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A universal, multifunctional, high-practicability superhydrophobic paint for waterproofing grass houses

Hai Zhu¹, Yu Huang^{1,2}, Shouwei Zhang¹, Shiwei Jin³, Xiaoding Lou¹ and Fan Xia¹

Abstract

Roof leakage is a common phenomenon on rainy days and makes residents uncomfortable. Superhydrophobic materials are promising candidates to protect grass houses from rainwater. However, mechanical weakness, chemical corrosion, and UV light sensitivity are the three main challenges restricting these nonwetting materials from wider application in real life. Herein, we developed an inorganic–organic superhydrophobic paint (IOS-PA) for preparing a waterproof grass house. IOS-PA not only showed mechanical robustness and chemical anticorrosion but also displayed self-healing properties, anti-icing properties, and high and low temperature (150 °C and -196 °C) resistance. Photocatalysis was also achieved with IOS-PA, as demonstrated by organic matter (Nile red, methyl blue, and methyl orange) degradation. Moreover, extremely long-term UV resistance, i.e., resistance to UV irradiation (365 nm, 5.0 ± 0.6 mW/cm^2) for 100 h and ambient sunlight for 8640 h (1 year), caused the conflicting properties of superhydrophobicity and photocatalysis to coexist in IOS-PA, further accomplishing self-cleaning for the removal of both dirt particles and organic contamination. Specifically, a grass house coated with IOS-PA exhibited favorable waterproof properties, indicating the potential to ensure comfortable living conditions for people living in undeveloped areas, even on rainy days. With a variety of excellent characteristics, IOS-PA, we believe, is advantageous for scalable production and practical application in reality.

Abbreviations

IOS-PA	Inorganic-organic superhydrophobic paint
PTFE	Polytetrafluoroethylene
NPs	Nanoparticles
PDMS	Polydimethylsiloxane
ER	Epoxyresin
PFOS	1H,1H,2H,2H-perfluorooctyltriethoxysilan
SEM	Scanning Electron Microscopes
TEM	Transmission electron microscopy
XPS	X-ray photoelectron spectroscopy
XRD	X-Ray Diffraction
WCA	Water contact angle
RA	Rolling-off angle

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MBMethyl blueMOMethyl orangeGDPGross Domestic Product

Introduction

According to *Monitoring Global Poverty*, published by the World Bank in 2017¹, extreme poverty still persists in remote and undeveloped areas in Africa and Asia. As a necessity, grass houses are the only places used for main living activities for a majority of residents^{2,3}. However, on rainy days, grass houses are easily wetted, and roof leaks bring tremendous inconveniences. Constructing waterproof grass houses is an effective approach to enable free and comfortable lives.

Due to their strong water repellency, bioinspired superhydrophobic materials have attracted wide attention⁴⁻⁹ and are anticipated to be usable for preparing nonwetting grass houses. To realize waterproof grass

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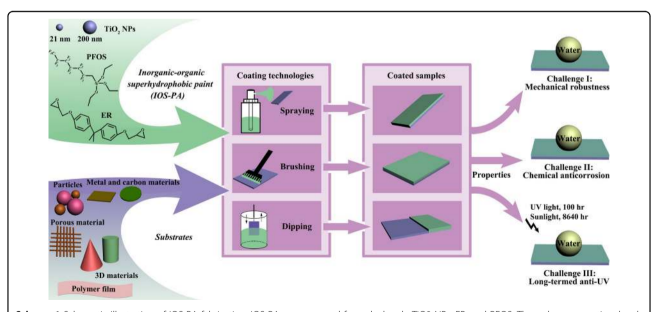
houses for real-world applications outdoors, three main challenges must be addressed: (I) mechanical weakness^{10,11}, (II) chemical corrosion¹², and (III) UV sensitivity¹³. Micro- and/or nanostructures provide surface roughness, but they wear easily under mechanical forces. To achieve mechanical superhydrophobicity, the flexible "coating + adhesive" method has been proven to be of significance and has indicated the potential for wide application^{14,15}. An all-organic water-repellent coating containing resin polymer and polytetrafluoroethylene nanoparticles (PTFE NPs) was developed that exhibited mechanochemical robustness resulting from the mechanical and chemical properties of the nanocomposite constituents¹⁶. UV light is a harmful factor for superhydrophobic materials^{17,18}. By mixing metal oxide photocatalysts with hydrophobic polymers, such as PTPE NPs¹⁹, polydimethylsiloxane (PDMS)^{20,21} and silicone nanofilaments²², (super)hydrophobic surfaces with longterm anti-ultraviolet properties have been prepared.

