Recent Developments in Artificial Super-Wettable Surfaces Based on Bioinspired Polymeric Materials for Biomedical Applications

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Abstract: Inspired by nature, significant research efforts have been made to discover the diverse range of biomaterials for various biomedical applications such as drug development, disease diagnosis, biomedical testing, therapy, etc. Polymers as bioinspired materials with extreme wettable properties, such as superhydrophilic and superhydrophobic surfaces, have received considerable interest in the past due to their multiple applications in anti-fogging, anti-icing, self-cleaning, oil–water separation, biosensing, and effective transportation of water. Apart from the numerous technological applications for extreme wetting and self-cleaning products, recently, super-wettable surfaces based on polymeric materials have also emerged as excellent candidates in studying biological processes. In this review, we systematically illustrate the designing and processing of artificial, super-wettable surfaces by using different polymeric materials for a variety of biomedical applications including tissue engineering, drug/gene delivery, molecular recognition, and diagnosis. Special attention has been paid to applications concerning the identification, control, and analysis of exceedingly small molecular amounts and applications permitting high cell and biomaterial cell screening. Current outlook and future prospects are also provided.

Keywords: nature; polymeric materials; super-wettable surfaces; bioinspired material; biomedicine

1. Introduction

Polymeric-materials-based superhydrophobic surfaces find significant applications [1] in the multidisciplinary areas of sciences with special emphasis on biomedicine, as illustrated in Figure 1. Such applications include wound dressing, implants, wettable surfaces, sensing, imaging, prosthetic materials, dental/bone materials, tissue engineering scaffolds, and drug carriers due to their desired physical and mechanical properties, low cost, and ease of modification for several functionalities. The development of biomedical materials currently needs extremely high standards in terms of functionality, bioactivity, and biocompatibility. Scientists have made sincere efforts [2] to partially or completely meet these criteria using a variety of techniques. The most promising among them have been biomimetic or bioinspired [3] since the compounds, architectures, and processes of living organisms display nature-based, time-tested approaches for obtaining desirable material properties.
Figure 1. Potentials of polymeric-materials-based super-wettable surfaces in diverse areas.

It is well known that early diagnosis is critical for many illnesses, including cancer, since it greatly improves the possibility of successful treatment. Unfortunately, the number of clinically important molecules for disease diagnostics is limited in the early phases of certain pathologies. Additionally, the identification of available molecular markers in biological fluid is extremely challenging due to the interference by numerous other complicated biomolecules found in body fluids [4]. Nanotechnology has delivered innovative methods to clinical practice, including the discovery of superhydrophobic surfaces. Such fields encourage important scientific developments with a wide range of opportunities for diagnostic and therapeutic applications in modern biology and medicine [5–8]. Artificial, superhydrophobic surfaces are biomimetic surfaces that facilitate naturally occurring structures such as those of lotus and rose petals. A droplet of water sitting on these surfaces must be approximately spherical with a water contact angle above 150°. This wetting property is attributed to the surface chemical effect since there is a rough surface fabric composed of micro-and nanostructures [9–11]. The scale of these constructions ranges between several hundred nanometers and tens of microns, the standard spatial region protected by advanced micro-manufacturing techniques. Scientists have developed powerful techniques to mimic superhydrophobic surfaces that exist naturally and to produce artificial surfaces with extreme wettable properties using both top-down and bottom-up approaches [9–11]. These activities have led to the development of modern, advanced superhydrophobic surfaces that can accommodate incredibly small concentrations of liquids quickly, tracking the locations and spatial configurations of a few droplets of microliters or even nanoliters [8]. Additionally, by integrating extremely precise micro- and nanostructures, it has enabled one to freely coordinate, analyze, and modify the materials found within droplets at both the micro scale and at nanoscale levels. In many disciplines, their multifaceted implementations have drawn considerable attention [5–9,11]. Various surface functions were combined with engineering surface topology and chemistry based on advances in nanotechnology, such as the self-cleaning capability of lotus leaves [12], anisotropic wetting of rice leaves [13], the efficiency of water collection by desert beetles [14], and liquid-repelling pitcher plants [15]. Their specific wetting property provides innovative, cell-based, micro-environmental solutions and cell substratum interactions that have been commonly used in bioinspired areas of intense wettability that cannot be accomplished using traditional tissue and cell culture platforms [16–21]. Although the use of extreme, wettable surfaces in medical applications has been of considerable interest, only a few studies are available to understand their potential role and
applications in the biomedicine sector [8,12,14,15,20,22–39]. This review presents the construction of polymer-based, super-wettable surfaces motivated by the superhydrophobic repellency of the lotus leaf, the anisotropic wetting capability of the butterfly wing and the rice leaf, the adhesion of the rose petal to water, desert beetle-patterned wettability, and the liquid slippery surfaces of the Nepenthes plant, and also discusses their potential applications in multiple areas. Current limitations and prospects for the biomedical evaluation of these polymeric-materials-based surfaces in contrast to concluding comments are also discussed.

2. Methodologies to Fabricate Super-Wettable Surfaces Using Various Polymeric Materials

The extreme wettability of certain living organisms has encouraged scientists and engineers working around the globe to develop surfaces that mimic natural properties. Consequently, several artificial, superhydrophobic surfaces were reported as advancements in the area of nanotechnology. The desired wettable properties in such artificial surfaces are usually accomplished by the engineering of micro- and nanostructures and surface energy. This section introduces some natural species with special wettable properties and also sheds light on their fabrication strategies.

2.1. Superhydrophobic Surfaces

Superhydrophobic surfaces are “water-repellant” or “non-wettable” surfaces that exhibit a water contact angle (CA) above 150°. Such non-wettable surfaces have gained substantial attention owing to their applicability in multiple areas over the last decade, such as self-cleaning [8,40,41], droplet manipulation [42], and corrosion-resistance [10,43–46]. Studies have confirmed that the non-wettable action of water droplets on non-wettable surfaces blends face chemistry and topology. Particularly, superhydrophobicity on the surface of nanostructured materials with low surface energy was achieved. Wenzel [47] and Cassie–Baxter [48] explained these surfaces thoroughly. A droplet forms a static CA (θ) on an ideal, solid, flat surface (Figure 2a). This CA is associated in Young’s equation with a surface energy of solid/gas (γ_{sl}), solid/liquid (γ_{sl}), and the interfaces of liquid/gas (γ_{lg}) [47].

\[ \gamma_{sl} = \gamma_{sl} + \gamma_{lg} \times \cos \theta \]  

(1)

![Figure 2](image_url)

**Figure 2.** Graphics of water droplet on rock-solid surfaces that are flat and rough. (a) Perfect flat surface with a droplet, (b) droplet situation in Wenzel state with a fully wet rough surface, and (c) droplet situation in Cassie–Baxter state with air pockets at the junction between an unwetted rough surface and a droplet.
When the solid surface is rough, the values of $\gamma_{sl}$, $\gamma_{sg}$, and $\gamma_{lg}$ can be taken into consideration since the direct interaction area between the droplet and the raw surface varies from the smooth surface. The whole rough surface is known to be in contact with the water droplet in the Wenzel model (Figure 2b). Here, the water CA in the Wenzel state ($\theta_{w^*}$) can be described as:

$$\cos \theta_{w^*} = R \times \cos \theta$$

(2)

The ratio between the actual surface area and the projected surface is the surface roughness factor ($R$). The water droplet is mounted on superhydrophobic surfaces due to its wide contact areas with the rock. It comprises the roughened surface’s topmost layer, producing air bags in the Cassie–Baxter model at the solid–liquid boundary (Figure 2c). Therefore, Young’s equation can be stated as follows:

$$\cos \theta_{c^*} = -1 + f (\cos \theta + 1)$$

(3)

where $\theta_{c^*}$ denotes the Cassie–Baxter CA and $f$ represents the ratio of actual droplet contact area to the total surface area. This decreased interaction region of surface-to-liquid leads to superhydrophobic surfaces that exhibit water-repellent and self-cleaning properties. Lotus leaves epitomize superhydrophobic natural surfaces repellent to water (Figure 3a). For the first time in 1997, Barthlott and Neinhuis reported that numerous leaves contain a naturally directed, epidermal, papillose cell structure coated with epicuticular hydrophobic wax (Figure 3b). In line with the Cassie–Baxter equation, they observed that, under a floating water droplet, the air can be trapped. Such discoveries revealed the use of hierarchical functional structures treated with energy materials and low surfaces to create superhydrophobic surfaces that resemble lotus leaves and are highly water-repellent. Various methods based on lotus leaves have been developed using diverse polymeric materials, such as wet chemical etching [49–56], electrochemical reagent [57–59], layer-by-layer repositioning, lithography [60–62], electrodynamic [63–65], or sol-gel [66–70] methods. The electrohydrodynamics technique was stated by Jiang et al. [71] as a multifaceted and efficient way of producing composite polystyrene (PS) films mixing pore microphones and cross-woven neurofibers. The fabricated PS-composite film demonstrated high hydrophobicity due to improved surface robustness, pore microspheres, and interwoven nanofibers. As shown in Figure 3c, the hierarchical structures of regularly arranged PS colloidal crystals decorated with a system of wet, self-assembly chemicals protected by carbon nanotubes (CNTs) were fabricated by Li et al. [72]. The acquired surfaces displayed superhydrophobicity with a low slide angle after the tuning of the surface with PFDTs (1H,1H,2H,2H-perfluoro-decyl trichlorosilane). Electrochemical reactions and photolithography provided a frequent series of hierarchical micro/nanostructures including structured, microprotrusion Cu covered with nanostructured Ag dendrites. These surfaces were manufactured in a broad pH range following the modification of PFDTs (Figure 3d).
was also studied. The rice leaf also exhibits a hierarchically ordered structure with waxy layers that were documented by Lai’s research team via simple, electrochemical methods for altering nanotubes. Seo et al. recently reported a vertically oriented silicon nanowire surface whose wettability was regulated from superhydrophilic to superhydrophobic under atmospheric oxygen after rapid thermal annealing (RTA). Reprinted with permission from [50,52,72,73]. Copyright 1997 Oxford University Press, copyright 2013 Elsevier, copyright 2005, 2007 American Chemical Society (ACS).

Species with water pinning properties on superhydrophobic surfaces have also been found in nature. For example, a rose petal displays high water CA with exciting features of water adhesion. As depicted in Figure 3e, the water droplet on the petals of rose does not move away even after the surface is tilted upright or upturned. A systematic series of nanoscale cuticular folds, as well as micropapillae, are found above each micropapillae on the surfaces of the rose petal (Figure 3f) [73–75]. Such a large periodic array generates a capillary action that, ultimately, facilitates the penetration of water droplets into grooves of micropapillae, thus, confirming the water-adhesion property. Employing a solvent-evaporation-triggered nanoimprint pattern transfer method, Feng and co-workers [74] replicated the micro/nanostructures of rose petals to mimic their liquid adhesion feature. In the replication process, a second template was obtained by negatively replicating the poly(vinyl alcohol) film (PVA). As shown in Figure 3g, the polystyrene film from this negative replica showed a water-adhesive, superhydrophobic property with a large water CA. Superhydrophobic, nanostructured TiO₂ films with tunable surface adhesion were documented by Lai’s research team via simple, electrochemical methods for altering nanotubes’ diameter and length [75]. The water-adhesive strength of superhydrophobic, porous nanostructures was manipulated by using capillary force and surface roughness. Seo et al. recently reported a vertically oriented silicon nanowire surface whose wettability was regulated from superhydrophilic to superhydrophobic under atmospheric oxygen after rapid thermal annealing (RTA) at 1000 °C [36]. Following the RTA cycle, the surfaces displayed a substantial shift (0 to 154°) in water CA, along with good water-adhesion characteristics. The change in surface chemistry from hydrophilic -Si-O-H groups to the hydrophobic -Si-O-Si- groups was the primary cause behind this dramatic transformation.

