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## **OPEN** Wetting theory for small droplets on textured solid surfaces

Donggyu Kim<sup>1</sup>, Nicola M. Pugno<sup>2,3,4</sup> & Seunghwa Ryu<sup>1</sup>

Conventional wetting theories on rough surfaces with Wenzel, Cassie-Baxter, and Penetrate modes suggest the possibility of tuning the contact angle by adjusting the surface texture. Despite decades of intensive study, there are still many experimental results that are not well understood because conventional wetting theory, which assumes an infinite droplet size, has been used to explain measurements of finite-sized droplets. Here, we suggest a wetting theory applicable to a wide range of droplet size for the three wetting modes by analyzing the free energy landscape with many local minima originated from the finite size. We find that the conventional theory predicts the contact angle at the global minimum if the droplet size is about 40 times or larger than the characteristic scale of the surface roughness, regardless of wetting modes. Furthermore, we obtain the energy barrier of pinning which can induce the contact angle hysteresis as a function of geometric factors. We validate our theory against experimental results on an anisotropic rough surface. In addition, we discuss the wetting on non-uniformly rough surfaces. Our findings clarify the extent to which the conventional wetting theory is valid and expand the physical understanding of wetting phenomena of small liquid drops on rough surfaces.

The contact angle is a material property determined by the surface tensions between substrate, liquid, and vapor<sup>1</sup>. Because the materials with extremely small or large contact angles, i.e., with super hydrophilicity or super hydrophobicity, are applicable in many ways, there have been a myriad of studies that have investigated tuning the contact angle via the surface roughness of the substrate based on the conventional wetting theory of rough surface with Wenzel (W), Cassie-Baxter (CB), and Penetrate (P, which is also referred to as hemi-wicking<sup>2</sup>) modes<sup>3-9</sup>.

However, the conventional wetting theory<sup>1,10-12</sup> assumes that the liquid droplet is much larger than the characteristic scale of the surface roughness, which frequently is not justifiable in many experiments. Therefore, the contact angle predictions using the conventional theory differ from experimental results when the droplet size is small<sup>13–19</sup>. The conventional theory considers a straight boundary between the liquid and the vapor regardless of the three-phase contact line location or the droplet size<sup>1,10-12,20</sup>. However, because the realistic contour of a liquid droplet forms part of a sphere<sup>21</sup>, the assumption does not hold when the droplet size to surface texture scale ratio is small. There have been several studies to overcome the limitations of the conventional wetting theories for the small droplet, based on wetting free energy calculations. As a pioneer of the subject, Marmur et al.<sup>12,13</sup> modelled the 2D circular droplet on the heterogeneous in CB mode or the saw-tooth-like rough surface in W mode, and clarified that the wetting on the rough surface can involve multiple local free energy minima with different contact angles and that the contact angle of the global free energy minimum approaches to that of the conventional theory only when the scale of the liquid droplet is much bigger than the scale of the periodicity of the rough surface. Shahraz et al.<sup>22</sup> computed the wetting free energies of every possible pinning points on the surface with periodic rectangular protrusions in W or CB mode, to predict the contact angles of the liquid droplet with finite volume at multiple pinning points. However, since the free energy landscape between the pinning points were not investigated, it cannot be guaranteed that all the pinning points investigated in the study are at local free energy minima.

In this work, we present a more generalized wetting theory by considering the entire free energy landscape including the transition configuration between the pinning points in three wetting modes, P, W, and CB, on a surface with rectangular protrusions as well as non-uniformly rough surfaces. First, we confirm that the pinning phenomena can be understood by the existence of multiple local minima of the free energy landscape separated

<sup>1</sup>Department of Mechanical Engineering, Korea Advanced Institute of Science and Technology (KAIST), 291 Daehak-ro, Yuseong-gu, Daejeon 34141, Republic of Korea. <sup>2</sup>Laboratory of Bio-Inspired and Graphene Nanomechanics, Department of Civil, Environmental, and Mechanical Engineering, University of Trento, Trento, Italy. <sup>3</sup>Center for Materials and Microsystems, Fondazione Bruno Kessler, Trento, Italy. <sup>4</sup>School of Engineering and Materials Science, Queen Mary University of London, Mile End Road, London, United Kingdom. Correspondence and requests for materials should be addressed to S.R. (email: ryush@kaist.ac.kr)



