Nature-inspired robust hydrochromic film for dual anticounterfeiting

Highlights
- Micro-structures are formed by self-assembly of polyurethane-polyelectrolyte colloids
- Information changes from one pattern to another within 3 s when exposed to water
- The hydrochromic films are mechanically robust in both dry and wet state

Pengxiang Si, Mingrui Liang, Manyou Sun, Boxin Zhao
zhaob@uwaterloo.ca

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Nature-inspired robust hydrochromic film for dual anticounterfeiting

Pengxiang Si,1 Mingrui Liang,1 Manyou Sun,1 and Boxin Zhao1,2,*

SUMMARY
Nature-inspired materials have been actively developed for anticounterfeiting applications. Among a variety of stimuli-responsive anticounterfeiting strategies, hydrochromic materials exhibit reversible color change in response to moisture or water and have the advantage of being easy to authenticate. However, the security level of current hydrochromic anticounterfeiting materials is not sufficient for practical applications since they only exhibit a single anticounterfeiting function, where the information switches between visible and invisible. To improve the security level and efficiency of hydrochromic anticounterfeiting materials, here we developed a robust dual hydrochromic material via the self-assembly of polyurethane (PU)-polyelectrolytes colloids with which the desired information can not only switch between visible and invisible but also transform from one pattern to another within 3 s without the need of any external instruments. The bio-inspiration, material design and demonstrated hydrochromic properties might have profound implications for using colloidal complexes to make advanced anticounterfeiting materials.

INTRODUCTION
Nature is a bountiful source of inspiration for modulating light-matter interactions (Choi et al., 2020; Liu et al., 2019a; Syurik et al., 2017; Völlers et al., 2016). Plant- or animal-derived materials containing unique nanostructures or microstructures are able to regulate light scattering, providing a wide range of structural color from whiteness to transparency (Jacucci et al., 2020). Scattering occurs when light propagates in a material with refractive index non-uniformities such as scattering centers or particles in bulk materials (Bohr, 1983). Thus, tuning the refractive index of a material via environmental stimuli enables dynamic color modification (Li and Yin, 2019). Hydrochromic materials exhibit reversible color change through humidity or water and have been widely used in displays, sensors, smart windows, and anticounterfeiting labels (Singh et al., 2017; Yu et al., 2020; Zhao et al., 2020). One example from nature is the skeleton flower (Diphylleia grayi) petals, which consist of numerous air cavities in dry weather, enabling light scattering and resulting in white color (Figure 1A, left). However, on a rainy day, those air gaps are filled with water that has a similar refractive index to its petals, minimizing light scattering and leading to a transparent appearance (Figure 1A, right) (Cai et al., 2018; Liu et al., 2019a; Yong et al., 2015; Yoo et al., 2020). Similarly, the tortoise beetle Charidotella egredia displays red color in dry weather and golden color on rainy days due to the filling of air or water in the nano-porous skin (Vigneron et al., 2007).

These interesting phenomena have inspired many researchers to develop a variety of hydrochromic materials for anticounterfeiting applications (Choi et al., 2020; Yu et al., 2018; Zeng et al., 2017). Compared with thermochromic, electrochromic, and photochromic strategies, hydrochromic anticounterfeiting materials are inexpensive, mass producible with printing technologies, and easy to authenticate by the naked eye without external instruments (Huang et al., 2019a; Huang et al., 2019b; Kang et al., 2018; Liu et al., 2019b; Peng et al., 2018; Wang et al., 2018; Yao et al., 2019). However, the security level of hydrochromic materials is not high enough for real uses because they only exhibit single anticounterfeiting function, with the information switching between visible and invisible by water stimuli (Choi et al., 2020; Singh et al., 2017; Yong et al., 2015; Yoo et al., 2020). To improve the safety of the secret information, Zhao et al. developed a thermal and hydrochromic double encrypted system. It requires a high temperature of 180°C enabled by an external instrument (Zhao et al., 2020). Note that organic phosphorescence materials could be used for anticounterfeiting; they had been used to realize multi-dimension/hydrochromic anticounterfeiting (Lei et al., 2020; Xiao et al., 2018).
Herein, inspired by *Diphylleia grayi*, we develop a robust dual anticounterfeiting hydrochromic material in which the information can not only switch between visible and invisible but also change from one pattern to another within 3 s without external instruments, providing opportunities to fabricate more complicated anticounterfeiting patterns for real applications. The designed dual anticounterfeiting hydrochromic material was fabricated using anionic polyurethane (PU) water dispersion and consisted of three layers which respond differently to water stimuli: the top layer changes from opaque to transparent; the middle layer remains transparent; and the bottom layer changes from transparent to opaque. Anionic PU water dispersion was selected as the matrix due to its high transparency, mechanical robustness, and solution processability. By controlling the deposition and self-assembly of PU colloidal particles during the film formation process, the obtained material exhibits different microscopic structures and transparency.

