Micro/Nano hierarchical peony-like Al doped ZnO superhydrophobic film: The guiding effect of (100) preferred seed layer

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In this communication, we present a versatile and controllable strategy for formation of superhydrophobic micro/nano hierarchical Al doped ZnO (AZO) films with a water contact angle (CA) of 170 ± 4°. This strategy involves a two-step layer-by-layer process employing an atomic layer deposition (ALD) technique followed by a hydrothermal method, and the resulting novel AZO surface layer consists of (100) dominant nano-rice-like AZO seed layer (the water CA of 110 ± 4°) covered with micro-peony-like AZO top. The growth mechanisms and superhydrophobic properties of the hierarchical AZO layer are discussed. It is believed that the present route holds promise for future success in the design and development of practical superhydrophobic materials.

Superhydrophobic materials and coatings with water contact angle larger than 150° have been extensively investigated due to their wide applications from housewares to industrial products1-2. For instance, solar cell panels, goggles and windows for photovoltaic devices generally demand their surfaces with properties of self-cleaning, anti-fog and superhydrophobicity3. The key points of superhydrophobic states include the surface energy of the chemical composition and the geometrical rough structure of solid surfaces4. Owing to the lower surface energy, the organic perfluorinated materials turn into the most common hydrophobic material, however those organic coatings are costly and unstable for the application. Therefore, increasing attentions are drawn on some inorganic materials with specific hierarchical rough surfaces5-6.

Al-doped ZnO (AZO) materials have attracted considerable attentions owing to their unique optical and electrical properties, such as wide band gap and high exciton binding energy7-9. AZO film can provide valence ions and higher carrier mobility, making it a desirable candidate for various photovoltaic and solar cell applications10-11. Therefore, the superhydrophobic AZO intelligent materials are potential and promising in the photovoltaic fields.

To fabricate the required superhydrophobic surface roughness of AZO film, various methods have been proposed in literatures. A two-step layer-by-layer approach were adopted in several research groups12,13. AZO seed layer was deposited on a substrate by solution method, sol-gel method, magnetron sputtering methods and others, which can lower the activation barrier for the following nucleation and enhance the film adhesive force with top layer. Micro-nano AZO structures were fabricated subsequently using the hydrothermal method, plasma etching, and vapor deposition on the seed layer14,15. It is worth fabricating the special and micro/nano hierarchical AZO structure surface, by enhancing the surface roughness, so that superhydrophobic AZO films materials can be achieved, without involving the perfluorinated material.

In most research works, the surface of ZnO micro-nanorods arrays grown based on (002) preferred seed layer is still hydrophilic, and thus the fluorination method needs to be used for formation of the superhydrophobic surface texture16. In contrast, other orientation dominant seed layer tends to induce more complex micro nanostructure for superhydrophobic surface17. In this case, the seed layer obviously plays a subtle guiding role in controlling

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the micro-structure for the subsequent growth process. Therefore, the growth method, orientation and guiding
effect of seed layer should be investigated to further discuss the morphology and the superhydrophobic feature
of the generated AZO film.

Herein, we present a facile strategy to fabricate superhydrophobic AZO film using an atomic layer deposition
(ALD) technique and hydrothermal methods. In our well-designed process, at first, nano rice-structured (100)
preferred AZO thin layer with a thickness of ~20 nm in-situ grows on a glass substrate using the ALD technique.
Subsequently, the micro-flower-structure AZO layer continuously grows on the seed layer using the hydrothermal
method. The resulting micro-nano hierarchical structure of AZO film is superhydrophobic and nearly transparent
without any further treatment. Our approach is much simpler and more feasible than the reported ones using
the expensive fluorochemicals to generate superhydrophobic surfaces. Owing to the flexibility of the ALD
technique, the facile two-step strategy introduced here is a promising candidate in the photoelectrical applica-
tions such as flexible photovoltaics and displays.

Results and Discussion

The strategy for preparing the hierarchically structured superhydrophobic AZO thin films is schematically pre-
sented in Figure S1. The AZO seed layer was formed on the silica and glass substrates through deposition using
the ALD method. Figure 1a,b show the 2D and 3D AFM images of AZO seed layer film with the thickness of
20 nm on the silica substrate. The seed layer consists of the arrays of uniform nano-rice-like protuberances with
acicular points, showing the gain size and roughness (rms) values of 10 nm and 3 nm, respectively. The seed layer
should be thick enough to allow subsequent growth of homogenous AZO crystal by the hydrothermal method.
In this study, we controlled the thickness of the seed layer around 20 nm, which presents good stability for the
subsequent treatment and high transparency. The novel AZO seed layer with the nano-rice array morphology
on the glass substrate shows hydrophobic against a spherical water droplet with a water CA of 110 ± 4°. Similar
features have been found in nature, such as lotus leaves and the cicada’s wings19. The transparency of the seed layer
is very close to the bare glass substrate (Insert Fig. 1a). The transmittance of the AZO 20 nm seed layer is about
95% (Figure S2).

