A Novel Simple Anti-Ice Aluminum Coating: Synthesis and In-Lab Comparison with a Superhydrophobic Hierarchical Surface

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Abstract: A simple process to obtain a slippery surface with anti-ice and ice-phobic properties has been developed and characterized in laboratory. The coating is realized by growing a nanostructured pseudo-boehmite on an aluminum substrate and applying an environmentally compatible final functionalization consisting of a fluorine-free oligomeric short-chain alkylfunctional silane. The resulting surface is conceptually similar to a slippery liquid infused porous surface (SLIP) material, but the porous infrastructure is inorganic and the process to generate it is very simple, rapid and economic. The coating performance in terms of hydrophobicity at room and low temperatures, ice nucleation temperature and ice adhesion forces were assessed in laboratory. Moreover, hydrophobicity persistence was tested after prolonged immersion in acid, basic and saline solutions while the durability of ice-phobic behavior was assessed by repeated shear stress tests. Results are compared with those of a superhydrophobic hierarchical aluminum surface obtained with a fluorinated siloxane. The novel coating shows very good anti-ice properties and relevant durability, with some differences from the fluorinated surface. The novel slippery coating process is promising for industrial applications, also taking into account its environmental compatibility, simplicity and low cost.

Keywords: superhydrophobic; slippery; ice-phobicity; durability; nanostructure; coating; aluminum

1. Introduction

Ice and snow accumulation can cause serious problems in many fields, for example aircrafts, roads and railways, wind turbines and photovoltaics [1]. The accretion of ice or snow sleeves on overhead power lines (OHL) may also cause the failure of conductors and guard wires and/or the collapse of the towers, provoking the disruption of the transmission networks and the interruption of the communication optical fibers embedded in the guard wires [2] with consequent severe economic impacts.

Recently, several superhydrophobic (SHP) coatings were studied for anti-icing applications: the excellent water repellency property sometimes leads to ice-phobicity; however, their application is still controversial [3–8]. Moreover, the durability aspects of SHP coatings and surfaces have to be taken into serious account. Many studies indeed evidence the quick deterioration of the SHP surfaces with loss of the hydrophobic and ice-phobic properties [8–10], while other works demonstrate that an accurate design of the SHP surfaces, based on a fine-tuning of both the morphology and the deposition method of the coating, gives rise to a real durable ice-phobic material [11]. However, the development of a
scalable and industrially feasible process, for instance for the fabrication of cables for overhead power lines, is still an open issue.

In a previous work, we studied a superhydrophobic surface obtained by applying a commercial fluoroalkyl silane (FAS) on a hierarchical micro-nano aluminum structure [12]. This coating maintained hydrophobic behavior even at low temperatures and evidenced amazing durability of ice-phobic properties after several freeze-thaw cycles in shear stress tests; however, the durability of this coating exposed in harsh environmental conditions was to be fully understood. Moreover, despite the very quick process to produce the ice-phobic material and the good results in terms of the ice-phobic property and its durability, the presence of polyfluorinated alkyl molecules may strongly limit the application of this coating for environmental problems [13].

Another interesting field of study concerns the development of slippery liquid infused porous surfaces (SLIPS) coatings composed of a micro/nanoporous structure with low surface energy, infused with a lubricant fluid [1,14,15]. These materials are designed according to two principles: (1) the lubricating fluid and the water have to be immiscible; (2) the chemical affinity between the lubricating fluid and the solid should be higher than that between the water and the solid [16]. Thanks to the presence of the infused fluid, locked in the micro-nanostructure, SLIPSs evidence a very good capability in repelling various liquids, in keeping very low contact angle hysteresis and in quickly restoring the lubricant after mechanical abrasion or damages [17]. They are also considered perspective materials for anti-icing applications because of their impressive stability in high humidity conditions [15] and their ability in reducing the ice adhesion strengths [1].

However, the anti-icing durability of SLIPS coatings is still a challenge [18]; in particular, icing/deicing cycles, water droplets and low temperatures, which can accelerate the lubricant depletion [19,20]. Moreover, to obtain SLIPS, complicated and multi-step synthetic methods are necessary, employing expensive starting materials and technologies [21–24].