To face much more complex environments in practice, in addition to addressing the above three challenges, various functions, such as a large coating area^{14–16}, selfhealing properties^{23,24}, anti-icing properties^{25–27}, highand low-temperature resistance^{11,28–30}, photocatalysis^{31,32} and self-cleaning properties^{14,33–35}, are strongly advocated for these nonwetting materials. However, although the aforementioned features have been achieved on diverse substrates in recent years, the combination of all of these features on a universal superhydrophobic surface or coating has yet to be reported.

In this study, by mixing dual-sized TiO₂ NPs with epoxy resin (ER) followed by grafting 1H,1H,2H,2H-perfluorooctyltriethoxysilan (PFOS), we prepared an inorganicorganic superhydrophobic paint (IOS-PA) (Scheme 1). IOS-PA featured mechanical robustness and anticorrosion arising from the mechanochemically stable ER. Self-healing properties, anti-icing properties, high-temperature (150 °C) and liquid nitrogen (-196 °C) resistance, and self-cleaning properties were also observed for IOS-PA. Furthermore, under UV illumination, IOS-PA showed efficient photodegradation of organic dyes, while super water repellency was still maintained even under exposure to UV light $(365 \text{ nm}, 5.0 \pm 0.6 \text{ mW/cm}^2)$ for 100 h and ambient sunlight for 8640 h (1 year). With its UV-resistant water repellency, mechanical robustness, chemical anticorrosion properties, self-healing properties, resistance to low- and hightemperature damage, anti-icing properties, photocatalysis, and self-cleaning properties, our multifunctional IOS-PA indicates the potential for necessary, meaningful, and practical applications in waterproofing houses in undeveloped areas.

Experimental procedures Materials

The epoxy resin and curing agent were obtained from Beijing Oriental Yuhong Waterproof Technology Co. Ltd.



Scheme 1 Schematic Illustration of IOS-PA fabrication. IOS-PA was prepared from dual-scale TiO2 NPs, ER, and PFOS. Through spray coating, brush coating, and dip coating, the paint could be coated on multiple substrates. The superhydrophobic coating showed mechanical robustness (Challenge I), chemical anticorrosion (Challenge II), and long-term UV resistance (Challenge III) at the same time. The superhydrophobicity remained stable even under exposure to UV light for 100 h and ambient sunlight for 8640 h (1 year). Moreover, self-healing, resistance to high and low temperatures, anti-icing, photocatalysis, and self-cleaning were also observed for IOS-PA.

P25 NPs with diameters of approximately 21 nm and $\rm TiO_2$ (anatase) NPs with diameters of approximately 200 nm (100 nm to 300 nm) were purchased from Evonki Degussa and Aladdin, respectively. PFOS was obtained from Innochem. Nile red powder was obtained from Aladdin. Methyl blue and methyl orange were purchased from Chinese Medicine. Grass houses made of straw and maize leaves were purchased from Taobao Shop.

Fabrication of IOS-PA

Six grams of epoxy resin was dissolved and stirred in 30 ml of acetone for 15 min, and 6 g of TiO_2 (21 nm) and 6 g of TiO_2 (200 nm) dissolved in 150 ml of acetone containing NPs were mixed and stirred for 15 min. Then, the above two solutions were mixed, stirred, and sonicated for two 15-min cycles. Then, 1.2 g of PFOS was added to the mixed solution, stirred, and sonicated for eight 15-min cycles. Finally, 0.6 g of curing agent and 0.2 g of PFOS were added to the above solution, stirred, and sonicated for two 10-min cycles. Thus, IOS-PA was successfully synthesized. Twenty-four substrates were coated with the super-water-repellent paint by spray coating.

Durability of IOS-PA

Mechanical destruction

Sandpaper abrasion and sand impact were used to test the mechanical robustness of our coated samples. The coated samples were placed face down on sandpaper in NO. A total of 320 or 200 g of weight was placed on the surface and moved for a distance of 10 cm with a velocity of approximately 7.5 cm/s. In every cycle, the water contact angle (WCA) on the surface was measured with $10 \,\mu\text{L}$ of water drops at five different positions. For the sand impact test, 15 g of sand particles held in a funnel were dropped onto the prepared surface 50 times from a height of 15 cm, and then, the WCA was tested.