**RESULTS AND DISCUSSION**

**Fabrication of hydrochromic films**

The schematic diagram for the preparation of PU/PEI and PU/PMANa hydrochromic films is presented in Figure 1B. The synthesis route of anionic PU water dispersion followed that of our previous work (Si et al., 2020a). Figure 1B1 displays a photo of the anionic PU water dispersion, which was an amphiphilic polymer containing a hydrophilic carboxyl head from 2,2-bis(hydroxymethyl) propionic acid (DMPA) and hydrophobic core from polypropylene glycol-2000 (PPG-2000) and isophorone diisocyanate (IPDI). The amphiphilic PU in water self-assembled into colloidal particles with an average diameter of 18 nm and Zeta potential of $\pm 38$ mV determined from dynamic light scattering (DLS) data, as seen in Figure 1C.

The arrangement of PU colloidal particles during the drying process could significantly affect the optical...
transparency of dried films. To fabricate a transparent PU substrate, the anionic PU water dispersion was cast onto a glass slide. As the water slowly evaporated at room temperature, PU colloidal particles diffused and entangled with each other to form a transparent polymer film (Figure 1B2).

Inspired by Diphylleia grayi, the fabrication of a thin layer of micro-particles on a transparent surface could effectively scatter visible light, leading to an opaque appearance (Sun and Bhushan, 2019). Cationic poly-electrolyte PEI was blended into anionic PU water dispersion with the protection of ammonium hydroxide to form a stable PU/PEI dispersion (Si et al., 2020a). During spray coating, the high pressure of the air flow divided PU/PEI dispersion into numerous microdroplets. As the ammonium hydroxide quickly evaporated, PU and PEI aggregated within these microwater droplets to form PU/PEI micro-particles on the PU substrate. Similar to Diphylleia grayi, the obtained PU film coated with the PU/PEI micro-particles was hydrochromic, changing between opaque in the dry state and transparent in the wet state (Figure 1B3, Video S1).

In contrast with PU/PEI film, anionic polyelectrolyte PMANa was mixed with anionic PU water dispersion to produce another type of hydrochromic film that changed from transparent to opaque via water stimuli. The PU/PMANa dispersion was directly cast onto PU substrate. The dried PU/PMANa film was visually transparent in the dry state and opaque in the wet state (Figure 1B4, Video S2). Both PU/PEI and PU/PMANa dispersions exhibited an average diameter less than 60 nm and Zeta potential less than −30 mV, indicating good colloidal stability for further processing (Figure 1C).

**Morphologies and swelling behavior of hydrochromic films**

To investigate the mechanism of the above two types of hydrochromic films, the morphologies of the PU/PEI and PU/PMANa hydrochromic films were examined with a scanning electron microscope (SEM).

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**Figure 2. The two types of hydrochromic films have different micro-structures and water swelling behavior**