In this work, we present a novel aluminum slippery surface (SLS) that is synthesized by boiling the aluminum substrate to grow a nanostructured pseudo-boehmite and by applying an oligomeric short-chain fluorine-free alkyl siloxane. We studied its wettability under different environmental conditions and its ice-phobic behavior; eventually, we assessed the durability of its properties. This surface is characterized by a light hydrophilic static contact angle, a very low sliding angle and low ice adhesion. SLS has similar anti-icing, wettability and even better durability behavior of SLIPS, despite the considerable differences in the morphology and in the chemistry: indeed, the under-layer structure is a porous hydrophilic inorganic oxide, no lubricant is infused and the final surface is dry.

The process to obtain the SLS is very intriguing because it is fast, environmentally friendly, economical, no special equipment and expensive chemicals are needed and thus could be easily transferred to industrial production. Although the process to grow the pseudo-boehmite is well known [25] and it is also very often used to build a nanostructure, as far as we know, a full characterization of durability and ice-phobicity of a surface similar to SLS is still missing.

Results and performance in terms of hydrophobicity at room and low temperatures, ice adhesion forces and durability are compared with those of the superhydrophobic boehmite surface studied in our previous work [12]. This study aims to directly compare results obtained in the same laboratory, applying the same ice-phobic characterization methods and the same aging protocols, given that, so far, a recognized protocol to assess the durability of the coatings is missing. Some important results are evidenced, indicating that slippery surfaces better tolerate critical and aggressive environmental conditions; however, the SHP coating keeps higher durability of ice-phobic properties.

2. Materials and Methods

2.1. Materials

Flat plates (20 mm × 70 mm × 2 mm) and bars (12 mm diameter × 100 mm length) of aluminum alloy (6082) were used as substrates.
Dynasylan® SIVO CLEAR EC was purchased from EVONIK (Essen, Germany); the product is composed of fluoroalkysilane (FAS) 2%, propan-2-ol 93% and dodecane 5%, and was used as received. Dynasylan® 9896 was purchased from EVONIK and it is an oligomeric short-chain alkylfunctional silane. It was used after dilution 1:1 in volume in ethanol. Hexane (95%), acetone (>99.5%) and acetic acid (≥99.7%) were purchased from Sigma Aldrich (St. Louis, MO, USA). Ethanol (≥99.9%), ethylene glycol (<98%), sodium chloride and sodium hydroxide were purchased from Carlo Erba (Milan, Italy). Diiodomethane (99%) was purchased from Alfa Aesar (Haverhill, MA, USA).

2.2. Preparation of Superhydrophobic Surface (SHP)

The aluminum alloy specimens were cleaned with basic soap, rinsed in an ultrasonic bath for 10 min with acetone and dried under nitrogen flux. For the preparation of the SHP samples, the aluminum alloy was sandblasted with micro glass beads in the 40–70 µm diameter range to generate a micrometric roughness. The sandblasting process was carried out for 10 s for each exposed area, with an applied pressure of 3.5 bar while keeping the sample 5 cm distant from the nozzle. Afterward, in order to generate a pseudo-boehmite nanostructured layer, the sample was boiled for 5 min at 100 °C in ultrapure water. Then, it was dried in air, rinsed with n-hexane and dried under nitrogen flux, as described in [12,26,27]. The obtained hierarchical substrate was dip-coated in the FAS bath (dipping–withdrawing speed: 0.7 mm/s, permanence time: 120 s) and cured at 70 °C for 1 h.

2.3. Preparation of Slippery Surface (SLS)

For the preparation of the slippery surface, the aluminum alloy specimens were cleaned with basic soap, rinsed in an ultrasonic bath for 10 min with acetone and dried under nitrogen flux. To obtain a porous nanostructured pseudo-boehmite, the aluminum alloy was boiled for 30 min at 100 °C in ultrapure water, dried in air, rinsed with n-hexane and dried under nitrogen flux. Dynasilan® 9896 was diluted in ethanol (1:1 in volume) and the solution (Dyn-50) was stirred for 1 h. The aluminum samples were dip-coated in ALK (dipping–withdrawing speed: 0.7 mm/s, permanence time: 120 s) and cured at 120 °C for 1 h.

