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A statistical model for the wettability of surfaces with heterogeneous pore geometries

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Abstract

We describe a new approach to modeling the wetting behavior of micro- and nano-textured surfaces with varying degrees of geometrical heterogeneity. Surfaces are modeled as pore arrays with a Gaussian distribution of sidewall reentrant angles and a characteristic wall roughness. Unlike conventional wettability models, our model considers the fraction of a surface’s pores that are filled at any time, allowing us to capture more subtle dependences of a liquid’s apparent contact angle on its surface tension. The model has four fitting parameters and is calibrated for a particular surface by measuring the apparent contact angles between the surface and at least four probe liquids. We have calibrated the model for three heterogeneous nanoporous surfaces that we have fabricated: a hydrothermally grown zinc oxide, a film of polyvinylidene fluoride (PVDF) microspheres formed by spinodal decomposition, and a polytetrafluoroethylene (PTFE) film with pores defined by sacrificial polystyrene microspheres. These three surfaces show markedly different dependences of a liquid’s apparent contact angle on the liquid’s surface tension, and the results can be explained by considering geometric variability. The highly variable PTFE pores yield the most gradual variation of apparent contact angle with probe liquid surface tension. The PVDF microspheres are more regular in diameter and, although connected in an irregular manner, result in a much sharper transition from non-wetting to wetting behavior as surface tension reduces. We also demonstrate, by terminating porous zinc oxide with three alternative hydrophobic molecules, that a single geometrical model can capture a structure’s wetting behavior for multiple surface chemistries and liquids. Finally, we contrast our results with those from a highly regular, lithographically-produced structure which shows an extremely sharp dependence of wettability on surface tension. This new model could be valuable in designing and evaluating processes for manufacturing liquid-repellent surfaces on an industrial scale.

Introduction

It has been well established that tuning a surface’s micro- and nano-structure can greatly increase the apparent contact angle of a liquid on the surface, rendering it superhydrophobic or superoleophobic [1–3]. One important way in which the apparent contact angle, and hence liquid repellence, can be increased is by reducing the fraction of a liquid’s surface that comes into contact with solid material—a relationship that was first modeled by Cassie and Baxter [4]. The aim of reducing contact area can be achieved by patterning a surface with micro- and nano-scale pillars or porous structures that support the liquid. Giving these surface features a re-entrant or under-cut shape where they meet the liquid (Ψ > 90° in figure 1(c)) is particularly valuable, because this directs the surface tension of the liquid out of the pores and enables the surface to repel the liquid, even including low-surface-tension oils and fluorinated liquids that will wet almost any smooth surface. When re-entrant features are combined with a low-surface-energy termination, such as a fluorinated molecule, the repellence is further enhanced.

There have been many experimental studies of how re-entrant geometries can impart high liquid-repellence to a surface. Geometrically highly regular hydrophobic and oleophobic surfaces have been made
lithographically. The microfabrication of re-entrant structures with this purpose was originally demonstrated by Tuteja, who patterned and etched SiO2 disks on a silicon substrate, and then under-cut the disks with an isotropic silicon etch [1]. After vapor-phase silanization, these structures were shown to deliver advancing contact angles in excess of 160° with octane. Liu subsequently extended this approach with doubly-reentrant structures that were formed by a multi-stage oxidation and etching process [5]. These structures offer re-entrant angles $\Psi$ in excess of 180° and apparent contact angles above 150° even for liquids with surface tension as low as 10 mN m$^{-1}$.

Although precisely fabricated, highly regular geometries have provided deeper understanding of how to engineer a liquid-repellent surface, many of the potential applications require large, non-flat surfaces to be modified at low cost [6]. The most lucrative of these applications include waterproof and stain-resistant clothing [7–9], food packaging and processing equipment [10], filters for separating oil and water [1, 11], anti-icing coatings for aircraft [12, 13], drag reduction in transport [14], antimicrobial coatings [15, 16], and the enhancement of heat-transfer by promoting dropwise condensation [17–19]. What these applications have in common is a requirement to augment pre-existing materials and geometries with the micro- and nano-structures that will enhance liquid repellence. Standard microfabrication techniques, including photolithography and plasma etching, have limited applicability in these situations since the techniques may require highly polished target surfaces, vacuum processing, and the use of specialized etchant gases.