(A) SEM image of PU/PEI hydrochromic film in a dry state.
(B) SEM image of PU/PEI hydrochromic film after soaking in water for 1 min and drying for 30 min.
(C) Cross-sectional SEM image of PU/PMANa hydrochromic film in a dry state.
(D) Cross-sectional SEM image of PU/PMANa hydrochromic film after soaking in water for 1 min and drying for 30 min.
(E) EDX data of PU/PMANa hydrochromic film.
(F) Water swelling mass ratio of PU/PEI and PU/PMANa hydrochromic films at different soaking times.
(G) Initial water contact angles on PU/PEI (left) and PU/PMANa (right) hydrochromic films, (each sample was measured for 3 times).
spray-coated PU/PEI particles had a diameter ranging from 400 nm to 20 μm in their dry state (Figure 2A). After soaking in water for 1 min and drying for 30 min, the PU/PEI hydrochromic films maintained the same microstructures as in their dry state, indicating that the PU/PEI particles are stable in water (Figure 2B). Compared with the PU/PEI hydrochromic film, the surface of the PU/PMANa hydrochromic film was smooth but had micro-domains inside the material (Figure 2C). These irregular domains were formed via self-assembly of PU and PMANa and had a diameter ranging from 400 nm to 10 μm when dry (Si et al., 2020b). These micro-domains on the cross-sectional surface of the dry PU/PMANa after being soaked in water for 1 min and dried for 30 min were found disappeared due to the high water affinity and solubility of PU/PMANa micro-domains (Figure 2D). The difference ratio of Na element from energy-dispersive X-ray spectroscopy (EDX) data indicated these micro-domains were PMANa dominated and the polymer matrix was PU dominated (Figure 2E). To further verify the capability of water absorbance, the swelling mass ratio and water contact angle of PU/PEI and PU/PMANa hydrochromic films were measured. The swelling mass ratios of PU/PEI and PU/PMANa hydrochromic films were 0.2 wt% and 90 wt%, respectively (Figure 2F). This could be explained by PEI physically cross-linking with the carboxyl groups on PU through electrostatic attraction to increase the hydrophobicity of PU, whereas the addition of PMANa would introduce more carboxyl groups to increase the hydrophilicity of PU. Therefore, the PU/PEI hydrochromic film had a higher initial water contact angle (90.5° ± 4.6°) than the PU/PMANa hydrochromic film (73.8° ± 2.8°) (Figure 2G). The water contact angle on the PU/PMANa film decreased within 180 s due to its hydrophilicity and good water absorbance (Figure S1A), whereas the contact angle of PU/PEI film maintained almost the same (Figure S1B). The significance of testing contact angle is to reveal the difference of wettability and water absorbance rate of PU/PEI and PU/PMANa film. The PU/PMANa absorbs water quickly but PU/PEI does not. Therefore, PU/PMANa turns white but PU/PEI remains transparent. Actually, PU/PEI film can eventually turn white after being soaked in water for 6 hr.

Optical transmittance of hydrochromic films

The optical transmittance of PU/PEI and PU/PMANa hydrochromic films in their dry and wet states was found to be dependent on coating thickness. We prepared two different thicknesses of 5 μm and 30 μm for each of the PU/PEI and PU/PMANa coating layers, which were measured by a 3D laser microscope (Figure 3A). As a control experiment, the neat PU exhibited high transmittance (above 95%) in both the dry and wet states without any hydrochromic behavior. The PU/PEI and PU/PMANa films with 5-μm coating thickness showed a transmittance change from 60% to 88% and 86% to 25% for their dry to wet states, respectively. The thicker films showed similar trends but a lower transmittance in both their dry and wet states (Figures 3B and 3C). The hydrochromic properties of PU/PEI and PU/PMANa films were reversible with no sign of degradation for 20 dry-wet cycles (Figure 3D). The sizes of PU/PEI particles and PU/PMANa micro-domains varied from 400 nm to 20 μm, which spanned beyond the wavelength of the visible light (400 nm–800 nm). The large particle size and the phenomenon of thickness-dependent transmittance collectively confirmed the Mie scattering mechanism of PU/PEI and PU/PMANa films (Rybin et al., 2009; Yao et al., 2019).