**Figure 1.** Assumptions and methods to achieve  $\mathbf{r} - \theta$  relation. (a) Schematic of liquid droplet and the 2D textured solid substrate. To simplify the problem, we assume that the liquid droplet has a vertical surface when the non-pinned tip is on the groove (b) Relation between n and  $\overline{L}$  when  $\overline{G} = 2$ , and  $\overline{W} = 1$  for two locally pinned states. (**c**-**e**) Relation between  $\theta$  and  $\overline{E}$ . Global minimum of free energy can be found at various roughness factors r. (f) Relation between r and  $\theta$ . The contour is constructed by contact angles of the global free energy minima at each roughness factor r.

by energy barriers. For all wetting modes (P, W, and CB), we show that the contact angle at the global minimum recovers the prediction of conventional wetting theory within 2° when the droplet size (diameter of the initial droplet) becomes at least 40 times larger than the characteristic scale of the surface roughness (periodicity of the texture). Second, we compute the free energy barriers between available pinning points in all wetting modes. For P and CB modes, the energy barrier between local minima tends to decrease as the contact angle becomes further apart from the contact angle at the global minimum energy and eventually becomes zero, which enables us to predict the ultimate bound of advancing and receding angles. Our theory is then applied to explain the measured contact angle on the surface with anisotropic roughness<sup>19</sup>. Finally, we calculate the free energy of the wetting on the non-uniformly rough surface and reaffirm in terms of free energy that the contact angle is determined by the roughness of the substrate near the three-phase contact line<sup>20,23</sup>, not by the overall roughness within the droplet-substrate contact area. When the effect of gravity is ignored<sup>22</sup>, the proposed theory is universally valid for any scale of droplets unless the droplet diameter is a few nanometers or smaller because we assume that the interfacial energy of rough surface can be written as the flat surface interfacial energy multiplied the ratio of true area to the apparent area. The length scale of the molecular interaction is known to be about one to two nanometers<sup>24</sup>. Our theory reproduces the conventional wetting theory in the limit of an infinite ratio between the droplet size and characteristic scale of surface roughness.

For the mathematical simplicity, we consider a two-dimensional (2D) finite-sized liquid droplet. As illustrated in Fig. 1a, we model the rough surface as periodic rectangular textures and model the boundary of the liquid droplet as an arc of a circle while ignoring the effect of gravity. In other words, our 2D droplet is a simplification of an infinitely long cylindrical droplet. Even though our theory cannot provide the exact quantitative prediction

|    | Curvature Radius R   | Free Energy E  |
|----|--|--|
| W  | $\overline{R}_W = \left( \tfrac{\pi - n \overline{G} \overline{H}}{\theta - \sin \theta \cos \theta} \right)^{\frac{1}{2}}.$ | $2\overline{R}_W(\theta - \cos \theta_e \sin \theta) - 2n\overline{H} \cos \theta_e (n \in \mathbb{N})$  |
|    |  | $2\overline{R}_{W}(\theta - \cos \theta_{e} \sin \theta) - 2n_{1}\overline{H} \cos \theta_{e} - \overline{H} \cos \theta_{e} + \overline{H}(n \not\in \mathbb{N})$ |
| СВ | $\overline{R}_{C} = \left( \frac{\pi}{\theta - \sin \theta \cos \theta} \right)^{\frac{1}{2}}.$                              | $2\overline{R}_{C}(\theta - \cos \theta_{e} \sin \theta) + n\overline{G}(1 + \cos \theta_{e}).$  |
| Р  | $\overline{R}_{p} = \left(\frac{\pi}{\theta - \sin\theta\cos\theta}\right)^{\frac{1}{2}}.$                                   | $2\overline{R}_p(\theta - \cos \theta_e \sin \theta) + n\overline{G}(-1 + \cos \theta_e).$   |

 Table 1. Expressions of curvature radius and free energies of three wetting modes.
 Overbar indicates

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of contact angle of realistic spherical-cap-like 3D droplets, it can still offer qualitative physical understanding about the wetting of small droplet on rough surface, such as the origin of pinning, the wetting on non-uniformly rough surface, and contact angle change upon droplet volume change. The transition between the pinning points in CB and P modes is modelled by assuming that the three-phase contact line horizontally slides between the top of grooves. In W mode, we assume that the liquid tip forms a vertical straight line when inside a groove. While such an assumption may cause a small numerical difference, the resulting free energy landscape would correctly capture the multiple local minima and the energy barriers between them qualitatively well (See Supplementary Information).