Traditional manufacturing processes—including stamping, embossing, injection or compression molding, or knurling—may have a role to play in delivering affordable liquid-repellent surfaces. However, re-entrant features are more challenging to manufacture at scale than, for example, simple vertical pillar arrays, because almost any mechanical process for imparting surface texture involves pressing the target material against a tool and then separating them, and so is not capable of creating re-entrant geometries.

Consequently, bottom-up fabrication processes have been widely explored. Prominent among them has been the controlled oxidation of metal surfaces and the precipitation of oxides and hydroxides to create porous films of, for example, anodic alumina [20–22], copper oxide [17, 18], zinc oxide [23–25], and brucite-type cobalt hydroxide [26]. The morphologies of grown and precipitated structures have ranged from porous films [20–22] to stacks of needles with deep sub-micron tip radii [17, 18, 26]. These structures are all irregular in shape and,

![Figure 1. Pore geometry and the effect of its variability on a liquid's apparent contact angle. The classical (a) Wenzel and (b) Cassie–Baxter models assume completely spatially uniform surface feature geometries with a fixed re-entrant angle $\Psi$, resulting in complete wetting (a) where $\theta_0 < \Psi$ or complete non-wetting (b) where $\theta_0 > \Psi$. Real surfaces exhibit a range of effective $\Psi$, such that the fraction of pores filled will depend on $\theta_0$.](image-url)
with the exception of anodic alumina, contain many re-entrant features. Ceramic surfaces are particularly suited to heat-transfer applications because they can be applied directly to a metallic component, the coatings themselves are much more thermally conductive than polymeric materials, and they can be grown in films less than a micrometer thick.

The ability to grow an oxide on to any shape of surface has enabled these structures to be combined with pre-textured surfaces such as woven metal meshes [27]. The oxide nanostructures together with the sub-millimeter features of the woven mesh yields structures with at least two length-scales of roughness, mimicking natural water-repellent materials, such as the lotus leaf [28], that are known to derive their high performance from their hierarchical geometry. Hierarchical yet irregular geometries have been achieved in other ways, including by gold-assisted electroless etching of silicon to create a multi-scale porous surface with overhanging features and a static water contact angle exceeding 160° [29].

Another potentially scalable approach to creating re-entrant structures is to deposit particles or fibers on to the target substrate, by spraying, painting, or dip-coating. For example, Campos sprayed precipitated silica nanoparticles in a fluoropolymer binder [30], Tarwal employed pyrolytic formation of ZnO from a heated, sprayed zinc acetate solution [31], Zhang smeared a polymer chalk of polyvinylbenzene on to a set of target surfaces [32], and Lu sprayed titania nanoparticles on to target substrates [33]. Electrospinning has also been widely employed (e.g. [34]). In all of these cases a porous surface with re-entrant microscopic geometries is created. Additionally, dip-coating of woven meshes with fluoropolymer solutions has been shown to be an effective way of combining a controlled re-entrant structure with a liquid-repellent surface chemistry [3].

Irregular, reentrant-featured surfaces certainly promise to be much less expensive to produce than regular, lithographically defined ones, and can be scaled more readily to larger areas [17], but their distributions of surface feature geometries—and thus their performance—are more challenging to characterize. The behavior of a liquid on a textured surface has conventionally been modeled either as completely wetting the surface (Wenzel mode [35], figure 1(a)) or as sitting entirely on top of the surface and not penetrating any pores (Cassie–Baxter mode [36], figure 1(b)). This binary distinction cannot, however, represent the true behavior of the many types of surfaces whose geometries are spatially heterogeneous, having, for example, pores with a range of wall angles. In these situations, some pores may wet while others remain empty, depending on their local geometries.