In addition to the superhydrophobicity of the leaves of the lotus and petals of the rose, the anisotropic wettability of natural species, such as butterfly wings and rice leaves, was also studied. The rice leaf also exhibits a hierarchically ordered structure with waxy nanobumps, as on the lotus leaf (Figure 4a(i)); however, the anisotropic wettability is attributed to the quasi-one-dimensional micropapillae configuration (Figure 4a(ii)) [11]. As shown in Figure 4a(iii), depending on orientation, this steering arrangement provides different energy barriers to moisture, which can be quickly rolled off vertical to the edge.
of the rice leaf. Several approaches to the development of man-made, superhydrophobic surfaces with anisotropic wetting properties have been developed [13,76–82]. Electrospinning nanofiber arrays developed by Wu’s research team were systematically aligned with poly(vinyl alcohol) (PVA) [78]. For obtaining uniform arrays over large areas, nanofibers were fabricated in between the two paralleled copper strips during the electrospinning process. In this method, on the aligned fiber arrays, the water droplet displayed anisotropic activity, as on a rice leaf. By using patterns on the fiber collector, systematic wetting behavior was established. Kang and co-workers recently reported three kinds of anisotropic microgroove array with various structures, such as rectangle, prism, and an overhang shape which was overhanging by a UV-mediated micromolding method and, thereafter, undergoing surface alteration with octafluorocyclobutane [24]. Surface wetting activities such as static CAs and SAs in the various microgrooves with different input types were altered. The highest fluid repellency and anisotropic wetting were observed in the overhang line arrays. Due to the incredibly low octafluorocyclobutane surface energy (~13 mJ/m²) and overhanging structure, the grooves instructed droplets of water and mineral oil.

Aside from anisotropic wetting, some natural surfaces with wetting in one direction also exist in nature. For example, the wings of Morpho butterflies exhibit wetting features in one direction owing to the presence of micro/nanostructures. As depicted in Figure 4b(ii,iii), these micro/nanostructures are frequently superimposed on the outer side of the wing, allowing the droplet to quickly move away from the wing as it spins strongly in a contrary direction [83]. Malvadkar and co-workers demonstrated a one-directional wetting pattern on tilted nanofilm using a nanorod array of poly(p-xylylene) (PPX) [84]. Oblique angle polymerization was adopted to develop the oblique PPX nanorods. Higher water CA and one-directional wetting were demonstrated by the resulting PPX nanorod film. As nanorods were pointed vertically upwards towards the surface, the droplets of water adhered to the surface. The droplets of water bounced away from the surface in the reverse direction.

Figure 4. (a) The anisotropic wetting of a rice leaf in two directions: (i) real picture and SEM photographs of the leaf of rice, (ii) rice leaf height map using optical profiler, (iii) the anisotropic wetting behavior in two directions. (b) The wetting activity of a butterfly wing in one direction: (i) real picture and SEM images of the wing of the butterfly, (ii) butterfly wing height map using optical profiler, (iii) the wetting activity in one direction. (c) The movement of a droplet on poly(p-xylylene) nanofilm. (d) Related time-lapse frames of droplet movement. Reprinted with permission from [83,84]. Copyright 2012 the Royal Society of Chemistry (RSC) and copyright 2010 Nature Publishing Group.
The gout motion in one direction was illustrated by the vibration of low amplitude on the half-pipe configurations covered with poly(p-xylylene) film (Figure 4c). In the PPX nanofilm-coated semi-pipe, a unidirectional movement of the droplet was noted (Figure 4d) when random vibration was applied to the droplet located in the half-pipe.

2.2. Decorated Wettability for Collection of Water

For their survival, organic species adapt themselves to live under harsh conditions. Animals and desert plants have unique features to retain water under dry conditions. Particularly, the Stenocara beetle collects and condenses fog water on its bumpy back in the Namib Desert. On the back surface [14], this water collection method is based on its hydrophobic and hydrophilic patterns. The Stenocara wings are protected by a random spectrum of hydrophilic and hydrophobic rough troughs and smooth peaks (Figure 5a(i)). Once completely submerged, the fog water settles down and condenses on the hydrophilic heights. The droplet rolls down the hydrophobic troughs in a critical dimension (Figure 5a(ii)). Numerous wettability-patterned surfaces on different organic and inorganic materials, such as oxides, polymers, and metals [85–98], have been created inspired by the patterned, hydrophilic beetles. Seo et al. [86] recently demonstrated arrays of superhydrophobic Si nanowire patterned hydrophilically to direct water droplets. The metal-assisted electroless engraving was employed to fabricate superhydrophobic Si nanowire followed by a self-assembled coating of monolayers. Subsequently, the hydrophilic guiding patterns were obtained under the illumination of UV light, as shown in Figure 5b(i), using shadow masks. Time-dependent images revealed that, due to high wettability in contrast with superhydrophobic Si nanowire arrays, a water droplet traveled alongside pre-determined hydrophilic pathways (Figure 5b(ii)). Based on the adhesive mechanism typically observed in mussel-adhesive proteins, Kang and co-workers [91] proposed a mask-less chemical method based on the solution. To partly surround the mussel-inspired polydopamine (pDA), the patterned polydimethyl siloxane (PDMS) mold-covered a synthetic membrane surface of the superhydrophobic porous oxide. Due to its hydrophilicity, the designed region became water-adhesive, and the surface was able to absorb and direct droplets of water in a pDA pattern. Li et al. [99] also developed superhydrophilic printed designs on a superhydrophobic surface. The phospholipid ink solution was printed on a porous, superhydrophobic surface. The great disparity in wettability between the superhydrophobic surface and printed spots allowed the sports area to be selectively wetted by aqueous solutions. This method provided a fast and convenient way to produce superhydrophilic shapes owing to its conformability with known methods of microfabrication such as inkjet printing, microcontact printing, and dip-pen nanolithography.

Figure 5. (a) Fused Stenocara beetle overwings showing water-capturing surfaces: (i) real picture and SEM images of unstained beetle wing regions (black) and wax-stained regions (color pictures);
(ii) wax droplets growing in a fog-laden wing in a time-dependent manner; (b) arrays of superhydrophobic Si nanowire arranged in a hydrophilic pattern for controlling water droplet; (i) water-guiding tracking by the construction of tilted Si NW arrays; (ii) images of water droplet channeled through the hydrophilic trajectory. Reprinted with permission from [14,86]. Copyright 2001 Nature Publishing Group and copyright 2011 American Chemical Society (ACS).

2.3. Smooth Liquid Surfaces

Many of the insects have two vital functioning properties that can be applied to multiple surfaces. Claws help to adhere to rougher textures, while adhesive pads [100,101] enable attaching to smooth surfaces. Nepenthes, the tropical, carnivore pitcher plant, exhibits a unique trapping organ that prevents the attachment of insects. The organ is specifically identified with its distinctive anisotropic and slippery qualities [15,102,103]. Within the lifting pitcher, the peristomic epidermal cells have stacks coated with hydrophobic wax and microstructures. Through secreted nectar and rainwater, the hydrophilic peristomic surface also encourages the production of fluid foil. Such physicochemical features prevent claws from being fastened into the peristome and adhesive pads. Encouraged by the Nepenthes pitcher plant’s exceptional repellent capability, Wong and co-workers developed (Figure 6a(i)) slippery surfaces based on nanostructured, porous material by penetrating liquid lubricants [25]. Such slippery films were produced with non-hazardous, perfluorinated lubricants based on oil and water. These films had improved liquid repellent properties with negligible CA hysteresis and low SAs in comparison with superhydrophobic, non-infused surfaces (Figure 6a(ii)). Moreover, they exhibited numerous features such as an instantaneous, repeatable, self-healing property, optical transparency, and pressure stability. They have numerous applications including marine anti-icing [104], anti-fouling coatings [105], and antibacterial [106]. Aside from the slippery surface applications, mechanical stimulus-responsive slippery surfaces have been developed. Using the elastic membrane of PDMS, Yao’s research team [26] (Figure 6b) developed a droplet-adhesion, controllable slippery surface. As shown in Figure 6c, the dynamic liquid interface kept switching between resting and stretching stages. The droplet sliding was promoted by the flat and smooth liquid interface in a relaxed state. The stretching, in contrast, revealed a rough surface (Figure 6d). In this prolonged situation, even a freshly formed oil drop was immobilized while, at its deposition location, the existing oil drop persisted. The two drops started to slide until the stress was released. The results of multiple deformation processes have been studied, such as bending, reversible swelling-drying, self-healing, and poking.
Figure 6. (a) (i) Fabrication of smooth film, (ii) the optical microscopic images of a drop of hexane sliding with a small angle; (b) the matrix formed on PDMS substrate and the pictures of lubricated, as well as dry substrates; (c) (i) topographical changes induced mechanically in the liquid slippery film by stretching and (ii) droplet motion; (d) dynamically slippery surface with the systematic movement of the oil droplet. Reprinted with permission from [25,26]. Copyright 2011, 2013 Nature Publishing Group.

2.4. Wettability of Switchable Superhydrophobic Surfaces

For transport with no loss and biological identification requiring minimal interaction of water substrates, reversible swapping of superhydrophobic adhesion properties is needed. New techniques for this action on the same substrate are supported by stimulating materials [5,42,107–118]. Li et al. first reacted a heat-sensitive side-chain liquid crystal polymer (LCP) on the rough surface that changed the properties of superhydrophobic adherence [111]. A water droplet was attached to the polymer-coated surface as the temperature approached a phase transition, while a hydrogen rearrangement of the polymer chains was carried out. The reverse process of this adhesion was found as the substratum cooled down to room temperature. Furthermore, raw surfaces were integrated with polymers for temperature, pH, and electrolyte sensors, allowing numerous adhesive adhesion switches [112]. Photoreactive compounds were also applied to accomplish adhesion-shifting surfaces. Wang et al. stated that UV-irradiated and heat treatment of switchable water adhesion [119] was observed by superhydrophobic TiO$_2$ nanotube films. To create hydrophilic regions evenly dispersed in superhydrophobic environments, these films were selectively illuminated by a mask. Such well-separated areas of lighting increased water adhesion and ensured superhydrophobia. Illuminated regions (80–180 °C) were regener-
ated by quick annealing. However, this technique was not sufficient for practical purposes since an improvement in temperature greatly sped up the dehydration of water droplets. Li and co-workers [107] reported a light-responsive raw surface employing an azobenzene liquid crystal polymer (LCP) with a separate change in polarity between cis and trans isomers under light irradiation (Figure 7). UV irradiation (365 nm) for 6 s was performed by azobenzene mesogens on the surface of the manufactured LCP film of azobenzene, which enhanced surface water adhesion by corresponding changes in polarity. As depicted in Figure 7a, under visible light (530 nm) irradiation for 30 s, the mesogens of azobenzene shifted to trans position which, ultimately, resulted in regaining the surface low adhesion to water. Thus, the wettability of the surface was changed dynamically from rolling to splintering, employing a combination of UV and visible light (Figure 7b).

Figure 7. (a) Photoswitchable adhesion with UV and visible light excitation; (b) dynamic adhesion capability of LCP film; (c) (i) the volume of Pd layers deposited expanded on arrays of Si NWs under air and hydrogen atmosphere, (ii) the angles of contact confirming superhydrophobicity under similar conditions; (d) pictures of moving water droplets under similar conditions. Reprinted with permission from [107,108]. Copyright 2012 the Royal Chemical Society (RSC) and copyright 2013 John Wiley and Sons.