2.4. Characterization Methods

Surface morphologies of all the samples and the thickness of the oxide layers were examined using a field emission scanning electron microscope (FE-SEM; Mira III, Tescan, Brno, Czech Republic). Energy dispersive X-ray spectroscopy (EDS) (Bruker, Billerica, MA, USA) micro-analysis was used to determine the composition of the materials. A Taylor Hobson stylus profilometer was used to measure surface roughness. Data were averaged over at least 5 runs for each sample. The Vickers hardness was estimated by the means of the FM 810E micro-hardness tester (Future-tech Corp., Kawasaki, Japan) applying a force of 1 g to the indenter. Data were averaged over 5 tests for each sample. The FT-IR characterizations were conducted with the FT-IR Alpha 1 (Bruker) spectrometer with ATR apparatus (diamond crystal was used as the internal reflection element). The gravimetric measure was performed with a microbalance Cubis MSU6 (Sartorius, Göttingen, Germany) with a sensitivity of 0.001 mg.

The static water contact angles (WCA) were measured with the DSA 30 Drop Shape Analyzer (Kruss, Hamburg, Germany) with the method of the sessile drop using 4 µL volume of water at 20 °C on at least 5 points for each sample. The WCA measurements at low temperatures were conducted by means of a Peltier plate in an environmental chamber at −4 °C, deposing the drop on the cooled samples. The measurements of dynamic contact angles (hysteresis (CAH), roll-off (RA) and sliding angles (SA)) at 20 °C were performed with the tilting plate method using 20 µL volume of water.
The measurements of the dynamic contact angles at $-4 \, ^\circ\text{C}$ were performed with the captive needle method in the Peltier chamber (Kruss, Hamburg, Germany). To measure the roll-off angles at $-4 \, ^\circ\text{C}$, the Peltier chamber was attached to a home-made tilting table and using a droplet of 20 $\mu\text{L}$ of water. Data were averaged on at least 3 measures for each sample.

The measurements of surface free energy (SFE) were performed with OWRK model measuring static contact angles with water, diiodomethane and ethylene glycol.

The ice nucleation temperature $T_n$ was measured in the following way: the sample was attached to a Peltier cell and covered by a transparent lid. A water droplet of 10 $\mu\text{L}$ was deposited at $20 \, ^\circ\text{C}$ on the sample; then, the temperature was cooled down with a cooling rate of $6 \, ^\circ\text{C}/\text{min}$ until the $T_n$ was reached and the drop started solidification [28].

The ice adhesion strength was evaluated by means of the shear force needed to extract a sample pole from an ice block. We adopted a home-made apparatus equipped with an electromechanical testing system (INSTRON 4507). The testing process and the apparatus are described in the Supplementary Materials section. No standard procedure was available for the ice-phobicity tests.

2.5. Durability Performance Tests

The durability study on the samples followed a protocol consisting of four tests (three in wet chemicals and one for ice shedding) and, for each test, different conditions were implemented according to [12,24,29] Hydrophobic and/or ice-phobic properties of the samples were measured after each test phase and compared to the pristine sample. The test frequencies and applied procedure allow distinguishing the different aging behaviors of the samples. Further information about the durability protocol is reported in the Supplementary Materials.

2.5.1. Resistance in Wet Chemicals

Freshly prepared SLS and SHP samples were used for each test. Different solutions were prepared to assess the durability of each sample in three different conditions: acid (acetic acid, pH = 3), basic (sodium hydroxide, pH = 8.5) and saline (sodium chloride 35 g/L). The as-prepared solutions should assess the resistance in different conditions: the acid solution simulated the pH of an acid rain event, the alkali environment recreated the pH of the seawater and the ionic solution recreated an average sea salt concentration. Samples were completely immersed in this aging solution for different times then rinsed with water and dried under nitrogen. The characterization was done by measuring WCA, contact angle hysteresis (CAH) and RA/SA to assess the variation of hydrophobic properties. Five and three replicated measurements were performed for WCA and for CAH/RA/SA, respectively. Tests were conducted at room temperature to avoid the combination of multiple effects.

