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Stood-up drop to determine receding contact angles

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The wetting behavior of drops on natural and industrial surfaces is determined by the advancing and receding contact angles. They are commonly measured by the sessile drop technique, also called goniometry, which doses liquid through a solid needle. Consequently, this method requires substantial drop volumes, long contact times, tends to be user-dependent, and is difficult to automate. Here, we propose the stood-up drop (SUD) technique as an alternative to measure receding contact angles. The method consists of depositing a liquid drop on a surface by a short liquid jet, at which it spreads radially forming a pancake-shaped film. Then the liquid retracts, forming a spherical cap drop shape (stood-up drop). At this quasi-equilibrium state, the contact angle (θ_{SUD}) closely resembles the receding contact angle measured by goniometry. Our method is suitable for a wide variety of surfaces from hydrophilic to hydrophobic, overcoming typical complications of goniometry such as needle-induced distortion of the drop shape, and it reduces user dependence. We delineate when the receding contact angle can be obtained by the stood-up method using volume-of-fluid (VoF) simulations that systematically vary viscosity, contact angle, and deposited drop volume. Finally, we provide simple scaling criteria to predict when the stood-up drop technique works.

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1 Introduction

Wetting¹ is ubiquitous in nature and technology, shaping phenomena such as drops rolling off plant leaves, animal skin, and insect wings, as well as applications including inkjet printing,² coatings,³ forensics,⁴ disease detection,⁵ and DNA analysis.⁶ Surface wettability is characterized by measuring contact angles. Primarily, three types of contact angles exist: Young–Dupré's (equilibrium) contact angle, θ_Y , advancing contact angle, θ_a , and receding contact angle, θ_r . Despite their relevance, measuring the different contact angles still remains

challenging, in particular measuring θ_r due to distortions of the drop shape by the needle. Therefore, a needle-free method to measure the receding contact angle would be desired.

At thermodynamic equilibrium, a stationary drop on a solid adopts θ_Y owing to the interfacial tension balance at the three phase contact line and follows Young–Dupré's equation:⁷

$$\cos \theta_Y = \frac{\gamma_{SG} - \gamma_{SL}}{\gamma_{LG}} \quad (1)$$

where γ_{SG} , γ_{SL} and γ_{LG} are the solid–gas, solid–liquid and liquid–gas interfacial tensions, respectively. It is important to point out that the Young–Dupré's contact angle is a theoretical concept, often attributed to the static contact angle of sessile drops.^{8–12} Although the static contact angle can classify surfaces as hydrophilic, hydrophobic and superhydrophobic, it provides little information on how drops adhere to, or roll or slide over a surface.

When the three-phase contact line moves, advancing of the liquid front yields θ_a , whereas retraction over an already wetted region yields θ_r . Both angles are affected by surface properties such as roughness and chemical heterogeneity,¹³ surface adaptation,¹⁴ and slide electrification.¹⁵ In practice, contact angles of resting drops lie within $[\theta_r, \theta_a]$; these bounds characterize wetting (θ_a) and de-wetting (θ_r) behavior on a given surface. The receding angle is indispensable for modeling drop

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removal. For example, the pull-off force per unit length is $\gamma_{LG}(1 + \cos \theta_r)$.^{16,17} Lower θ_r promotes residual drops during withdrawal ($\theta_r < 90^\circ$), and θ_r correlates with practical adhesion in anti-icing^{18,19} and anti-biofouling^{20,21} applications. Moreover, θ_r predicts instabilities of receding lines, the occurrence and duration of rebound,²² and drop-surface friction,²³ making it central to drop dynamics on solids.^{23–26}

Standard measurements of θ_a and θ_r use sessile-drop goniometry: a needle injects liquid into a sessile drop to increase volume (advancing), then withdraws to decrease volume (receding). Despite algorithmic advances,^{27–32} this protocol has drawbacks. The needle perturbs the free surface (capillary rise) and especially biases θ_r at small volumes; needle position can alter the contact-line speed. Typical devices require samples of a few cm^2 and volumes of $\sim 30\text{--}50 \mu\text{L}$ to reach θ_r , limiting heterogeneous or small specimens. The inflow/outflow cycle and setup time render the method time-consuming and user-dependent.³³ An alternative approach has also been proposed, in which a droplet is deposited through a pre-drilled hole in the substrate.³⁴ Although this method can prevent needle induced effects, it represents an idealized configuration,³⁵ which is time consuming and can induce surface damage in fragile samples. This can alter the local topography, compromising the integrity of heterogeneous or patterned surfaces.

Here we introduce the stood-up drop (SUD) method for receding-angle metrology. A short liquid jet (jetting time of a few ms) deposits a thin lamella that spreads radially and then retracts to a spherical-cap drop. After a quasi-equilibrium is reached, the apparent contact angle θ_{SUD} closely approximates θ_r . The measurement completes in ~ 10 ms and requires no needle, mitigating deposition artefacts and facilitating automation. We validate SUD by directly comparing θ_{SUD} with θ_r across hydrophilic–hydrophobic substrates, and we delineate when θ_{SUD} recovers θ_r using direct numerical simulations based on a volume-of-fluid (VoF) technique that systematically varies viscosity, static contact angle, and deposited volume. Finally, we summarize simple scaling criteria that predict the SUD regime and practical volume limits, and we highlight conditions under which SUD can fail (*e.g.*, violent oscillations or detachment on highly hydrophobic surfaces). Together, these results position SUD as a fast, needle-free alternative for robust receding-angle characterization.

2 Materials and experimental methods

2.1 Experimental setup: stood-up drop (SUD) device

To form drops from a liquid jet, a device designed by KRÜSS company was used. The device generates a liquid jet by applying a constant pressure P_{app} to a liquid reservoir (Fig. 1a), which is connected to a valve and nozzle. The distance between sample and nozzle was kept at 4 mm. The applied pressure P_{app} was varied between 100 and 700 mbar. The time at which the valve is open (jetting time τ) can be controlled by software. It should not exceed a few milliseconds to ensure the formation of a thin lamella. If not stated otherwise, $\tau = 1$ ms is used. A

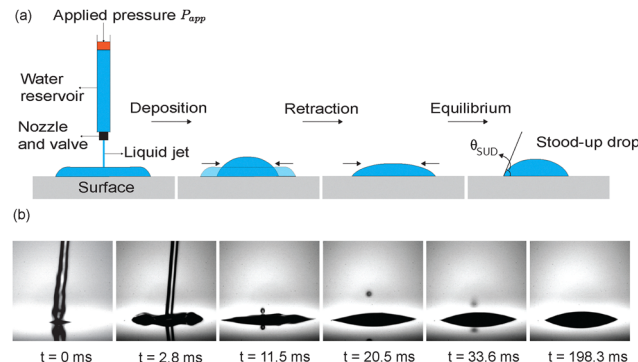


Fig. 1 (a) Experimental setup for SUD technique. A liquid jet is generated at certain applied pressure P_{app} for a very short jetting time τ . Once the liquid impacts the sample surface, it spreads radially and then retracts. During the first retraction phase the drop can vibrate, before the contact line smoothly recedes. The liquid forms a drop with a spherical-cap shape, whose contact angle is (θ_{SUD}). The drop profile in the first column represents the initial configuration of the drop for the numerical simulations. The configuration resembles a pancake shape, similar to that observed in experiments post-impact. The second to fourth columns sketch the retraction of the drop from an initial (lighter shade) to the quasi-equilibrium state (darker shade) as time progresses. (b) Experimental snapshots of a liquid jet impacting on a Si wafer surface at $P_{\text{app}} = 350$ mbar and jetting time $\tau = 5$ ms. See also Video S1.

liquid jet is generated immediately after the valve opening. After spreading and retraction of the liquid, a stationary drop is formed (Fig. 1b). Here, the contact angle at this stage (θ_{SUD}) is measured and compared with θ_r obtained by Goniometry technique. We determined θ_{SUD} values by a polynomial fitting method, which was designed by a python code (SI). This code determines the average of θ_{SUD} values from the average of contact angles at the left and right side of drops. Our results consider the θ_{SUD} average of the last 50 frames of each video. For a subset of all investigated samples no high-speed recordings were performed. In such cases, the SUD contact angle was determined using the KRÜSS ADVANCE software after drop deposition using the KRÜSS DS3251 SUD dosing unit.