Here, therefore, we introduce a new model to characterize geometrically heterogeneous surfaces accurately. Our model takes account of the average wall angle of surface structures as well as the degree of variability of wall angle and the average roughness of the structures. The new model’s crucial contribution is the introduction of a parameter that captures the variability of pore geometry. It thus goes beyond the derivations [36–39] and modifications [40, 41] that have previously been used to specialize the Cassie–Baxter model for particular regular geometries such as woven meshes. All of these previously proposed models are entirely deterministic and regard a surface as being capable of being defined by a small set of fixed dimensional parameters. In contrast, the approach that we describe here incorporates pore-to-pore variation and relates a liquid’s apparent contact angle to the particular fraction of pores that are filled at a given time.

**Modeling framework**

We model a surface as having many equal-volume pores with sidewalls whose re-entrant angles $\Psi$ near their tips approximately follow a Gaussian distribution with mean $\Psi_0$ and standard deviation $\sigma_0$ (figure 1(c)). Let $P(\Psi)$ be the cumulative distribution function for $\Psi$:

$$P(\Psi) \approx \int_{-\infty}^{\Psi} \frac{1}{\sigma_0 \sqrt{2\pi}} e^{-\left(\frac{\Psi' - \Psi_0}{2\sigma_0}\right)^2} \, d\Psi'. \quad (1)$$

In our model, a pore will instantaneously and completely fill when $\theta_0$, the contact angle of the liquid on a flat surface, is less than the re-entrant angle: $\theta_0 < \Psi$. In that situation, the surface tension of the liquid will pull the liquid down into the pore until it fills. Conversely, we assume that a pore remains completely empty when $\Psi > \Psi$ and the surface tension holds the liquid out of the pore. We assume that a liquid droplet making contact with the surface is large enough to encounter many pores. Below the droplet, the proportion of pores filled, $\phi_F$, for a given value of $\theta_0$ is:

$$\phi_F = 1 - P(\theta_0). \quad (2)$$

We then model the overall apparent contact angle, $\theta^*$, of a droplet of liquid on the surface structure using a generalized Cassie–Baxter model [4]. The general Cassie–Baxter model expresses the cosine of the apparent contact angle on a composite surface as the area-weighted sum of the contact angle cosines for each of the components, $i$, of the surface:
\[ \cos \theta^* = \sum_i f_i r_i \cos \theta_{0,i} \]  

where \( f_i \) is the fraction of the surface’s projected area that is occupied by component \( i \), \( r_i \) is the roughness of the component (i.e. the ratio of total surface area to projected area) and \( \theta_{0,i} \) is the contact angle of the liquid on a flat surface with the same surface chemistry as component \( i \).

Our specific model assumes that the projected surface area of the porous film is divided into three components (figure 1(c)). Firstly, a fraction \( \phi_Y \) of the area sits at the tips of the material between the pores and is always in contact with the liquid. The remaining area, the fraction \((1 - \phi_Y)\), is divided between a proportion \( \phi_P \) of pores that are filled, and a proportion \((1 - \phi_P)\) of unfilled pores in which a liquid–air interface exists. The pores are assumed to have a roughness \( r \) such that the total surface area of a pore is \( r \) times the projected area of one pore opening. This roughness plays a role only when a pore is filled with liquid. The general Cassie–Baxter model thus reduces to:

\[ \cos \theta^* = \phi_Y \cos \theta_0 + (1 - \phi_Y) [\phi_P \cos \theta_0 - (1 - \phi_P)]. \]

Equation (4) encapsulates the essential contribution of our new model: that the apparent contact angle of a liquid on a surface with a range of pore geometries can be written as a function of the fraction of pores, \( \phi_P \), filled by a given liquid. That fraction is, in turn, a function of the distribution of pore angles, as expressed in equation (1), and the contact angle \( \theta_0 \) of the liquid on a flat surface (equation (2)).

When a pore is not filled with liquid, the local effective contact angle is taken to be 180°, since in the complete absence of any solid surface, a volume of liquid naturally forms a sphere. Thus, the second term inside the square brackets of equation (4), corresponding to unfilled pores, is \( -(1 - \phi_P) \), since the cosine of the contact angle for the liquid–air interface is \( -1 \). In cases where equation (4) would imply that \( \cos \theta^* > 1 \), \( \cos \theta^* \) is set to equal 1 and complete wetting is assumed to take place. By measuring the apparent contact angles, \( \theta^* \), of four or more liquids with a range of surface tensions, we can fit estimates for \( \Psi_0 \), \( \sigma_0 \), \( \phi_Y \) and \( r \), and thereby characterize a surface.