We find that the free energy has local minima when one end of the droplet is fixed at either the left or right corner of the step. Two cases are illustrated in the inset of Fig. 1b. Because a hydrophobic surface prefers to reduce the contact area between the droplet and the substrate at a given contact angle, Case I has the lower free energy for the hydrophobic surfaces ( $\theta_e > 90^\circ$ ), and Case II has the lower free energy for the hydrophilic surfaces ( $\theta_e < 90^\circ$ ) (see Supplementary Information for more details).  $\theta_e$  refers to the equilibrium contact angle on a flat surface determined by Young's equation<sup>1</sup>,  $\sigma_{LV} \cos \theta_e = \sigma_{SV} - \sigma_{SL}$ , where  $\sigma_{LV}, \sigma_{SV}$  and  $\sigma_{SL}$  denote the liquid-vapor, solid-vapor and solid-liquid interfacial energies, respectively. For the two cases, the number of grooves below the liquid droplet n can be expressed with length of the baseline L, width of the step W, and width of the groove G, as described below.

$$n = \begin{cases} \max\left(n_{1}, n_{1} + \frac{\overline{L} - n_{1}(\overline{G} + \overline{W}) - \overline{W}}{\overline{G}}\right) \text{ for Case I} \\ \min\left(n_{1} + 1, \frac{\overline{L} - n_{1}(\overline{G} + \overline{W})}{\overline{G}}\right) \text{ for Case II} \\ n_{1} = \left\lfloor\frac{\overline{L}}{\overline{G} + \overline{W}}\right\rfloor \end{cases}$$
(1)

 $n_1$  is the number of groove-step pairs which fully covered with liquid, hence a natural number, and n is defined to account for partially filled grooves (real number). The overbar indicates the dimensionless variables. In what follows, all length and energy variables are normalized against the radius of the circular droplet  $R_0$  (i.e. the volume of droplet is  $\pi R_0^2$ ) and  $\sigma_{LV} R_0$ , respectively. The values of n are visualized in Fig. 1b. for the case when  $\overline{G} = 2$  and  $\overline{W} = 1$ . As illustrated in the figure, n continuously increases when the three-phase contact line is on the groove and is conserved when the contact line is on the step, as  $\overline{L}$  increases. Thereafter, the radius of the curvature and the free energy of each wetting modes can be derived in terms of n, geometrical factors, and  $\theta$ , which are presented in Table 1. Details on the numerical calculations are described in the Supporting Information. Because the current study considers only 2D droplet cases, the effect of the line tension<sup>25</sup> can be ignored. However, one needs to incorporate the line tension effect when dealing with 3D droplets on textured surfaces<sup>26,27</sup>.

Based on the free energy expression as a function of  $\theta$ , we can find the allowable contact angles with free energy minima at a given roughness factor<sup>28</sup>  $r = \frac{\overline{G} + \overline{W} + 2\overline{H}}{\overline{G} + \overline{W}}$ , i.e., the ratio of the true surface area to the projected area. For example, Fig. 1c–e show the relationship between the free energy and the contact angle for the W mode when r = 1.2, 1.5 and 1.8, respectively when  $\overline{G} = \overline{W} = 0.05$  (i.e., a droplet size of  $2R_0$  is 20 times that of the characteristic scale of the surface roughness, G + W). The vertical dotted line is the contact angle determined by the conventional wetting theory on the Wenzel mode,  $\cos \theta = r \cos \theta_e$ . The contact angles at the global free energy minima under different surface roughnesses, which are highlighted by the blue circle, green triangle, and red square, do not match perfectly with the prediction from the conventional wetting theory. Thereafter, one can predict the contact angle as a function of the surface roughness factor r by connecting the contact angles at the global free energy minima, as illustrated in Fig. 1f. The conventional wetting theory prediction (red dotted curve),  $\cos \theta = r \cos \theta_e$ , is also presented in comparison. The contact angles for the CB and P modes can be obtained in a similar way as functions of  $f = \frac{\overline{W}}{\overline{G + W}}$ , which is the fraction of the step area from the projected area.