Mie scattering effect of hydrochromic films

Figure 4 shows a schematic diagram of the scattering effect in PU/PEI and PU/PMANa hydrochromic films in their dry and wet states. The dried PU/PEI hydrochromic composite film was in a scattering state because the path of incident light was disturbed by the discrete PU/PEI micro-particles adhered on the PU surface. In the wet state, water wetted both PU/PEI particles and PU substrate and water filled in the spaces among the PU/PEI particles, replacing the air; the water wetting minimized the overall scattering effect since the refractive index of water (n = 1.33) is closer to PU (n = 1.49) than to air (n = 1) (Figures 4A and 4B). Therefore, the fabrication of PU/PEI particles on the top of PU surface so as to have the air gaps among them was critical to achieve a hydrochromic effect. In contrast, the PU/PEI film showed high transmittance of 95% when the fabrication method was solution casting instead of spray coating and no change of transmittance was observed after soaking in water for 1 min. In contrast to the PU/PEI hydrochromic composite film which transmittance increased in wet state, the transmittance of the PU/PMANa hydrochromic composite film significantly decreased when exposed to water. This is because that the PU/PMANa micro-domains were formed inside the PU matrix due to the self-assembly of colloidal solution. The similarity between the refractive indices of PMANa (n = 1.48) and PU (n = 1.49) minimized the scattering effect to achieve a high transparency in the dry state. When soaked in water, the hydrophilic PU/PMANa particles quickly absorbed water, leading to the spreading of water...
inside the film. These PU/PMANa micro-domains acted as scattering centers that decreased the transmittance of the film (Figures 4C and 4D). Notably, it is the water adsorbed in the micro-domains that act as the scattering center for the PU/PMANa composite film. In dry state, the PU/PMANa and PU have similar refractive indices so that the composite film is transparent. In wet state, the absorbed water in the micro-domains has a different refractive index than the PU, subsequently acting as a scattering center while not affecting the film integrity (Figure 2D). These invisible micro-domains once dried could still absorb water. Therefore, the hydrochromic phenomenon of PU/PMANa composite film is reversible.

Mechanical properties of hydrochromic films
To be more practical for potential applications in anticounterfeiting, the transmittance contrast of hydrochromic films between their dry and wet states needs to be maximized. Therefore, PU/PEI and PU/PMANa films with 5 μm thickness coating layers were selected to fabricate dual anticounterfeiting material. In addition to optical properties, the mechanical properties of anticounterfeiting material are also critical for practical applications. From the tensile stress-strain curves, the PU/PEI hydrochromic film showed a tensile strength of 9.1 MPa, breaking elongation of 1600% and toughness of 84.5 MJ/m$^3$. The PU/PMANa hydrochromic film was not as tough as the PU/PEI hydrochromic film because the non-crosslinkable and intrinsically rigid PMANa blocked the diffusion and entanglement of PU colloidal particles. After soaking in water for 1 min, the tensile strength decreased and breaking elongation increased for both PU/PEI and PU/PMANa hydrochromic films, indicating that the water molecules served as a plasticizer of the polymers. Due to the difference between their water swelling ratios, the PU/PMANa hydrochromic film had a much larger toughness decrease of 20.2 MJ/m$^3$ than the PU/PEI hydrochromic film with 1.1 MJ/m$^3$ (Figure 5A). The adhesive properties of PU/PEI and PU/PMANa hydrochromic films were characterized through a 180° peeling test to evaluate the bonding strength between the PU/PEI or PU/PMANa coatings and the PU substrate. The test was applied under both dry and wet
states. After peeling, there was no damage either at the tape/PU/PEI (or tape/PU/PMANa) interface or the PU/PEI/PU (or PU/PMANa/PU) interface, indicating the strong adhesion between the PU/PEI (or PU/PMANa) coatings and the PU substrate. The peel force was higher for the wet state because water softened the polymer and increased its contact area with the tape (Figure 5B). Furthermore, scratch tests were performed to produce PU/PEI and PU/PMANa hydrochromic film surface damage under controlled loading. Figures 5C and 5D exhibited typical scratch track of damages on PU/PEI and PU/PMANa hydrochromic films under an optical microscope with an increasing load from 2 g to 5000 g in dry and wet states. The value of critical load was used to determine the maximum load that PU/PEI and PU/PMANa hydrochromic films can tolerate prior to delamination. The delamination of PU/PEI and PU/PMANa films started at about 1000 g and 500 g, respectively, suggesting good scratch resistance in both dry and wet states. Overall, the toughness, adhesion, durability, and failure resistance of the PU/PEI and PU/PMANa hydrochromic films were robust for practical applications.

**Dual anticounterfeiting hydrochromic film**

As a proof of concept, we developed a robust dual anticounterfeiting hydrochromic material by integrating PU/PEI and PU/PMANa coatings on each side of a PU film. The transparent PU substrate of 200 μm thickness is the middle layer, the “cloud” pattern made with PU/PEI colloidal dispersion was on the top side, and the “PU” letters made using PU/PMANa colloidal dispersion were on the bottom side of PU film (Figure 6A). The obtained dual anticounterfeiting hydrochromic material exhibited “cloud” patterns in the dry state. After soaked in water for 3 s, the “cloud” patterns changed to “PU” letters (Figure 6B and Video S3). In addition to a PU substrate, we have performed preliminary tests, showing that the dual anticounterfeiting material can also be fabricated on other transparent substrates such as glass, polystyrene, polyethylene terephthalate, etc. Compared to the previous single anticounterfeiting hydrochromic materials on which information can only appear or disappear, our designed material successfully accomplished a complete information change on the same area of material with fast responsive time and without external instruments. More complicated two-dimensional patterns and coded information could be created by a variety of combinations of hydrochromic PU/PEI, PU/PMANa, and non-hydrochromic materials on the same area of materials, providing a dual anticounterfeiting function.
Conclusion

