Role of natural convection in the dissolution of sessile droplets

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The dissolution process of small (initial (equivalent) radius $R_0 < 1\,\text{mm}$) long-chain alcohol (of various types) sessile droplets in water is studied, disentangling diffusive and convective contributions. The latter can arise for high solubilities of the alcohol, as the density of the alcohol–water mixture is then considerably less than that of pure water, giving rise to buoyancy-driven convection. The convective flow around the droplets is measured, using micro-particle image velocimetry (μPIV) and the schlieren technique. When non-dimensionalizing the system, we find a universal $Sh \sim Ra^{1/4}$ scaling relation for all alcohols (of different solubilities) and all droplets in the convective regime. Here $Sh$ is the Sherwood number (dimensionless mass flux) and $Ra$ is the Rayleigh number (dimensionless density difference between clean and alcohol-saturated water). This scaling implies the scaling relation $\tau_c \propto R_0^{5/4}$ of the convective dissolution time $\tau_c$, which is found to agree with experimental data. We show that in the convective regime the plume Reynolds number (the dimensionless velocity) of the detaching alcohol-saturated plume follows $Re_p \sim Sc^{-1}Ra^{5/8}$, which is confirmed by the μPIV data. Here, $Sc$ is the Schmidt number. The convective regime exists when $Ra > Ra_t$, where $Ra_t = 12$ is the transition $Ra$ number as extracted from the data. For $Ra \leq Ra_t$ and smaller, convective transport is progressively overtaken by diffusion and the above scaling relations break down.

Key words: convection, drops and bubbles, plumes/thermals

1. Introduction

Conventional wisdom says that oil and water do not mix. However, some oily liquids, e.g. long-chain alcohols, are slightly soluble in water (see table 1). When a droplet of such an alcohol is placed in a bath of water, it will slowly dissolve, analogous to the much studied topic of sessile droplet evaporation (Cazabat & Guéna 2010; Erbil 2012; Lohse & Zhang 2015), or the dissolution of small gas bubbles

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Figure 1. (Colour online) (a) Snapshot of a dissolving long-chain alcohol droplet (1-hexanol) in water. The dotted black line indicates the location of the silicon substrate, which mirrors part of the droplet. The footprint radius $R_{fp}$ and the contact angle $\theta$ are indicated. A movie of the entire dissolution process may be found online in the supplementary movies available at http://dx.doi.org/10.1017/jfm.2016.158. (b) Evolution of the aforementioned parameters in time, along with the volume $V$ of the droplet and the equivalent radius $R = (3V/(2\pi))^{1/3}$. All parameters have been normalized by their initial values: $V_0 = 750$ nl, $R_{fp,0} = 0.825$ mm, $R_0 = 0.708$ mm and $\theta_0 = 72^\circ$.

(Peñas López, Parrales & Rodríguez-Rodríguez 2015). Figure 1 shows an example of a sessile 1-hexanol droplet in water dissolving in the stick–jump mode (Dietrich et al. 2015; Zhang et al. 2015): unavoidable chemical and geometrical inhomogeneities in the substrate cause the contact line of the droplet to be pinned during dissolution, until the contact angle $\theta$ has decreased to a critical depinning value, which was found to be $62^\circ \pm 2^\circ$. At this point, the contact line depins and the contact angle quickly increases to a value $\theta = 66^\circ \pm 1^\circ$, resulting in a simultaneous decrease of the footprint radius $R_{fp}$. For the 1-hexanol droplet in figure 1, the dissolution time $\tau$ was approximately 3 h. Considering this long dissolution time, it may seem plausible to assume that mass transport away from the droplet is governed by diffusion. Equivalent to the diffusion-driven mass transport from small gas bubbles (Epstein & Plesset 1950) or small sessile droplets (Popov 2005; Stauber et al. 2014; Lohse & Zhang 2015; Zhang et al. 2015), the relevant time scale would in this case then be given by