We then consider how the contact angle changes with the droplet size by varying the dimensionless variable  $\overline{G}$ , which is the ratio of the groove width to the initial droplet radius. For a given  $\overline{G}$ , the roughness factor<sup>28</sup> r and the step fraction f can be tuned by changing the height or width of the steps,  $\overline{H}$  or  $\overline{W}$ . The predicted contact angles for W, CB, and P modes are presented in Fig. 2a–d with varying values of r and f. When  $\overline{G} = 0.5$ , i.e., the droplet size is 2 times the characteristic scale of the surface roughness, our theory and the conventional theory<sup>10–12</sup> predict different contact angles because the wetting free energy landscape of a small droplet is extremely different from that of a large droplet. Hence, the conventional theory should not be used. However, in case of  $\overline{G} = 0.05$ , the contact



Figure 2. Results of contact angle and wetting mode prediction from the proposed theory. (a–d) Convergence curve of each wetting modes ((a) Wenzel (hydrophilic), (b) Wenzel (hydrophobic) (c) Cassie-Baxter, (d) Penetrate). Conventional wetting theory should be used when  $\overline{G} < 0.025$  (e) Wetting mode selection with respect to  $\theta_e$  when  $\overline{G} = 0.1$ , r = 1.5 and f = 0.5.

angle from our theory converges to that from the conventional theory within the range of  $\pm 10^{\circ}$ . When  $\overline{G}$  reaches 0.005, the contact angle from our theory recovers the prediction of conventional theory<sup>10-12</sup> almost perfectly, which is expected. Considering the typical resolution of the contact angle measurements  $(1\sim 2^{\circ})^{15}$ , we suggest the conventional theory<sup>10-12</sup> should be applied in the case when  $\overline{G} \leq 0.025$  (see Supplementary Information for details), i.e., when the droplet size is at least about 40 times bigger than the characteristic scale of the surface roughness.

We can predict the most stable wetting mode for the substrates with different Young's angle  $\theta_e$  by comparing the free energies of the three modes<sup>29</sup>. As depicted in Fig. 3d, we find that in accordance with the conventional theory, on a hydrophilic surface ( $\theta_e < 90^\circ$ ), the contact angle is given by the higher contact angle between the predictions based on the P mode and W mode and on the hydrophobic surface ( $\theta_e < 90^\circ$ ) by the lower contact angle between the predictions based on the CB mode and W mode. The choice between the W mode and the CB mode is made by comparing the free energy values in Table 1. To select between the W mode and the P mode, we use the critical contact angle theory<sup>2</sup> instead of the direct free energy comparison because the initial free energy of the P mode differs from the other modes. The critical contact angle,  $\theta_C$  is determined when the free energy variation to fill an additional groove is 0. If the contact angle of the flat surface  $\theta_e$  is smaller than  $\theta_C$  the free energy variation to fill another groove becomes negative, spreading occurs, and the Penetrate mode is selected. In our work, the critical contact angle can be expressed with geometric factors as  $\theta_C = \arccos\left(\frac{\overline{G}}{\overline{G}+2\overline{H}}\right)$ . As illustrated in Fig. 2d, for the case of  $\overline{G} = 0.1$ , r = 1.5, and f = 0.5, one can find that the most stable wetting mode curve follows a similar path as that of the conventional theory.

We then confirm the origin of the pinning effect is the occurrence of multiple local minima and calculate the free energy barriers in terms of geometric factors for the three wetting modes. In the W mode, a drastic free energy change occurs when the three-phase contact line is located near the corner of the step because the additional





| Wetting Mode  | $\Delta \overline{\mathrm{E}}_{1}$   | $\Delta \overline{E}_2$  |
|---------------|--|--|
| Wenzel        | $\overline{H}(1 + \cos \theta_e)$  | $\overline{H}(1 - \cos \theta_e)$  |
| Cassie-Baxter | $2\overline{R}^{A}_{C}\!\theta^{A}+\overline{W}\cos\theta_{e}-2\overline{R}^{B}_{C}\!\theta^{B}$ | $2\overline{R}^C_C\!\theta^C + \overline{G} - 2\overline{R}^B_C\!\theta^B$                       |
| Penetrate     | $2\overline{R}_{P}^{A}\!\theta^{A}+\overline{G}-2\overline{R}_{P}^{B}\!\theta^{B}$               | $2\overline{R}_{P}^{C}\!\theta^{C}-\overline{W}\cos\theta_{e}-2\overline{R}_{P}^{B}\!\theta^{B}$ |

Table 2. Dimensionless energy barriers for each wetting mode.