In the methods described above, to regulate the superhydrophobic adhesion of the surface, some constraints in biological applications were found due to their chemistry. UV irradiation, for instance, may affect enzymes and biological cells. Wu et al. showed that, by varying its curvature as another approach to tuning adhesion forces on a superhydrophobic surface, a superhydrophobic periodic PDMS micropillar array could be amended [42]. A water droplet was mounted in a trapped condition on a smooth, pillar-array film that had a large force of adhesion. As the surface curvature increased, the adhesion decreased gradually. When the curvature rose further, the droplet was removed from the series. This process allowed droplets of water to be transferred in situ without loss. A new
approach was put in motion recently to address the drawbacks of chemical processes using gas-responsive materials. There is also a modern approach to address the limitations of gas-responding materials using the above-mentioned techniques. Fast gas-powered adhesion to a superhydrophobic Si NW palladium (Pd)-coated layer was demonstrated by Seo et al. [108]. Fast adhesive switching resulted in a morphological phase transformation in the Pd-coated Si NW arrays. H atoms resulting from H\textsubscript{2} decomposition were applied under the H\textsubscript{2} atmosphere to the as-deposited Pd sheet, resulting in volume expansion (Figure 7c(i)). When environmental parameters were transferred from hydrogen atmosphere to air, the extended palladium layer automatically collapsed towards its original volume. Although the air and H atmospheres were low and extremely sticky, these Si NW arrays were superhydrophobic in the air and H\textsubscript{2} (Figure 7c(ii),d). Droplets were also rolled off and pinned off by switching among these ambient and H\textsubscript{2} requirements. Thus, Section 2 summarizes the development of bioinspired, superhydrophobic surfaces with unique wettable properties. A detailed check of wettability and liquid adhesion is given by the micro/nanostructure and energy surface. These unusual wetting properties of organic areas are likely to be used in numerous biomedical applications and, using traditional methods, cannot be achieved using cell culture pipelines or flat Petri dishes.

3. Extreme Wetting Surfaces Inspired by Nature and Their Applications in Biomedicine

For various biomedical uses, analysis, practical cell culture biomedical devices, and lab-on-one-chip systems may be used directly on bioinspired surfaces with intense weaving functionality. This segment provides new advances in the platforms that are bioinspired.

3.1. Cell Configuration for Studying Cellular Interactions

The topology and chemistry of surfaces are very dependent on protein adsorption and attachment to cells. With patterned superhydrophilic and superhydrophobic surfaces [17,18,120–129], cell interactions have been thoroughly investigated. Piret et al. noted the selective adhesion of Chinese hamster ovary K1 cells to micropatterns on a surface of a superhydrophobic area [20]. Likewise, Ishizak’s research team demonstrated that physicochemical properties of surface (e.g., robustness, weight resistance) influence cell interactions and cell–cell adhesion [16]. Specifically, the mouse 3T3 fibroblast cells adhered to the superhydrophilic areas immediately after seeding, while the cells scarcely adhered to superhydrophobic planes. The preference for protein absorption was due to the variation in cell attachment in superhydrophobic areas. The cultured cells modified their shapes and directions of adhesion selectively, based on the distance between superhydrophobic areas. Moreover, there was direct contact between cells in adjacent locations under a designated cell gap of 250 µm cell–cell. In terms of tissue and organism growth in distinct but neighboring compartments, multiple-cell-type cultivation proved to be important for imitating and analyzing different biological actions, such as intercellular connectivity and cell signaling. Multiple cell patterning forms were applied to a porous, hydrophilic/hydrophobic boundary substratum by Efremov et al. [124]. These hydrophilic substrates with a superhydrophobic border were developed in two stages: firstly, a hydrophilic nanoporous layer of the polymer was generated through a UV polymerization reaction. Afterward, the hydrophilic polymer coating was placed over a hydrophobic, photographic material. With a photomask, the UV was radiated again to make the superhydrophobic boundaries. The resulting surface was used in a reciprocal culture medium through pre-patterned cell co-cultivation for cell migration and signaling. The movement of MLTy-mCherry and HeLa-GFP cell lines through the thin, superhydrophobic periphery to different geometries was tracked within three days of co-culture. Cell movement was avoided by the thin boundary and could be utilized without intermingling to cultivate different types of cells correctly. At the same time, Wnt proteins were also used for the cell signaling process, a crucial class of proteins issued by the association of border areas. Precisely, the Wnt ligand was cultured that expressed zebrafish fibroblast Pac-2 cells that neighbored each pattern.
without direct cell interaction. In the tissue’s extracellular space, the proteins migrated independently without any interaction with the cell. Interactions between cell–cell and cell–biomaterial on flat, 2D surfaces have been extensively studied, but experiments have been more useful in 3D, as micro-environments are best imitated in vivo [130–134]. In hydrophilic, patterned, superhydrophobic substrates between cells and polymers, Salgado et al. [27] analyzed 3D interactions. UV/ozone irradiation was carried out by a superhydrophobic PS substrate using a square-patterned photomask. By comparing wettability in several 3D hydrogel sizes, the hydrophilic patterns could be repositioned (Figure 8a(ii)). Fibroblast, L929, and MC3T3-E1 pre-osteoblast cells were employed for testing 3D cell–hydrogel interactions. Among 24 various hydrogels, alginate (Alg) with various amounts of chitosan (Chi), hyaluronic acid (HA), gelatin (G), and collagen (Coll) were combined. The cytotoxicity of various compositions of hydrogel (total 24) was studied for 24 h (Figure 8a) and 5 µL of cell-mixed polymer/crosslinker solutions was deposited on the substratum. Followed by cell cultivation for 24 h, all the materials were evaluated for cell quantification and cell viability and assessed for cell metabolic activity. The fluorescence imaging studies of live (green) and dead (red) cells in hydrogels, after 24 h of culture, are shown in Figure 8b. Dead cells were comparatively low in a 40% Alg containing HA/G mixture but high in a 40% Alg–Coll mixture. It was observed that, for L929 fibroblast cells, 70% of the Alg hydrogels containing HA and Coll and hydrogels containing Chi demonstrated high and low viability. For Coll- and HA-style hydrogels with 70% Alg, L929 cells were of high viability but with low compatibility. Additionally, even though the bone native tissue and pre-bone (osteoid) represent the most abundant protein structure, the presence of Coll enhanced pre-osteoblast cell viability.

![Image](image_url)

**Figure 8.** (a) (i) 3D hydrogel array of patterned hydrophilic marks; (ii) pictures before and after immersion in culture medium for 24 h; (b) fluorescence imaging of live (green) and dead (red) cells after cell culture for 24 h using 24 diverse hydrogels. Reprinted with permission from [27]. Copyright 2012 Oxford University Press.

### 3.2. Functional 3D Cell Spheroids

Cell culture in 3D conditions makes it possible to study interactions between the cell and extracellular matrix (ECM) more easily. Particularly, 3D cell spheroids offer cells with larger micro-environments in vivo [135,136]. Moreover, they also hold significant interest due to improved therapeutic capacities relative to cells grown on two-dimensional substrates. Traditionally, with classical dish cultivation, the hanging drop technique, and spinner flask culture [137–140], 3D cellular spheroids have been developed. By regulating their cell size and viability, hanging drop cultures can be associated with the hydrophilic patterned superhydrophobic surface [141–143]. Employing mussel-inspired adhesive polymer pDA [144], Lee et al. performed photolithography on the fluorosilane-coated super-
hydrophobic substrate to acquire the wettability-patterned surface. The hanging drop technique was employed for culturing rat islet cells (ICs) and human mesenchymal stem cells (MSCs) on the pDA-patterned surface. The active surface acquisition of 3D MSC spheroids increased the secretion of its vascular endothelial growth factor (VEGF) to an approximately 300% higher level compared to the level reached by the spinner flake culture. IC spheroids were also shown to improve glucose stimulus sensitivity by about 200%. In contrast with those cultivated with the technique of hanging drops on traditional Petri dishes, the uniformity, feasibility, and functionality of spheroid cells were increased. Such observations indicate that, by increasing cell–cell behavior, a spherical gout form increases cell–cell and cell–matrix interactions. Thus, it is helpful on superhydrophobic surfaces without hydrophilic architecture for hanging drop spheroid culture. The superhydrophobic surface, 3D micro-environments dramatically decrease the cell cultivation media volume and avoid any soil contact. Neto and co-workers used a superhydrophobic surface patterned in micro-morphology for spheroid cultivation [20]. By applying force, the superhydrophobic surface of PS was primed and the surface was dented into sharp tips. Water droplets displayed a strong CA on the indentation as the water was fixed into the indentation. The preserved superhydrophobic features contributed to the creation of spherical water droplets on the surface and contact areas between the droplets and reduction of the surface. On the superhydrophobic surface of the mechanically modified PS, 3D spheroids were developed (Figure 9a(i)). Droplets-based cells continued, suspended from the indented base, despite the surface being rotated upside-down. Using different cell densities or drug concentrations, a cell spheroid combinatory study was performed and established using gravitational power. Mouse-immortalized lung fibroblast cell line L929 droplets were dispensed and cultured at two densities of thirty and forty thousand cells/droplet on the inverted superhydrophobic surface. The compound was subjected to varying levels of anti-cancer drug doxorubicin after 24 h of cell culture to determine the dose-dependent reaction of the tumor spheroid. In Figure 9a(ii,iii), the results of drug screening achieved from the stained fluorescent images are demonstrated. The cell ratios in both spheroids between live (green) and dead (red) decreased as the dose decreased. Additionally, such ratios were larger at the lower density of spheroid. In comparison, dead cells collected mainly in the inner spheroid (Figure 9a(iii)) since it was more difficult to determine nutrients and waste releases in larger and denser spheroids.

Figure 9. (a) (i) Sticking drop culture on a superhydrophobic surface with hollow shapes; (ii) the ratios of life (green) and dead (red) cells after introducing different doxorubicin concentrations; (iii) fluorescence imaging of L929 spheroids in doxorubicin-treated spheroids. (b) (i) Culture medium droplets on Sn NWs coated with Pd and exposed with H2 at various tile angles; (ii) post four days of culture at different cell densities and medium sizes, live/dead cell staining, size distribution, and VEGF protein secretion from spheroids. (n = 3, **: p < 0.01 compared to the density of 1.25 × 10⁵ cells/mL, ###: p < 0.01 compared to the density of 2.5 × 10⁵ cells/mL). Reprinted with permission from [20,28]. Copyright 2014 John Wiley and Sons.
The superhydrophobic surface with reversible adhesion properties was demonstrated by Seo et al. [138] to reduce cell-interface interactions during hanging drop spheroid culture. The gas-triggered adhesion to the hydrogen-sensitive surface of the superhydrophobic surface is shown in Figure 7c,d by deposition of Pd to Si NWs. Pd-coated Si NW arrays are adhesion swapping features that, using an attachment system without patterns, are specifically applied to 3D spheroid forming. The adhesion switch properties of Pd-coated Si NW arrays are directly applied to 3D spheroid formation using the hanging drop technique without any patterning process. Due the adhesion properties of Pd-coated Si NW arrays after exposure to $\text{H}_2$, droplets of variable sizes containing human adipose-derived stem cells (hADSC) were adhered and maintained under ambient air, as shown in Figure 9b(i). Figure 9b(ii) indicates the viability of spheroids after four days of culture and the controllability of spheroid size by different cell densities ($1.25, 2.5$, and $5.0 \times 10^5$ cells/mL) and different medium volumes (10, 15, and 20µL). It was demonstrated that the secretion of VEGF from hADSC spheroids depended on the size of the spheroids. Improved cell density paracrine behavior and culture medium volume adaptation were observed in hADSC spheroids. A decreased distribution and substantially improved VEGF secretions of hADSC spheroids were observed relative to traditional spinner bottles and Petri dishes. In contrast to the Petri-dish strategies, this drop-out strategy increased the efficacy of hADSC, paracrine seclusion, mitochondrial metabolic processes, apoptosis, and ECM. Additionally, an angiogenic potential was assessed as an operative assay for hADSC spheroids. The conditioned media obtained in Pd-coated Si NW pads from hADSC spheroids increased the proliferation of human endothelial cells and accelerated capacity formation.