$$\tau_d = \frac{R_0^2 \rho_d}{2D \Delta c}, \tag{1.1}$$

where $R_0$ is the initial equivalent radius of the droplet, $D$ is the diffusion constant of the alcohol in water, $\rho_d$ is the density of the droplet material, and $\Delta c$ is the difference between the saturated concentration $c_s$ at the droplet interface and the (undersaturated) concentration $c_\infty < c_s$ far away from the drop. However, for the 1-hexanol droplet with an initial radius $R_0 = 0.7$ mm, one finds $\tau_d \approx 11$ h, which is much longer than the 3 h observed experimentally. In previous work (Dietrich et al. 2015), we hypothesized that this discrepancy is caused by the neglect of buoyancy-driven convection of the slightly lighter alcohol–water mixture near the droplet interface. The same idea has been put forward in the context of slowly growing CO$_2$ bubbles in small supersaturations (Enríquez et al. 2014) and evaporating droplets (Shahidzadeh-Bonn et al. 2006). Also in these cases, the rate of mass transport in excess of the diffusion-limited prediction could be explained by assuming a contribution from buoyancy-driven convection. On the other hand, even for millimetre-sized droplets, there also seem to be circumstances under which the diffusive time scale is accurate (Picknett & Bexon...
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Table 1. Properties of the alcohols used in this work, measured at 25 °C: chemical composition; density \( \rho \) of the pure alcohol; diffusion constant \( D \) of the alcohol in water; saturation solubility \( c_s \) of the alcohol in water; molal volume \( \phi^0 \) of the alcohol in water at infinite dilution; density difference \( \Delta \rho = \rho_{\text{H}_2\text{O}} - \rho_{\text{H}_2\text{O,sat}} \) between clean water and water saturated with the alcohol; interfacial tension \( \gamma \) of the water–alcohol interface; and diffusive time scale \( \tau_d \) calculated according to (1.1), for \( R_0 = 0.7 \text{ mm} \). Data were obtained from Crittenden & Hixson (1954)‡, Kinoshita, Ishikawa & Shinoda (1958)*, Høiland & Vikingstad (1976)♣, Stephenson, Stuart & Tabak (1984)♠, Demond & Lindner (1993)♭, Hao & Leaist (1996)†, Romero, Suárez & Jiménez (2007)* and Yalkowsky, He & Jain (2010)♦. The diffusion constants for 2-heptanol and 3-heptanol were assumed to be equal to the 1-heptanol value; and the diffusion constant of 1-octanol was obtained by extrapolating data from Hao & Leaist (1996). No values for \( \gamma \) for 2-heptanol and 3-heptanol could be found.

<table>
<thead>
<tr>
<th>Alcohol</th>
<th>Composition</th>
<th>( \rho ) (kg m(^{-3}))</th>
<th>( D ) (10(^{-9}) m(^2) s(^{-1}))</th>
<th>( c_s ) (kg m(^{-3}))</th>
<th>( \phi^0 ) (cm(^3) mol(^{-1}))</th>
<th>( \Delta \rho ) (kg m(^{-3}))</th>
<th>( \gamma ) (mN m(^{-1}))</th>
<th>( \tau_d ) (10(^4) s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1-Pentanol</td>
<td>C(_5)H(_11)OH</td>
<td>811</td>
<td>0.888†</td>
<td>22‡</td>
<td>102.62♣</td>
<td>3.42</td>
<td>4.4♣</td>
<td>1.0</td>
</tr>
<tr>
<td>1-Hexanol</td>
<td>C(_6)H(_13)OH</td>
<td>814</td>
<td>0.83‡</td>
<td>5.9‡</td>
<td>118.65♣</td>
<td>0.92</td>
<td>6.8♣</td>
<td>4.0</td>
</tr>
<tr>
<td>1-Heptanol</td>
<td>C(_7)H(_15)OH</td>
<td>822</td>
<td>0.800†</td>
<td>1.67*</td>
<td>136.95*</td>
<td>0.29</td>
<td>7.7♣</td>
<td>14.5</td>
</tr>
<tr>
<td>2-Heptanol</td>
<td>C(_7)H(_15)OH</td>
<td>817</td>
<td>0.800</td>
<td>3.5*</td>
<td>134.39♣</td>
<td>0.53</td>
<td>—</td>
<td>7.6</td>
</tr>
<tr>
<td>3-Heptanol</td>
<td>C(_7)H(_15)OH</td>
<td>818</td>
<td>0.800</td>
<td>4.0*</td>
<td>133.30♣</td>
<td>0.57</td>
<td>—</td>
<td>5.8</td>
</tr>
<tr>
<td>1-Octanol</td>
<td>C(_8)H(_17)OH</td>
<td>827</td>
<td>0.780</td>
<td>0.5‡</td>
<td>148.41*</td>
<td>0.07</td>
<td>8.52♣</td>
<td>48.1</td>
</tr>
</tbody>
</table>
This reflects the existence of a threshold for convection. However, the details of this threshold remain unclear. The low velocities and small refractive index differences involved often inhibit direct observation of the buoyant flow in the surrounding medium. As far as we are aware, direct visualization attempts of the external flow have only been undertaken in the context of evaporating droplets, using either schlieren (Kelly-Zion, Batra & Pursell 2013a), infrared spectroscopy (Kelly-Zion et al. 2013b) or interferometry (Dehaeck, Rednikov & Colinet 2014) techniques, and only very recently by tracing tiny oil droplets in air (Somasundaram, Anand & Bakshi 2015).