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boundary may form or disappear. The A' and B' points at the end of the step in Fig. 3a include an additional boundary line while A and B do not. As depicted in Fig. 3a, the free energy differences between A' and  $A(\Delta \overline{E}_1)$  or B' and  $B(\Delta \overline{E}_2)$  act as the primary free energy barrier and form a local free energy minimum. The amount of the energy barrier can be expressed with  $\overline{H}$  and  $\theta$ , as shown in Table 2. When the substrate is hydrophobic, the local minimum point is located on B, and one can notice that this corresponds to the experimentally observed three-phase contact line location<sup>19</sup> when pinning occurs. A similar discussion can be repeated for the Cassie-Baxter or the Penetrate mode. By calculating the free energy of each point of Fig. 3b and c, the energy barriers can be formulated, as summarized in Table 2. The superscripts A and B refer to the points A and B in the figures, respectively, and the subscripts C and P refer to the wetting modes. Interestingly, while the Wenzel mode always has local free energy minima because of the additional liquid-vapor boundary line, the Cassie-Baxter or the Penetrate mode do not possess local minimum points when  $\theta < \theta_e$  or  $\theta > \theta_e$ , respectively. The disappearance of the energy barrier indicates the ultimate bound of advancing and receding angles which are determined by available local free energy minima. The maximum advancing (receding) angle can be obtained from experiments will be the highest (lowest) contact angle at the pinning points with nonzero energy barrier. We speculate that the conventional Wenzel state has very large contact angle hysteresis because of the non-vanishing energy barrier between local minima<sup>30</sup>.

Now, we apply the proposed theory to understand the wetting experiments on the surface with anisotropic roughness<sup>19</sup>. An experiment performed by Chen *et al.*<sup>19</sup> used morphologically patterned surface to measure the contact angle of the surface with anisotropic roughness made of PDMS ( $\theta_e = 114^\circ$ ) along both perpendicular and parallel directions. The width of step and groove were 23 µm and 25.6 µm each. The height of the step was 30 µm. The measurement of the contact angle were conducted with water droplet of volume from 0.59 mm<sup>3</sup> to 5.679 mm<sup>3</sup>. They confirmed that droplets are in the CB mode, and measure the number of pillars filled by or the base line length of the droplet. It was reported that the contact angle at the direction perpendicular to the grooves is likely to be similar to the advancing contact angle<sup>31,32</sup>, and that the advancing contact angle is chosen as the maximum contact angle among the local free energy minima<sup>33</sup>. However, since there exists an external energy perturbation such as ambient vibration and the inertia associated with the droplet spreading, we expect that the measured advancing contact angle must be smaller than the maximum contact angle among local minima. We compute the energy barrier at each local minima for different droplet volumes, and compared the energy barrier between B







**Figure 5.** Relation between  $\theta$  and  $\overline{E}$  of the Wenzel mode when the surface has non-uniform roughness. The contour contains slices of free energy contours of fixed n (red dotted). The global free energy point is highlighted with the green square symbol (a) The global free energy minimum point might be located on the  $n = n_{MAX}$  curve or (b) the global free energy point of the surface with uniform roughness.

state and C state in Fig. 3b plays a main role in determining the advancing contact angle in CB mode. The energy barrier can be calculated by the formula in Table 2. Since the energy barrier formula is non-dimensional, we multiply  $R_0\sigma_{LV}$  and the baseline length parallel to the groove direction to dimensionalize the formula to compare the energy barriers for different droplet volume. The energy barrier between B state and C state, at each local minimum is plotted in Fig. 4a. Each dot in the curves refers to the energy barrier at each pinning point and all open symbols refer to the experimental result for different droplet volume. In the figure, one can notice that the energy barriers for different droplet volumes are similar. The average energy barrier is represented with the black dotted horizontal line which is located close to all open symbols. One may consider the black dotted line as the typical external energy perturbation in the series of experiments reported in the study. The contact angle for specific liquid volume can be predicted by capturing the pinning point which has the smallest distance from the average energy barrier (black dotted in Fig. 4b). The predicted contact angle agrees well with experimental results and show the same trend with experiments. One may perform a similar analysis to predict the receding contact angle or the contact angle hysteresis of other experiments.