Chemical corrosion

In the chemical corrosion experiment, the WCAs and rolling-off angles (RAs) of our coating samples were tested after immersion in solutions of pH = 1 for 4 h, pH = 7 for 8 h, and pH = 14 for 8 h.

Self-healing

Treatment with O_2 plasma (1 min) and boiling water (5 min) was conducted to assess the self-healing properties of our coating. Superhydrophobicity was regained after heating in a 150 °C oven for 5 min.

Anti-icing

A water drop was placed on a fresh surface or coated surface at -10 °C. Then, the process of liquid water becoming ice was observed, and the freezing time was recorded.

Low- and high-temperature treatment

The superhydrophobic coating surface was placed in an oven at 150 °C for 100 h, and the surface WCA and RA were measured every 10 h. Two hundred milliliters of liquid nitrogen, serving as an extremely cold substance, was poured on the coated samples, and then the WCA and RA were measured after drying at room temperature for 10 repeats.

Photocatalytic activity and UV resistance

To show the photocatalytic performance, three organic dyes, Nile red, methyl blue, and methyl orange, were used with stirring at 500 rpm. Nile red was dissolved in ethanol at a concentration of $10 \,\mu\text{g/L}$. MB/water ($40 \,\mu\text{g/L}$) and MO/water (40 µg/L) were prepared before mixing with ethanol (volume, water:ethanol = 2:1). First, 0.06 g of our coating particles was added to these three solutions and then placed under UV illumination $(5.0 \pm 0.6 \text{ mW/cm}^2)$ before dark treatment for 30 min. The degradation of the dyes was judged by the disappearance of color and ultraviolet spectrophotometry. Moreover, organic dyes were dropped onto the coated samples, in which were then exposed to UV light. The colors were decomposed with almost no observable color. Our coated samples were placed under UV light $(5.0 \pm 0.6 \text{ mW/cm}^2)$ and ambient sunlight at the Xuehai Building, China University of Geosciences (Wuhan), from Nov. 1st, 2018, to Nov. 1st, 2019, to test the UV resistance.

Instrumentation

Surface nanostructures were observed by a HITACHI SU8010 field-emission scanning electron microscope (FESEM, 20 kV). The complex structures between the ER and TiO₂ NPs were demonstrated by transmission electron microscopy (Tecnai G2 20 S-TWIN). To analyze the chemical composition and surface element content of the samples, X-ray photoelectron spectroscopy (XPS, ESCA-LAB Xi + , Thermo Fisher Scientific) with an Al K α line as the excitation source was adopted. X-ray diffraction (XRD, Bruker D8 Advance) was performed using a Dmax-3β diffractometer with nickel-filtered Cu Kα radiation to distinguish the crystal forms of titanium dioxide (rutile and anatase). WCA and RA measurements were performed by a DSA100 contact angle analyzer with 10 µL water droplets at five different positions to obtain the average contact angle values. Water droplets with a volume of 10 µL and a height of 15 mm were dripped at a velocity of 0.25 m/s to show the bouncing phenomena on the coated samples. UV intensity was tested by an ultraviolet radiometer (HANDY). The concentration of Nile red in the presence of our coating and UV light was measured with an ultraviolet spectrophotometer (SHI-MADZU). By using a paramagnetic resonance spectrometer (ESR/EPR, Bruker A300), superoxide and hydroxyl

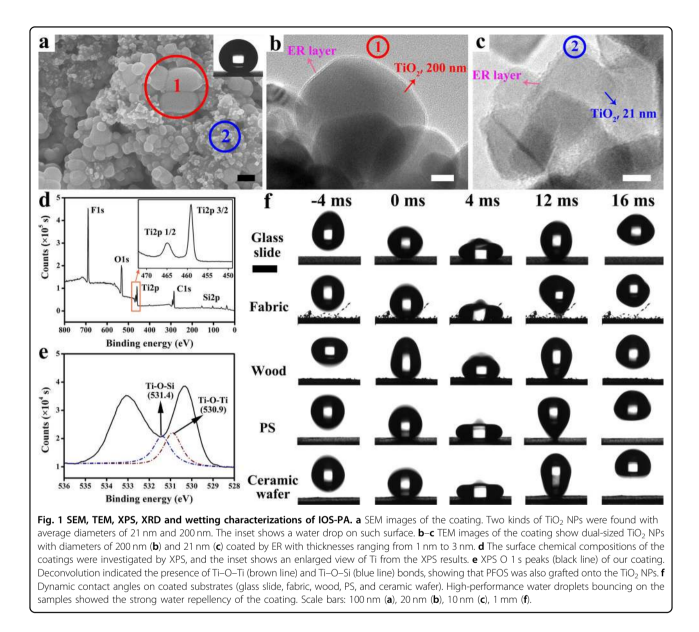
free radicals from the ER, PFOS-TiO₂ NPs, and PFOS-ER-TiO₂ NPs were studied in an ethanol solution for UV illumination for 10 min.