In this work we combine qualitative schlieren imaging with quantitative micro-particle image velocimetry (µPIV) to directly visualize the concentration field and flow around slowly dissolving droplets of various types of long-chain alcohols in clean water. We show that, above a transition solutal Rayleigh number, which corresponds to the buoyancy of the alcohol–water mixture, which is lighter than the surrounding clean water, the solute is mainly transported away in a single steady plume above the droplet. Knowledge of the flow structure allows us to derive scaling laws for both the dissolution rate and the plume velocity in the convective regime, which are in good agreement with experiments. Finally, as the droplet shrinks and its Rayleigh number drops below the transition value, a transition occurs in which convection dies out and is overtaken by diffusion.

2. Experimental procedure
2.1. Materials and preparation
As shown in table 1, the solubility of long-chain alcohols strongly depends on their length, while other properties, such as density and diffusion coefficient, are relatively insensitive to this. By increasing the number of carbon atoms in the chain from five (pentanol) to eight (octanol), one decreases the solubility (and thereby the buoyant force of the water–alcohol mixture) by two orders of magnitude. This makes these alcohols very suitable to study the possible transition between diffusion and convection. Alcohols with purities of \( \geq 98 \% \) (Sigma-Aldrich) were used. The density of the alcohol–water mixture (table 1) was calculated for a mixture at 100 \% saturation, using the molal volume \( \phi V \) at infinite dilution (Høiland & Vikingstad 1976; Romero et al. 2007). The molal volume \( \phi V \) gives the volume occupied by one mole of solute in the solvent. The assumption is made that \( \phi V \) is independent of the solute concentration, which introduces a negligible error in \( \Delta \rho \) of \( <1 \% \) when compared to the direct density measurements given by Romero et al. (2007).

A sketch of the experimental set-up is provided in figure 2. All measurements were conducted in a cubical glass tank of \( 5 \text{ cm} \times 5 \text{ cm} \times 5 \text{ cm} \). The container was cleaned using isopropyl alcohol and water, and then filled with 100 ml of clean water. This water was obtained from a Reference A+ system (Merck Millipore, at 18.2 MΩ cm) several hours before the measurement and stored in a clean flask to equilibrate and thus reduce thermal convective currents. After the tank was filled, a single droplet was dispensed from a glass syringe with a Teflon plunger, fitted in a motorized syringe pump. The droplet was placed on a hydrophobized silicon wafer \( \approx 1 \text{ cm} \times 1 \text{ cm} \) (P/Boron/(100), Okmetic), placed at the bottom of the tank. Hydrophobization was achieved by coating the wafer with a self-assembled monolayer of PFDTDS (1H,1H,2H,2H-perfluorodecyltrichlorosilane 97 \%, ABCR GmbH, Karlsruhe, Germany), following the procedure described earlier (Karpitschka 2012). Prior to each experiment, the samples were cleaned by insonication in acetone for 10 min and dried under a stream of nitrogen. After the droplet was placed on the substrate, the needle was removed and the tank was closed.
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Figure 2. (Colour online) Experimental set-up, showing the glass tank (1), with the substrate and the droplet in place (not to scale). The droplet was deposited under water using a syringe (2) fitted in a motorized syringe pump (not drawn). A dichroic mirror (3) was used to couple the laser beam (4) into the long-distance microscope (5). A trigger-delay box (6) synchronized the laser pulses with the camera exposure. Parts (3), (4) and (6) were used in the μPIV measurements only. The assembly used for schlieren, consists of a positive lens (7) and a knife edge (8), located at the focal point of the lens. The parallel LED light source (9) was used in all experiments, with the exception of the PIV measurements.