The assumption of a Gaussian distribution of the angle \( \Psi \) is approximate because in engineered surfaces \( \Psi \) is usually constrained to lie between \(-90^\circ \) and \( 180^\circ \); \( \Psi \) cannot, in reality, extend towards \( \pm \infty \) as a Gaussian distribution would imply. Nevertheless, for the values of \( \Psi_0 \) and \( \sigma_0 \) that we fit for actual surfaces, a negligible portion of the tails of a Gaussian distribution lie outside a realistic range of \( \Psi \).

**Methods**

To demonstrate the model’s ability to describe the behavior of real surfaces, we have fabricated and characterized three porous, liquid-repellent coatings. All materials for fabricating and characterizing these coatings were obtained from Sigma-Aldrich.

The first test coating was a nanoporous zinc oxide that was grown hydrothermally on to aluminum. The porous zinc oxide surfaces were synthesized by immersing an aluminum scanning electron microscopy (SEM) sample-mounting peg into a 25 mM equimolar aqueous solution of zinc nitrate and hexamine at 70 °C for 90 min. The ZnO-coated substrates were then oxygen plasma-treated (60 W, 200 mTorr, 2 min) to clean them and impart hydroxyl surface termination.

The zinc oxide surfaces were modified in one of three ways to impart hydrophobic surface chemistry. In the first modification, the substrate was placed, immediately after plasma treatment, in a vacuum desiccator with 100 µl of 1H, 1H, 2H, 2H-Perfluorooctyltrichlorosilane (PFOS). The desiccator was then pumped for 20 min to vaporize the silane followed by letting it settle for 40 min before venting the chamber. Upon removal, the substrates were rinsed with deionized water to remove unreacted perfluorosilane, dried, and finally annealed at 120 °C for 1 h on a hot plate. In the second modification, the substrate was dipped in a 3% perfluorooctyl polyhedral oligomeric silsesquioxane (PFO-POSS) solution in Vertrel™ XF (Dupont) for 3 min and then used as coated. In the third modification, the substrate was dipped in 1% (3-aminopropyl)triethoxysilane (APTES) solution in deionized water for 10 min, and then rinsed with deionized water and heated to 120 °C on a hot plate for 1 h.

As a second test-case for the model, a film composed of approximately 600 nm diameter polyvinylidene fluoride (PVDF) spheres was synthesized by spinodal decomposition from a ternary solvent system. A 20 wt% polymer solution was created by dissolving PVDF pellets into dimethylformamide in an 80 °C water bath for 3 h.

1.7 wt% deionized water was then loaded into the solution. The solution was vortex-mixed for 5 min; then 50.0 µl of the solution was drop-cast on to an aluminum SEM peg, and spun at 1200 rpm for 30 s. Directly after spinning, the sample was submerged in deionized water at 90 °C for 30 min. After removal, the sample was air-dried then pumped under dynamic vacuum at room temperature overnight to ensure that the pores were free of solvent and water.
As a third test-case, a porous polytetrafluoroethylene (PTFE) film was created via templating with sacrificial poly(styrene-co-divinylbenzene) (PS) microspheres, according to the procedure of van der Wal [42]. Briefly, PTFE and PS colloid suspensions in water were mixed to give a solid volume fraction of 60%, with 67% of the solid volume being the sacrificial PS. 100 μl of this suspension was blade-cast on to an aluminum SEM peg and placed on a hot plate at 120 °C for 1 h to evaporate the water. A further hot-plate treatment at 290 °C for 30 min was used to sint the PTFE particles together and partially decompose the sacrificial PS. The PS was then removed by washing in acetone.