The survey XPS spectrum of the AZO seed layer is shown in Fig. 1c, which shows the representative peaks of
Zn and O. Figure 1d–f demonstrate the high resolution XPS spectra of the Zn 2p, O1s, and Al 2p core levels and
their Gaussian-resolved results for the AZO seed, respectively. The Gaussian-resolved result for Zn 2p spectra in
Fig. 1d shows two main peaks centered at 1020.68 eV and 1043.78 eV, which were assigned to Zn 2p3/2 and Zn
2p1/2 states respectively19. The Gaussian-resolved result for O1s spectra in Fig. 1e shows two components of oxygen
varying in chemical states which are different in binding energy. The lower binding energy component, cen-
tered at 529.8 eV, is attributed to O2− ions on the wurtzite structure of hexagonal Zn2+ ion array20. Meanwhile, the
high binding energy component located at 531.1 eV is associated with O2− in the oxygen deficient regions within
the matrix of ZnO21. Figure 1f indicates the presence of Al3+ in the seed layer according to the Al 2p peak, which
can be Gaussian-resolved with two peaks centered at 74.08 eV and 74.8 eV. The 74.08 eV obviously suggested that
the Al3+ had substitutionally incorporated into Zn2+ sites in the seed layer. In addition, the higher energy 74.8 eV is associated
with the stoichiometric Al2O3, indication of generation of Al2O3 within the ALD growth process22. It can be found
that the atomic concentration of Al is 3.43% for the AZO seed layer. In addition, the distribution of Al dopants
in the surface and the whole sample were also investigated using energy-dispersive x-ray spectroscopy (EDS) in
Figure S2 and inductively coupled plasma atomic emission spectroscopy (ICP-AES), respectively. The Al atomic
dopants of 3.41% and 3.01% had been obtained for the surface layer and longitudinal direction in the ALD AZO
film. The Al dopant concentration for the whole sample in ICP-AES was slightly lower than that in the surface in
the XPS and EDS analysis. This behavior can be attributed to the ALD AZO film growth process, in which each
cycle ended up with 19 Zn: 1 Al: O sequence. Therefore, the Al dopant was more abundant in the surface layer
and the similar results have also been obtained by other groups23.

The seed layer with specific nanostructures, excellent transparency and ion doping modification were suc-
cessfully obtained in the first step. Then, upper AZO micro-flower thin layers were fabricated via the short time
hydrothermal method.

Figure 2a,b are SEM top-images of the hierarchical AZO hydrothermal film at low and high magnifications. It
can be seen that the diameters of AZO micro-flowers are about 2 ~ 3 μm, showing a uniform peony-like structure
with dense and thin nano-petals. The unique micro/nano flower-like AZO layer presents excellent superhydro-
phobicity and good transparency, as indicated by the clear letters underneath the glass slide coated with this layer
(Insert Fig. 2b). Figure S3 illustrates the transmittance spectra of the AZO seed layer and the hydrothermal layer.
The average transmittance values of the AZO hydrothermal layer are about 80%, which is suitable for the practical
optical device. The optical band gap (Eg) of the AZO seed layer and the hydrothermal layer were estimated by
the Tauc relation (αhv)(2) = B(hν − Eg). Where α, h, and Eg is the optical absorption coefficient, the photon energy
and the energy gap, respectively. Based on the transmittance spectra, the (αhv)(2) versus hν graph was plotted in
Figure S4, in which a straight line is fitted for the straight region. The extrapolation of this straight line to x axis
gives the value of the band gap for AZO films. The results indicate that AZO seed and hydrothermal layers both
exhibit higher Eg values ~3.7 eV than undoped ZnO (3.36 eV), which is attributed to Moss-Burstein effect, caused
by an increase in free electron concentration due to Al doping24.