Results and discussion

To synthesize the superhydrophobic paint, epoxy resin/ acetone (6 g/30 ml) and dual-scale TiO₂ NPs/acetone (6 g:6 g/100 ml) were mixed, stirred, and sonicated. Subsequently, 1.2 g of PFOS was added to the above solution to decrease the surface free energies of ER and the TiO₂ NPs. Finally, 0.6 g of curing agent and 0.2 g of PFOS were added to this composite solution to achieve IOS-PA. In this work, spray coating was used to prepare superhydrophobic samples in all subsequent experiments unless otherwise specified. Various substrates were coated by IOS-PA, including "0D" particles (MnO₂, CuO, SiO₂, and sand) (Supplementary Fig. S1), 2D carbon-based substrates (fabric, wood, A4, and filter paper) (Supplementary Fig. S2), 2D silicon-based substrates (ceramic wafers and glass slides) (Supplementary Fig. S3), 2D metal substrates (Zn and Al plates) (Supplementary Fig. S4), 2D polymer films (polyvinyl chloride (PVC), polyethylene (PE) and polystyrene (PS) films and culture dishes) (Supplementary Fig. S5), 3D materials (tetrahedral, cylindrical, and circular cone shapes) (Supplementary Fig. S6), porous substrates (stainless steel mesh and sponges) (Supplementary Fig. S7) and natural leaves (maple, apricot, and loquat leaves) (Supplementary Fig. S8). The coating of a large area by IOS-PA was accomplished on a PS plate with dimensions of $0.5 \times 1 \text{ m}^2$ (Supplementary Fig. S9 and Movie S1). It has been previously proven that surface superhydrophobicity can be achieved with an optimal mass ratio of TiO_2 (21 nm): TiO_2 (200 nm) = 1:1. ^[10] Herein, a mass ratio of ER:TiO₂ (21 nm):TiO₂ (200 nm) = 1:1:1 was adopted since the coexistence of superhydrophobicity (Supplementary Fig. S10-S12), photocatalysis (Supplementary Fig. S13), and UV resistance (Supplementary Fig. S14) could be perfectly maintained (Supplementary Table S1, Fig. S15). Two kinds of TiO₂ NPs were observed by scanning electron microscopy (SEM) (Fig. 1a) and were confirmed to be coated with ER according to transmission electron microscopy (TEM) images (Fig. 1b-c). As indicated by X-ray photoelectron spectroscopy (XPS) analysis (Fig. 1d), PFOS was grafted onto ER (Supplementary Fig. S16–S17) and the TiO₂ NPs (Fig. 1e and Supplementary Fig. S16) to provide a low surface energy. Anatase and rutile TiO₂ NPs in our coating were also demonstrated by X-ray diffraction (XRD) (Supplementary Fig. S18). Thus, the textured nanostructures and low surface energy of IOS-PA endowed the substrates with superhydrophobic properties, displaying high WCAs >150° and low RAs <10° (Supplementary Fig. S19 and Movie S2). To demonstrate water droplet bouncing phenomena and dynamic contact angles, five kinds of coated substrates (glass slide, fabric, wood, PS, and ceramic wafer) were utilized. Due to the air films formed over rough structures^{36–38}, water droplets (release height = 15 mm, $\nu = 0.25$ m/s) completely left the surfaces without wetting (Fig. 1f). Moreover, the coated surface repelled the impact of water drops released from a height of approximately 80 cm with different dripping velocities ($\nu = 0.25$ m/s, 1 m/s, 2 m/s, and 3 m/s) (Supplementary Movie S3). Although raindrops with a velocity of approximately 9 m/s hit our coated surface, strong water-repellent properties were still observed (Supplementary Figs. 20–21, Movie S4).