2.5.2. Resistance to Ice Shedding

The resistance of the surfaces to ice removal was tested by repeating the shear stress test on the same samples several times; ten bars for each sample were tested and the average shear stress was calculated. The variation of the shear force required to remove ice after several (up to 20) icing-deicing cycles allows assessing the durability of ice-phobic properties.

3. Results and Discussion

3.1. Morphological and Chemical Evaluation

The hierarchical surface of SHP is shown in Figure 1. At low magnification, the micrometric roughness generated by the sandblasting process is visible; the resulting roughness ($R_a$) is 0.94 $\mu\text{m}$. At higher magnification, the typical grass-like aspect of the pseudo-boehmite layer is observable [30]. A thickness of about 300 nm was estimated for the oxide layer [31].
Table 1. EDX semi-quantitative analysis of the SLS sample with that of a non-nanostructured aluminum sample similarly covered with Dyn-50 (SLS-1), a silicon content (attributable to the Dyn-50) much higher in SLS than in SLS-1 can be appreciated (Table 1).

The roughness ($R_a$) of the SLS aluminum substrate was 0.30 µm, and after the boiling process a boehmite layer of about 500 nm of thickness was obtained [31]. Differently than SHP, in the SEM images of SLS (Figure 2), the nano-grass-like pseudo-boehmite is partially hidden; only some pinnacles are visible. In fact, the Dyn-50 completely fills up the nanovoids in the pseudo-boehmite layer and the surface is almost totally covered by a much thicker layer than the FAS one. The thickness is calculated at about 1 µm according to the gravimetric analysis and the Landau–Levich model [33].

In this sample, the role of the porous pseudo-boehmite is dual: firstly, similarly to SHP, it improves the adhesion of the coating thanks to Al-O-Si bonds, secondly, the porosity of the nano-oxide layer allows to host a large amount of Dyn-50. In fact, by comparing the EDX semi-quantitative analysis of the SLS sample with that of a non-nanostructured aluminum sample similarly covered with Dyn-50 (SLS-1), a silicon content (attributable to the Dyn-50) much higher in SLS than in SLS-1 can be appreciated (Table 1).
The FT-IR spectra (Figure 3) further confirm that SLS is hosting a bigger amount of Dyn-50 than SLS-1. This is well visible in the peaks at 2960–2872 cm$^{-1}$, referable to the stretching mode of C–H bonds in the siloxane. In the SLS spectrum, the peaks related to the pseudo-boehmite layer are also visible at 780–470 cm$^{-1}$ together with the weak peaks at 901 and 864 cm$^{-1}$ attributable to the stretching of Si–O–Al bonds [34]. It is also important to underline the presence of the signals due to Si–O–Si (1060 cm$^{-1}$) indicating the presence of siloxane chains.

![Figure 3. FT-IR spectra of SLS and SLS-1: (2960–2872 cm$^{-1}$) ν C–H; (1463–1297 cm$^{-1}$) ω C–H; (1215 cm$^{-1}$) ν Si–CH$_2$; (1060 cm$^{-1}$) ν Si–O–Si.](image)

Thanks to the gravimetric analysis, it was estimated that SLS hosts almost 70% more of Dyn-50 than SLS-1 (about 149 μg/cm$^2$ and 87 μg/cm$^2$ of siloxane were calculated, respectively).

### 3.2. Hydrophobic Properties at Room and Low Temperatures

The static (WCA) and dynamic angles (CAH and RA/SA) of SHP and SLS were collected at room and low temperatures and are shown in Table 2 together with the surface energy data (SFE).

#### Table 2. Wettability and surface free energy at room and low temperatures.

| Sample | Room Temperature | Low Temperature |
|--------|------------------|-----------------|
|        | WCA (°) | CAH (°) | RA/SA (°) | SFE (mJ/m$^2$) | WCA (°) | CAH (°) | RA/SA (°) |
| SHP    | 170.2 ± 2.7 | 4.2 ± 0.7 | 2 ± 1 | 0.11 ± 0.03 | 143.2 ± 2.7 | 32.3 ± 2.7 | >90 |
| SLS    | 86.7 ± 1.5 | 2.0 ± 0.1 | 2.0 ± 0.2 | 38.50 ± 0.02 | 81.5 ± 0.9 | 8.6 ± 0.8 | 16.0 ± 2.0 |

The SHP sample evidences very good superhydrophobic properties with high WCA and very low RA and CAH; however, at low temperatures, the surface loses its super-hydrophobicity and the dynamic properties highly degenerate. This behavior is typical of SHP surfaces [14] and it is mainly due to the humidity condensation from the surrounding air on the surface that drives the transition from the Cassie–Baxter to the Wenzel state [35].