The liquid impact process is recorded by a high-speed camera (Photron UX10) at up to 8000 fps. Simultaneous high speed-video experiments were performed from the top of the surface, to study the shape of the drop contact area and pinning of the three-phase contact line. Experiments were repeated at least three times for each surface (in some cases five times), with their wetting nature ranging from hydrophilic to hydrophobic. For each droplet, the contact angle was obtained by averaging the left and right contact angles, and then the mean and standard deviation were calculated across repeated experiments.

2.2 Receding contact angle measurements

Receding contact angles θ_r were measured by the sessile drop method/Goniometer (KRÜSS, DSA100). First, a $0.5 \mu\text{L}$ drop of Milli-Q water (18 megaohm) is deposited on the sample. Then, the baseline is positioned at the contact line between drop and surface, while the needle is adjusted to be in the middle of the drop. Afterwards, the volume is increased up to $50 \mu\text{L}$ at a flow



rate of $0.5 \mu\text{L s}^{-1}$. Then, the drop is deflated at the same flow rate. A tangent fit method was used to determine receding contact angles by the ADVANCE software of the device. Experiments were repeated at least three times (up to five times in some cases) on different spots of the surface. At each time step, the mean of the left and right contact angles was taken, and the average and standard deviation of these values were then determined over the plateau region of the contact angle vs. drop base diameter plot.

2.3 Image processing of contact angle measurements

For automated contact angle measurements using image processing of the high speed recordings, we adapted the open source 4S-SROF toolkit to match our requirements.³⁶ The 4S-SROF toolkit effectively utilizes the OpenCV library³⁷ to manipulate the images, such as separate the drop from its background. We chose morphological transformations for noise reduction, and they proved to be superior to the median filter, guaranteeing the accuracy of our advancing angle measurements.³⁶ We calculated the advancing and receding angle using the tangent fitting method for the final 10 pixels of the drop near the substrate. The Savitzky–Golay filter³⁸ was employed in specific cases to eliminate unwanted noise and enhance the smoothness of the final diagram, facilitating easier interpretation.

2.4 Surface preparation

Most investigated surfaces are commercially available or provided by customers from Krüss. These surfaces were cleaned by water and ethanol before usage.

2.4.1 PFOTS coatings. Si wafer, 1 mm thick SiO_2 slides ($76 \times 25 \times 5 \text{ mm}^3$, Thermo Fisher Scientific) and glass slides were coated with perfluoroctadecyltrichlorosilane (PFOTS). After O_2 plasma cleaning at 300 W for 10 min (Femto low-pressure plasma system, Diener electronic), samples were placed in a vacuum desiccator containing a vial with 0.5 mL of 1H,1H,2H,2H-perfluoro-octadecyltrichlorosilane (97%, Sigma Aldrich). The desiccator was evacuated to less than 100 mbar, closed and the reaction was allowed to proceed for 30 min.

2.4.2 PDMS brushes. Plasma-activated silicon wafers (Si-Mat, Kaufering, Germany, plasma activation for 10 min at 300 W) were placed in a desiccator together with 1,3-dichlorotetramethyl-disiloxane (96%, Alfa Aesar, 50 μL in 2.150 cm^3) and the reaction was allowed to proceed at ambient temperature (21°C) and ambient humidity (40–60%) for 3 h. Non-bonded oligomers were removed by washing the samples in toluene ($\geq 9.8\%$, Thermo Fisher Scientific), isopropanol ($\geq 99.8\%$, Thermo Fisher Scientific), and Milli-Q water for a minimum of 30 s each.

2.4.3 Teflon AF on ITO. Teflon AF1600 samples were prepared by dip-coating of indium tin oxide (ITO) coated glass slides. The slides were immersed into 1 wt% Teflon AF1600 (Sigma Aldrich) in FC-317 (Sigma Aldrich with a speed of 90 mm min^{-1}). After being immersed for 10 s, the substrates were withdrawn from the solution at a constant speed of 10 mm min^{-1} . Finally, the coated substrates were annealed at 160°C in a Heidolph vacuum oven for 24 h.

3 Simulation methodology

3.1 Governing equations

In this section, we present the governing equations that describe the retraction dynamics of a liquid drop on a solid surface. Following the one-fluid formulation, both the drop and ambient gas are treated as a single continuum, and the interfacial boundary condition is enforced *via* the localized surface tension force \mathbf{f}_γ .³⁹ Specifically, the drop's retraction arises from the conversion of interfacial free energy into kinetic energy, countered by viscous dissipation as captured by the Navier–Stokes equation,

$$\rho \left(\frac{\partial \mathbf{v}}{\partial t} + \nabla \cdot (\mathbf{v} \mathbf{v}) \right) = -\nabla p + \nabla \cdot (2\eta \mathcal{D}) + \mathbf{f}_\gamma + \rho \mathbf{g}, \quad (2)$$

with mass conservation that enforces a divergence-free velocity field $\nabla \cdot \mathbf{v} = 0$. Here, \mathbf{v} and p are the velocity and pressure fields respectively, and t represents time. The variables η and ρ are the fluid viscosity and density, respectively. \mathcal{D} is the symmetric part of the velocity gradient tensor ($\mathcal{D} = (\nabla \mathbf{v} + (\nabla \mathbf{v})^T)/2$).⁴⁰ Lastly, \mathbf{g} is the acceleration due to gravity.

3.2 Non-dimensionalization of the governing equations

To non-dimensionalize the governing equations, we use the characteristic scales of the system: the liquid density ρ_L , the surface tension at the liquid–gas interface γ_{LG} , and the maximum post-impact spreading thickness of the drop H . These parameters define the characteristic inertiocapillary timescale τ_γ and the characteristic inertiocapillary velocity v_γ ,⁴¹

$$\tau_\gamma = \frac{H}{v_\gamma} = \sqrt{\frac{\rho_L H^3}{\gamma_{\text{LG}}}}, \quad (3)$$

and

$$v_\gamma = \sqrt{\frac{\gamma_{\text{LG}}}{\rho_L H}}, \quad (4)$$

respectively. Using these characteristic scales, and using \tilde{x} to represent the dimensionless form of the respective variable x , we rewrite the Navier–Stokes equation eqn (2) in its dimensionless form

$$\left(\frac{\partial \tilde{\mathbf{v}}}{\partial \tilde{t}} + \tilde{\nabla} \cdot (\tilde{\mathbf{v}} \tilde{\mathbf{v}}) \right) = -\tilde{\nabla} \tilde{p} + \tilde{\nabla} \cdot (2\text{Oh} \tilde{\mathcal{D}}) + \tilde{\mathbf{f}}_\gamma + \text{Bo} \hat{\mathbf{g}} \quad (5)$$

where $\tilde{p} = pH/\gamma_{\text{LG}}$ is the dimensionless pressure, $\tilde{\mathbf{f}}_\gamma \approx \tilde{\kappa} \tilde{\nabla} \psi$ is the dimensionless surface tension force, where κ is the interfacial curvature $\kappa = \nabla \cdot \hat{\mathbf{n}}$ with $\hat{\mathbf{n}}$ as the normal to the interface marked between the two fluids: liquid with $\psi = 1$ and gas with $\psi = 0$. Lastly, the Ohnesorge number and Bond number

$$\text{Oh} = \frac{\eta}{\sqrt{\rho_L \gamma_{\text{LG}} H}}, \quad (6)$$

$$\text{Bo} = \frac{\rho_L g H^2}{\gamma_{\text{LG}}} \quad (7)$$

characterize the dimensionless viscosity of the retracting drop and the dimensionless gravitational acceleration, respectively. The non-

dimensionalization also gives two additional control parameters, the density ratio $\rho_L/\rho_G = 1000$ and the Ohnesorge number ratio $Oh_L/Oh_G = 50$ between liquid and gas phases. These ratios remain constant throughout the paper. Additionally, the Bond number based on the formulation in eqn (7) is $O(10^{-3})$. At Bond numbers of this magnitude, it is expected that gravity influences the macroscale dynamics very weakly, and even more so, it is dominated by the capillary and viscous stresses near the contact line. Therefore, throughout the paper we set $Bo = 0$, with the exception of Fig. 3, where the Bond number is set to be $Bo = 5.7 \times 10^{-3}$.