Durable superhydrophobic materials are required for practical applications. Mechanical robustness, chemical anticorrosion properties, self-healing properties, antiicing properties, and high- and low-temperature resistance functioned well on IOS-PA. To display the mechanical stability of IOS-PA, sandpaper abrasion and sand impact approaches were applied. A coated glass slide was placed face down on sandpaper (No. 320) (Supplementary Fig. S22), a weight of 200 g was placed on the reverse side, and then a force was used to push the coated surface and weight for a distance of 10 cm with a velocity of approximately 7.5 cm/s (Supplementary Fig. S23–S24). After 50 repeats, the WCA of the sample was still larger than 150° (Fig. 2a), showing abrasion-resistant superhydrophobicity. Regarding sand impact destruction, 15 g of sand with diameters ranging from 0.5 mm to 2 mm fell from a height of 15 cm to hit the surface (Supplementary Fig. S25–S26), and super water repellency of the samples was sustained with WCAs > 150° after 50 cycles (Fig. 2b). Upon mechanical damage, the original surface morphology changed but still had textured structures, thus offering sufficient roughness to ensure super-nonwetting behavior, mainly due to the favorable adhesive and mechanically robust properties of ER^{14,16,39-41}. In addition to the glass slide-based superhydrophobic surface, the coated fabric, PS, wood, and ceramic wafer substrates exhibited stable superhydrophobic behavior after sandpaper abrasion and sand impact.

After storage in acidic (pH = 1) solution for 4 h and saline (pH = 7) and alkaline (pH = 14) solutions for 8 h, high WCAs and low RAs remained on coated samples (Fig. 2c–e and Supplementary Fig. S27). Thus, IOS-PA exhibited stable water repellency against chemical corrosion, originating from the inherent chemical inertness of $ER^{16,39-41}$. Moreover, low-surface-energy compounds on nonwetting materials are often sensitive to outside stimuli, and thus, self-healing water-repellent materials are required^{23,24}. IOS-PA manifested superior self-healing ability against oxygen plasma (Fig. 2f and Supplementary Fig. S28) and boiling water vapor treatments (Supplementary Fig. S29). Self-healing proceeded for 10 cycles of these treatments with no obvious morphology variations,



and super water repellency could be regained by treatment at 150 °C for 5 min due to the thermostable properties of the material (Fig. 2g and Supplementary Fig. S30). IOS-PA also featured favorable anti-icing properties, attributed to the thermal barriers on the textured nanostructures and the small contact region between the surface and liquid²⁶. Taking the glass slide as an example (Supplementary Fig. S31a), it took approximately 30 s for a water droplet (volume: $10 \,\mu$ L) to freeze into ice on the bare surface (-10 °C) (Supplementary Movie S5). However, for the superhydrophobic sample under the same conditions, the freezing time was 120 s (Supplementary Movie S6). Additionally, ice formation delays were observed on the other four coated samples, i.e., fabric, PS, wood, and the ceramic wafer (Supplementary Fig. S31b). An ice cube easily rolled away from the surface at room temperature (Supplementary Fig. S31c). Notably, even when the coated filter paper (Supplementary Movie S7) and sponge (Supplementary Movie S8) were exposed to extremely cold liquid nitrogen (LN), they still manifested strong water repellency (Supplementary Fig. S32). The WCA and RA on the surface remained almost unchanged after liquid nitrogen treatment (Fig. 2h), indicating outstanding resistance of the composite coating to low temperature.

Under UV illumination, excited superoxide and hydroxyl free radicals from TiO₂ NPs can degrade organic substances (Fig. 3a–b and Supplementary Fig. S33)^{18,19,31,32}. To demonstrate the photocatalytic properties of IOS-PA, Nile red solutions without and with our coating (PFOS-ER-TiO₂ NPs) were placed under UV irradiation (365 nm, $5.0 \pm 0.6 \text{ mW/cm}^2$). As illustrated in Supplementary Fig.