3.3. Biomedical Devices

In clinical applications for implantable biomedical instruments, the adhesion of harmful biomedical materials to the surface must be avoided. Thus, if the implanted system requires prevention of inflammation or infection in cell/tissue culture systems, an antibacterial surface property is highly desirable. Different methods for the development of antibacterial surfaces have been reported [145–154]. A superhydrophobic surface inspired by biological structures has become a favorable antibacterial surface due to the anti-fouling capability that could circumvent surface bacteria from adhering [155–159]. Privett et al. [155] also produced xerogels which contained silica colloids, fluoroalkoxy silane, and silane backbone. The superhydrophobic characteristics of the coated surface of the xerogel were facilitated by fluorinated silica nanoparticles for low surface energy and a hierarchical structure. The antibacterial properties of the xerogel were described by a conventional cell flow assay using Gram-negative *Pseudomonas aeruginosa* (*P. aeruginosa*) and Gram-positive *Staphylococcus aureus* (*S. aureus*). Microbial adhesion to the superhydrophobic xerogel surface was decreased by 98% and 99% relative to the blank area. The antibacterial properties of surfaces with Gram-negative bacteria were reported by Feschau et al. [29], using superhydrophobic polycarbonate (PC) and polyethylene (PE) and PS substrates. Using a basic casting method, the superhydrophobic PC, PE, and PS surfaces were acquired from a micro/nanostructured PDMS casting. A total of 10 µL of *E. coli* was cultured to test standardized PS, PC, and PE surface anti-bacterial properties. Then, a bacterial *E. coli* solution was cultivated for 24 h (Figure 10). Fewer than 100 colony-forming units (CFUs) were observed on PS- and PE-based superhydrophobic surfaces after 24 h, whereas no bacterial growth was observed on the PC substrate. Moreover, 100 CFUs were observed each for flat PS and PC, while 25,800 were observed for PE. These findings suggest that, relative to flat surfaces, superhydrophobic surfaces efficiently reduce adhesion of bacteria to <0.1%. Because of the wide variety of anti-fouling properties, liquid slippery surfaces often have tremendous potential for various medical applications in comparison to different liquids and environmental pressures. Over a wide range of temperatures, heats, surface tensions, and multiple conditions they retain repellency [159]. Epstein’s research team [160] showed the importance of a slippery surface to avert binding biofilm. On a smooth, slippery surface, the bacteria were presented, which could not be anchored on the
mobile interface compared to solid interface. Irrespective of the principal solid, porous structure, separate biofilm accumulations were stopped by a slippery surface over more than a week. In contrast to common, polyethylene glycol, anti-fouling surfaces, it also decreased bacterial attachment by 96–96.6%. To minimize the morbidity and mortality caused by thrombosis, Leslie et al. [30] used slippery properties on tubing and catheters of indwelling medical devices. They developed a tethered perfluorocarbon coating (TP) at the top of the tube followed by its coating with a liquid perfluorodecalin (LP) surface to attain anti-thrombogenic and non-adhesive surfaces. The thin, moving liquid layer permitted the tethered-liquid perfluorocarbon (TLP) surface to resist fluids effectively also after the surface contacted an immiscible liquid, such as blood (Figure 10b(i)). The surface was cleaned almost immediately from a new, whole human droplet of blood (Figure 10b(ii)). Compared to the uncoated surfaces of acrylic and polysulfone, they showed (Figure 10c(i)) a decrease in the adhesion and polymerization of the TLP surface. The slithery condition also decreased the adhesion of platelets in contrast to uncoated surfaces. These findings demonstrated that the smooth surface decreased fibrin polymerization and inhibited adherence, as well as plates activation. To realize an arteriogenic shunt for in vivo analysis of the anti-thrombogenic effect of the slippery surface, polycarbonate connectors, TLP-treated polyurethane cannulae, and medicinal, polyvinyl chloride (PVC)-based heart pulmonary perfusion tubes were studied. Compared with control tubes, Figure 10c(ii) displays polymer TLP-treated tubes that minimize occlusive thrombosis post flow for 8 h. Such noteworthy, anti-fouling properties can, therefore, be used in several additional applications.

**Figure 10.** (a) The growth of bacteria on structured and smooth PS and PC substrates; (b) (i) the smooth surfaces after coating with tethered-liquid perfluorocarbon (TLP) showing blood repellency; (ii) the images of the slippery surface showing slipping of a blood droplet; (c) (i) fluorescence images of fibrinogen on polysulfone or acrylic surfaces in the presence and absence of TLP coating; (ii) the images of polycarbonate connectors, polyurethane cannulae, and PVC tubing in the presence and absence of TLP coating. Reproduced from [29] under CC BY license. Reprinted with permission from [30]. Copyright 2014 Nature Publishing Group.

3.4. **Lab-on-a-Chip Based on Open Channel Droplets**

In terms of fast penetration into the sample fluid, energy consumption and low samples, quick chemical and biological reactions [161–171], such as microfluidic droplet systems, have many advantages compared to traditional closed-channel, microfluidic applications. A gout-based microfluidic system [163] was introduced on a superhydrophobic, porous oxide membrane by You et al. To direct the water droplets, they inserted hydrophilic PDAs on superhydrophobic surfaces. A square pDA for more complex droplet handling was also planned in the middle of the pDA micro-line. A fluid droplet traveled along the micro-lines by momentum, stopping at the square, which showed adequate surface energy to catch the droplet. Such an immobilized droplet was pushed downwards by inserting a second droplet. With this droplet capacity to combine, preparation of monodispersed gold nanoparticles and rapid, structural protein changes were introduced. For chemical and
biological organic solvent-based reactions and experiments, however, superhydrophobic hydrophilic patterned surfaces could not be employed. For compatibility with different solvents, a pDA, micropatterned slippery surface was introduced by You et al. [172]. To attain a fluid-guiding feature, PDA micro-lines were engineered on a nanostructured surface before lubricant infiltration. Multiple solvents, for example, gasoline, dimethyl sulfoxide, water, dimethylformamide, 1,2-dichloroethane, n-hexane, toluene, acetone, and diesel oils, were compatible with the produced slippery device. Any solvent with a surface tension larger than that of the lubricant could be depleted by gravitational force along the micro-lines by the infused lubricant located on top of the pDA micro-line. They implemented the square shape pattern for the gout mixture in the pDA micro-line intersection with the same aforementioned process (Figure 11a(i)) [163]. The chemical reactions using an organic solvent were carried out on the slippery surface of the pDA micro-model due to the broad compatibility of solvents. The organic reaction was performed between benzaldehyde and ortho-phenylenediamine to create 2-arylbenzimidazole. The droplet of THF solution composed of two reactants, as well as a THF, was dropped on the oily surface, as shown in Figure 11a(ii). By fast and homogeneous mixing, the reaction rate (70.3%) improved relative to the standard bulk reaction using a vial (52.1%). Furthermore, the surface was cleared and used again after ultrasonication.

![Figure 11](image-url)

**Figure 11.** (a) (i) The movement of the droplet and its mixing on a slippery system fabricated with pDA (ii) THF-based chemical reaction on the slippery surface; (b) (i) regulation of motions of water (movement, mixing, and examination) on PDMS substrate appended with micropillar arrays; (ii) the dimple structure’s SEM picture; (iii) water droplet image on the substrate of PDMS; (c) (i) the measurement system for SERS; (ii) formation of minor, interfering lipidoid-RNA complex; (d) SERS measurement spectra (in situ/ex situ) for R6G/Ag NP droplet mixture and an evaporated R6G/Ag NP droplet; (e) fluorescence imaging and flow cytometry studies of transfected GFP-Hela cells. Scale bar-200 μm (n = 3, **p < 0.01, compared to the conventional group). Reprinted with permission from [172,173]. Copyright 2014 American Chemical Society (ACS) and copyright 2015 Nature Publishing Group.

Decorated droplet-handling systems only had minimal control over fluid processes, for instance, start and stop motion. Newly developed by Seo et al., the superhydrophobic
PDMS substrate contained micropillar arrays for programmed water droplet manipulation (Figure 11b(i)) [173]. When applying the substrate, the vacuum pressure on the suspended PDMS substratum was expanded (Figure 11b(ii)) to form a local, pale structure. Decreased performance and superhydrophobicity of the micropillar array was observed (Figure 11). In the interface region between the substrate and water droplets, the dimple arrangement could be used to monitor the motions of water droplets individually. The curvature was positive at the boundary of the pillars adjacent to the PDMS, and the width was greater than the flat substratum distance. The lower number of micropillars decreased the adhesiveness of the water and allowed isolation from the droplet substratum. The flow of water drops on the substratum also could not be conveniently regulated through the vacuum dimple structure; however, the moving direction could also be readily built without any additional pattern. Surface-enhanced Raman spectroscopy (SERS) analyzed the substrate’s analytical efficiency (Figure 11c(ii)). Rhodamine 6G (R6G) in situ/ex situ SERS tests were carried out at different concentrations (Figure 11d). In situ, individual water droplets consisting of Ag nanoparticles and R6G molecules were determined by the SERS and combined at the Raman detection point before the collection of data. The lower limit of detection for R6G/Ag NP was calculated to be $10^{-5}$ M, owing to disseminated R6G and Ag molecules in a mixed droplet. The mixture was evaporated and its components condensed to resolve this detection cap within an area of a certain length. The R6G molecules were also observed at $10^{-15}$ M because of the aggregation of Ag NPs and R6G for ex situ SERS calculation. To manufacture regular, intracellular, gene transfer nanoparticles, they used the platform. In the lipid (ND98) complexes with green fluorescent protein (GFP) siRNA, a basic method of mixing and blending outlets was used (siGFP). On the platform, homogenous mixing of droplets and the efficiency of transfections to GFP-HeLa cells from the platform’s lipidoid-siRNA complexes (78.8% ± 0.5%), in contrast to classical manual pipetting mixing (70.0%), could be achieved, as shown in Figure 11e.

3.5. Fluorescent Intelligence Sensing Based on Bioinspired Super-Wettable Patterns

In analytical chemistry, fluorescence is a popular detection method. In a light-absorption process, fluorescence is a luminescence process induced by an appropriate molecule under light excitation. Fluorescence intensity is measurable at specific excitation/emission wavelengths. Under dilute concentrations, the amplitude of fluorescence would be equal to the fluorophore concentration. Several groups recently focused their efforts on the fabrication of bioinspired super-wettable microchips for profound fluorescence-based detection. These examples will be highlighted in the following text: mainly their features such as anchoring and enriching capabilities for biosensing. The analytes were stored in microdroplets that were enclosed inside the superhydrophilic microwells with the superhydrophobic substrate serving as a wall to keep the microdroplets from spreading. Huang and co-workers recorded sensitive detection of metal ions by integrating the fluorescence technique, as seen in Figure 12a, by using the anchoring capacities of bioinspired micro-patterns. A hydrophilic, photonic-crystal, colloidal microchip was created by assembling hydrophilic, photonic-crystal, colloidal particles on a hydrophilic–hydrophobic patterned substrate. The developed microchip improved fluorescence detection in multiple channels selectively, as well as performed highly effective testing under discriminative conditions using twelve metal ions [31]. Such a well-performing, patterned microchip demonstrates a modern use of bioinspired microchips in detection and their crucial role in the development of sophisticated fluorescent devices and advanced discriminative analysis.
Figure 12. Super-wettable fluorescence finding. (a) High-performance metal-ion recognition using a bioinspired photonic-crystal microchip; (b) ultra-trace DNA identification using super-wettable microchips; (c) example of free prostate-specific antigen (f-PSA) detection on homogeneous super-wettable microchips in a schematic diagram; (d) a green super-wettable miRNA biochip built on TiO₂ substrate. Reprinted with permission from [31,33,37,39]. Copyright 2013, 2015 Wiley-VCH and copyright 2018 Elsevier B.V.