We then apply our theory to analyse a surface with non-uniform roughness that has a rough center and flat periphery. In other words, we compute the free energy landscape when there exists an upper bound  $n_{MAX}$  in the number of filled grooves, n. For example, we compute the free energies of the wetting states when  $\overline{G} = \overline{W} = 0.05$ and  $\theta_e = 120^\circ$  as functions of contact angle  $\theta$  with  $n_{MAX} = 14$  and 19, as depicted in Fig. 5a and b, respectively. Figure 5a illustrates the situation where the area of the rough central region is small enough that the three-phase contact line is located on the flat region. We find that the global free energy minimum is located at the Young's angle  $\theta_e$ . It is the case for any  $n_{MAX} \le 14$  because the free energy curve with a fixed n has a minimum at  $\theta_e$ , as depicted by the red dotted curves in Fig. 5a. On the contrary, Fig. 5b shows a case where the area of the rough region is large enough and the contact line is located within the rough center. In this case, the local minimum associated with  $n = n_{MAX} = 19$  has a higher free energy than the global minimum. Hence, the contact angle prediction becomes identical to the surface with uniform roughness. Our results agree with the previously proposed wetting theory for a non-uniform rough surface<sup>20,23</sup>, which proposed that the contact angle is determined by the roughness condition near the three-phase contact line. Our work offers an extended theory that enables us to predict whether the contact line will be located on the rough center or on the flat region. We note that our theory can be generalized to study the wetting on surfaces with a more complex non-uniform roughness or be used to build a surface to fix the droplet to the specific location<sup>34</sup>, complementing the pioneering works by Johnson and Dettre<sup>35,36</sup>.

In conclusion, we have developed a wetting theory that can predict the wetting angle and wetting mode when the liquid droplet is not much larger than the surface texture scale. Because the conventional theory assumes a much larger size of the droplet compared to the texture scale, there have been limitations in how to analyse experiments that investigate the wetting of small liquid drops. Our theory suggests that conventional theory should be used when the droplet size is about 40 times larger than the characteristic scale of the surface roughness and provides a deeper physical understanding regarding the wetting of smaller liquid droplets on non-uniform rough surfaces.

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#### **Author Contributions**

D.K. and S.R. designed the research, developed the theoretical models, and analyzed the results. D.K. carried out numerical calculations. N.M.P. contributed to the theoretical analysis. D.K. and S.R. wrote the manuscript. All authors discussed the results and commented on the manuscript.

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### **Supplementary Information**

### Wetting theory for small droplets on textured surfaces

Donggyu Kim<sup>1</sup>, Nicola M. Pugno<sup>2</sup>, and Seunghwa Ryu<sup>\*,1</sup>

#### Affiliations

<sup>1</sup> Department of Mechanical Engineering, Korea Advanced Institute of Science and

Technology (KAIST), 291 Daehak-ro, Yuseong-gu, Daejeon 305-701, Republic of Korea

<sup>2</sup> Department of Civil, Environmental, and Mechanical Engineering, University of Trento, Trento, Italy

\* Corresponding author email : <u>ryush@kaist.ac.kr</u>

Supplementary Figure 1: Free energy comparison between circular boundary and vertical boundary



**Supplementary Figure S1.** Free energy comparison between circular boundary and vertical boundary. (a) Schematic of circular (A) and vertical (B) boundary. (b) Relative error between free energy of two boundary assumptions. The surface was set by  $\bar{G} = \bar{W} = 0.5$ , r=1.5. (c) Relative error between free energy of two boundary assumptions. The surface was set by  $\bar{G} = \bar{W} = 0.005$ , r=1.5. Rather than the droplet volume, the Young's angle mainly affects the error arising from the vertical boundary assumption.



#### Supplementary Figure 2: Free energy curve of a droplet in Wenzel mode

**Supplementary Figure S2**. Convex curvature of Case II is always on the right side of Case I. Because  $\overline{\mathbf{E}}$  decreases for (a) a hydrophilic substrate and increases for (b) a hydrophobic surfaces as  $\boldsymbol{\theta}$  increases, the former has a minimum at Case II and the latter has a minimum at Case I.