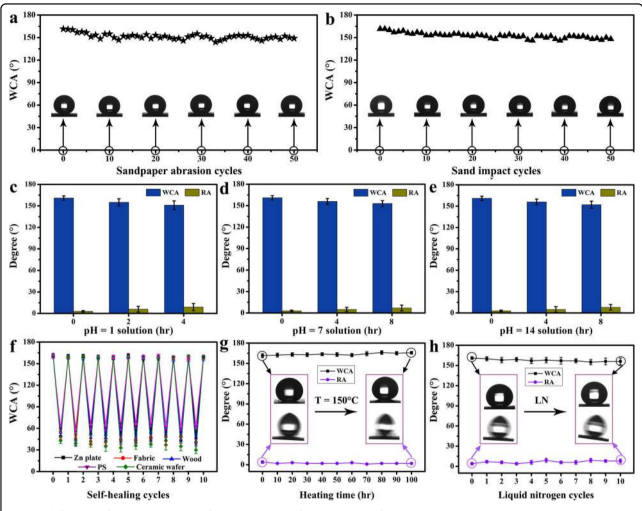
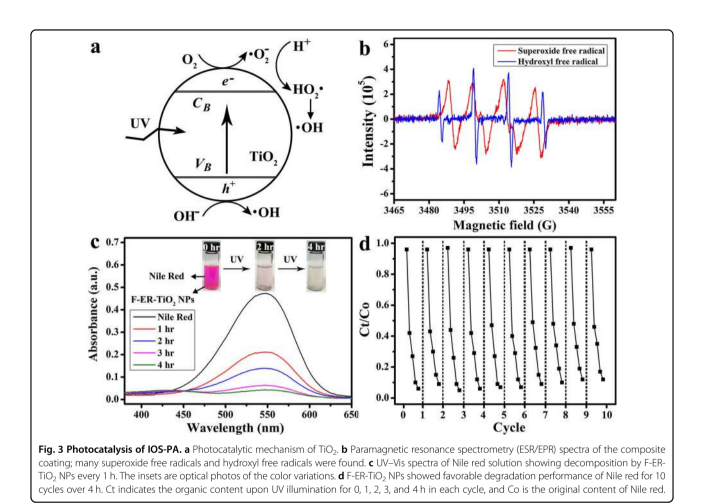


Fig. 2 Multifunction of IOS-PA. a The WCAs of the paint-coated surfaces were tested after each abrasion cycle, and stable superhydrophobicity was obtained with almost all WCAs larger than 150°. **b** After sand impact for 50 cycles, the WCAs of the coatings remained high, also showing super water repellency. When placed in pH = 1 (**c**), pH = 7 (**d**), and pH = 14 (**e**) solutions for 4, 8, and 8 h, respectively, the coating still manifested super water repellency with high WCAs and low RCAs. **f** The WCAs on five coated samples (glass slide, fabric, wood, PE film, and ceramic wafer) changed upon oxygen plasma treatment and heating in an oven. After plasma treatment, the surface changed from superhydrophobic to hydrophilic, and then superhydrophobicity was obtained again after treatment at 150 °C for 5 min in an oven. The self-healing cycles could be repeated ten times. **g** Super water repellency was still obtained even after treatment at 150 °C for 100 h. **h** Coated paper was treated with extremely cold liquid nitrogen (LN) and dried at room temperature. The WCA and RA of the coated fabric were tested after 10 cycles of liquid nitrogen treatment.

S34, the bare Nile red solution did not show obvious color variation, and a high chemical concentration was still observed under UV irradiation. For PFOS-ER-TiO₂ NPs, the red dye was almost completely decomposed in 4 h, as further proven by the UV–Vis spectra (Fig. 3c). After 10 repeats, efficient degradation of Nile red was still stably achieved (Fig. 3d). Moreover, eight other kinds of coated samples were used to show photocatalytic activity, i.e., fabric, filter, A4 paper, a culture dish, PE, PVC and PS films and natural loquat leaf; all Nile red-contaminated samples remained clean after UV irradiation (Supplementary Fig. S35). Even in ambient sunlight, organic dye decomposition was observed on the coated substrates (Supplementary Fig. S36). In our composite coating, this

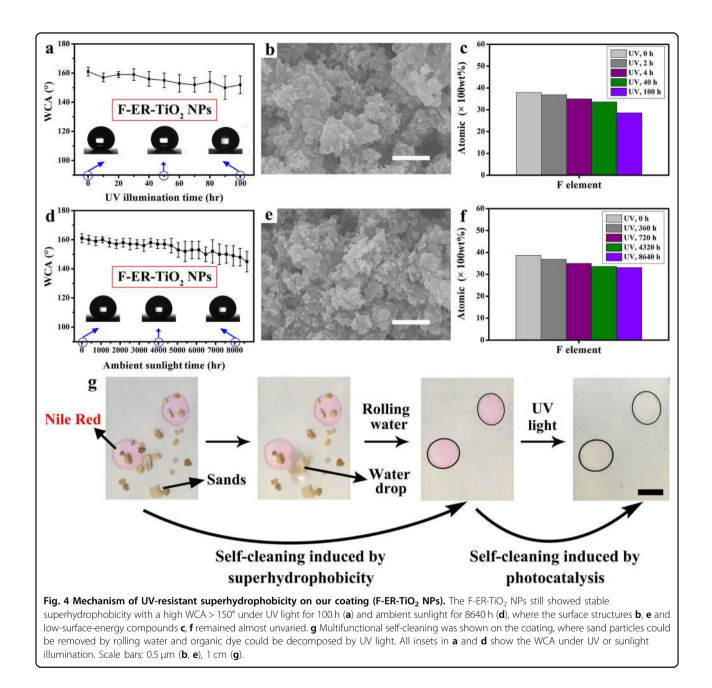
high-performance photocatalytic effect was obtained solely because of the existence of dual-sized TiO₂ NPs (Supplementary Fig. S34 and S37) since ER showed no photocatalytic ability (Supplementary Fig. S38). In addition to Nile red decomposition, methyl blue (MB) and methyl orange (MO) were degraded by IOS-PA (Supplementary Fig. S39–S40). The coated surfaces showed robust superhydrophobic properties when exposed to mechanical destruction, chemical corrosion, and liquid nitrogen. Here, when subjected to the same damage, favorable photocatalytic activity was still observed for our coating (Supplementary Fig. S41).