The “analyte solution concentration in the hydrophilic region increases as the microdroplet evaporates, which is ideal for capturing the analyte for ultra-trace detection. In Figure 12b [33], ultra-trace fluorescent DNA detection is demonstrated by using the super-wettable microchips’ enrichment ability. UV-based etching of the octadecyl trichlorosilane (OTS), functionalized, superhydrophobic, nanodendritic Si coating through a photomask and deposition of candle soot were employed to create this super-wettable microchip. The analyte was captured from an extremely diluted solution by constant evaporation and emission signals were eventually intensified to attain the recognition of DNA with a low detection limit (10⁻¹⁶ M). Aggregation-induced emission enhancement (AIEE) and evaporation-induced enhancement were accomplished concurrently by incorporating AIE molecules onto a super-wettable microchip [33]. As a result of the synergetic enhancement, a miRNA biosensor with enhanced efficiency (detection limit-1 pM, linear range-10⁻⁶ to 10⁻¹² M) can be achieved. The super-wettable microchip’s overall fluorescence strength was just half that of industrial hydrophilic or hydrophobic glass substrates. Due to the coffee rings, however, unequal signal propagation occurred on the hydrophilic and hydrophobic surfaces, possibly limiting repeatability. The sensitive and precise identification of biomarkers included excellent spots in the superhydrophilic microwells. The coffee-ring effect could be reduced in super-wettable microchips, and the homogeneity of superhydrophilic spots could be increased, as seen in Figure 12c [37]. The superior Marangoni effect in the superhydrophilic spot and the reduced outward flow, owing to the large hydrodynamic flow resistance of 3D Si nanodendritic structure, contributed to this phenomenon. Figure 12d [39,174,175] displays the sustainable super-wettable biochips for miRNA identification. The nanodendritic TiO₂ nanostructure was made hydrothermally on FTO-coated glass substrates, then modified with OTS and exposed to UV light via a photomask. By using its anchoring and enrichment abilities, super-wettable fluorescence detection provides a sensitive technique that could be especially useful in biomarker sampling, diagnosis, and disease monitoring. However, the need for large instruments and
complicated biomolecule fluorescent labeling can restrict their use in point-of-care (PoC) detection. Integration of bioinspired micropatterns with compact fluorescence instruments for point-of-care research will be a potential priority.

Aside from the above-discussed papers, recent years have witnessed several other polymeric-materials-based, super-wettable surfaces for applications in diverse research areas [176–185]. To summarize, some common polymeric-material-based, super-wettable surfaces utilized for various applications are illustrated in Table 1.

### Table 1. Some common polymeric materials used in the fabrication of super-wettable surfaces.

| Polymeric Materials                    | Fabrication Method                  | Application          | Wetting Property | Reference |
|----------------------------------------|-------------------------------------|----------------------|------------------|-----------|
| Hyaluronic acid (HA)                   | Phase separation                    | Tissue engineering   | Hydrophobic      | Ref [27] |
| Polyethylene (PE)                      | Shrink-induced mold method          | Antibacterial        | Superhydrophobic | Ref [29] |
| Polycarbonate (PC)                     | Shrink-induced mold method          | Antibacterial        | Superhydrophobic | Ref [29] |
| Polystyrene (PS)                       | Electrohydrodynamics (EHD)          | Tissue engineering   | Superhydrophobic | Ref [71] |
| 1H,1H,2H,2H-perfluoro-decyl trichlorosilane (PFDTS) | Wet chemical self-assembly | Conductive stainless steel | Superhydrophobic | Ref [72] |
| Poly(vinyl alcohol) (PVA)              | Solvent evaporation                 | Petal effects        | Hydrophilic-Superhydrophobic | Ref [74] |
| Poly(p-xylene) (PPX)                   | Soft lithography                    | Self-cleaning        | Superhydrophobic | Ref [91] |
| Polydimethyl siloxane (PDMS)           | Soft lithography                    | Water adhesion       | Hydrophilic-Superhydrophobic | Ref [91] |
| Polydopamine (PD)                      | Soft lithography                    | Water adhesion       | Hydrophilic-Superhydrophobic | Ref [91] |
| Polytetrafluoroethylene (PTFE)         | Sol-gel strategy                    | Water adhesion       | Superhydrophobic | Ref [176] |
| Polyvinylpyrrolidone (PVP)             | Sol-gel strategy                    | Self-cleaning        | Superhydrophobic | Ref [176] |
| Polyamide (PA)                         | Dip-coating method                  | Water and oil separation | Superhydrophobic | Ref [181] |
| Silane-modified polymer (SMP)          | Spray-coating method                | Water and oil separation | Superhydrophobic | Ref [181] |

### 4. Conclusions and Future Prospects

This review systematically described the design and development of functional bioinspired surfaces based on smart polymeric materials with extreme wettable properties for biomedical and engineering applications. The specific values of properties were identified, and representative living organisms with bioinspired surfaces were investigated, from the leaf of lotus to the pitcher plant. The importance of such features in the area of biomedical science was also highlighted. Furthermore, various functional approaches, such as high-performance cell assays and droplet-based lab-on-chips were also described. In spite of the significant advancements in the respective research area, some key issues remain that are required to be overcome to apply bioinspired, polymeric-materials-based surfaces for wide usage, especially as cutting-edge biomedical platforms. For instance, the long-term stability and durability of the surfaces with extreme wetting characteristics must be enhanced. To provide nutrients and bioactive molecules for cell culture, cultural substrates should provide a culture medium for in vitro cell/tissue culture, but continuous interaction with
the culture medium and surface results in adsorption of protein and penetration of liquid to the structured surface causing a change in the surface wetting properties. Therefore, the construction of bioinspired and long-lasting surfaces based on new polymeric materials with good stability and durability can produce more micro-environments in cell culture in living conditions and can be employed as an innovative method for investigating cellular activity in contrast to traditional procedures. The stimuli-responsive and self-healing composites that can restore degraded, bioinspired surfaces could be a promising material for achieving a platform with long-lasting stability. In addition, the advancement of bioinspired, intelligent surfaces made up of smart, polymeric materials in the controlling and release of drugs could be crucial to the faster and successful diagnosis of diseases and drug discovery applications. Moreover, there is also a great demand to build and introduce fast and cost-effective manufacturing methods for bioinspired surfaces based on advanced polymeric systems, adapting traditional methodologies in mass production such as spray coating and the latest 3D bioprinting technology. Such practical innovations could have a powerful effect on the biomedical industry in the forthcoming days after circumventing the existing aforementioned challenges.

Author Contributions: All authors discussed the contents of the review manuscript. A.A. designed the paper outline and wrote the text of the manuscript; C.Z., M.A. and A.A. performed the literature search; Modification and the proofreading of the manuscript have been done by S.H., A.K., S.H.M. and A.W. The manuscript was prepared under the supervision of S.H. and S.H.M. All authors have read and agreed to the published version of the manuscript.

Funding: This research was funded by the China Postdoctoral Science Foundation (grant no. 2020M673368) and the Fundamental Research Funds for the Central Universities (grant no. xjh012020001).

Institutional Review Board Statement: Not applicable.

Informed Consent Statement: Not applicable.

Data Availability Statement: The data presented in this study are available on request from the corresponding author.

Conflicts of Interest: The authors declare no conflict of interest.

References
1. Zhu, H.; Huang, Y.; Lou, X.; Xia, F. Bioinspired super wetting surfaces for biosensing. VIEW 2021, 2, 20200053. [CrossRef]
2. Li, S.; Fan, Y.; Liu, Y.; Niu, S.; Han, Z.; Ren, L. Smart Bionic Surfaces with Switchable Wettability and Applications. J. Bionic Eng. 2021, 3, 473–500. [CrossRef] [PubMed]
3. Shin, S.; Seo, J.; Han, H.; Kang, S.; Kim, H.; Lee, T. Bioinspired Extreme Wetting Surfaces for Biomedical Applications. Materials 2016, 3, 116. [CrossRef]
4. Zhang, Y.; Xia, H.; Kim, E.; Sun, H. Recent developments in superhydrophobic surfaces with unique structural and functional properties. Soft Matter 2012, 8, 11217–11231. [CrossRef]
5. Liu, X.; Liang, Y.; Zhou, F.; Liu, W. Extreme wettability, and tunable adhesion: Biomimicking beyond nature? Soft Matter 2012, 8, 2070–2086. [CrossRef]
6. Bixler, G.; Theiss, A.; Bhushan, B. LSuperhydrophilic TiO$_2$ surface without photocatalytic ee, S. Anti-fouling properties of microstructured surfaces bioinspired by rice leaves and butterfly wings. J. Colloid Interface Sci. 2014, 419, 114–133. [CrossRef] [PubMed]
7. Yao, X.; Song, Y.; Jiang, L. Applications of bioinspired special wettable surfaces. Adv. Mater. 2011, 23, 719–734. [CrossRef] [PubMed]
8. Nature Publishing Group. A Chinese nano-society? Nat. Mater. 2005, 4, 355. [CrossRef] [PubMed]
9. Zorba, V.; Chen, X.; Mao, S. Superhydrophilic TiO$_2$ surface without photocatalytic activation. Appl. Phys. Lett. 2010, 96, 093702. [CrossRef]
10. Wang, P.; Zhang, D.; Qiu, R. Liquid/solid contact mode of super-hydrophobic film in aqueous solution and its effect on corrosion resistance. Corros. Sci. 2012, 54, 77–84. [CrossRef]
11. Bai, H.; Wang, L.; Ju, J.; Sun, R.; Zheng, Y.; Jiang, L. Efficient water collection on integrative bioinspired surfaces with star-shaped wettability patterns. Adv. Mater. 2014, 26, 5025–5030. [CrossRef] [PubMed]
12. Barthlott, W.; Neinhuis, C. Purity of the sacred lotus, or escape from contamination in biological surfaces. Planta 1997, 202, 1–8. [CrossRef]
13. Feng, L.; Li, S.; Li, Y.; Li, H.; Zhang, L.; Zhai, J.; Song, Y.; Liu, B.; Jiang, L.; Zhu, D. Super-hydrophobic surfaces from natural to artificial. **Adv. Mater.** 2002, **14**, 1857–1860. [CrossRef]

14. Parker, A.; Lawrence, C. Water capture by a desert beetle. **Nature** 2001, **414**, 33–34. [CrossRef]

15. Bohn, H.; Federle, W. Insect aquaplaning, Nepenthe’s pitcher plants capture prey with the peristome, a fully wettable water-lubricated anisotropic surface. **Proc. Natl. Acad. Sci. USA** 2004, **101**, 14136–14143. [CrossRef] [PubMed]

16. Ishizaki, T.; Saito, N.; Takai, O. Correlation of cell adhesion behaviors on superhydrophobic superhydrophilic and micropatterned superhydrophobic/superhydrophilic surfaces to their surface chemistry. **Langmuir** 2010, **26**, 8147–8154. [CrossRef] [PubMed]

17. Geyer, F.; Ueda, E.; Liebel, U.; Grau, N.; Levkin, P. Superhydrophobic-superhydrophilic micropatterning: Towards genome-on-a-chip cell microarrays. **Angew. Chem. Int. Ed.** 2011, **50**, 8424–8427. [CrossRef]

18. Piret, G.; Galopin, E.; Cofinnyer, Y.; Boukherroub, R.; Legrand, D.; Slomianny, C. Culture of mammalian cells on patterned superhydrophilic/superhydrophobic silicon nanowire arrays. **Soft Matter** 2011, **7**, 8642–8649. [CrossRef]

19. Oliveira, M.; Neto, A.; Correia, C.; Rial Hermida, M.; Alvarez Lorenzo, C.; Mano, J.F. Superhydrophobic chips for cell spheroids high-throughput generation and drug screening. **ACS Appl. Mater. Interfaces** 2014, **6**, 9488–9495. [CrossRef] [PubMed]

20. Neto, A.I.; Correia, C.R.; Custódio, C.A.; Mano, J. Biomimetic miniaturized platform able to sustain arrays of liquid droplets for high-throughput combinatorial tests. **Adv. Funct. Mater.** 2014, **24**, 5096–5103. [CrossRef]

21. Neto, A.I.; Custodio, C.A.; Song, W.; Mano, J.F. High-throughput evaluation of interactions between biomaterials, proteins and cells using patterned superhydrophobic substrates. **Soft Matter** 2011, **7**, 4147–4151. [CrossRef]