In summary, inspired by *Diphylleia grayi*, we have demonstrated a mechanically robust hydrochromic film based on PU/PEI and PU/PMANa colloidal dispersions for dual anticounterfeiting, wherein the information can not only switch between visible and invisible but also change from one pattern to another within fast responsive time and without external instruments. Water was utilized to tune the refractive index of hydrochromic films in order to modify the light scattering pattern of PU/PEI and PU/PMANa micro-structures, resulting in transmittance change of the material. The water-transparent PU/PEI and water-opaque PU/PMANa were coated on the top and bottom surfaces of a PU substrate, respectively. After soaked in water, the coated “cloud” patterns made by PU/PEI visually transformed into “PU” letters made by PU/PMANa within 3 s. The designed dual anticounterfeiting hydrochromic films are mechanically robust, quick to respond, inexpensive, environmentally friendly, mass producible, printable, non-destructive, and easy to authenticate with high security. Two-dimensional patterns and codes could be fabricated by a variety of combinations of hydrochromic PU/PEI, PU/PMANa, and non-hydrochromic materials on the same area of materials, showing great potential in fabricating more complicated anticounterfeiting hydrochromic materials in the future.

Limitations of the study

The transparency contrast between dry and wet states of the PU/PEI layer needs to be further improved in the future. Additionally, the anticounterfeiting material might lose the hydrochromic phenomenon after being continuously soaked in water for longer than one day due to the dissolving of PMANa and hydrolysis of PU.

STAR METHODS

Detailed methods are provided in the online version of this paper and include the following:

![Figure 5. The hydrochromic films are mechanically robust](image-url)}
**KEY RESOURCES TABLE**

**RESOURCE AVAILABILITY**
- Lead contact
- Materials availability
- Data and code availability

**METHOD DETAILS**
- Synthesis of anionic PU water dispersion
- Preparation of PU/PEI hydrochromic film
- Preparation of PU/PMA Na hydrochromic film
- Preparation of hydrochromic film for dual anticounterfeiting
- Characterization

**SUPPLEMENTAL INFORMATION**
Supplemental information can be found online at https://doi.org/10.1016/j.isci.2021.102652.

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**AUTHOR CONTRIBUTIONS**
Conceptualization, design, methodology, and analysis, P.S. and B.Z.; experiments, P.S., M.S., and M.L.; manuscript writing, P.S. and B.Z.

**DECLARATION OF INTERESTS**
The authors declare no competing interests.

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**Figure 6.** The dual anticounterfeiting hydrochromic film is designed by the combination of three layers
(A) Schematic illustration of dual anticounterfeiting hydrochromic film.
(B) Photo of dual anticounterfeiting hydrochromic film: switchable “cloud” patterns and “PU” letters between dry and wet states.
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STAR★METHODS

KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| Chemicals, peptides, and recombinant proteins | Sigma-Aldrich | CAT#81380 |
| Polypropylene glycol-2000 (PPG-2000) | Sigma-Aldrich | CAT#81380 |
| Dimethylolpropionic acid (DMPA) | Sigma-Aldrich | CAT#106615 |
| Isophorone diisocyanate (IPDI) (98%) | Sigma-Aldrich | CAT#317624 |
| Ethylene diamine (EDA) (99.5%) | Sigma-Aldrich | CAT#391085 |
| Triethylamine (TEA) (99%) | Sigma-Aldrich | CAT#0086 |
| Reagent acetone | Sigma-Aldrich | CAT#179124 |
| Branched polyethylenimine (PEI) (average Mw = 25000) | Sigma-Aldrich | CAT#408727 |
| Ammonium hydroxide solution (28–30%) | Sigma-Aldrich | CAT#221228 |
| Poly (methacrylic acid, sodium salt) solution (PMANa, Mw = 4000-6,000, 40 wt % in H2O) | Sigma-Aldrich | CAT#674044 |

RESOURCE AVAILABILITY

Lead contact
Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, Boxin Zhao (zhaob@uwaterloo.ca).