The SLS evidences a weak hydrophilic behavior at room temperature, comparable with that of bare aluminum (WCA about 85°; SFE about 33 mJ/m$^2$) [36]. Despite this, the dynamic properties are consistently better with respect to the bare aluminum. The very low hysteresis and sliding angles are

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**Table 1.** EDX atomic concentration in SLS and SLS-1 samples.

| Sample | Si (at.%) | Al (at.%) | O (at.%) | C (at.%) |
|--------|-----------|-----------|----------|----------|
| SLS    | 4         | 56        | 17       | 23       |
| SLS-1  | 1         | 67        | 8        | 24       |
comparable to those of SLIPS [37] but, notwithstanding this similarity, SLS cannot be classifiable as a
SLIPS. Indeed, the porous surface in SLS is composed of hydrophilic pseudo-boehmite while SLIPS
typically have a hydrophobic porous structure, as previously discussed in [16]; moreover, the surface
is dry and there is not a liquid intra-layer between the solid and the repellent fluid [38].

At low temperatures, the wettability tests of SLS evidence a little reduction of the static angle but
the SA and the hysteresis keep low values. This behavior indicates that the SLS coating better tolerates
the cold conditions and the humidity condensation phenomena, similarly to SLIPS coatings [37].
Probably, this is due to the absence of impurities and surface defects, which are very likely to be present
on a rough superhydrophobic surface exposed to condensed humidity. On the smooth surface of SLS,
the receding angle, which is the parameter related to the adhesion of the drops [39], keeps a good
value, contributing to maintaining a low hysteresis even at low temperatures. As a matter of fact,
the underlying pseudo-boehmite in SLS seems to have an important role in decreasing the hysteresis
and sliding angle both at room and low temperatures: indeed, the un-boiled SLS-1 sample showed
worse dynamic properties than SLS at room temperature (CAH 9.6° and SA 12.3°) and even worse at
−4 °C (CAH 11.8 and SA 26.3°). Further investigation to explain this phenomenon will be done.

3.3. Ice-Proof Performances

Although a protocol to measure ice-proof performances is not defined yet, two main parameters
can be studied to evaluate the properties of a surface with respect to the ice accretion and adhesion: the
ice nucleation temperature ($T_n$) and the ice adhesion strength on the surface. The ability of a surface
in lowering the $T_n$ of a supercooled water droplet is defined in literature as anti-icing property [40];
thanks to this, the surface can prevent or hinder the ice formation. In parallel, the ability of the surface
to lower the adhesion of the ice can be referred to as ice-phobicity and it defines how easily the ice can
be removed from the surface [22].

Considering these definitions, we studied both anti-icing and ice-phobic properties of the coated
samples compared to the bare aluminum ones.

3.3.1. Anti-Icing Properties

According to the literature [41], ice nucleation is a heterogeneous process in which ice nuclei
forms at the interface with the solid substrate rather than inside the water droplet for energetic reasons.
The heterogeneous nucleation is therefore strictly related to the underlying surface structure, and in
particular to the presence of sites favoring the nucleation and to the contact area between water and
solid surface. A method to measure the influence of the surface structure on the ice nucleation process
is to analyze the temperature $T_n$ at which the drop starts to solidify, as described above.

In our experimental setting, the water droplets on the bare aluminum froze at −7.8 °C while the
$T_n$ for SHP and SLS were −12.1 and −12.4 °C, respectively (Figure 4).

![Figure 4. Icing nucleation temperature $T_n$ on different surfaces.](image-url)
The experimental results demonstrate that both the SHP and SLS surfaces can hinder the formation of frozen droplets, providing interesting and comparable anti-icing properties, although attributable to different mechanisms.