The micrographs of the hierarchical AZO film were taken by TEM and high-resolution TEM (HRTEM).
Figure 2c depicts bright field TEM and HRTEM images of certain nano-petals of the AZO flower. From the
HRTEM observation, it can be found that there are AZO polycrystalline grains with the distances of crystalline
planes of 0.28 nm (Insert Fig. 2c), which are corresponding to the interplanar spacing of (100) planes of AZO. The
selected area electron diffraction (SAED) image is shown in Fig. 2d. The presence of diffused rings and regular
spots further confirms the above results.
Besides the transparency and surface morphologies, the crystallographic structure and dominant orientation of the seed layer are significant and critical for guiding growth of AZO top layer during the second step. Owing to the island-like growth mechanism of AZO during the ALD process, heat treatment was not adopted. The crystallographic structures of the as-prepared AZO seed layer (lower) and the AZO hydrothermal film (upper) were characterized by using XRD, as shown in Fig. 3a. The results indicate that the AZO seed layer possesses a wurtzite crystal structure according to the standard data (JCPDS, No. 99–1111) with (100) preferred plane, different from (002) dominant ZnO thin layer prepared using other methods like the sol-gel and magnetron sputtering methods. There are no extra peaks related to aluminum metal and those from other zinc aluminum phase, indicating that the Al ions substituted the Zn sites without changing the wurtzite structure. In the second hydrothermal step, the AZO top layer sequentially grows and evolves restrictedly guided by the (100) crystallographic planes of the former seed layer. Because of the different hierarchical structures and surfaces tensions of the two layers, the diffraction peaks of the AZO hydrothermal film were slightly shifted compared with those of the AZO seed layer (Insert Fig. 3a).

It can be seen that the (002) peak of the AZO hydrothermal film can be decomposed to two peaks at 34.43° and 34.53° by Gaussian fitting. As shown in Fig. 3b, 34.43° is the typical (002) diffractive peak of ZnO wurtzite
crystal. The (002) peak splitting at 34.53° can be inferred the lattice disorder of the AZO crystal. The morphology of the peony-like AZO structure with dense and thin nano-petals (Fig. 2a) is a possible reason for the disorder of crystal structure27.

The chemical states of the compositional elements in AZO hydrothermal film were investigated by the XPS and the survey and representative Al 2p spectra are shown in Figure S5. The XPS results reveal that the alumina concentration was equal to 3.08% after hydrothermal process.

The formation mechanism of the unique micro/nano flower structure of AZO film was proposed as below. Normally, because (002) polar plane of ZnO has the highest atomic packing density, and thus c axis is the thermodynamically favorable growth direction of the ZnO with wurtzite structure under normal crystallization conditions. Therefore the aligned nanorods array structure was common micro-topography for the ZnO or AZO material (Fig. 4a). The preferred orientation and the guiding effects of the AZO seed layer play a significant role during the process of growth and evolution of the followed AZO film. In this work, the (100) dominant plane of the AZO seed layer is special and critical factor for the subsequent growth. In fact, orientation control of ALD thin film growth is much complex and affected by many factors such as pressure, precursor and temperature etc28.

In ALD method, methyl and ethyl groups have been involved in AZO film growth process (Fig. 4b). Due to the adsorption of the ethyl groups on (002) crystallographic plane of AZO, the c-axis crystal growth of AZO may be suppressed17,29. Furthermore, the methyl groups aggravate such suppression on (002) plane and thus reinforce the preferential growth of (100) crystallographic plane. As a result, the as-prepared AZO seed layer using the ALD technique exhibits (100) dominant plane, which is in agreement with the XRD patterns of the seed layer. In the process of ALD AZO film growth, methyl and ethyl groups play a role in suppression for the c-axis crystal of AZO, but will totally be replaced by H2O in every reaction cycles and then are removed by the purge gas. In order to confirm the results, the FT-IR measurements at the middle and far infrared wave bands has been adopted to explore the bonding features in the resulting ALD AZO film. Figure 5a presents the FT-IR spectrum of AZO seed layer at middle infrared range. The bands at 3430 and 1630 cm⁻¹ were assigned to the stretching and bending vibration of surface -OH groups. The peak at 434 cm⁻¹ and the weak band observed at 1075 cm⁻¹ corresponds to

Figure 2. FE-SEM top-images of the as-prepared AZO films at high magnification (a) and low magnification (b). Inset in (b) is digital image of AZO films on glass substrate. (c) TEM image and high-resolution TEM image (insert of Fig. 2c,d) selected area electron diffraction (SAED) of AZO petal.
Figure 3. (a) X-ray diffraction of the AZO seed layers and the AZO hydrothermal films; Inset shows the zoom-in view of 31–37 deg. (b) The magnification and Gaussian fitting results of (002) diffractive peak of the AZO hydrothermal films.