We have used six probe liquids to characterize these surfaces; these are, in descending order of surface tension, water (73 mN m⁻¹ [43]), glycerol (64 mN m⁻¹ [43]), thioglycol glycol diacrylate with a molar mass of 700 g (~43 mN m⁻¹ [44]), dipropylene glycol (34 mN m⁻¹ [43]), and methanol (23 mN m⁻¹ [43]). Sessile contact angle measurements were made by pipetting 5 μl of the liquid on to the surface, and then, after waiting for approximately 15 s, imaging the droplet from the side using a 4X objective (Olympus Plan Achromat), a 30 mm focal-length achromatic doublet (Thorlabs, Newton, NJ), and a 1280 × 1024-pixel CMOS image sensor (Thorlabs, model DCC1645C). Images were analyzed to extract contact angles using ImageJ [45] with the low-bond asymmetric drop shape analysis algorithm described by Stalder [46, 47]. Each material/probe liquid combination was measured with five separate droplets.

Values of θl for each liquid–solid pairing were determined by directly measuring the liquid’s contact angle on a flat silicon wafer treated with the same surface termination as the textured surface. Then, contact angles between the probe liquids and the textured surfaces were measured. Each liquid–solid pairing was characterized with five separate contact angle measurements made using different droplets.

For each model calibration, an upper bound for φT is estimated by considering θ* for the highest-surface-tension liquid tested—in this case water—and assuming that no pores are filled in that case. For the ZnO surfaces, an upper bound on φT was computed using the results for the PFO-POSS termination, the least-wetting treatment. Equation (4) then reduces and rearranges to:

\[ \phi_T \approx \frac{\cos \theta^* + 1}{\cos \theta_0 + 1}. \]  

(5)

This value of φT is an upper bound because it is possible that a few pores are in fact filled with water during the observation of θ*, so that not all the liquid–solid contact area is located within the tips of the structure. By using our physical understanding of the liquid–solid interaction to establish an upper bound on φT prior to the fitting procedure, we eliminate the possibility of extracting an unrealistically high value for φT during fitting. This approach helps to increase confidence in the values that are extracted for all four model parameters. With θ0 and the upper bound on φT established, a least-squares error minimization routine (Microsoft Excel’s Solver plug-in with the generalized reduced gradient algorithm) is then used to fit Ψ0, σ0, φT and r.

Results

Table 1 records the means and standard errors of the contact angles measured across all five surfaces and six probe liquids. As shown in figure 2(a), the zinc oxide pore morphology is highly irregular, and pores are surrounded by thin walls of material that are aligned approximately perpendicularly to the surface of the sample but with considerable angular variation. Figure 2(b) shows that a single set of geometrical parameters fits the data closely for all six probe liquids and all three surface terminations. Our model finds that pores are, on average, moderately re-entrant, with Ψ0 = 51°, and with a broad spread of pore geometry: σ0 = 25° (table 2). The gradual increase of cos θ* with cos θ0 indicates that a progressively larger proportion of the pores becomes filled as the contact angle of the probe liquid on a smooth surface reduces (i.e. as cos θ0 increases).

In figure 3, we re-plot the contact angle data from the ZnO with PFOSS surface termination, together with results for the PVDF and PTFE surfaces. These three surfaces show quite different relationships between cos θ* and cos θ0. The PVDF surface exhibits a much sharper non-wetting-to-wetting transition than we observe with the zinc oxide. From our contact angle measurements, we extract for the PVDF an effective average sidewall angle of Ψ0 = 57° and a low pore sidewall angle variability of σ0 = 1° (table 2). This apparent tight distribution could be explained by the reasonably uniform distribution of PVDF sphere diameter (figure 4(a)) that results from the precipitation kinetics. The curvatures locally encountered by a liquid droplet at its interface with the solid are likely to be spatially uniform, even though the pores within the film are irregular in shape. On the other hand, a high within-pore roughness, r, of 11.2 is fit, and we attribute this result both to the micro-scale roughness created by the irregular clustering of microspheres and to the nano-scale surface texture of individual microspheres that is seen in a higher-magnification electron micrograph of the material (figure 4(b)).