The efficacy of the superhydrophobic surface in reducing ice formation is related to the small contact area between the liquid and the solid. Indeed, the water droplet floats on the superhydrophobic surface and it comes into contact with this only for a small area. This leads to a reduction in both heat exchange and the ice nucleation rate [42–44]. Moreover, it is well reported that in SHP coatings, the air pockets further hinder the freezing process by acting as heat insulators [45,46], delaying the heat exchange.

The anti-icing behavior of SLS is comparable to the SHP in terms of Tn. Two main factors are to be considered. First, the surface is almost completely impregnated with the Dyn-50, which creates a heat insulation effect similar to the air pockets in an SHP surface [47]. Second, the presence of hydrogen bonds favors the ice nucleation and the SLS does not contain chemical groups capable of forming hydrogen bonds. Moreover, it is reported that the interaction between a hydrophilic surface and water can prevent the first layer of water molecules from arranging in ice structure [48].

### 3.3.2. Ice-Phobic Properties

The shear stress force values and the calculated adhesion reduction factor (ARF) are displayed in Figure 5. The ARFs were calculated as the ratio between the shear stress values of bare aluminum alloy and of the samples.

![Figure 5. Shear stress values for bare aluminum, SHP and SLS samples (adhesion reduction factors are also reported). In the box, the home-made equipment used for shear stress measurements.](image)

Both superhydrophobic and slippery surfaces evidenced a good reduction of ice adhesion with respect to the bare aluminum alloy; noteworthy, the SLS has better ice-phobic properties.

As described in several studies [12,39,49,50], the ice-phobic behavior of the SHP surface can be attributed to the high water repellency and to the existence of an “air-cushion” on the surface of the sample that maintain it partially unwetted during the ice formation or that considerably reduce the contact area of ice on the surface. The ice-phobic results obtained for SHP are comparable or better than those of superhydrophobic materials reported in the literature [51–53].
The SLS evidenced an even higher ARF than the SHP. This remarkable result is comparable to the shear stress values of several SLIPS coatings [28,38,54,55].

On a molecular scale, the presence of the non-polar allylic groups completely covering the surface gives rise to very weak intermolecular attraction forces between surface and ice. Thus, the electrostatic van der Waals forces and the hydrogen bonds, between ice and the sample, contribute little to the ice adhesion, minimizing the shear force [51,56]. On a microscopic scale, the SLS is a smooth surface because the nanoporous layer is filled with the coating: this leads to a minor surface area in contact with the ice and, consequently, to a decreasing of the ice adhesion [7,51].

Moreover, as it is widely evidenced in the literature [39,57], the low adhesion of the ice is strongly related to the low CAH: in particular, the hysteresis of SLS kept a low value even at low a temperature, and this gave rise to the low ice adhesion.

3.4. Durability of Wettability Properties

The results of the wettability after aging in harsh environments are reported in Figures 6–8 (for more details, see the Supplementary Materials).

![Figure 6. Static contact angles of SHP and SLS after aging in acid, saline and basic solutions.](image-url)
Figure 7. Hysteresis of SHP and SLS after aging in acid, saline and basic solutions.

Figure 8. Roll-off/slide-off angles of SHP and SLS after aging in acid, saline and basic solutions: values of 90° mean that the drops did not roll off.
As a general consideration, the hysteresis and the roll-off or slide-off angle are strongly affected by the degradation of the coatings, and the increase of their values is a clue of the deterioration of the surfaces [58].

In general, the SLS sample showed better durability than the SHP surface: the CAH and SA values of SLS slowly degraded in harsh environments. In acid and saline conditions, they remained substantially steady after 20 days, while a sudden degradation is observed for SHP surfaces. The test in basic solution is the most critical for both samples as the timescale of degradation is particularly rapid. Anyhow, the samples maintained a rather good RA/SA value (about 10°) up to 16 h.

After the exposition in wet chemical solutions, the FT-IR spectra of the samples evidence some differences with respect to the pristine ones (Figure 9).

![FT-IR of SLS (a) and SHP (b) surfaces after aging in harsh environment.](image)

Figure 9. FT-IR of SLS (a) and SHP (b) surfaces after aging in harsh environment.