## Supplementary Figure 3: Convergence to conventional theory with respect to $\overline{G}$ for each wetting mode



Supplementary Figure S3. Convergence to conventional theory due to  $\overline{G}$  for each wetting mode.  $\overline{W}$  and  $\overline{H}$  are set to satisfy the roughness factors (r, f) in the legend. The contact angle difference between the proposed theory and the conventional theory ( $\Delta \theta$ ) converges to  $1 \sim 2^{\circ}$  when  $\overline{G} \leq 0.025$ 

## Supplementary Note 1: Free energy comparison between circular boundary and vertical boundary

It was reported that the liquid droplet forms the part of the circle when the effect of the gravity is neglected<sup>1</sup>. However, it is mathematically very hard to model the liquid droplet on periodic rectangular protrusion with circular boundary with the constant droplet volume constraint. Therefore, we used vertical liquid boundary (path B in **Fig. S1a**) to model the droplet in Wenzel mode (CB mode and P mode do not need the assumption) rather than the circular liquid boundary (path A in **Fig. S1a**). To check the validity of our assumption, we compare the free energy of two boundary for a simple case where the three-phase contact line touches the bottom of the groove.

The free energy of each liquid boundary can be formulated as below.

$$\bar{E}_{C} = \bar{R}_{W} \alpha$$
$$\bar{E}_{V} = \bar{H} + (\bar{R}_{W} \sin(\theta + \alpha) - \bar{R}_{W} \sin(\theta)) \cos\theta_{e}$$

Here,  $\bar{E}_C$  and  $\bar{E}_V$  refer to the free energy of the circular and the vertical boundary. The free energy can be formulated by  $E = \sigma_{LV}A_{LV} + (\sigma_{SL} - \sigma_{SV})A_{SL}$ .  $\alpha$  refers to the angle between circular boundary within the groove as depicted in **Fig. S1a**. The relative error between two energy barriers can be defined by  $\frac{\bar{E}_C - \bar{E}_V}{\bar{E}_V}$ . The relative error between two boundaries are calculated for the surface with r=1.5, when  $\bar{G} = \bar{W} = 0.5$  (small droplet) and  $\bar{G} = \bar{W} =$ 0.005 (large droplet). The region with high contact angle ( $\theta > 140^\circ$ ) when the droplet volume is small is not investigated because  $\alpha$  cannot be decided in the case because the three-phase contact line cannot reach the floor of the groove. As shown in **Fig. S1b and Fig. S1c**, the relative error is smaller than 15% unless the surface is highly hydrophilic ( $\theta_e < 60^\circ$ ) or highly hydrophobic ( $\theta_e > 120^\circ$ ). Because the droplet on the highly hydrophilic (hydrophobic) surface prefers P (CB) mode, we can conclude that our assumption is reasonable within 15% of errors.

#### Supplementary Note 2: The choice between Case I and Case II along wetting modes

1-1. Cassie-Baxter & Penetrate mode

From Young's relation<sup>2</sup>, one can say the free energy from same area of liquid - solid boundary is always larger than that from the liquid – vapor boundary for any substrate when  $\theta > 0^\circ$ . Because Case II always contains larger n than in Case I, regardless of L (**Fig. 2b**), Case I contains less area of the liquid-vapor boundary. Consequently, Case I become more stable if the same  $\theta$  is assumed. In the case of Penetrate mode, on the contrary, Case II become more stable because the liquid-vapor boundary is substituted by a liquid-liquid boundary, which contains 0 surface energy.

#### 1-2. Wenzel mode

If the same amount of n is assumed, Case I always has a larger  $\overline{L}$  or smaller  $\theta$  than Case II. As we noted in the text, the  $\overline{E} - \theta$  graph contains convex contours of conserved n, where the minimum is at  $\theta_e$  (red dotted in **Figs. S1a, S1b**). Because Case I has a larger  $\theta$  for conserved n, one can notice the convex contour of Case II is always on the right side of Case I for conserved n. Recalling that the Wenzel mode has a larger equilibrium contact angle than  $\theta_e$ , when the substrate is hydrophilic, the free energy of the liquid decreases as  $\theta$  increases. Therefore, the local free energy minimum near the equilibrium contact angle is on the right side of the contour, Case II. If a hydrophobic substrate is assumed, by the opposite logic, the local free energy minimum is on Case I.