22. Ueda, E.; Levkin, P.A. Emerging applications of superhydrophilic-superhydrophobic micropatterns. **Adv. Mater.** 2013, **25**, 1234–1247. [CrossRef] [PubMed]

23. Oliveira, M.B.; Mano, J.F. High-throughput screening for integrative biomaterials design: Exploring advances and new trends. **Trends Biotechnol.** 2014, **32**, 627–636. [CrossRef]

24. Kang, S.M.; Lee, C.; Kim, H.N.; Lee, B.; Lee, E.; Kwak, M.K.; Suh, K.Y. Directional oil sliding surfaces with hierarchical anisotropic groove microstructures. **Adv. Mater.** 2013, **25**, 5756–5761. [CrossRef]

25. Wong, T.S.; Kang, S.H.; Tang, S.K.Y.; Smythe, E.J.; Hatton, B.D.; Grinthal, A.; Aizenberg, J. Bioinspired self-repairing slippery surfaces with pressure-stable omniphobicity. **Nature** 2011, **477**, 443–447. [CrossRef] [PubMed]

26. Yao, X.; Hu, Y.; Grinthal, A.; Wong, T.S.; Mahadevan, L.; Aizenberg, J. Adaptive fluid-infused porous films with tunable transparency and wettabillity. **Nat. Mater.** 2013, **12**, 529–534. [CrossRef]

27. Salgado, C.L.; Oliveira, M.B.; Mano, J.F. Combinatorial cell-3D biomaterials cytompatibility screening for tissue engineering using bioinspired superhydrophobic substrates. **Integr. Biol.** 2012, **4**, 318–327. [CrossRef] [PubMed]

28. Seo, J.; Lee, J.S.; Lee, K.; Kim, D.; Yang, K.; Shin, S.; Mahata, C.; Jung, H.B.; Lee, W.; Cho, S.W.; et al. Switchable water-adhesive, superhydrophobic palladium-layered silicon nanowires potentiate the angiogenic efficacy of human stem cell spheroids. **Adv. Mater.** 2014, **26**, 7043–7050. [CrossRef]

29. Frescauf, L.R.; McLane, J.; Sharma, H.; Khine, M. Shrink-induced superhydrophobic and antibacterial surfaces in consumer plastics. **PLoS ONE** 2012, **7**, e40987. [CrossRef] [PubMed]

30. Leslie, D.C.; Waterhouse, A.; Berthet, J.; Valentin, T.M.; Watters, A.L.; Jain, A.; Kim, P.; Hatton, B.D.; Nedder, A.; Donovan, K.; et al. A bioinspired omniphobic surface coating on medical devices prevents thrombosis and biofouling. **Nat. Biotechnol.** 2014, **32**, 1134–1140. [CrossRef] [PubMed]

31. Huang, Y.; Li, F.; Qin, M.; Jiang, L.; Song, Y. A multi-stopband photonic-crystal microchip for high-performance metal-ion recognition based on fluorescent detection. **Angew. Chem. Int. Ed. Engl.** 2013, **52**, 7296–7299. [CrossRef] [PubMed]

32. Hou, J.; Zhang, H.; Yang, Q.; Li, M.; Song, Y.; Jiang, L. Bioinspired photonic-crystal microchip for fluorescent ultratrace detection. **Angew. Chem. Int. Ed. Engl.** 2013, **53**, 5791–5795. [CrossRef] [PubMed]

33. Xu, L.P.; Chen, Y.; Yang, G.; Shi, W.; Dai, B.; Li, G.; Cao, Y.; Wen, Y.; Zhang, X. Ultratrace DNA Detection Based on the Condensing-Enrichment Effect of Superwettatable Microchips. **Adv. Mater.** 2015, **27**, 6878–6884. [CrossRef] [PubMed]

34. Hussain, S.; Zhao, H.; Zhou, L.; Zhou, X.; Iyer, P.K.; Lv, F.; Liu, L.; Wang, S. An optoelectronic device for rapid monitoring of creatine kinase using cationic conjugated polyelectrolyte. **Adv. Mater. Technol.** 2019, **4**, 1900361. [CrossRef]

35. Hussain, S.; Lv, F.; Qi, R.; Senthilkumar, T.; Zhao, H.; Chen, Y.; Liu, L.; Wang, S. Förster resonance energy transfer mediated rapid and synergistic discrimination of bacteria over fungi using a cationic conjugated glycopolymer. **ACS Appl. Bio Mater.** 2020, **3**, 20–28. [CrossRef]

36. Hussain, S.; Malik, A.H.; Iyer, P.K. FRET-assisted selective detection of flavins via cationic conjugated polyelectrolyte under physiological conditions. **J. Mater. Chem. B** 2016, **4**, 4439–4446. [CrossRef]

37. Chen, Y.; Xu, L.P.; Meng, J. Superwettatable microchips with improved spot homogeneity toward sensitive biosensing. **Biosens. Bioelectron.** 2018, **102**, 418–424. [CrossRef] [PubMed]

38. Hussain, S.; Malik, A.H.; Iyer, P.K. Highly precise detection, discrimination, and removal of anionic surfactants over the full pH range via cationic conjugated polymer: An efficient strategy to facilitate illicit-drug analysis. **ACS Appl. Mater. Interfaces** 2015, **7**, 3189–3198. [CrossRef] [PubMed]

39. Wu, T.; Xu, T.; Chen, Y.; Yang, Y.; Xu, L.; Zhang, X.; Wang, S. Renewable superwettatable biochip for miRNA detection. **Sens. Actuators B** 2018, **258**, 715–721. [CrossRef]

40. Zimmermann, J.; Reifler, F.A.; Fortunato, G.; Gerhardt, C.; Seeger, S. A simpl one-step approach to durable and robust superhydrophobic textiles. **Adv. Funct. Mater.** 2008, **18**, 3662–3669. [CrossRef]
41. Wang, L.; Zhang, X.; Li, B.; Sun, P.; Yang, J.; Xu, H.; Liu, Y. Superhydrophobic and ultraviolet-blocking cotton textiles. *ACS Appl. Mater. Interfaces* **2011**, *3*, 1277–1281. [CrossRef] [PubMed]

42. Wu, D.; Wu, S.; Chen, Q.; Zhang, Y.; Yao, J.; Yao, X.; Niu, L.G.; Wang, J.; Jiang, L.; Sun, H.B. Curvature-driven reversible in situ switching between pinned and roll-down superhydrophobic states for water droplet transportation. *Adv. Mater.* **2011**, *23*, 545–549. [CrossRef] [PubMed]

43. Yin, Y.; Liu, T.; Chen, S.; Liu, T.; Cheng, S. Structure stability and corrosion inhibition of super-hydrophobic film on aluminium in seawater. *Appl. Surf. Sci.* **2008**, *255*, 2978–2984. [CrossRef]

44. Grignard, B.; Vaillant, A.; de Coninck, J.; Piens, M.; Jonas, A.M.; Detrembleur, C.; Jerome, C. Electrospinning of a functional perfluorinated block copolymer as a powerful route for imparting superhydrophobicity and corrosion resistance to aluminium substrates. *Langmuir* **2011**, *27*, 335–342. [CrossRef] [PubMed]

45. Cui, Z.; Zhang, F.; Wang, L.; Xu, S.; Guo, X. In situ crystallized zirconium phenylphosphonate films with crystals vertically to the substrate and their hydrophobic, dielectric, and anticorrosion properties. *Langmuir* **2010**, *26*, 179–182. [CrossRef] [PubMed]

46. Liu, H.; Szunerits, S.; Xu, W.; Boukherroub, R. Preparation of superhydrophobic coatings on zinc as effective corrosion barriers. *ACS Appl. Mater. Interfaces* **2009**, *1*, 1150–1153. [CrossRef] [PubMed]

47. Wenzel, R.N. Resistance of solid surfaces to wetting by water. *Ind. Eng. Chem.* **1936**, *28*, 998–994. [CrossRef]

48. Cassie, A.B.D.; Baxter, S. Wettability of porous surfaces. *Trans. Faraday Soc.* **1944**, *40*, 546–551. [CrossRef]

49. Young, T. An essay on the cohesion of fluids. *Philos. Trans. R. Soc. Lond.* **1805**, *95*, 65–87.

50. Neinhuis, C.; Barthlott, W. Characterization and distribution of water-repellent, self-cleaning plant surfaces. *Philos. Trans. R. Soc. Lond.* **1997**, *352*, 667–677. [CrossRef]

51. Kim, T.; Tahk, D.; Lee, H.H. Wettability-controllable super water- and moderately oil-repellent surface fabricated by wet chemical etching. *Langmuir* **2009**, *25*, 6576–6579. [CrossRef] [PubMed]

52. Seo, J.; Lee, S.; Han, H.; Chung, Y.; Lee, J.; Kim, D.; Kim, Y.W.; Lim, S.; Lee, T. Reversible wettability control of silicon nanowire surfaces from superhydrophilicity to superhydrophobicity. *Thin Solid Films* **2013**, *527*, 179–185. [CrossRef]

53. Piret, G.; Coffinier, Y.; Roux, C.; Melnyk, O.; Boukherroub, R. Biomolecule and nanoparticle transfer on patterned and heterogeneously wetted superhydrophobic silicon nanowire surfaces. *Langmuir* **2008**, *24*, 1670–1672. [CrossRef] [PubMed]

54. Coffinier, Y.; Piret, G.; Das, R.; Boukherroub, R. Effect of surface roughness and chemical composition on the wetting properties of silicon-based substrates. *Comptes Rendus Chim.* **2013**, *16*, 65–72. [CrossRef]

55. Dupré, M.; Enjalbal, C.; Cantel, S.; Martinez, J.; Megouda, N.; Hadjersi, T.; Boukherroub, R.; Coffinier, Y. Investigation of silicon-based nanostucture morphology and chemical termination on laser desorption ionization mass spectrometry performance. *Anal. Chem.* **2012**, *84*, 10637–10644. [CrossRef] [PubMed]

56. Nguye, T.P.N.; Dufour, R.; Thomy, V.; Senez, V.; Boukherroub, R.; Coffinier, Y. Fabrication of superhydrophobic and highly oleophobic silicon-based surfaces via electroless etching method. *Appl. Surf. Sci.* **2014**, *295*, 38–43. [CrossRef]

57. Yu, L.Y.; Shen, H.M.; Xu, Z.L. PVDF-TiO<sub>2</sub> composite hollow fiber ultrafiltration membranes prepared by TiO<sub>2</sub> sol-gel method and blending method. *J. Appl. Polym. Sci.* **2009**, *113*, 1763–1772. [CrossRef]

58. He, G.; Wan, K. The super hydrophobicity of ZnO nanorods fabricated by electrochemical deposition method. *Appl. Surf. Sci.* **2011**, *257*, 6590–6594. [CrossRef]

59. Thomas, Y.R.J.; Benayad, A.; Schroder, M.; Morin, A.; Pauchet, J. New method for super hydrophobic treatment of gas diffusion layers for proton exchange membrane fuel cells using electrochemical reduction of diazonium salts. *ACS Appl. Mater. Interfaces* **2011**, *3*, 15068–15077. [CrossRef] [PubMed]

60. Tadanaga, K.; Morinaga, J.; Matsuda, A.; Minami, T. Superhydrophobic-superhydrophilic micropatterning on flowerlike alumina coating film by the sol-gel method. *Chem. Mater.* **2010**, *22*, 590–592. [CrossRef]

61. Kim, H.C.; Kreller, C.R.; Tran, K.A.; Sisodiya, V.; Angelos, S.; Wallraff, G.; Swanson, S.; Miller, R.D. Nanoporous thin films with hydrophilicity-contrasted patterns. *Chem. Mater.* **2004**, *16*, 4267–4272. [CrossRef]