A degradation of the SLS due to the Dyn-50 layer reduction after the aging tests is visible from the decreasing intensity of the peaks around 2900 cm\(^{-1}\) (aliphatic C–H bonds stretching), whereas the samples aged in acid and saline solutions (Figure 9a, green and red lines, respectively) showed only a smaller intensity reduction. These results are in good agreement with the wettability analysis of the aged samples.

The FT-IR spectra of SHP samples are less explicative because of the very weak signals of the FAS layer coating. Anyhow, the signals from the underlying pseudo-boehmite layer are well visible (780–470 cm\(^{-1}\)) and their increasing intensities after aging point out the loss of the FAS, especially severe for the basic solution, if considering the short immersion times (Figure 9b).

All the collected data evidenced that the loss of the wettability properties is due in both cases to the significant depletion of the siloxane coatings. According to the literature, this could be attributed to the hydrolysis reaction of Si–O–Al bonds [50,59,60] catalyzed by acids, bases or nucleophilic chlorine ions. In the SHP, this reaction is likely to occur forthwith because the FAS monolayer is widely exposed to the reacting agents and because the fluorinated chains increase the electrophilic nature of silicon atoms. On the contrary, in SLS, this reaction is strongly hindered by the thick coating that protects the interface: the Si–O–Al bonds are exposed to the water only after the hydrolysis of Si–O–Si bonds occurs. Furthermore, the better durability of SLS with respect to SHP can be attributed to the thicker layer of Dyn-50 deposed on the substrate, not only for its role in protecting the interface but also because it has a kind of self-healing properties. When the first functionalized Si-O-Si layer is hydrolyzed, a second layer equal to the first is still available to ensure the wettability properties. In order to confirm this, a durability study on the SLS-1, which is coated with a layer of Dyn-50 of the same order of thickness as SLS, was carried out. As expected, better durability of this sample with respect to the SHP was evidenced for the acid and saline solutions, while the basic solution rapidly degraded the sample. Data highlighting the differences in surface deterioration are reported in Table 3. Moreover, by comparing the hysteresis and sliding angles of SLS and SLS-1, a better endurance of the SLS sample is shown, especially for the tests in acid ad saline solutions. This confirms that the pseudo-boehmite
layer, grafting the coating with Si–O–Al chemical bonds, gives rise to a more durable surface with respect to the mere material deposition on the substrate.

Table 3. Significant wettability results after durability test in wet chemicals for SLS, SLS-1 and SHP.

| Sample | Acid Solution | Saline Solution | Basic Solution |
|--------|---------------|-----------------|----------------|
|        | 7 Days | 15 Days | 24 h | 7 Days | 15 Days | 24 h | 7 Days | 15 Days | 24 h |
|        | CA (°) | CAH (°) | RA/SA (°) | CA (°) | CAH (°) | RA/SA (°) | CA (°) | CAH (°) | RA/SA (°) |
| SLS    | 89.1   | 6.7    | 6.1    | 92.8   | 6.6    | 8.2    | 100.6 | 26.4    | 32.8 |
| SLS-1  | 88.6   | 26.3   | 44.5   | 99.5   | 38.9   | 56.9   | 99.4  | 33.4    | 39.8 |
| SHP    | 144.6  | 90.9   | no roll | 87.1   | 46.2   | no roll | 150.9 | 41.7    | 19.7 |

The durability of the slippery sample is comparable to that of some SLIPS described in the literature [22,24]; the remarkable stability of the SLS surface evidenced also a role as a protective layer for the aluminum substrate under the three tested aging environments. This behavior is considered very important for prospective applications in outdoor environments.

3.5. Durability of Ice-Phobic Properties

Shear stress tests were repeated twenty times on the same bars for sample SLS and compared to the results obtained for SHP after the same test procedure [12]. The shear stress values are shown in Figure 10.

![Figure 10. Durability of ice-phobic properties of SLS and SHP with repeated shear stress measurements. Pale areas represent the standard deviation for each value.](image)

As reported in a previous study, the SHP evidenced remarkable durability for 20 shear stress cycles and showed better performances with respect to several superhydrophobic coatings described in the literature [12,61]. This behavior is thought to be attributable to the presence of the micro-roughness which enhances the hardness of the sample that is, consequently, less exposed to ice abrasion. In order to confirm this, the hardness of a sandblasted and of a pristine aluminum alloy sample was tested with a microhardness tester and the resulting marks were measured by SEM analysis. The hardness of the pristine aluminum was calculated to be about 70 hv, while on the sandblasted sample, no indentation
was evidenced, and this could indicate harder material. For comparison purposes, we measured a tumbled finished specimen, assuming that the sandblasting treatment could give rise to a surface with similar mechanical properties. The hardness was about 169 hv and we can assume that the sandblasted specimen should have a similar or higher value.