The PTFE structure, meanwhile, contains a broad distribution of pore sizes (figure 4(c)) by virtue of the wide distribution of sacrificial PS sphere diameters used: we estimate that these diameters range from 1 to 10 μm. We
Figure 2. Calibration of the surface wettability model for zinc oxide nanoporous surfaces terminated with PFOS, PFO-POSS and APTES. The pores in the hydrothermally grown ZnO have highly heterogeneous sizes and sidewall angles, as shown in a scanning electron micrograph (a). Inset: 5 μl droplet (diameter ≈ 2.1 mm) on the surface. A single set of fit parameters accurately describes the dependence of \( q^* \cos \theta \) on \( q \cos \theta_0 \) for this surface geometry with all three surface terminations (b). Error bars correspond to ± one standard error of the mean, based on five measurements. In many cases, error bars are smaller than symbols. Probe liquids are detailed in the results section. On a given surface, the higher the liquid’s surface tension, the lower the value of \( q \cos \theta_0 \). The model fit is shown as a dashed line.

Table 1. Measured apparent contact angles (in degrees) for the five surfaces and six probe liquids tested. The mean of five separate droplet measurements and the standard error of the mean are shown. Where the contact angle is recorded as ‘<10’, the surface was observed to wet completely, and contact angles were thus not measured.

<table>
<thead>
<tr>
<th>Probe liquid</th>
<th>Surface material</th>
<th>PFOS–ZnO</th>
<th>Mean</th>
<th>Std. err.</th>
<th>Mean</th>
<th>Std. err.</th>
<th>Mean</th>
<th>Std. err.</th>
<th>Mean</th>
<th>Std. err.</th>
<th>Mean</th>
<th>Std. err.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Water</td>
<td></td>
<td>173.8</td>
<td>0.8</td>
<td>175.8</td>
<td>0.6</td>
<td>61.6</td>
<td>1.5</td>
<td>163.3</td>
<td>0.6</td>
<td>169.2</td>
<td>0.9</td>
<td></td>
</tr>
<tr>
<td>Glycerol</td>
<td></td>
<td>169.6</td>
<td>0.5</td>
<td>170.8</td>
<td>0.4</td>
<td>34.8</td>
<td>1.9</td>
<td>158.1</td>
<td>0.7</td>
<td>169.0</td>
<td>0.9</td>
<td></td>
</tr>
<tr>
<td>Thiodiglycol</td>
<td></td>
<td>162.7</td>
<td>0.6</td>
<td>163.1</td>
<td>0.8</td>
<td>&lt;10</td>
<td>—</td>
<td>152.8</td>
<td>0.4</td>
<td>&lt;10</td>
<td>—</td>
<td></td>
</tr>
<tr>
<td>Polyethylene glycol diacrylate</td>
<td></td>
<td>151.5</td>
<td>1.5</td>
<td>158.6</td>
<td>0.3</td>
<td>&lt;10</td>
<td>—</td>
<td>147.3</td>
<td>1.0</td>
<td>&lt;10</td>
<td>—</td>
<td></td>
</tr>
<tr>
<td>Dipropylene glycol</td>
<td></td>
<td>123.7</td>
<td>0.8</td>
<td>145.2</td>
<td>0.5</td>
<td>&lt;10</td>
<td>—</td>
<td>137.2</td>
<td>1.6</td>
<td>&lt;10</td>
<td>—</td>
<td></td>
</tr>
<tr>
<td>Methanol</td>
<td></td>
<td>62.3</td>
<td>1.7</td>
<td>107.4</td>
<td>0.9</td>
<td>&lt;10</td>
<td>—</td>
<td>108.5</td>
<td>1.4</td>
<td>&lt;10</td>
<td>—</td>
<td></td>
</tr>
</tbody>
</table>

Contact angles on smooth surfaces (\( \theta_0 \))