3.3 Numerical setup & domain description

We perform direct numerical simulations using the volume-of-fluid (VoF) method, implemented using the open-source language Basilisk C.⁴² The simulations solve the governing equations presented in Section 3.1, under the assumption of axial symmetry (see Fig. 2). The bottom boundary of the domain represents the substrate and enforces a no-penetration condition for the normal velocity component. The tangential component is influenced by an implicit slip model, which emerges due to the interpolation of fluid properties in the VoF scheme, despite the no-slip condition being imposed.⁴³ The implicit slip is further constrained to satisfy the prescribed contact angle θ at the first grid cell. The pressure boundary condition at this interface follows a zero-gradient condition (see Fig. 2).

At the top and right domain boundaries, free-slip conditions are applied to the velocity field, while a Dirichlet zero condition is enforced for pressure. The domain size, denoted as L_D , is chosen such that these boundaries remain sufficiently distant to avoid influencing the contact line dynamics. Heuristically L_D is chosen to be $D_{\max}/(2H) + 10$, where D_{\max} is the maximum spreading diameter following jet impingement. To enhance computational efficiency and accuracy, adaptive mesh refinement (AMR) is applied in regions exhibiting large velocity gradients. The refinement depth is selected to ensure a minimum of 40 grid cells across the drop thickness, *i.e.*, $H > 40\Delta$, where Δ denotes the smallest grid cell size.

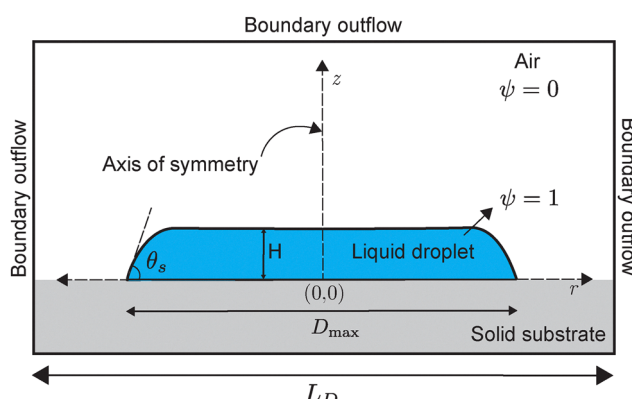


Fig. 2 The initial configuration of the drop for the numerical simulations. The configuration resembles a pancake shape, similar to that observed in experiments post-impact.

3.4 Initial conditions

The numerical simulations commence with the drop at rest, immediately after reaching the “pancake” configuration following impact (Fig. 2). At this stage, the drop assumes a pancake-like shape, characterized by a thin, radially spread film with an aspect ratio $\Gamma = D_{\max}/H$. Since the drop spreading is directly controlled by the kinetic energy provided by the jet,⁴⁴ we also assume that there is sufficient kinetic energy to overcome pinning and reach the pancake configuration. This assumption is justified by the drop deposition protocol in experiments where the liquid jet is turned off once a drop is created at the surface. Furthermore, the flow timescales involved in the impact and spreading process are distinct. The transition from an impacting jet to a flattened drop occurs over the kinematic timescale D_0/V_0 where the subsequent retractions occurs on the inertio-capillary timescale, $\tau_\gamma = \sqrt{\rho_L H^3/\gamma_{LG}}$. Here, we focus on the retraction phase only where the influence of left-over kinetic energy from impact is assumed minimal.⁴⁵

The dimensions of the spread drop are determined by the impact conditions, with the base diameter D_{\max} and thickness H depending on the volume of the deposited drop which depends on the nozzle flow rate q and jetting time τ , and the balance of inertial and capillary forces during the spreading phase. The volume of the drop Ω is related to the base diameter and thickness, up to the leading order, as

$$\Omega \sim q\tau \sim D_{\max}^2 H \quad (8)$$

The jetting time is kept constant at $\tau = 1$ ms during the experiments, resulting in drops with a volume of 0.5 μL . However, we do vary the volume of the drop in our simulations, ranging from 0.25–3 μL . The system is modeled under axial symmetry in a three-dimensional framework, ensuring that the retraction dynamics are captured without assuming any two-dimensional simplifications. We note that in the experiments, the drop retraction is not always axisymmetric. The initial state provides a well-defined starting point for investigating the subsequent retraction process, which is governed by surface tension and viscous effects.

4 Results and discussion

4.1 SUD mechanism

Initially, a liquid jet, ejected from a nozzle at a controlled pressure (see Section 4.2.1), impacts the surface with a speed of approximately 3 m s^{-1} , as measured from the last few frames of the high-speed videos (Fig. 1b and 3a, $t = 0^+$, ms). Subsequently, the liquid spreads radially, converting kinetic energy into surface energy and viscous dissipation, forming a pancake-shaped liquid film until reaching a maximum diameter D_{\max} . Analogous to impacting drops,⁴⁶ the impact and spreading duration scales with the inertio-capillary timescale. Consequently, we turn off the jet after roughly 1 ms, (Video S1).

The pancake spreading state serves as the initial condition for our simulations, assuming that the internal flow at this moment doesn't influence subsequent retraction dynamics



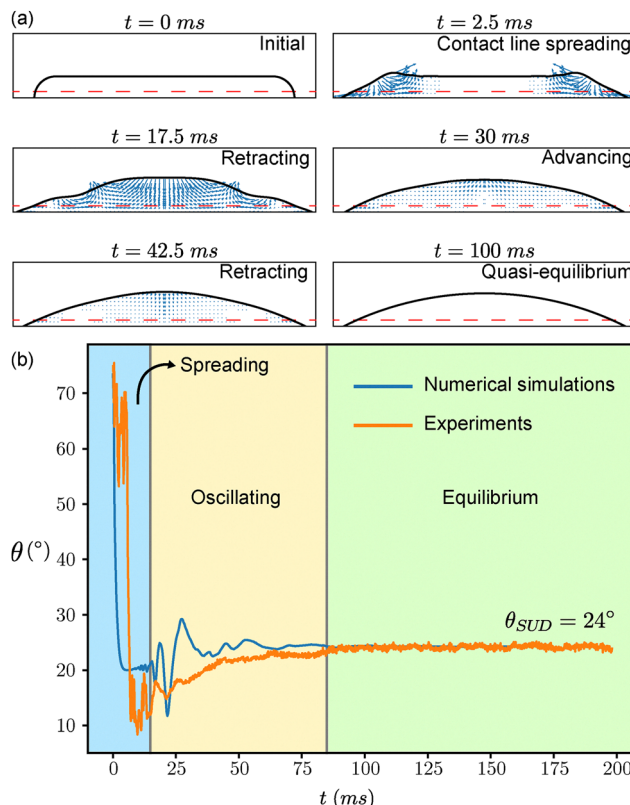


Fig. 3 (a) Snapshots of a simulated drop with static contact angle $\theta_s = 27^\circ$ and volume of $0.5 \mu\text{L}$. The arrows represent the velocity profile inside the drop to visualize whether the drop is in the retracting or advancing phase. The dashed red line represents the height at which the contact angles are measured, in line with experimental conditions. (b) Comparison of measured contact angle from numerical simulations and experiments, as a function of time for the simulation shown in (a). $\tau = 1 \text{ ms}$.