The SLS evidenced a very good and constant ice-phobic behavior for as long as 12 tests, after which the degradation began to appear, followed by a strong and sudden increase of the ice adhesion during the further tests. This trend can be attributed to the progressive depletion of Dyn-50 layers up to the complete removal of the coating, with a consequent abrupt surge of ice adhesion. The proposed mechanism of the coating depletion is double: the breakage of the Si–O–Si bonds is due to the reaction with water during the process of ice formation [50,59,60], similarly to what was mentioned for wet chemical tests (Section 3.4) and the mechanical abrasion of the coating layer due to the ice during the process of ice detachment.

The partial removal of the coating from the surface also exposes the acicular nanolayer to the ice and the nanostructures indented into the ice are very likely to be damaged during ice removal [3,9,50,62]. Further confirmation of this comes from SEM images and EDX analysis of SLS rods after 20 shear stress tests (Figure 11). In contrast with the pristine sample (Figure 2), the pseudo-boehmite layer is visible but it is inhomogeneous: it clearly comes out in some zones (Figure 11a) but appears less porous and not nanostructured in others (Figure 11b). This is attributed to ice friction. Moreover, from the EDX, it is confirmed that the Dyn-50 is almost completely removed (Si content < 1 at.%).

![SEM image of SLS rod after 20 shear stress tests: the boxes evidenced (a) acicular nano layer and (b) highly deteriorated oxide layer.](image)

**Figure 11.** SEM image of SLS rod after 20 shear stress tests: the boxes evidenced (a) acicular nano layer and (b) highly deteriorated oxide layer.

### 4. Conclusions

In this study, we present an economical, environmentally friendly and fast process to synthetize a dry and fluorine-free slippery aluminum surface with ice-phobic properties. A comparison between a superhydrophobic coated aluminum (SHP) and the slippery surface (SLS) was carried out. They evidenced similar anti-icing behavior in lowering the nucleation temperature of freezing water; however, the SLS demonstrated a lower adhesion to the ice. Durability tests were carried out both on hydrophobic and ice-phobic behavior: SLS maintains its wettability properties in wet chemicals for a long time (up to 20 days) while SHP completely lost the super-hydrophobicity in few hours in all the tested environments. On the contrary, SHP demonstrated longer ice-phobic durability (up to twenty repeated shear stress tests) than SLS, which lost the ice-phobic property after 12 tests. The very good
The durability of ice-phobicity is related to the hardness of the SHP, enhanced by the sandblasting process. On the other side, the SLS evidenced a loss of coating and a deterioration of the nanostructure. SLS surfaces are promising materials because of their very good ice-phobic and anti-icing properties and remarkable durability in harsh environments even if further study is necessary in order to enhance their ice-phobic durability. A three-step process combining the sandblasting, the hydrothermal treatment and the Dyn-50 coating could lead to a better ice-phobic durability, although an increased surface area is expected to slightly enhance the adhesion between the sample and the ice. A deeper understanding of these conflicting effects (enhanced durability vs enhanced ice adhesion) is crucial to achieving a more durable ice-phobic surface.

Supplementary Materials: The following are available online at http://www.mdpi.com/2079-6412/10/2/111/s1, Figure S1: home-made equipment for shear stress measures, Table S1: Protocol for shear stress measures, Figure S2: sample treatment protocol for the durability testing in wet chemicals, Table S2: Schedule of the wettability measures after the immersion in wet chemicals, Figure S3: durability in acid solution: static contact angles, Figure S4: durability in acid solution: hysteresis, Figure S5: durability in acid solution: roll-off and sliding angles. Figure S6: durability in saline solution: static contact angles, Figure S7: durability in saline solution: hysteresis angles, Figure S8: durability in saline solution: roll-off and sliding angles.

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