Materials availability
This study did not generate new unique reagents.

Data and code availability
This study did not generate/analyze [datasets/code]

METHOD DETAILS

Synthesis of anionic PU water dispersion
50 g of PPG-2000, 3.4 g of DMPA and 16 mL of IPDI were added into a three-neck flask equipped with a water-cooled condenser and N₂ tubes. The reaction took place for 4 hr at 110°C under mechanical stirring; the resulting polymer was cooled down to 40°C, and then 38 mL of reagent acetone was added into the flask under mechanical stirring; then, 100 mL of amine water solution containing 0.018 mol EDA and 0.022 mol TEA was added into flask under vigorous stirring at room temperature. Acetone was removed by vacuum oven for 20 min. Anionic PU water dispersion was obtained and condensed to 30 wt % solid content.

Preparation of PU/PEI hydrochromic film
2 mL anionic PU water dispersion was cast on 75 mm by 25 mm glass slides. The transparent PU film was obtained after drying naturally at room temperature for 24 hr. Branched PEI was dissolved in DI water with a concentration of 30 mg/mL. 0.3 mL ammonium hydroxide solution and 0.3 mL branched PEI water solution were added into 2 mL of the anionic PU water dispersion. The obtained PU-PEI solution was spray coated on transparent PU film via air brush and then dried at room temperature for 30 min.

Preparation of PU/PMANa hydrochromic film
2 mL anionic PU water dispersion was cast on 75 mm by 25 mm glass slides. The transparent PU film was obtained after drying naturally at room temperature for 24 hr. PMANa solution was diluted to 4 wt % and then 2 mL PMANa solution was mixed with 2 mL anionic PU water. 0.5 mL PU/PMANa solution was cast on PU film followed by drying at room temperature for 2 hr.
Preparation of hydrochromic film for dual anticounterfeiting

2 mL anionic PU water dispersion was cast on 75 mm by 25 mm glass slides. The transparent PU film was obtained after drying naturally at room temperature for 24 hr. PU/PEI solution was spray coated on PU film via a stencil with a “cloud” pattern to produce a thin layer “cloud” pattern of PU/PEI with around 5 μm thickness. After drying for 30 min at room temperature, the film was peeled off from the glass slide and then flipped over to attach on the glass slide. The “PU” letters were handwritten via a glass pipette filled with PU/PMANa solution followed by drying at room temperature for 2 hr. The “cloud” pattern was visible in the air and changed to “PU” letters after soaking the obtained hydrochromic film in water for 3 s. After drying, the “PU” letters changed back to the “cloud” pattern.

Characterization

Dynamic light scattering (DLS) (Malvern Zetasizer Nano ZSP) was used to measure the average diameter and zeta potential of the PU, PU/PEI and PU/PMANa colloidal dispersions.

UV-vis diffuse reflectance spectroscopy (UV-Vis) (Lambda 1050) was performed to measure the visible light (400 nm–800 nm) transmittance of PU/PEI and PU/PMANa films in the dry and wet states. Tensile tests were conducted with a universal mechanical tester (UMT) (T1377, Center for Tribology Inc.) at a speed of 2 mm/s using a 10 kg load cell. 180-degree peeling tests were completed with UMT at a speed of 0.5 mm/s using a 1 kg load cell. A backing Scotch tape with 19 mm width was adhered on samples for peeling. Scratch tests were performed with UMT by using a 1 mm diameter stainless steel ball indenter that was fixed on a 10 kg load cell. The indenter was moved horizontally along the film surface for 20 mm at 0.5 mm/s while the normal force linearly increased from 2 g to 5000 g. A digital microscope (AD41132T, Dino-Lite) was used to observe the scratch track. A 3D laser microscope (OLSS5000, Olympus) was applied to measure the thickness of PU/PEI and PU/PMANa coatings on PU film. The morphology of PU/PEI and PU/PMANa films was observed using a SEM (ZEISS Ultra) at 10 kV accelerating voltage. 5 μL of DI water was used for measuring the water contact angle of PU/PEI and PU/PMANa films.