2.2. Imaging

The droplet was illuminated from one side using a collimated light-emitting diode (LED) light source (Thorlabs, wavelength $\lambda = 625$ nm) and imaged onto a charge-coupled device (CCD) camera (Pixelfly USB, PCO Germany), with a long-distance microscope providing a magnification up to 16×. The images were recorded at a rate of one frame per second (f.p.s.), and post-processed using a Matlab code to extract the droplet profile with subpixel accuracy (van der Bos et al. 2014). Since all droplets were smaller than the capillary length ($\sqrt{\gamma/(\rho_{H_2O} - \rho_{alcohol})} g \approx 2$ mm), the droplet profile could be accurately fitted to a spherical cap to obtain the radius of curvature and contact angle. With this method, droplets could be traced until $V < 0.05V_0$.

2.3. Schlieren

The concentration gradients developing around the dissolving droplets were qualitatively visualized using the schlieren technique (Settles 2001). For this, a positive lens and a knife edge were placed at the camera side of the tank, as shown in figure 2. After passing through the tank, the parallel light from the LED source is focused onto the edge of a sharp knife placed perpendicular to the beam. To be sensitive to both horizontal and vertical concentration gradients, the knife edge was placed at an angle of 45°. In the resulting image, solute-rich regions are visible as local changes in light intensity.

2.4. μPIV measurements

For the μPIV measurements, the water in the tank was seeded with red fluorescent tracer particles (Fluoro-Max, Thermo Fisher Scientific, 3 μm diameter). A pulsed green laser, (Nd:YAG, $\lambda = 532$ nm) was coupled into the microscope by a dichroic mirror. The focal plane of the microscope was centred at the droplet. Therefore,
only tracer particles in the $\approx 100 \, \mu m$ thick focal plane were imaged, producing a two-dimensional velocity field around the symmetry axis of the droplet. The red light ($\lambda = 612 \, \text{nm}$) emitted by the fluorescent particles was recorded by the CCD camera at 8 f.p.s. A BNC 575 pulse/delay generator was used to synchronize the laser pulse and the camera exposure. The obtained images were then post-processed in ImageJ to remove static features and to enhance the contrast. Consecutive image pairs were analysed with JPIV, using an interrogation window of $32 \times 32$ pixels, corresponding to $\approx 70 \, \mu m \times 70 \, \mu m$. The concentration of tracer particles was kept low to avoid excessive absorption and blurring by out-of-focus particles. Because of this low particle density, the velocity fields from multiple image pairs were combined for improved accuracy (Raffel et al. 2007).

3. Visualization results

Figure 3 shows snapshots of the $\mu$PIV and schlieren measurements for 1-pentanol ($a,c$) and 1-heptanol droplets ($b,d$). The dissolving 1-pentanol droplet generates a clear plume originating from its apex, while fresh liquid is drawn in from the sides. The tip of the plume ends in a vortex ring. As this ring moves away from the droplet as the experiment proceeds, the effect of the transient flow generated by the vortex
Figure 4. (Colour online) Sketch of a sessile droplet dissolving at a rate $\dot{m}_d$ and the convective plume. A concentration boundary layer of thickness $\delta_c$ develops over the droplet interface; since $Sc \gg 1$, the velocity boundary layer has a thickness $\delta_v > \delta_c$. The boundary layers merge into plumes with diameters $2\delta_c$ and $2\delta_v$, moving at a vertical velocity $v_p$. The boundary layer widths $\delta_c$ and $\delta_v$ will change as a function of height, but are linked via the relation $\delta_v/\delta_c \sim \sqrt{Sc}$ (Bejan 1993).

ring diminishes rapidly, leaving behind a single plume (see figures 3c and 11). For 1-heptanol, with its lower solubility, the plume is far less pronounced. The particle velocities are significantly lower and the contrast of the schlieren image had to be strongly enhanced to see the plume at all (see figure 3d). The weak 1-heptanol plume seems to be affected by a small mean flow in the cell, possibly caused by thermal convection due to changes in room temperature or the illumination. As we will show later on, this mean flow seems to have little influence on the dissolution behaviour. Appendix A contains additional µPIV results, including time-resolved velocity fields around a 1-pentanol droplet, the velocity field around an insoluble sessile droplet, and the flow around a dissolving sessile droplet placed on a vertical substrate. A movie, showing the motion of the bulk around the dissolving droplet, is available online in the supplementary movies.

