (1)$$

where θ_{CA} is the contact angle; f_1 and f_2 are the area fractions of the solid and the air at the interface, respectively; and θ_1 and θ_2 are the contact angles measured on the flat solid and air surfaces, respectively. Then, equation (1) becomes:

$$\cos \theta_{\rm CA} = f_1 \cos \theta_1 + (1 - f_1) \cos \theta_2 \qquad (2)$$

Here, equation (2) can be rewritten when $\theta_2 = 180^\circ$ as:

$$\cos \theta_{\rm CA} = f_1 \cos \theta_1 + f_1 - 1 \qquad (3)$$

The relationship between the calculated contact angle and the area fraction of solid at the composite interface in the Cassie-Baxter state is shown in Fig. 5b. The calculated contact angle gradually increased as the area fraction of solid decreased, and a similar curved relation between the experimental result and the Cassie-Baxter model was obtained. The maximum ACA value obtained experimentally remained at approximately 165° on the surface that contained the minimum number density of the pyramidal structures (Fig. 5a), and additional low-density pyramidal structures may result in an increased superhydrophobic contact angle of more than 165°. However, the amorphous ANWs formed by long-term anodizing were completely bent due to their own weight; thus, the pyramidal structures were not present on the surface (Fig. 3). Therefore, there may be a limitation to increasing the superhydrophobicity by decreasing the number density of the pyramidal structures formed by anodizing in pyrophosphoric acid.

As the shape of the contact area on each of the pyramidal structures that supported the water droplet was assumed to be a circle, the real area fraction of the pyramidal structures, f_{1-real} , can be calculated as follows:

$$f_{1-\text{real}} = \rho \pi r^2 \qquad (4)$$

where *r* is the radius of each circular contact area. The number density of the pyramidal structures formed after anodizing for 22 min was 2.98×10^{13} m⁻² (Fig. 3), and the ACA value was measured to be approximately 152° (Figs. 2a and 5a). The value of *r* that satisfied these experimental results was calculated to be approximately 50 nm using equation (4). Fig. 6 shows a) a bright-field (BF) STEM image and b) a high-resolution TEM image of the pyramidal ANW structures formed by anodizing for 22 min. A porous oxide structure measuring approximately 100 nm in thickness was formed on the aluminum substrate, and pyramidal structures consisting of sub10-nm scale ANWs formed by the chemical dissolution of aluminum oxide were observed on the porous oxide. We previously found that the ANWs grew and become longer with the anodizing time, whereas the thickness of the porous layer was almost unchanged due to the rapid dissolution of pore walls [35]. Therefore, only the ANWs grows longer length during anodizing. The average diameter of the ANWs was measured to be 7.1 nm. Although the size of the outermost part of the pyramidal structures increased due to the bundling of multiple ANWs, their width was measured to be approximately 10-15 nm, which differed from the calculated value (50 nm). Therefore, the shape of the pyramidal structures was expected to be quite pressed and transformed by the weight of the supported water droplet during the contact angle investigations.

We demonstrated the slipping behavior of a water droplet on the superhydrophobic surfaces with low CAH values that were produced by the long ANW structures and high CAH values that were produced by the pyramidal ANW structures. Fig. 7 shows video snapshots of the sliding behavior of a water droplet on the 5°-tilted superhydrophobic specimens. Here, a 12.5 µL water droplet was continuously dropped on the superhydrophobic surface from a needle. When a water droplet was dropped onto the long ANW structures (Fig. 7a), the droplet became stuck on the nanostructured surface and did not slip. Although the size of the water droplet increased due to a continuous water supply and the shape gradually changed to an ellipsoid, the large droplet also became stuck to the surface. In contrast, as the water droplet was applied to the pyramidal ANW structures (Fig. 7b), the droplet immediately slipped down to the right side without any sticking.

These superhydrophobic specimens with different CAH values were vertically placed on a laboratory table, and a 10 μ L water droplet was splayed toward the specimen using a micropipette (Fig. 8). In the case of the long ANW structures (Fig. 8a), the water droplet released from the pipette became stuck on the 90°-tiled superhydrophobic surface without any splashing due to their low CAH value. Such sticking behavior occurred on the whole anodized surface (right, Fig. 8a). Conversely, the water droplet applied to the pyramidal ANW structures was dispersed everywhere and did not stick due to the high CAH value, and the surface remained clean (Fig. 8b). Based on Figs. 7 and 8, even though similar superhydrophobic aluminum surfaces were covered with ANWs, completely different slipping behaviors were easily obtained by controlling the anodizing time and the corresponding CAH value.

4. Conclusions

We investigated the influence of the ANW morphology, including a pyramidal ANW structure, on the water wettability and slipping behavior on a hydrophobic surface fabricated via a two-step aluminum anodizing process. The ANWs immediately formed pyramidal ANW structures during the early stage of anodizing, and their pyramidal structures increased in size due to joining of the surrounding nanowires. The advancing contact angle increased with the number density of pyramidal ANW structures due to a reducing area fraction of the solid/liquid interface, and a highest superhydrophobic contact angle measuring approximately 165° was obtained on a low-density pyramidal structure with a density of 8.1×10^{11} m⁻². Further anodizing led to complete nanowire bending and the disappearance of the pyramidal structures due to the growth of long ANWs; thus, the advancing contact angle decreased. The pyramidal ANW structure exhibited high slipping value indicated by a CAH < 10°, whereas the bent nanowire structure exhibited a lower slipping value indicated by a CAH > 100°. Superhydrophobic surfaces with a range of water slipping properties can be fabricated by controlling the anodizing time and the corresponding CAH value.

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Figure Captions

Figure 1 a) Current density-time curves during aluminum anodizing at a constant voltage of 40 V in a 0.3 M oxalic acid solution at 290 K (first anodizing process) and a 74 wt% pyrophosphoric acid solution at 283 K (second anodizing process). b) SEM image of the aluminum nanobowl surface fabricated by anodizing in oxalic acid and subsequent oxide removal.

Figure 2 a) Changes in the advancing contact angle (ACA) and the receding contact angle (RCA) of a water droplet with anodizing time during second anodizing in pyrophosphoric acid. b) The change in the contact angle hysteresis (CAH) with the anodizing time.

Figure 3 SEM images of the anodized aluminum surface in a 74 wt% pyrophosphoric acid solution at 283 K from 5 min to 180 min.

Figure 4 a) AFM images of the pyramidal alumina nanowire (ANW) structures formed by anodizing for 22 min, 25 min, and 60 min. b) The change in the height of pyramidal structures with the anodizing time.c) The change in the number density of pyramidal structures with anodizing time.

Figure 5 a) The relationship between the advancing contact angle and the number density of pyramidal structures. b) The change in the calculated contact angle with the area fraction of solid at the interface of the Cassie-Baxter state.

Figure 6 a) A BF-SETM image and b) high-resolution TEM image of the pyramidal ANW structure fabricated by the second anodizing process for 22 min.

Figure 7 Video snapshots of a 12.5 μ L water droplet dropped from a needle onto the 5°-tilted superhydrophobic surfaces with a) a high CAH value and b) a low CAH value.

Figure 8 Video snapshots of a 10 μ L water droplet splayed from a micropipette onto 90°-tiled superhydrophobic surfaces with a) a high CAH value and b) a low CAH value.



Figure 1.



Figure 2.



Figure 3.



Figure 4.



Figure 5.



Figure 6.



Figure 7.



Figure 8.