(Fig. 3a) in our simulations. For hydrophilic substrates, the contact line initially spreads to achieve its maximum spreading state. Afterwards, the pancake-shaped drop recoils, converting surface energy back into kinetic energy and viscous dissipation (Fig. 3a), finally forming a static cap-spherical drop characterized by the stood-up contact angle θ_{SUD} (Fig. 3, $t \approx 100 \text{ ms}$, Video S1).

In experiments, during spreading, the contact angle significantly decreases until reaching a minimum (Fig. 3b, blue region for both experiments and simulations). The minimum is caused by the large impact pressure, resulting in the formation of a very thin lamellae. As the contact line recedes, first the contact angle oscillates (yellow region). The release of the stored surface energy can cause the oscillations. Oscillations of the contact angle level off after approximately 80 ms at a larger angle.

The temporal development of the contact angle is well reproduced in numerical simulations (Fig. 3b). Oscillations are more pronounced on hydrophobic surfaces (Fig. S1) but they ultimately cease due to viscous dissipation, resulting in a plateau at θ_{SUD} (Fig. 3b, blue line). Since no solid needle is required, effects on the drop shape due to the deposition protocol can be ruled out. This facilitates the fitting procedure to calculate contact angles.

To investigate possible pinning during retraction, we recorded the impacting jet from the top of the substrate using high-speed imaging. No pinning points or residual drops were observed (Fig. 4). The final SUD drop area was clearly circular, Video S2.

4.2 Further experimental results

4.2.1 Influence of pressure on the observed SUD contact angle. To analyze the influence of pressure on θ_{SUD} , we generate SUD drops at three different applied pressures P_{app} (100, 350, and 700 mbar) on PMMA. Our experiments show that θ_{SUD} remains largely unchanged across this pressure range (Fig. S2). As P_{app} increases, the liquid impacts the surface with higher speed and kinetic energy, causing the drop to spread further and reach larger maximum diameters D_{max} , thus increasing the aspect ratio Γ – an effect analogous to increasing the Weber number We for impacting drops.⁴⁶ Nonetheless, as the drop retracts, memory effects can be excluded. The impacting jet is always turned off before the retraction phase starts. As soon as the drop comes to rest the contact angle converges to a constant value of θ_{SUD} and is independent of P_{app} .⁴⁷

We stress that increasing P_{app} increases both the flow rate (volume of liquid ejected per unit time during valve opening) and the maximum attainable SUD drop volume. This volume depends on P_{app} , the jetting time τ , and the nozzle diameter. At the end of the retraction phase, a SUD behaves like a spherical cap with volume (Fig. 4):

$$\Omega = \frac{\pi}{6} H_c (3r^2 + H_c^2), \quad (9)$$

where H_c and r are the height and footprint radius of the cap. However, the SUD volume variation is insignificant in our experiments due to the short τ used. For instance, on a PMMA surface, the SUD volume increases by $\sim 0.5 \mu\text{L}$ when P_{app} changes from 350 to 700 mbar. An ejected volume that is too large can prevent the SUD state (see Section 4.3.1).

In previous work,⁴⁷ we showed that the flow rate—defined by the combination of P_{app} and the jet diameter—is the key

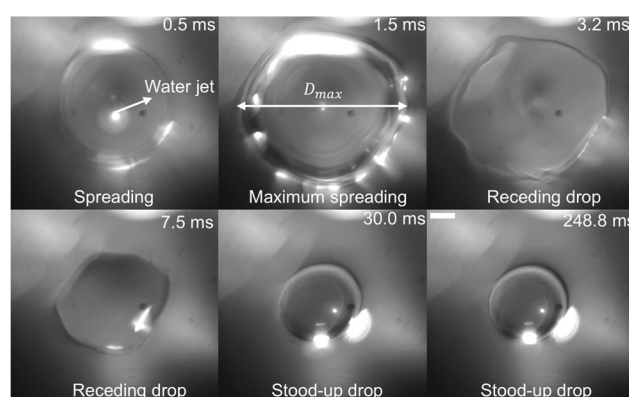


Fig. 4 Top view snapshots of the impacting liquid jet on a PMMA surface. Frame rate: 4000 fps. No pinning points are observed. Pressure: 350 mbar. Jetting time: $\tau = 1 \text{ ms}$. Scale bar represents 0.5 mm. White dots correspond to specular reflections.

factor determining the largest drop volume that still achieves the SUD state. If, for a given flow rate, the final drop volume is below a critical threshold, the contact line recedes after dosing, and the drop's rest angle is the receding contact angle θ_r . From experiments on six substrates,⁴⁷ we identified a phenomenological correlation between the flow rate and the maximum drop volume ensuring SUD dosing.

4.2.2 Comparison between θ_{SUD} and θ_r . We measured the corresponding θ_{SUD} for different surfaces, from hydrophilic to hydrophobic and compared the values with θ_r obtained by the goniometer technique. Our results reveal a good agreement between θ_r and θ_{SUD} (Fig. 5). Agreement is slightly worse for a few samples showing particularly low contact angle hysteresis, such as polycarbonate.

Remarkably, SUD technique is suitable for a hydrophilic surface (bare glass slide), overcoming the difficulties of measuring contact angles for these surfaces by the sessile-drop method, caused by evaporation and drop pinning. As the SUD drop forms in a few milliseconds, evaporation does not affect θ_{SUD} values. Moreover, the high-speed of the liquid jet for both spreading and the first period of the retraction phase, avoid efficiently the influence of pinning points on the drop dynamics. These pinning points can prevent the contact line motion during inflation/deflation of drops using goniometry. Determining θ_r for a very hydrophilic glass slide using goniometry has proven challenging. However, a contact angle of $\theta_r = 4.5^\circ$ has been determined previously by capillary bridges.⁴⁸ This is in good agreement with our θ_{SUD} values of $\sim 6^\circ$ (Fig. S3).

Therefore, SUD method arises as a reliable alternative to determine θ_r for surfaces with low wettability.

For the case of hydrophobic surfaces like PFOTS and PDMS surfaces, the equilibrium state at which the SUD is formed takes longer time. This occurs due to the capillary oscillations generated during the retraction phase, when the surface energy is converted back to kinetic energy and viscous dissipation.

Drop oscillations can lead to asymmetric contact line motion and the drops can even slide owing to this asymmetry or that of the surface orientation. The effect is more pronounced when the hydrophobicity increases, as shown for the case of Teflon AF on ITO (Fig. S1). The SUD method fails when: (1) insufficient viscous dissipation allows drops to oscillate violently and settle in an advancing phase or (2) rapid receding motion generates enough upward momentum for the drop to detach from the surface. Both situations are common on highly hydrophobic surfaces, where bubble entrainment during rapid retraction can further complicate the dynamics.^{49–52} Under these conditions, the SUD state is not the result of a final receding of the contact line, and thus the measured contact angles no longer correspond to θ_r . Therefore, the SUD technique requires that the final motion of the contact line is a clear, dominant receding event, which is essential for reliably resembling the receding contact angle.