<table>
<thead>
<tr>
<th>Surface material</th>
<th>Probe liquid</th>
<th>PFOS–ZnO</th>
<th>Mean</th>
<th>Std. err.</th>
<th>Mean</th>
<th>Std. err.</th>
<th>Mean</th>
<th>Std. err.</th>
<th>Mean</th>
<th>Std. err.</th>
<th>Mean</th>
<th>Std. err.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Water</td>
<td></td>
<td>105.3</td>
<td>0.2</td>
<td>118.2</td>
<td>0.3</td>
<td>53.7</td>
<td>0.7</td>
<td>122.6</td>
<td>1.3</td>
<td>79.0</td>
<td>2.0</td>
<td></td>
</tr>
<tr>
<td>Glycerol</td>
<td></td>
<td>102.9</td>
<td>1.6</td>
<td>112.7</td>
<td>0.7</td>
<td>47.7</td>
<td>0.5</td>
<td>114.6</td>
<td>1.0</td>
<td>65.9</td>
<td>0.6</td>
<td></td>
</tr>
<tr>
<td>Thiodiglycol</td>
<td></td>
<td>96.6</td>
<td>0.5</td>
<td>104.3</td>
<td>0.4</td>
<td>36.2</td>
<td>1.1</td>
<td>107.2</td>
<td>0.4</td>
<td>57.8</td>
<td>1.0</td>
<td></td>
</tr>
<tr>
<td>Polyethylene glycol diacrylate</td>
<td></td>
<td>83.5</td>
<td>0.7</td>
<td>92.3</td>
<td>0.5</td>
<td>27.1</td>
<td>1.4</td>
<td>96.5</td>
<td>0.3</td>
<td>35.0</td>
<td>1.1</td>
<td></td>
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<tr>
<td>Dipropylene glycol</td>
<td></td>
<td>70.2</td>
<td>1.0</td>
<td>83.5</td>
<td>0.7</td>
<td>16.0</td>
<td>2.5</td>
<td>90.8</td>
<td>0.6</td>
<td>18.7</td>
<td>0.5</td>
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<tr>
<td>Methanol</td>
<td></td>
<td>53.2</td>
<td>0.8</td>
<td>65.2</td>
<td>1.8</td>
<td>&lt;10</td>
<td>—</td>
<td>65.4</td>
<td>1.5</td>
<td>20.2</td>
<td>1.1</td>
<td></td>
</tr>
</tbody>
</table>

Table 2. Summary of the model parameters fit for three heterogeneous, nanoporous surfaces and for one regular, lithographically-produced surface. The fitting parameters are: mean reentrant angle (\( \Psi \)), standard deviation of the reentrant angle (\( \sigma_\Psi \)), within-pore surface roughness factor (\( r \)), and top-of-surface solid fraction (\( \phi_T \)).

<table>
<thead>
<tr>
<th>Surface</th>
<th>( \Psi )</th>
<th>( \sigma_\Psi )</th>
<th>( r )</th>
<th>( \phi_T )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Porous ZnO</td>
<td>51°</td>
<td>25°</td>
<td>3.5</td>
<td>0.0051</td>
</tr>
<tr>
<td>PVDF microspheres</td>
<td>57°</td>
<td>1°</td>
<td>11.2</td>
<td>0.014</td>
</tr>
<tr>
<td>PTFE templated by sacrificial PS microspheres (using the method of van der Waal [42])</td>
<td>54°</td>
<td>38°</td>
<td>1.4</td>
<td>0.092</td>
</tr>
<tr>
<td>Regular, reentrant disk array [3]</td>
<td>27°</td>
<td>0°</td>
<td>1.2</td>
<td>0.051</td>
</tr>
</tbody>
</table>
extract surface characteristics of ψ₀ = 54° and φ₀ = 38°, making this appear to be the most geometrically heterogeneous of the surfaces we have examined.

Discussion

We have used our statistical model to extract three very different wettability characteristics from the three porous surfaces. None of these trends could have been captured with a simple model of completely regular, straight-sidewalled features; such a model can comprehend only a single, critical value of \( q \) below which the surface is assumed to be in Cassie–Baxter mode, and above which pore wetting is assumed to occur spontaneously and completely.

It is conceivable that the observed shapes of the \( q^* \) versus \( q \) plots could be reproduced without considering pore-to-pore variability by developing instead a tailored geometrical model for sidewall profile whose angle varied with depth in such a way that every single pore would progressively fill at the same rate as \( q \) increased. However, a very particular profile would need to be proposed for each and every material characterized, in order to capture fully the various transitions from completely non-wetting to completely wetting behavior that are shown in figures 2 and 3. Our model of pore angle variability, with only four fitting parameters, is a much more realistic and versatile way to explain how irregular surfaces wet.