62. Kavalenka, M.N.; Hopf, A.; Schneider, M.; Worgull, M.; Höscher, H. Wood-based microfibred superhydrophobic and underwater superoleophobic surfaces for oil/water separation. *RSC Adv.* **2014**, *4*, 31079–31083. [CrossRef]

63. Zhu, Y.; Zhang, J.; Zheng, Y.; Huang, Z.; Feng, L.; Jiang, L. Stable superhydrophobic and conductive polyaniline/poly styrene films for corrosive environments. *Adv. Funct. Mater.* **2006**, *16*, 568–574. [CrossRef]

64. Sarkar, M.K.; Bal, K.; He, F.; Fan, J. Design of an outstanding super-hydrophobic surface by electro-spinning. *Appl. Surf. Sci.* **2011**, *257*, 7003–7009. [CrossRef]

65. Nuraie, N.; Khan, W.S.; Lei, Y.; Ceylan, M.; Asmatulu, R. Superhydrophobic electrospun nanofibers. *J. Mater. Chem. A* **2013**, *1*, 1929–1946. [CrossRef]

66. Tadanaga, K.; Katata, N.; Minami, T. Formation process of super-water-repellent Al<sub>2</sub>O<sub>3</sub> coating films with high transparency by the sol-gel method. *J. Am. Ceram. Soc.* **1997**, *80*, 3213–3216. [CrossRef]

67. Shirtcliffe, N.J.; McHale, G.; Newton, M.I.; Perry, C.C. Intrinsically Superhydrophobic organosilica sol-gel foams. *Langmuir* **2003**, *19*, 5626–5631. [CrossRef]

68. Tsai, M.Y.; Hsu, C.C.; Chen, P.H.; Lin, C.S.; Chen, A. Surface modification on a glass surface with a combination technique of sol-gel and air brushing technique. *Appl. Surf. Sci.* **2011**, *257*, 8640–8646. [CrossRef]
69. Mahidak, S.; Mahidak, D.B.; Kavale, M.S.; Parale, V.G.; Wagh, P.B.; Barshilia, H.; Gupta, S.; Hegde, N.D.; Rao, A.V. Thermally stable and transparent superhydrophobic sol-gel coatings by spray method. *J. Sol. Gel Sci. Technol.* 2012, 63, 580–586. [CrossRef]

70. Latthe, S.S.; Terashima, C.; Nakata, K.; Sakai, M.; Fujishima, A. Development of sol-gel processed semi-transparent and self-cleaning superhydrophobic coatings. *J. Mater. Chem. A* 2014, 2, 5548–5553. [CrossRef]

71. Jiang, L.; Zhao, Y.; Zhai, J. A lotus-leaf-like superhydrophobic surface: A porous microsphere/nanofiber composite film prepared by electrohydrodynamics. *Angew. Chem. Int. Ed.* 2004, 11, 4438–4441. [CrossRef]

72. Li, Y.; Huang, X.J.; Heo, S.H.; Li, C.C.; Choi, Y.K.; Cai, W.P.; Cho, S.O. Superhydrophobic bionic surfaces with hierarchical microsphere/SWCNT composite arrays. *Langmuir* 2007, 23, 2169–2174. [CrossRef]

73. Sun, M.; Luo, C.; Xu, L.; Ji, H.; Ouyang, Q.; Yu, D.; Chen, Y. Artificial lotus leaf by nanocasting. *Langmuir* 2005, 21, 8978–8981. [CrossRef]

74. Feng, L.; Zhang, Y.; Xi, J.; Zhu, Y.; Wang, N.; Xia, F.; Jiang, L. A superhydrophobic state with high adhesive force. *Langmuir* 2008, 24, 4114–4119. [CrossRef] [PubMed]

75. Lai, Y.; Gao, X.; Zhuang, H.; Huang, J.; Lin, C.; Jiang, L. Designing superhydrophobic porous nanostructures with tunable water adhesion. *Adv. Mater.* 2009, 21, 3799–3803. [CrossRef]

76. Zhao, Y.; Lu, Q.; Li, M.; Li, X. Anisotropic wetting characteristics on submicrometer-scale periodic grooved surface. *Langmuir* 2007, 23, 6212–6217. [CrossRef] [PubMed]

77. Xia, D.; Brueck, S.R.J. Strongly anisotropic wetting on one-dimensional nanopatterned surfaces. *Nano Lett.* 2008, 8, 2819–2824. [CrossRef] [PubMed]

78. Wu, H.; Zhang, R.; Sun, Y.; Lin, D.; Sun, Z.; Pan, W.; Downs, P. Biomimetic nanofiber patterns with controlled wettability. *Soft Matter* 2008, 4, 2429–2433. [CrossRef] [PubMed]

79. Gao, J.; Liu, Y.; Xu, H.; Wang, Z.; Zhang, X. Mimicking biological structured surfaces by phase-separation micromolding. *Langmuir* 2009, 25, 4365–4369. [CrossRef]

80. Zhang, X.; Zhang, J.; Ren, Z.; Li, X.; Zhang, X.; Zhu, D.; Wang, T.; Tian, T.; Yang, B. Morphology and wettability control of silicon cone arrays using colloidal lithography. *Langmuir* 2009, 25, 7375–7382. [CrossRef]

81. Wu, S.Z.; Wu, D.; Yao, J.; Chen, Q.D.; Wang, J.N.; Niu, L.; Fang, H.H.; Sun, H.B. One-step preparation of regular micropearl arrays for two-direction controllable anisotropic wetting. *Langmuir* 2010, 26, 12012–12016. [CrossRef] [PubMed]

82. Wu, D.; Wang, J.N.; Wu, S.Z.; Chen, Q.D.; Zhao, S.; Zhang, H. Three-level biomimetic rice-leaf surfaces with controllable anisotropic sliding. *Adv. Funct. Mater.* 2011, 21, 2927–2932. [CrossRef]

83. Wu, D.; Wang, J.N.; Wu, S.Z.; Chen, Q.D.; Zhao, S.; Zhang, H. Three-level biomimetic rice-leaf surfaces with controllable anisotropic sliding. *Adv. Funct. Mater.* 2011, 21, 2927–2932. [CrossRef]

84. Malvadkar, N.A.; Hancock, M.J. An engineered anisotropic nanofilm with unidirectional wetting properties. *Adv. Mater.* 2008, 20, 6212–6217. [CrossRef] [PubMed]

85. Bixler, G.D.; Bhushan, B. Bioinspired rice leaf and butterfly wing surface structures combining shark skin and lotus effects. *Soft Matter* 2012, 8, 11271–11284. [CrossRef]

86. Hussain, S.; Malik, A.H.; Afroz, M.A.; Iyer, P.K. Ultrasensitive detection of nitroexplosive-picric acid via a conjugated polyelectrolyte in aqueous media and solid support. *Chem. Commun.* 2015, 51, 7207–7210. [CrossRef] [PubMed]

87. Chen, X.; Hussain, S.; Hao, Y.; Tian, X.; Gao, R. Review-Recent advances of signal amplified smart conjugated polymers for optical detection on solid support. *ECS J. Solid State Sci. Technol.* 2021, 10, 037006. [CrossRef]
98. Zhai, L.; Berg, M.C.; Cebeci, F.; Kim, Y.; Milwid, J.M.; Rubner, M.F.; Cohen, R.E. Patterned superhydrophobic surfaces Toward a synthetic mimic of the namib desert beetle. *Nano Lett.* 2006, 6, 1213–1217. [CrossRef] [PubMed]

99. Li, J.S.; Ueda, E.; Nallapaneni, A.; Li, L.X.; Levkin, P.A. Printable superhydrophilic-superhydrophobic micropatterns based on supported lipid layers. *Langmuir* 2012, 28, 8286–8291. [CrossRef] [PubMed]

100. Zhang, L.; Wu, J.; Hedhili, M.N.; Yang, X.; Wang, P. Inkjet printing for direct micropatterning of a superhydrophobic surface, toward biomimetic fog harvesting surfaces. *J. Mater. Chem. A* 2015, 3, 2844–2852. [CrossRef]

101. Beutel, R.G.; Gorb, S.N. Ultrastructure of attachment specializations of hexapods (arthropoda) Evolutionary patterns inferred from a revised ordinal phylogeny. *J. Zool. Syst. Evol. Res.* 2001, 39, 177–207. [CrossRef]

102. Gorb, E.; Kastner, V.; Peressadko, A.; Arzt, E.; Gauze, L.; Rowe, N.; Gorb, S. Structure and properties of the glandular surface in the digestive zone of the pitcher in the carnivorous plant nepenthes ventrata and its role in insect trapping and retention. *J. Exp. Biol.* 2004, 207, 2947–2963. [CrossRef] [PubMed]

103. Bauer, U.; Federle, W. The insect-trapping rim of nepenthes pitchers. *Plant Signal. Behav.* 2009, 4, 1019–1023. [CrossRef] [PubMed]

104. Xiao, L.; Li, J.; Mieszkin, S.; Di Fino, A.; Clare, A.S. Slippery liquid-infused porous surfaces showing marine antibiofouling properties. *ACS Appl. Mater. Interfaces* 2013, 5, 10074–10080. [CrossRef]

105. Wilson, P.W.; Lu, W.; Xu, H.; Kim, P.; Kreder, M.J.; Alvarenga, J.; Aizenberg, J. Inhibition of ice nucleation by slippery liquid-infused porous surfaces (SLIPS) for snow and ice protection. *Adv. Mater.* 2015, 27, 6704–6711. [CrossRef]

106. Li, C.; Cheng, F.; Lv, J.; Zhao, Y.; Liu, M.; Jiang, L.; Yu, Y. Light-controlled quick switch of adhesion on a micro-arrayed liquid crystal polymer superhydrophobic film. *Soft Matter* 2012, 8, 3730–3733. [CrossRef]

107. Seo, J.; Lee, S.; Han, H.; Jung, H.B.; Hong, J.; Song, G.; Cho, S.M.; Park, C.; Lee, W.; Lee, T. Gas-driven ultrafast reversible switching of super-hydrophobic adhesion on palladium-coated silicon nanowires. *Adv. Mater.* 2013, 25, 4139–4144. [CrossRef] [PubMed]

108. Zhu, Y.; Li, J.; He, H.; Wan, M.; Jiang, L. Reversible wettability switching of polyaniline-coated fabric, triggered by ammonia gas. *Macromol. Rapid Comm.* 2007, 28, 2230–2236. [CrossRef]

109. Cheng, Z.; Feng, L.; Jiang, L. Tunable adhesive superhydrophobic surfaces for superparamagnetic microdroplets. *Adv. Funct. Mater.* 2008, 18, 3219–3225. [CrossRef]

110. Li, C.; Guo, R.; Jiang, X.; Hu, S.; Li, L.; Cao, X.; Yang, H.; Song, Y.; Ma, Y.; Jiang, L. Reversible switching of water-droplet mobility on a superhydrophobic surface based on a phase transition of a side-chain liquid-crystal polymer. *Adv. Mater.* 2009, 21, 4254–4258. [CrossRef]

111. Liu, X.; Ye, Q.; Yu, B.; Liang, Y.; Liu, W.; Zhou, F. Switching water droplet adhesion using responsive polymer brushes. *Langmuir* 2010, 26, 12377–12382. [CrossRef]

112. Ying, G.; Sun, T. Chirality-triggered wettability switching on a smart polymer surface. *Adv. Mater.* 2011, 23, 1615–1620. [CrossRef]

113. Liu, X.; Cai, M.; Liang, Y.; Zhou, F.; Liu, W. Photo-regulated stick-slip switch of water droplet mobility. *Soft Matter* 2011, 7, 3331–3336. [CrossRef]

114. Li, C.; Zhang, Y.; Ju, J.; Cheng, F.; Liu, M.; Jiang, L.; Yu, Y. In situ fully light-driven switching of superhydrophobic adhesion. *Adv. Funct. Mater.* 2012, 22, 760–763. [CrossRef]