A particular case arises on highly hydrophobic pillared surfaces. At sufficiently high impact pressures, the liquid jet may impale the surface, leading to a localized Wenzel state at the impact spot. If a clean receding phase is achieved at a given pressure without liquid detachment, the SUD state can still be

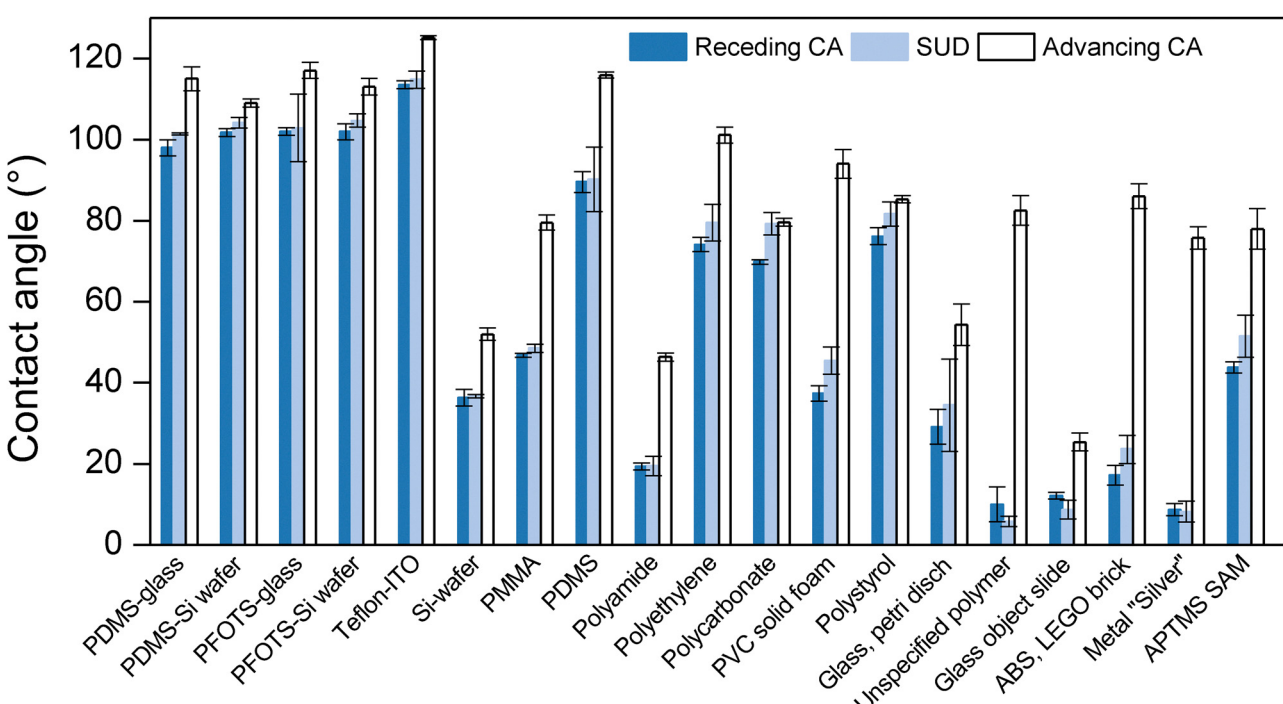


Fig. 5 Comparison between the receding (θ_r) and advancing (θ_a) contact angles measured by goniometry technique (dark blue and white rectangles) and the stood-up contact angles (light blue rectangles) (θ_{SUD}). Values of (θ_r) and (θ_{SUD}) are very close, apart from a few samples. For the following samples: PDMS/PFOTS on Si wafer and glass, Teflon on ITO, and Si wafer, θ_{SUD} was measured by the tangent fitting method described in Section 2.3, while the rest of the surfaces by the Young–Laplace fit provided by the goniometer software.



reached. However, at higher pressures the contact line dynamics may be influenced by local liquid penetration, as previously reported for droplets impacting superhydrophobic nanostructured surfaces.⁵³ This scenario would represent an additional limitation of the SUD method, with underlying physics that lies beyond the scope of the present study.

In the following section, we quantitatively analyze when the SUD method works by systematically exploring the parameter space defined by the Ohnesorge number Oh and the aspect ratio Γ at three representative static contact angles of $\theta_s = 60^\circ$ (hydrophilic), $\theta_s = 90^\circ$, and $\theta_s = 120^\circ$ (hydrophobic).

4.3 Simulation results for the full $Oh-\Gamma$ parameter space

This section delineates the regions in the $Oh-\Gamma$ parameter space, for hydrophobic ($\theta_s > 90^\circ$), $\theta_s = 90^\circ$, and hydrophilic ($\theta_s < 90^\circ$) surfaces, where SUD is a viable technique. In the SUD regime, the viscous dissipation is sufficient to ensure an over-damped retracting drop that stays on the surface. Additionally, the total volume of the drop (keeping all other material and

flow properties fixed) presents additional constraints on this technique, which is discussed at the end of this section.

4.3.1 When does the SUD technique work? For drops smaller than the gravito-capillary length ($H \ll l_c \equiv \sqrt{\gamma_{LG}/\rho_{LG}}$), the amount of viscous dissipation during the retraction phase largely dictates whether a drop remains attached to the surface after contact line recoiling (SUD regime), with or without oscillations or detaches from the surface (non-SUD regime).⁴⁵

Detached drops often fall back onto the surface. This leads to pronounced oscillations for drops already in a spherical cap shape, which opens the possibility of the final drop at rest to be the result of a wetting process and not of a de-wetting process. Consequently, the contact angle might not resemble the receding angle but rather a contact angle between advancing and receding angle. To systematically characterize the conditions classifying stable SUD formation *versus* detachment, we explored the parameter space spanned by the Ohnesorge number Oh and aspect ratio Γ across multiple static contact angles θ_s , as illustrated in Fig. 6.