To contrast our randomly distributed surface geometries with a more regular geometry, we have used the new model to interpret previously published data from a highly regular array of singly reentrant surface structures that was produced lithographically by Liu and Kim [5]. These structures consist of a square array of silicon oxide disks of diameter \( \sim 20 \) μm that are supported by narrower silicon pillars and coated with \( \sim 150 \) nm of C₄F₈ hydrophobic polymer. As expected, this surface exhibits a far sharper transition from non-wetting to wetting as \( q \) increases (figure 3) than do our three more variable porous surfaces. Our least-squares fitting procedure extracts an average re-entrant angle of \( \psi_0 = 27° \)—which is plausible if the etching of the silicon oxide disks involves some sidewall tapering—together with zero angle variability \( \sigma_0 \), which is consistent with highly repeatable lithographic production. For this particular surface, therefore, it is reasonable to suppose that the inter-disk regions are either completely empty or completely filled according to the liquid used. Consequently, while our model-fitting procedure extracts a roughness value between the structures of \( \tau = 1.2 \), this value has no practical relevance because as soon as any inter-structure void fills, wetting is complete and \( q^* \) becomes 1.

We have thus shown that our model can be used to characterize the relationship between a liquid’s contact angle on a flat surface and its apparent contact angle on a range of porous or otherwise randomly distributed material geometries. The model predicts a one-to-one relationship between the contact angle on a flat surface, \( \theta_0 \), and the apparent contact angle. It does not capture the possible effects of ‘breakthrough’, where a load—such as a droplet’s Laplace pressure or a randomly occurring inertial force—causes the liquid surface to deform into the pore, touch its base, and wet the geometry. Rather, our model assumes that as \( \cos \theta_0 \) increases, the sidewalls
of the pores progressively guide liquid into an increasing fraction of the pores, so that the increase in \( \cos \theta^* \) with \( \cos \theta_0 \) is gradual and predictable, and relies only on the action of the liquid’s surface tension within the pores.

For the ZnO, PVDF and PTFE surfaces that we have demonstrated here, our model is accurate because the pores are mostly sub-micrometer in size and thus highly robust against breakthrough. According to the model of, for example, Chhatre et al [3], we estimate our surfaces’ robustness—essentially, the ratio of required breakthrough pressure to a typical droplet’s Laplace pressure—to be around 1000. In other words, it is very unlikely that a spontaneous Cassie–Baxter-to-Wenzel transition occurs in these structures. For materials with

Figure 4. Scanning electron micrographs of the PVDF foam shows microspheres with uniformly distributed diameters (a) and nanoscale surface roughness (b). The PTFE film exhibits large variation of pore size (c). Insets in (a) and (c) show 5 μl droplets (diameter ≈ 2.1 mm) on the respective surfaces.
considerably larger pores, it may become necessary to incorporate robustness as an additional factor. Additionally, our model does not consider contact angle hysteresis or the relationship between pore geometry and pinning. Both of these considerations may be relevant in some applications, particularly those involving condensation of the liquid that is to be repelled.

Concluding remarks

We have demonstrated a new approach to modeling the dependence of a liquid’s apparent contact angle on the geometrical characteristics of a porous surface on which it sits. Our model is simple to calibrate, requiring only contact-angle measurements to be made, and provides a framework for experimentally quantifying the variability of pore geometries on surfaces. We have demonstrated the calibration procedure for three geometrically heterogeneous porous surfaces that we have produced, as well as for one previously reported, spatially regular structure. These four materials show considerable differences between wetting characteristics that can be explained by considering pore geometry variation. All three of our coating methods involve inexpensive reagents and simple apparatus, and thus promise to be more affordable than precisely engineered, geometrically regular, patterned surfaces. The coatings can be applied to non-flat surfaces without special handling.

In addition to providing new insight into the importance and role of pore shape variability on droplet contact angles, our modeling technique offers a rapid way to screen candidate surface deposition and synthesis processes. One of the method’s key strengths is that once a particular surface geometry (e.g. that of porous zinc oxide) has been characterized, the \( \cos \theta_0 \) versus \( \cos \theta^* \) relationship can be used to predict the effect of modifying the surface chemistry (e.g. PFOS or APTES) applied to that geometry, provided that the surface modification does not alter the geometry. Thus, our approach could readily be applied to the screening of multiple candidate liquid-repellent coating processes to quantify their performance.

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