## Supplementary Note 3: Convergence to conventional theory with respect to $\overline{G}$ for each wetting modes

As depicted in **Figs. 2a-c**, the predicted contact angle from the proposed theory ( $\theta_P$ ) converges to the contact angle from the conventional theory ( $\theta_T$ ) when  $\bar{G} \leq 0.025$ . We are not able to find a closed form expression for the difference between the two contact angles ( $\Delta \theta = \theta_P - \theta_T$ ), but find that the envelope of the oscillating  $\Delta \theta$  curve decreases with  $\bar{G}$  for all wetting modes (**Fig. S2**). The contact angle difference converges with vibration and becomes  $1 \sim 2^\circ$  range when  $\bar{G} \leq 0.025$ .

#### Supplementary Note 4: Free energy and curvature radius calculation about $\theta$

The curvature radius of the liquid droplet of each wetting mode (W,CB,P) can be formulated with a circular trace of the boundary of the liquid and the constant volume constraint as Shahraz et al. <sup>1</sup> reported. In Wenzel mode, the sum of the area of the circular part and of the groove should be constrained. Therefore, the curvature radius of the Wenzel mode  $\overline{R}_W$  can be formulated as follows.

$$\overline{R}_{W} = \left(\frac{\pi - n\overline{G}\overline{H}}{\theta - \sin\theta\cos\theta}\right)^{\frac{1}{2}}$$

After the curvature radius is formulated, the free energy of the droplet can be calculated by summing up the free energy from the liquid-vapor boundary and the liquid-solid boundary as follows.

$$\mathbf{E} = \sigma_{\rm LV} \mathbf{A}_{\rm LV} + (\sigma_{\rm SL} - \sigma_{\rm SV}) \mathbf{A}_{\rm SL}$$

 $A_{LV}$  or  $A_{SL}$  refer to the area of the boundary between the liquid and the vapor or solid and the liquid. Employing the curvature radius formula, the free energy of the droplet in Wenzel mode can be shown.

$$\overline{E}_{W} = 2\overline{R}_{W}(\theta - \cos\theta_{e}\sin\theta) - 2n\overline{H}\cos\theta_{e} \ (n \in \mathbb{N})$$
$$\overline{E}_{W} = 2\overline{R}_{W}(\theta - \cos\theta_{e}\sin\theta) - 2n_{1}\overline{H}\cos\theta_{e} - \overline{H}\cos\theta_{e} + \overline{H} \ (n \notin \mathbb{N})$$

Because an additional boundary of the liquid-vapor boundary exists when  $n \notin \mathbb{N}$  in Wenzel mode, the free energy expression either differs when  $n \in \mathbb{N}$  or not. Similar analysis can be repeated for the Cassie-Baxter mode and Penetrate mode to include the curvature radius or free energy of the liquid in CB mode or P mode. In particular, in the case of P mode, we only model the liquid forming the droplet to construct  $R_0$  and set the initial state of the free energy with a rough surface in which grooves are filled with liquid for convenience. Then the free energy and the curvature radius of the penetrate mode can be formulated similar with CB mode as follows.

$$\overline{R}_{C,P} = \left(\frac{\pi}{\theta - \sin\theta\cos\theta}\right)^{\frac{1}{2}}$$

$$\overline{E}_{C,P} = \overline{R}_{C,P}(\theta - \cos\theta_e \sin\theta) + n\overline{G}(1 + \cos\theta_e)$$

Because the  $n - \theta$  relation is known if the liquid tip condition (Case I or Case II) is suggested, a set of ( $\overline{R}$ , n,  $\overline{E}$ ) can be shown with respect to a specific  $\theta$ . In the case of CB mode and P mode, because  $\overline{R}_{C,P}$ ,  $\overline{E}_{C,P}$ , and n are explicitly expressed by  $\theta$ , a pair of ( $\overline{R}_{C,P}$ , n,  $\overline{E}_{C,P}$ ) can be easily obtained for a specific  $\theta$ . In W mode,  $\overline{R}_W$ ,  $\overline{E}_W$ , and n are in an implicit relation; therefore we adopted the bisection method to numerically calculate the set of ( $\overline{R}_W$ , n,  $\overline{E}_W$ ). With this process, the free energy of the droplet can be numerically obtained from a specific  $\theta$  and can be used to find the contact angle of the minimum free energy.

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