In the outside environment, long-term sunlight illumination often damages superhydrophobicity because



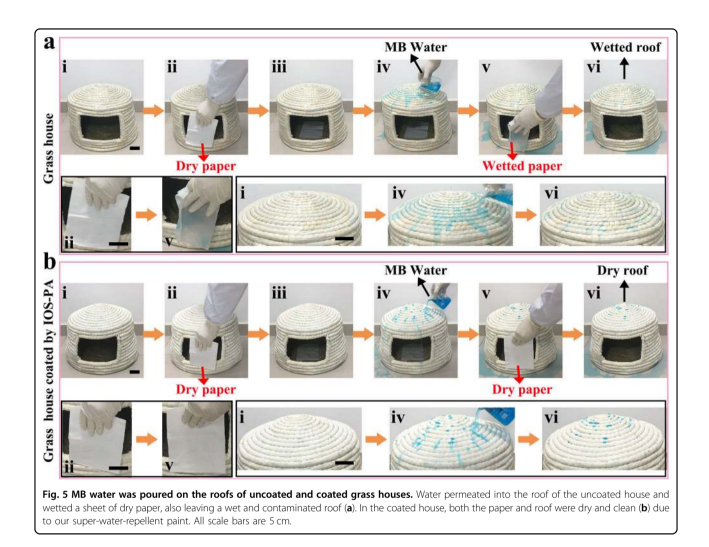
interfacial compositions and structures are sensitive to UV light; thus, UV-resistant superhydrophobicity is required for outdoor applications. Specifically, for photocatalysts, (super)hydrophobicity could not be achieved for a long time since the low-surface-energy compounds that were used to reduce surface energy were easily degraded by excited free radicals under UV light^{17,18}. Therefore, water-repellent photocatalysts often require dark storage or protection from light. For our coating, superhydrophobicity with a high WCA > 150° and low RA < 20° (Supplementary Fig. S42) was achieved under UV irradiation (365 nm, $5.0 \pm 0.6 \text{ mW/cm}^2$) for a period of 100 h (Fig. 4a) with unaffected surface structures (Fig. 4b) and a high content of F (Fig. 4c). When the coated samples were exposed to ambient sunlight for as long as 8640 h (1 year), they also possessed strong water repellency (Fig. 4d and Supplementary Fig. S42), resulting from their stable surface roughness (Fig. 4e) and chemical composition (Fig. 4f). In IOS-PA, fluorosilane was grafted onto ER and the TiO₂ NPs to obtain a low surface energy (Supplementary Fig. S43), which combined with the sufficient roughness derived from the dual-sized TiO₂ NPs to achieve super-nonwetting properties. However, for TiO₂ NPs directly modified with PFOS (PFOS-TiO₂ NPs), superhydrophobicity changed into superhydrophilicity under both UV light and sunlight illumination (Supplementary Fig. S44a–S44c and S45, Table S2) because PFOS was decomposed by the TiO₂ NPs. However, fluorinated ER (PFOS-ER) with hydrophobicity (Supplementary Fig. S46) exhibited almost unchanged wetting behavior (Supplementary Fig. S44d) even in the presence of UV light and TiO₂ NPs, where almost no obvious variations were found in the surface structure (Supplementary Fig. S44e and S46) or chemical composition (Supplementary Fig. S44f). Hence, PFOS on ER was protected from superoxide and hydrogen free radicals excited by TiO₂ NPs, which contributed to the durable low surface energy of IOS-PA and thus long-term super water repellency.