115. Liu, Y.; Wang, X.; Fei, B.; Hu, H.; Lai, C.; Xin, J.H. Bioinspired, stimuli-responsive, multifunctional superhydrophobic surface with directional wetting, adhesion, and transport of water. *Adv. Funct. Mater.* 2015, 25, 5047–5056. [CrossRef]

116. Verplank, N.; Coffinier, Y.; Thomy, V.; Boukherroub, R. Wettability switching techniques on superhydrophobic surfaces. *Nanoscale Res. Lett.* 2007, 2, 577–596. [CrossRef]

117. Gras, S.L.; Mahmud, T.; Rosengarten, G.; Mitchell, A.; Kalantar-Zadeh, K. Intelligent control of surface hydrophobicity. *ChemPhysChem* 2007, 8, 2036–2050. [CrossRef]

118. Wang, D.; Liu, Y.; Liu, X.; Zhou, F.; Liu, W.; Xue, Q. Towards a tunable and switchable water adhesion on a TiO2 nanotube film with patterned wettability. *Chem. Commun.* 2009, 7018–7020. [CrossRef]

119. Alves, N.M.; Shi, J.; Oramas, E.; Santos, J.; Tomás, H.; Mano, J.F. Bioinspired superhydrophobic poly(l-lactic acid) surfaces control bone marrow derived cells adhesion and proliferation. *J. Biomed. Mater. Res. A* 2009, 9, 480–488. [CrossRef]

120. Oliveira, S.M.; Song, W.; Alves, N.M.; Mano, J.F. Chemical modification of bioinspired superhydrophobic polystyrene surfaces to control cell attachment/proliferation. *Soft Matter* 2011, 7, 8932–8941. [CrossRef]

121. Ballester Beltran, J.; Rico, P.; Moratal, D.; Song, W.; Mano, J.F. Role of superhydrophobicity in the biological activity of fibronectin at the cell-material interface. *Soft Matter* 2011, 7, 10803–10811. [CrossRef]

122. Auda, P.; Ueda, E.; Levkin, P.A. Facile and multiple replication of superhydrophilic-superhydrophobic patterns using adhesive tape. *ACS Appl. Mater. Interfaces* 2013, 5, 8053–8057. [CrossRef]

123. Efremov, A.N.; Stanganello, E.; Welle, A.; Scholpp, S.; Levkin, P.A. Micropatterned superhydrophobic structures for the simultaneous culture of multiple cell types and the study of cell-cell communication. *Biomaterials* 2013, 34, 1757–1763. [CrossRef] [PubMed]

124. Oliveira, S.M.; Alves, N.M.; Mano, J.F. Cell interactions with superhydrophilic and superhydrophobic surfaces. *J. Adhes. Sci. Technol.* 2014, 28, 843–863. [CrossRef]

125. Li, H.; Yang, Q.; Li, G.; Li, M.; Wang, S.; Song, Y. Splitting a droplet for femtoliter liquid patterns and single cell isolation. *ACS Appl. Mater. Interfaces* 2015, 7, 9060–9065. [CrossRef] [PubMed]
155. Privett, B.J.; Youn, J.; Hong, S.A.; Lee, J.; Han, J.; Shin, J.H.; Schoenfisch, H. Antibacterial fluorinated silica colloid superhydrophobic surfaces. *Langmuir* 2011, 27, 9597–9601. [CrossRef] [PubMed]

156. Fadeeva, E.; Truong, V.K.; Stiesch, M.; Chichkov, B.N.; Crawford, R.; Wang, J.; Ivanova, E.P. Bacterial retention on superhydrophobic titania surfaces fabricated by femtosecond laser ablation. *Langmuir* 2011, 27, 3012–3019. [CrossRef]

157. Liu, T.; Dong, L.; Liu, T.; Yin, Y. Investigations on reducing microbiologically influenced corrosion of aluminum by using super-hydrophobic surfaces. *Electrochim. Acta* 2010, 55, 5281–5285. [CrossRef]

158. Ye, W.; Shi, Q.; Hou, J.; Jin, J.; Fan, Q.; Wong, S.; Xu, X.; Yin, J. Superhydrophobic coating of elastomer on different substrates using a liquid template to construct a biocompatible and antibacterial surface. *J. Mater. Chem. B* 2014, 2, 7186–7191. [CrossRef]

159. Grinthal, A.; Aizenberg, J. Mobile interfaces. Liquids as a perfect structural material for multifunctional, antifouling surfaces. *Chem. Mater.* 2014, 26, 698–706. [CrossRef]

160. Epstein, A.K.; Wong, T.S.; Belisle, R.A.; Boggs, E.M.; Aizenberg, J. Liquid-infused structured surfaces with exceptional anti-biofouling performance. *Proc. Natl. Acad. Sci. USA* 2012, 109, 13182–13187. [CrossRef]

161. Balu, B.; Berry, A.D.; Hess, D.W.; Breedveld, V. Patterning of superhydrophobic paper to control the mobility of micro-liter drops for two-dimensional lab-on-paper applications. *Lab Chip* 2009, 9, 3066–3075. [CrossRef] [PubMed]

162. Mertaniemi, H.; Jokinen, V.; Sainiemi, L.; Franussa, S.; Marmur, A.; Ikkala, O.; Ras, R.H.A. Superhydrophobic tracks for low-friction, guided transport of water droplets. *Adv. Mater.* 2011, 23, 2911–2914. [CrossRef] [PubMed]

163. You, I.; Kang, S.M.; Lee, S.; Cho, Y.O.; Kim, J.B.; Lee, S.B.; Nam, Y.S.; Lee, H. Polydopamine microfluidic system toward a two-dimensional, gravity-driven mixing device. *Angew. Chem. Int. Ed.* 2012, 51, 6126–6130. [CrossRef] [PubMed]

164. Sousa, M.P.; Mano, J.F. Superhydrophobic paper in the development of disposable labware and lab-on-paper devices. *ACS Appl. Mater. Interfaces* 2013, 5, 3731–3737. [CrossRef]

165. Elsharkawy, M.; Schutzius, T.M.; Megaridis, C.M. Inkjet patterned superhydrophobic paper for open-air surface microfluidic devices. *Lab Chip* 2014, 14, 1168–1175. [CrossRef] [PubMed]

166. Zhao, Y.; Xu, Z.; Niu, H.; Wang, X.; Lin, T. Magnetic liquid marbles, Toward “lab in a droplet”. *Adv. Funct. Mater.* 2015, 25, 437–444. [CrossRef] [PubMed]

167. Kim, D.; Seo, J.; Shin, S.; Lee, S.; Lee, K.; Cho, H.; Shim, W.; Lee, H.; Lee, T. Reversible liquid adhesion switching of superamphiphobic Pd-decorated Ag dendrites via gas-induced structural changes. *Chem. Mater.* 2015, 27, 4964–4971. [CrossRef]

168. Jonsson Niedziolka, M.; Lapierre, F.; Coffinier, Y.; Parry, S.J.; Zoueshtiagh, F.; Foat, T.; Thomy, V.; Boukherroub, R. EWOD driven cleaning of biocaptes on hydrophobic and superhydrophobic surfaces. *Lab Chip* 2011, 11, 490–496. [CrossRef]

169. Lapière, F.; Preit, G.; Drobecq, H.; Melnyk, O.; Coffinier, Y.; Thomy, V.; Boukherroub, R. High sensitive matrix-free mass spectrometry analysis of peptides using silicon nanowires-based digital microfluidic device. *Lab Chip* 2011, 11, 1620–1628. [CrossRef]

170. Lapière, F.; Harnois, M.; Coffinier, Y.; Boukherroub, R.; Thomy, V. Split and flow: Reconfigurable capillary connection for digital microfluidic devices. *Lab Chip* 2014, 14, 3589–3593. [CrossRef] [PubMed]

171. Li, L.; Breedveld, V.; Hess, D.W. Hysteresis controlled water droplet splitting on superhydrophobic paper. *Colloid Polym. Sci.* 2013, 291, 417–426. [CrossRef] [PubMed]

172. You, I.; Lee, T.G.; Nam, Y.S.; Lee, J.; Han, J.; Shin, J.H.; Schoenfisch, H. Antibacterial fluorinated silica colloid superhydrophobic surfaces. *Langmuir* 2011, 27, 9597–9601. [CrossRef] [PubMed]

173. Seo, J.; Lee, S.K.; Lee, J.; Seung, I.J.; Kwon, H.; Cho, S.W.; Ahn, J.H.; Lee, T. Path-programmable water droplet manipulations on an adhesion controlled superhydrophobic surface. *Sci. Rep.* 2015, 5, 12326. [CrossRef] [PubMed]

174. Chen, X.; Liu, C.; Xu, Z.; Pan, Y.; Liu, J.; Du, L. An effective PDMS microfluidic chip for chemiluminescence detection of cobalt (II) in water. *Microsyst. Technol.* 2012, 19, 99–103. [CrossRef]

175. Sima, F.; Xu, J.; Wu, D.; Sugio, K. Ultrafast laser fabrication of functional biochips: New avenues for exploring 3D micro- and nano-environments. *Microachines* 2017, 8, 40. [CrossRef]

176. He, J.; Zhao, Y.; Yuan, M.; Hou, L.; Abbas, A.; Xue, M.; Ma, X.; He, J.; Qu, M. Fabrication of durable polytetrafluoroethylene superhydrophobic materials with recyclable and self-cleaning properties on various substrates. *J. Coat. Technol. Res.* 2020, 17, 755–763. [CrossRef]

177. He, J.; Zhang, Y.; Zhou, Y.; Wang, J.; Zhao, Y.; Ma, L.; Abbas, A.; Qu, M. A facile approach to fabricate the durable and buoyant superhydrophobic fabric for efficient oil/water separation. *Fibers Polym.* 2019, 20, 1003–1010. [CrossRef]

178. Stetsyshyn, Y.; Raczkowska, J.; Budkowski, A.; Awiuki, K.; Kostruba, A.; Nastys, S.; Harhay, K.; Lyckhovskyy, E.; Ohar, H.; Nastishin, Y. Cholesterol-based grafted polymer brushes as alignment coating with temperature-tuned anchoring for nematic liquid crystals. *Langmuir* 2016, 32, 11029–11038. [CrossRef]

179. Pan, Z.; Cheng, F.; Zhao, B. Bioinspired polymeric structures with special wettability and their applications. *Polymers* 2017, 9, 725. [CrossRef] [PubMed]

180. Raczkowska, J.; Stetsyshyn, Y.; Awiuki, K.; Lekak, M.; Marzec, M.; Harhay, K.; Ohar, H.; Ostapiv, D.; Sharand, M.; Yaremchuk, I.; et al. Temperature-responsive grafted polymer brushes obtained from renewable sources with potential application as substrates for tissue engineering. *Appl. Surf. Sci.* 2017, 407, 546–554. [CrossRef] [PubMed]
182. Wang, R.; Wu, F.; Yu, F.; Zhu, J.; Gao, X.; Jiang, L. Anti-vapor-penetration and condensate microdrop self-transport of superhydrophobic oblique nanowire surface under high subcooling. *Nano Res.* 2021, 14, 1429–1434. [CrossRef]

183. Wang, D.; Sun, Q.; Hokkanen, M.J.; Zhang, C.; Lin, F.-Y.; Liu, Q.; Zhu, S.-P.; Zhou, T.; Chang, Q.; He, B.; et al. Design of robust superhydrophobic surfaces. *Nature* 2020, 582, 55–59. [CrossRef] [PubMed]

184. Zeng, J.; Zhang, Y.; Zeng, T.; Aleisa, R.; Qiu, Z.; Chen, Y.; Huang, J.; Wang, D.; Yan, Z.; Yin, Y. Anisotropic plasmonic nanostructures for colorimetric sensing. *Nano Today* 2020, 32, 100855. [CrossRef]

185. Xu, T.; Xu, L.-P.; Zhang, X.; Wang, S. Bioinspired superwettable micropatterns for biosensing. *Chem. Soc. Rev.* 2019, 48, 3153–3165. [CrossRef] [PubMed]