Figure 4. Schematic of plausible mechanism of (100) dominant planes in the seed layer and the formation of AZO architectures for (a) AZO nanorods arrays and (b) AZO flower-like structure.
the Zn—O and Al-O bond stretching vibration. In the spectrum, the C-H stretching mode of methyl and ethyl group in the region around 2900 cm$^{-1}$ is not observed. In order to obtain some information about AZO seed layer, the FT-IR spectrum of AZO film at the far infrared range is shown in Fig. 5b, where the sharp Zn—O peak also can be obtained at 434 cm$^{-1}$. The significant spectroscopic bands at 643, 595, and 461 cm$^{-1}$ appear, which are identified to be the characteristic absorption bands of Al-O. The bands at 505 and 372 cm$^{-1}$ were assigned to the oxygen deficiency in ZnO and the Si-O bond of the substrate material, respectively.

According to the synergistic reaction mechanism of the precursors, a seed layer with the desired dominant orientation can be obtained controllably, which further guides the self-assemble growth of the AZO top layer using the hydrothermal method. Actually, the nanorods array structure and nano flowers were two kinds of typical morphology for the hydrothermal AZO, which can be obtained under condition of the low and high growth temperature, respectively. In ZnO wurtzite structure, each Zn$^{2+}$ ion is surrounded by four O$^{2-}$ ions and vice versa. The asymmetric distribution of the zinc and oxygen atoms along the c axis determines the positive and negative sides of the crystal. The (002) plane is the positive side, which is the exposed surface of zinc atoms, and (00–2) plane is the negative side with the exposed surface of oxygen atoms. The growth units Zn(OH)$_4^{2-}$ were negatively charged, which should be easily superimposed on the positive side of ZnO crystals. When the temperature of reaction was low, HMTA was able to slowly release OH$^-$ groups, thus making the low-concentration of OH$^-$ assist in forming nanorods easily and the OH$^-$ hardly affect the morphology. However the higher hydrothermal temperature speeds up the releasing rate of OH$^-$ groups from HMTA, and more OH$^-$ groups can dissolve the (002) plane of ZnO crystal, then the ZnO nanosheets and flower begin to form.

In the hydrothermal growth, for the lower temperature around 95 °C, the nano-rods array AZO structure had been fabricated and presents hydrophobicity with CA ~ 130° (Figure S6). The (002) dominant planes generally lead to the nanorods array structure along [0001] direction, and resulting film still needs the further fluorination method to build the superhydrophobic surface. In contrast, the AZO nucleates and assembles along (100) planes instead of (002) planes, and thus the unique complex flower-like superhydrophobic structures can be formed around 140 °C.

Additionally, HMTA concentrations in solution are an important factor for the hierarchical structure in the hydrothermal process. In this present work, a 1:1 ratio of zinc nitrate hexahydrate (Zn(NO$_3$)$_2$·6H$_2$O) and hexamethylenetetramine (HMTA) were employed in the reaction nutrient solution to meet the reaction requirement. Four different HMTA concentrations 25 mM/L, 50 mM/L, 75 mM/L and 100 mM/L were used to participate the AZO hydrothermal structure growth (Figure S7). The results indicate that high HMTA concentrations will increase the nucleation rate of the AZO crystal grains. Smaller contact angle ~140° was obtained for lower HMTA concentrations of 25 mM/L. On the other hand, lower optical transmittance ~60% happened in higher

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**Figure 5.** The FT-IR at thin film reflecting measurement mode spectra of the AZO ALD seed layer at middle infrared range (a) and at far infrared range (b).
concentrations of 75 mM/L and 100 mM/L. For the balance of the superhydrophobic and transparent properties of the films, 50 mM/L were chosen as optimal concentrations of HMTA.

The schematic pictures of the AZO seed layer and the hydrothermal film with hydrophobic properties are shown in Fig. 6. The CA value of 110 ± 4° was obtained for AZO seed layer, which can be attributed to the lower surface energies of ALD film. Owing to the peony-like morphology with a combination of micro-sized flowers and petals in nano-sized, the AZO hydrothermal film displays the excellent superhydrophobic property with the CA of 170 ± 4° without any other treatment, which is better than other reported values of 158° and 160° using fluoridation method on inorganic materials5,16.

The different hydrophobic properties between the AZO seed layer and the AZO hydrothermal film can be explained by classical wetting theories34. The hydrophobic property of rice-like AZO seed layer was a good example of Wenzel model (Fig. 6a), in which liquids are assumed to completely fill the grooves of rough surfaces, and the apparent contact angle $\theta_W$ follows equation: $\cos \theta_W = r \cos \theta_Y$ ($\theta_Y$ is the contact angle, and $r$ is the surface roughness factor, defined as the ratio between the actual surface area and the apparent surface area of the rough surface).