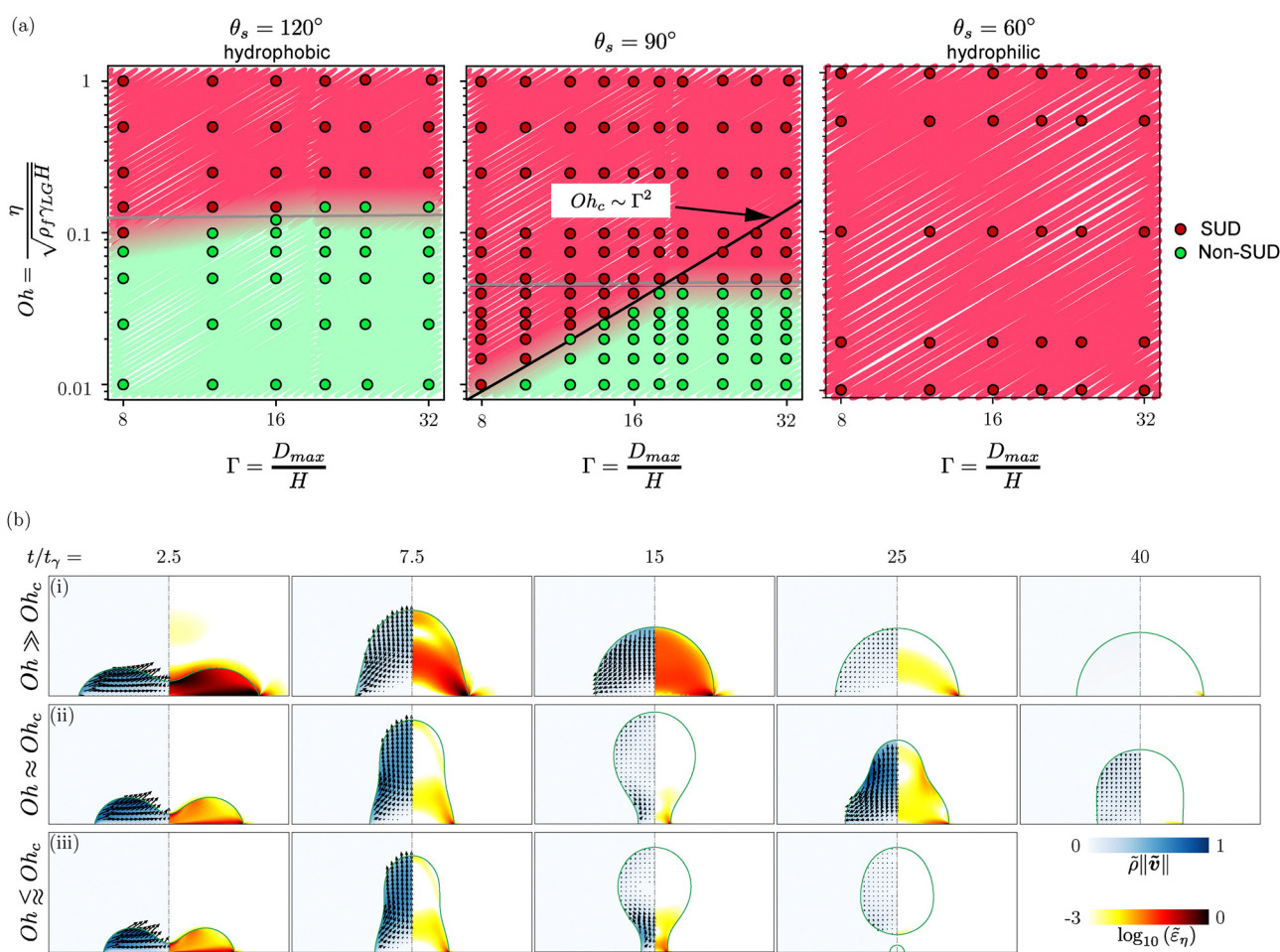


Fig. 6 (a) The $Oh-\Gamma$ parameter space illustrating regions where SUD is a viable technique for $\theta = 120^\circ$ (left, hydrophobic), $\theta = 90^\circ$ (middle), and $\theta = 60^\circ$ (right, hydrophilic). (b) Numerical simulations of drop evolution for $\theta = 90^\circ$, $\Gamma = 12$ at $Oh =$ (i) $0.75 \gg Oh_c$, (ii) $0.025 \approx Oh_c$ – a narrow neck forms at $\tilde{t} \sim t/t_\gamma = 15$, but the drop escapes pinch off and continues surface oscillations, and (iii) $0.022 \ll Oh_c$ – a narrow neck develops which eventually pinches off. Here, $Oh_c(\Gamma, \theta)$ represents the Ohnesorge number dictating the SUD to non-SUD transition. The left half of every simulation snapshots represents the dimensionless momentum and the right half shows the dimensionless rate of viscous dissipation per unit volume normalized using the inertio-capillary scales, represented on a \log_{10} scale to differentiate the regions of maximum dissipation. The black arrows depict the velocity vectors inside the drop.



We observe three distinct mechanisms governing the transition between SUD and non-SUD regimes, dependent on the static contact angle:

1. For highly hydrophobic surfaces (large θ_s), contact line dissipation is minimal, and the transition is primarily governed by bulk dissipation overcoming the excess surface energy—analogous to the bouncing-to-non-bouncing transition on superhydrophobic surfaces.⁴⁵ In this regime, the critical Ohnesorge number Oh_c remains approximately constant; drops with $Oh > Oh_c$ dissipate sufficient energy to remain in the SUD regime, while those with $Oh < Oh_c$ detach.

2. For intermediate wettability ($\theta_s \approx 90^\circ$), there is a balance between dissipation and released energy as illustrated by the numerical snapshots in Fig. 6(b). For sufficiently large Oh , dissipation in both bulk and contact line produces SUD behavior, with the drop undergoing overdamped oscillations as it dissipates energy (Fig. 6(b-i)). Near the critical threshold Oh_c , drops may undergo several oscillation cycles before eventually stabilizing (Fig. 6(b-ii)). The SUD method fails when insufficient viscous dissipation allows drops to oscillate violently and drops settle in an advancing phase. Below Oh_c , insufficient viscous dissipation leads to detachment, with the liquid neck pinching off as shown in Fig. 6(b-iii).

3. For hydrophilic surfaces (small θ_s), contact line dissipation dominates and effectively suppresses detachment across the entire parameter range investigated – encompassing the parameter space relevant for experimental applications. Consequently, as shown in Fig. 6(a) – ($\theta_s = 60^\circ$), all simulated conditions for hydrophilic surfaces result in stable SUD behavior, making the technique particularly robust for such surfaces.

We develop a theoretical model balancing surface energy and viscous dissipation to quantify these transitions. The initial surface energy of the pancake-shaped drop at $t = 0$ is $E_{\text{surf}} = \gamma_{\text{LG}} A_{\text{surf}}$. To leading order in Γ , the surface area scales as

$$E_{\text{surf}} \sim \gamma_{\text{LG}} H^2 \Gamma^2. \quad (10)$$

During retraction, viscous dissipation⁴¹

$$E_{\text{diss}} = 2 \int_0^t \int_{\Omega} \eta (\mathcal{D} : \mathcal{D}) d\Omega dt \quad (11)$$

energizes energy from the system. With velocity gradients scaling as v_{γ}/λ (where λ is the characteristic length scale) and dissipation occurring over volume Ω_{diss} , we obtain

$$E_{\text{diss}} \sim \eta \left(\frac{v_{\gamma}}{\lambda} \right)^2 \tau_{\gamma} \Omega_{\text{diss}} \sim \eta v_{\gamma} H \left(\frac{\Omega_{\text{diss}}}{\lambda^2} \right) \quad (12)$$

The dominant velocity gradient arises from shear flow with $\lambda \sim H$, giving v_{γ}/H .

The location of viscous dissipation depends critically on the Ohnesorge number. At high Oh , velocity gradients develop throughout the drop immediately-analogous to the high-viscosity limit in Taylor–Culick retraction.^{54,55} Here, dissipation occurs over the entire bulk volume:

$$\Omega_{\text{diss}} \sim H^3 \Gamma^2 \quad (13)$$

Substituting into eqn (12) yields

$$\begin{aligned} E_{\text{diss,bulk}} &\sim \eta v_{\gamma} H \left(\frac{H^3 \Gamma^2}{H^2} \right) \\ &\sim \eta V_{\gamma} H^2 \Gamma^2 \\ &\sim \gamma_{\text{LG}} H^2 \Gamma^2 Oh \end{aligned} \quad (14)$$

When bulk dissipation dominates and balances the released surface energy, we get

$$Oh_c \sim 1, \quad (15)$$

where Oh_c is the critical Ohnesorge number for SUD behavior.^{45,56}

At low Oh , viscous effects remain localized near the retracting contact line, with the central film region remaining nearly stationary initially. Dissipation concentrates in a boundary layer of volume:^{57,58}

$$\Omega_{\text{diss}} \sim \lambda^3 \quad (16)$$

The boundary layer thickness λ is bounded by the drop thickness H (Fig. 6). With $\lambda \sim H$, eqn (12) gives

$$\begin{aligned} E_{\text{diss,CL}} &\sim \eta v_{\gamma} H \left(\frac{H^3}{H^2} \right) \\ &\sim \eta v_{\gamma} H^2 \sim \gamma_{\text{LG}} H^2 Oh \end{aligned} \quad (17)$$

When contact line dissipation dominates and balances the surface energy,

$$Oh_c \sim \Gamma^2 \quad (18)$$

These criteria accurately demarcate the SUD and non-SUD regimes in Fig. 6(a) – ($\theta_s = 90^\circ$). At very large Oh values, the transition becomes controlled primarily by bulk dissipation, resulting in a constant Oh_c marked by the gray line for $\theta_s = 120^\circ$ and 90° . Lastly, below a critical static contact angle (exemplified by $\theta_s = 60^\circ$ in Fig. 6a), the system features only SUD behavior.