Lotus-inspired superhydrophobic surfaces^{4,10,14,33–35} and (super)hydrophilic metal oxide photocatalysts^{19–22} are two kinds of self-cleaning materials. The first kind of surface can remove dirt (sand, dust, particles, etc.) with the aid of strong water repellency. The latter has been applied to decompose organic contamination by excited free radicals. However, in the presence of both dirt and organic contamination, the above self-cleaning surfaces



did not perform adequately. Fortunately, the coexistence of superhydrophobicity and photocatalytic properties induced by long-term UV resistance is useful for sustaining a clean surface under such double-polluted conditions. Here, self-cleaning for lotus-inspired dirt removal and organic dye degradation were realized in our coating (Fig. 4g). In Nile red-contaminated areas, super water repellency was also found (Supplementary Fig. S47), where sand with sizes ranging from 0.5 mm to 2 mm was freely removed by rolling water droplets (Supplementary Movie S9). Then, organic contaminants were degraded under UV irradiation. Therefore, IOS-PA-coated samples can always remain clean on account of these two kinds of self-cleaning mechanisms.

On rainy days, grass houses are easily wetted, causing considerable trouble for residents in undeveloped regions. Here, we used IOS-PA to coat a simulated grass house (Supplementary Fig. S48). To assess the nonwetting performance, clean and dry papers were placed in grass houses without and with coated IOS-PA (Supplementary Fig. S49). When MB water was poured on the uncoated grass house (Fig. 5a, Supplementary Movie S10), the



original dry paper became wetted, resulting in a wet and MB-contaminated roof. In contrast, because of the strong water repellency of IOS-PA (Fig. 5b, Supplementary Movie S11), the paper always remained in the dry state, and a waterproof roof was observed.

The costs of the components (ER, curing agent, TiO_2) (21 nm), TiO₂ (200 nm), PFOS, and acetone) of IOS-PA are shown in Supplementary Table S3. Eighty milliliters of IOS-PA is sufficient to coat a substrate with a surface area of 1 m², and the cost per unit area for our paint is approximately $2.70 \text{ }^{\text{s}}/\text{m}^2$, or $17.46 \text{ }^{\text{s}}/\text{m}^2$. In addition, the cost of a grass house is approximately \$20. For a house roof with an area of 100 m^2 , the total cost to accomplish waterproofing is approximately 290 (270 + 20). As reported by the World Bank, the most poverty-stricken country possesses a per capita GDP of approximately \$303. Therefore, our reported IOS-PA is affordable for residents in poverty-stricken regions. Furthermore, with its mechanochemical robustness, self-healing properties, high- and low-temperature resistance, anti-icing properties, photocatalytic properties, ultralong UV resistance, and self-cleaning properties, the application of IOS-PA to grass houses exhibits promising and wide potential for real-world applications.

Conclusion

Multifunctional IOS-PA was fabricated by incorporating dual-sized TiO₂ NPs, ER, and PFOS together, showing superior mechanical robustness and chemical stability. Moreover, self-healing properties, high- and lowtemperature (150 °C and -196 °C) resistance, and antiicing properties were also observed for our IOS-PA. Photodegradation of organic matter (Nile red, MB and MO) was observed on IOS-PA. Moreover, the conflicting properties of photocatalysis and superhydrophobicity were achieved in our paint with long-term functionality, resulting in ultralong UV resistance under UV irradiation $(365 \text{ nm}, 5.0 \pm 0.6 \text{ mW/cm}^2)$ for 100 h and ambient sunlight exposure for 8640 h (1 year). Moreover, IOS-PA showed self-cleaning properties with both dirt removal and organic decomposition. Based on these advantages, grass houses coated with IOS-PA also manifested

favorable waterproof performance in real life. We believe the reported IOS-PA has the potential for practical production in the near future, as well as daily use for waterproofing items such as roofs, clothes, umbrellas, and shoes.

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Author contributions

H.Z., Y.H., and F.X. conceived and designed the overall experiments. H.Z. and S. Z. fabricated the IOS-PA. H.Z. and Y.H. conducted the durability tests. H.Z. and S.J. assessed the XPS data, EPR data, and photocatalysis performance. H.Z., Y.H., X.L., and F.X. wrote and revised the paper. All authors reviewed the manuscript.

Conflict of interest

The authors declare no competing interests.

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