On the other hand, the polar (002) planes are metastable and have high surface energy, while the nonpolar (100) planes are stable and have lower surface energy, which is beneficial to the fabrication of the superhydrophobic surface. According to the experimental results and theories models, the high ratio of (100)/(002) AZO hydrothermal film with the micro-sized flowers and nano-sized petals presents an excellent superhydrophobic property, which could be a model for future design and development of the superhydrophobic surfaces.

The durability of the superhydrophobic film is important for the practical applications. The CA of the AZO hydrothermal film was measured every month after storage under ambient condition for a period of time. The superhydrophobicity of the film remained excellent, as indicated by the constant CA values for 5 months. The durable superhydrophobicity results from the stable chemical composition and structure of the inorganic material. The studies on the superhydrophobicity of the AZO films with different hierarchical structures are ongoing and the results will be reported soon.

Conclusion

A simple, inexpensive and controllable two-step layer-by-layer process using the ALD and hydrothermal methods was applied to fabricate transparent and durably superhydrophobic (CA = 170 ± 4°) inorganic micro/nano hierarchical films. The bottom homogeneous AZO seed layer controlled the subsequent growth of AZO film along (100) preferred planes with micro/nano scale peony-like structure. This is the first report about fabrication of the transparent and superhydrophobic coatings through the guiding effect of the ALD seed layer instead of using the expensive fluorinated reagents. Due to the controllable crystalline phase and special morphology of the AZO layer obtained by the ALD and hydrothermal methods, the current strategy is suitable for numerous other inorganic oxide materials and substrates with special shapes, and promises a potential foreground in various areas.
Methods

Preparation of AZO seed layer by ALD. Substrates were cleaned by acetone, alcohol and distilled water, respectively. An ALD system (TALD-150 Kmerico Jiaxing) was employed to deposit AZO films on substrates. DEZn, TMAl and distilled water (H2O) were used as precursors of Zn, Al, and O, respectively. DEZn, TMAl and H2O were fed into the chamber through separate inlet lines and purged by N2 gas in ALD cycles. The pulse sequence of ZnO was DEZn - N2 purge - H2O - N2 purge, and the dose time and purge time was 0.02-0.015-25 s. The Al doping sequence in ZnO was DEZn - N2 purge - TMAl - N2 purge - H2O - N2 purge, and the dose time and purge time was 0.01-0.02-0.015-25 s. The sequence ratio of Zn:Al was 19:1 and the growth rate of AZO is 1.5 Å per cycle at 150 °C and 0.15 Torr.

Preparation of AZO micro flower structure composite film via hydrothermal method. The substrates with AZO seed film were placed flat on the bottom of a 50 mL Teflon-lined stainless steel autoclave. 0.5652 g Zn (NO3)2·6H2O, 0.0375 g Al (NO3)3·9H2O (5% atomic concentration doped) and 0.2668 g C6H12N4 (HMTA) were dissolved in 40 mL of deionized water and poured into the autoclave. Hydrothermal synthesis was carried out at 140 °C for 4 h. The autoclave was cooled down to room temperature and the sample was washed repeatedly with ethanol, deionized water and dried with nitrogen.

Materials characterization. The surface morphology and root mean square (RMS) roughness of the AZO seed films were investigated by an atomic force microscopy (AFM) (Asylum Research MFP-3D-SA). X-ray Photoelectron Spectroscopy (XPS, TSC K-Alpha, AlKa) was adopted to investigate the elemental states of Zn and Al. The concentration of Al was determined by using inductively coupled plasma atomic emission spectroscopy (ICP-AES, Perkin-Elmer, Optima 2100DV). The morphology of the composite films was observed in a scanning electron microscope (E-SEM; Quanta 250, FEG) and transmission electron microscopy (TEM; Tecnai G2). Crystal properties of the AZO seed layer film and films were characterized by grazing-incidence x-ray diffraction (GI-XRD) using Cu-Kα radiation of 0.15406 nm. The fourier transform infrared spectra (FTIR) were recorded using a Thermo Fisher IS 50 spectrophotometer in the wave number range of 0–4000 cm−1 at room temperature. WCA measurement was carried out with a CA101 contact angle goniometer using a 4-μL water droplet.

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Author Contributions
Y.L., X.H.W. and W.Q. devised the original concept, designed the experiment, discussed the interpretation of results and co-wrote the paper. Y.L., Y.K. and G.W. performed the experiments. J.F.W. and J.Z.W. analysed the dates. Q.K.L. made the preliminary simulation. J.Z. and B.H. polished the manuscript. All authors reviewed the manuscript.

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