A drop detaching from the surface obviously poses serious challenges for measurements using the SUD technique. Such detachment can cause the drop to exit the camera frame, entrain bubbles during pinch-off and redeposition,⁵² or significantly increase the time required to reach equilibrium. Furthermore, detached drops often fall back onto the surface, resulting in uncontrolled secondary deposition unlike the controlled jetting process discussed previously.

Indeed, a key takeaway from the numerical simulations is that SUD technique remains viable for hydrophilic surfaces across the entire parameter space, even at extreme combinations of small Oh and large Γ . For hydrophobic surfaces, however, the technique only works within the restricted range of large Oh and small Γ . Lastly, we stress that apart from the non-SUD cases discussed in this section, the technique can also fail when surfaces are superhydrophobic, causing the jet to bounce instead of depositing a drop, or when drop breakup occurs (Fig. S4). Analysis of lift-off during jet impingement lies beyond our current scope.

4.3.2 Other volume limitations. In addition to directly affecting the Ohnesorge number, very small volume of the drop can cause practical issues in measuring the contact angle, such as having a limited resolution to measure the contact angle close to the surface relative to the size of the drop. On the other hand, at very large volumes, liquid jets do not reach the SUD regime as the spreading behavior is favored by the excess of volume. This prevents the onset of a receding phase.

5 Conclusions and outlook

In this work, we have introduced and validated the stood-up drop (SUD) technique as an effective method for measuring receding contact angles. Microliter-sized water drops, formed after an impacting liquid jet spreads and recoils on a surface, exhibit a contact angle (θ_{SUD}) that remarkably resembles the receding contact angle (θ_r) measured by traditional goniometry. Our numerical simulations, performed using the volume-of-fluid method, corroborate our experimental findings and provide a theoretical framework for understanding this technique and its limitations.

The SUD method offers several key advantages over conventional goniometry. First, it eliminates the need for a solid needle, thereby avoiding distortion of the drop shape during measurement. Second, it dramatically reduces measurement time from minutes to milliseconds, enhancing experimental efficiency. Third, it performs exceptionally well on hydrophilic surfaces, where traditional techniques often struggle due to evaporation and pinning issues. Fourth, it requires smaller sample sizes ($\approx 0.5 \mu\text{L}$), making it suitable for testing smaller or heterogeneous surfaces. Fifth: the contact angle after SUD dosing θ_{SUD} describes the smallest possible contact angle of a drop at rest on the investigated surface. This in combination with the already established liquid needle dosing³³ describing the largest possible contact angle of a drop on the investigated surface allow for the easiest way to determine contact angle hysteresis. By systematically exploring the parameter space defined by the Ohnesorge number (Oh) and aspect ratio (Γ), we have established clear boundaries for the applicability of this technique. For hydrophilic surfaces, the SUD method remains viable across the entire parameter space investigated. For hydrophobic surfaces, its applicability is restricted to combinations of sufficiently large Oh and small Γ , where viscous dissipation prevents detachment. This framework allows researchers to determine *a priori* whether the SUD technique will provide reliable measurements for their specific systems. While the SUD method exhibits exceptional performance across a wide range of surfaces, it has limitations for highly hydrophobic materials. In these cases, insufficient viscous dissipation can lead to violent oscillations or detachment, preventing the formation of a stable equilibrium state dominated by receding dynamics. Additionally, volume constraints must be considered, as tiny drops present resolution challenges, while large drops deviate from the spherical cap assumption due to gravitational deformation.

Looking forward, the SUD technique opens several promising avenues for research. The temporal dynamics of the surface tension-driven receding process varies significantly between substrates (e.g., 2–3 seconds on PMMA *versus* approximately 30 ms on other samples). This difference remains poorly understood but might provide valuable insights into a surface's dewetting properties beyond the steady-state contact angle measurements. We have shown that the stood-up drop technique works excellently on smooth surfaces. Future work could explore these dynamic aspects to develop a more comprehensive characterization of surface wettability, especially on more complex substrates such as sticky or textured surfaces. Moreover, the rapid and accurate determination of receding contact angles enabled by the SUD method could enhance our ability to predict contact line instabilities such as drop pinning and splitting. This has practical implications for various applications including spray coatings, self-cleaning surfaces,^{59,60} anti-icing materials,⁶¹ and biofouling-resistant coatings,⁶² where the receding contact angle offers better correlation with practical adhesion work than the advancing contact angle.^{1,63} The SUD technique could also be extended to investigate more complex fluids with non-Newtonian properties^{64,65} or to study the temperature dependence of receding contact angles¹ – both relevant for industrial applications. Additionally, combining the SUD method with simultaneous measurements of contact line dynamics might reveal further insights into the fundamental physics of wetting and dewetting processes.⁶⁶

Author contributions

D. V., T. W. and D. D. planned the experiments. D. D. and F. W. performed the experiments, using a code written by S. S. A. S. supported with the preparation of the samples. V. S., D. L. and A. B. planned the simulations. A. B. ran the simulations. D. D., F. W. and A. B. analyzed the data. D. V., V. S., T. W., D. D. and A. B. designed the structure of the manuscript. D. D., A. B., D. V., T. W., V. S., R. B., M. K., H.-J. B. and D. L. wrote the manuscript. All authors discussed the measurements and approved the manuscript.

Conflicts of interest

Thomas Willers is employed by KRÜSS GmbH.

Data availability

The codes used for the numerical simulations are available at <https://github.com/comphy-lab/Retracting-Droplet>. The experimental data is available from the corresponding authors upon reasonable request.

The data supporting this article have been included as part of the supplementary information (SI). The supplementary information contains additional images and movies captured from experiments for hydrophilic as well as hydrophobic substrates. See DOI: <https://doi.org/10.1039/d5sm00985e>.

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Notes and references

- 1 D. Bonn, J. Eggers, J. Indekeu, J. Meunier and E. Rolley, *Rev. Mod. Phys.*, 2009, **81**, 739.
- 2 D. Lohse, *Annu. Rev. Fluid Mech.*, 2022, **54**, 349–382.
- 3 H. B. Eral, D. t Mannetje and J. M. Oh, *Colloid Polym. Sci.*, 2013, **291**, 247–260.
- 4 D. Attinger, C. Moore, A. Donaldson, A. Jafari and H. A. Stone, *Forensic Sci. Int.*, 2013, **231**, 375–396.
- 5 T. A. Yakhno, O. A. Sedova, A. Sanin and A. Pelyushenko, *Tech. Phys.*, 2003, **48**, 399–403.
- 6 D. Brutin, *Droplet wetting and evaporation: from pure to complex fluids*, Academic Press, 2015.
- 7 T. Young, *Abstracts of the Papers Printed in the Philosophical Transactions of the Royal Society of London*, 1832, pp. 171–172.
- 8 J. W. Drellich, L. Boinovich, E. Chibowski, C. Della Volpe, L. Holysz, A. Marmur and S. Siboni, *Surf. Innovations*, 2019, **8**, 3–27.
- 9 A. Marmur, *Soft Matter*, 2006, **2**, 12–17.
- 10 D. Cwikel, Q. Zhao, C. Liu, X. Su and A. Marmur, *Langmuir*, 2010, **26**, 15289–15294.
- 11 A. Marmur, *Eur. Phys. J.: Spec. Top.*, 2011, **197**, 193.
- 12 A. Marmur, C. Della Volpe, S. Siboni, A. Amirfazli and J. W. Drellich, *Surface Innovations*, 2017, **5**, 3–8.
- 13 C. McPhee, J. Reed and I. Zubizarreta, *Developments in Petroleum Science*, Elsevier, 2015, vol. 64, pp. 313–345.
- 14 H.-J. Butt, R. Berger, W. Steffen, D. Vollmer and S. Weber, *Langmuir*, 2018, **34**, 11292–11304.
- 15 A. Stetten, A. Golovko, S. Weber and H.-J. Butt, *Soft Matter*, 2019, **15**, 8667–8679.
- 16 B. Samuel, H. Zhao and K.-Y. Law, *J. Phys. Chem. C*, 2011, **115**, 14852–14861.
- 17 A. Dupré and P. Dupré, *Théorie Mécanique de la Chaleur*, Gauthier-Villars, 1869.
- 18 A. J. Meuler, J. D. Smith, K. K. Varanasi, J. M. Mabry, G. H. McKinley and R. E. Cohen, *ACS Appl. Mater. Interfaces*, 2010, **2**, 3100–3110.
- 19 K. Golovin, S. P. Kobaku, D. H. Lee, E. T. DiLoreto, J. M. Mabry and A. Tuteja, *Sci. Adv.*, 2016, **2**, e1501496.
- 20 T. H. Duong, J.-F. Briand, A. Margaignan and C. Bressy, *ACS Appl. Mater. Interfaces*, 2015, **7**, 15578–15586.
- 21 Z. Yi, C.-J. Liu, L.-P. Zhu and Y.-Y. Xu, *Langmuir*, 2015, **31**, 7970–7979.
- 22 C. Antonini, F. Villa, I. Bernagozzi, A. Amirfazli and M. Marengo, *Langmuir*, 2013, **29**, 16045–16050.
- 23 N. Gao, D. Geyer, F. Pilat, S. Wooh, D. Vollmer, H.-J. Butt and R. Berger, *Nat. Phys.*, 2018, **14**, 191–196.
- 24 H.-J. Butt, I. V. Roisman, M. Brinkmann, P. Papadopoulos, D. Vollmer and C. Semprebon, *Curr. Opin. Colloid Interface Sci.*, 2014, **19**, 343–354.
- 25 V. Liimatainen, M. Vuckovac, V. Jokinen, V. Sariola, M. Hokkanen, Q. Zhou and R. Ras, *Nat. Commun.*, 2017, **8**, 1798.
- 26 D. Daniel, J. Timonen, R. Li, S. Velling and J. Aizenberg, *Nat. Phys.*, 2017, **13**, 1020–1025.
- 27 T. Huhtamäki, X. Tian, J. T. Korhonen and R. H. Ras, *Nat. Protoc.*, 2018, **13**, 1521–1538.
- 28 J. T. Korhonen, T. Huhtamäki, O. Ikkala and R. H. Ras, *Langmuir*, 2013, **29**, 3858–3863.
- 29 A. Kalantarian, *Development of Axisymmetric Drop Shape Analysis-No Apex (ADSA-NA)*, University of Toronto, 2011.
- 30 M. Hoofar and A. Neumann, *J. Adhes.*, 2004, **80**, 727–743.
- 31 S. Srinivasan, G. H. McKinley and R. E. Cohen, *Langmuir*, 2011, **27**, 13582–13589.
- 32 Z. N. Xu, *Rev. Sci. Instrum.*, 2014, **85**, 125107.
- 33 M. Jin, R. Sanedrin, D. Frese, C. Scheithauer and T. Willers, *Colloid Polym. Sci.*, 2016, **294**, 657–665.
- 34 D. Y. Kwok, R. Lin, M. Mui and A. W. Neumann, *Colloids Surf. A*, 1996, **116**, 63–77.
- 35 J. Drelich, *Surf. Innovations*, 2013, **1**, 248–254.
- 36 S. Shumaly, F. Darvish, X. Li, A. Saal, C. Hinduja, W. Steffen, O. Kukharenko, H.-J. Butt and R. Berger, *Langmuir*, 2023, **39**, 1111–1122.
- 37 G. Bradski, *Dr Dobb's Journal: Software Tools for the Professional Programmer*, 2000, **25**, 120–123.
- 38 W. H. Press and S. A. Teukolsky, *Comput. Phys.*, 1990, **4**, 669–672.
- 39 G. Tryggvason, R. Scardovelli and S. Zaleski, *Direct Numerical Simulations of Gas-Liquid Multiphase Flows*, Cambridge University Press, 2011.
- 40 G. K. Batchelor, *An introduction to fluid dynamics*, Cambridge University Press, 2000.
- 41 V. Sanjay, PhD Thesis – Research UT, graduation UT, University of Twente, Netherlands, 2022.
- 42 S. Popinet and Collaborators, Basilisk, 2013–2025, <https://basilisk.fr>, Last accessed: July 22, 2025.
- 43 M. Renardy, Y. Renardy and J. Li, *J. Comput. Phys.*, 2001, **171**, 243–263.
- 44 E. J. Watson, *J. Fluid Mech.*, 1964, **20**, 481–499.
- 45 V. Sanjay, P. Chantelot and D. Lohse, *J. Fluid Mech.*, 2023, **958**, A26.



46 V. Sanjay and D. Lohse, *Phys. Rev. Lett.*, 2025, **134**, 104003.

47 T. Willers and O. Katrin, *European Patent*, EP4109072B1, 2022.

48 N. Nagy, *Langmuir*, 2019, **35**, 5202–5212.

49 D. Bartolo, C. Josserand and D. Bonn, *Phys. Rev. Lett.*, 2006, **96**, 124501.

50 T. V. Nguyen and M. Ichiki, *Microsyst. Nanoeng.*, 2020, **6**, 36.

51 B. Zhang, V. Sanjay, S. Shi, Y. Zhao, C. Lv, X.-Q. Feng and D. Lohse, *Phys. Rev. Lett.*, 2022, **129**, 104501.

52 V. Sanjay, B. Zhang, C. Lv and D. Lohse, *J. Fluid Mech.*, 2025, **1004**, A6.

53 X. Deng, F. Schellenberger, P. Papadopoulos, D. Vollmer and H.-J. Butt, *Langmuir*, 2013, **29**, 7847–7856.

54 N. Savva and J. W. Bush, *J. Fluid Mech.*, 2009, **626**, 211–240.

55 H. Deka and J.-L. Pierson, *Phys. Rev. Fluids*, 2020, **5**, 093603.

56 A. Jha, P. Chantelot, C. Clanet and D. Quéré, *Soft Matter*, 2020, **16**, 7270–7273.

57 P. G. D. Gennes, X. Hua and P. Levinson, *J. Fluid Mech.*, 1990, **212**, 55–63.

58 D. Bartolo, C. Josserand and D. Bonn, *J. Fluid Mech.*, 2005, **545**, 329–338.

59 D. L. Schmidt, R. F. Brady, K. Lam, D. C. Schmidt and M. K. Chaudhury, *Langmuir*, 2004, **20**, 2830–2836.

60 K. M. Wisdom, J. A. Watson, X. Qu, F. Liu, G. S. Watson and C.-H. Chen, *Proc. Natl. Acad. Sci. U. S. A.*, 2013, **110**, 7992–7997.

61 F. Wang, Y. Zhuo, Z. He, S. Xiao, J. He and Z. Zhang, *Adv. Sci.*, 2021, **8**, 2101163.

62 K. Sun, Y. Gizaw, H. Kusumaatmaja and K. Votchovsky, *Soft Matter*, 2025, **21**, 585–595.

63 P.-G. De Gennes, *Rev. Mod. Phys.*, 1985, **57**, 827.

64 M. Jalaal, B. Stoeber and N. J. Balmforth, *J. Fluid Mech.*, 2021, **914**, A21.

65 H. L. França, M. Jalaal and C. M. Oishi, *Phys. Rev. Res.*, 2024, **6**, 013226.

66 J. H. Snoeijer and B. Andreotti, *Annu. Rev. Fluid Mech.*, 2013, **45**, 269–292.