The two different techniques used in figure 3 reveal that the convective plume displays two different features, as illustrated in figure 4. Firstly, the schlieren images visualize the plume-shaped region that contains dissolved alcohol. The concentration profile is characterized by a width $2\delta_c$, which increases as $\delta_c \propto \sqrt{Dz/v_p}$, with $z$ the height above the droplet, and $v_p$ the plume flow speed. Secondly, the buoyant force on the (lighter) water–alcohol mixture results in a flow, as visualized by the µPIV. The velocity profile of this flow (also drawn in figure 4) is characterized by a width $2\delta_v$. The liquid viscosity $\nu$ also causes the velocity profile to broaden for increasing height, namely with the same height dependence as the concentration profile, i.e. $\delta_v \sim \sqrt{vz/v_p}$. Therefore, the ratio $\zeta$ between the widths of the concentration and velocity profiles is fixed, $\zeta = \delta_v/\delta_c \sim \sqrt{Sc}$ (Bejan 1993), where $Sc$ is the Schmidt number, $Sc \equiv \nu/D$. In the current system, $Sc \approx 1200$, so $\delta_v \approx 30\delta_c$ is expected.

To obtain a theoretical description for the velocity and concentration profiles at $Sc = 1200$, we solved equations (II.8) and (II.9) in the paper of Fujii (1963), who described the analogous case of a thermal plume above a heat source. Here, we
followed the numerical procedure described by Vázquez, Pérez & Castellanos (1996), and used equation (II.15) in Fujii’s paper as a condition in the solving procedure, to obtain the velocity and concentration profiles at $Sc = 1200$, as shown in figure 5(a) by the black and pink lines, respectively. The lateral coordinate $\tilde{X}$ in the theory is scaled by $\sqrt{L\epsilon}$, with $L = \sqrt{2\pi\nu^3/(g\beta_c \dot{m}_d)}$, where $g$ is the acceleration of gravity, $\beta_c \equiv (\partial \rho/\partial c)/\rho_b$ is the solutal expansion coefficient, and $\dot{m}_d$ is the mass loss rate of the droplet. The high $Sc$ number in the current system results in distinct shapes for the velocity and concentration profiles. The measured value for $\zeta$ therefore depends on the definition of the plume width, as shown in figure 5(a). The anticipated $\zeta = 30$
is retrieved when evaluated at a relative amplitude of 0.1, corresponding to the 90% boundary layer definition.

To compare our measurements to the theoretical profiles, a cross-sectional intensity profile was measured in the schlieren image (figure 3c) at a height of 300 µm above the apex of the droplet (as illustrated by the dashed line in figure 3c). This intensity profile is plotted as the solid red line in figure 5(b), where we have to keep in mind that it represents the first derivative of the concentration profile, as it is the result from a schlieren measurement. To allow for an easy comparison, we plotted the theoretical concentration profile (pink solid line) from figure 5(a) and its derivative (red dashed line) in figure 5(b), and fitted the derivative to match the experiment. From this fit, a conversion factor was obtained to translate the dimensionless lateral coordinate of the theory to the length scale of the experiment. We executed the same procedure to obtain the velocity profile from the µPIV data in figure 3(a), again at 300 µm above the droplet. This profile is plotted as the black circles in figure 5(c). The theoretical velocity profile (black solid line) is superimposed on the measurement, where the previously found conversion factor was used to match the lateral coordinate. Comparison of the experimental and theoretical profiles in figure 5(c) reveals that, while the central part of the plume shows fair agreement with the theoretical profile, the general shape of the plume is much narrower than expected from theory. The cause of this discrepancy is not understood as of yet. Possibly the vortex, substrate and droplet influence the plume shape, and the expected profile can be recovered when measured at higher distance above the droplet.

As mentioned before, the plume changes over time. To visualize the evolution of the plume, cross-sections are taken in the µPIV data at subsequent times, at 300 µm above the 1-pentanol droplet. These cross-sections are plotted in figure 5(d), and show that the plume properties are linked to the droplet size: both the width and the maximum velocity of the plume steadily decrease as the droplet shrinks. At \( t = 3000 \) s, the droplet has dissolved completely.

4. Dissolution rate and plume velocity

The µPIV and schlieren images show that the convective flow around small droplets takes the form of a thin boundary layer over the droplet interface, culminating in a single plume rising from its apex. A schematic drawing of this flow and the concentration profile is shown in figure 4. If we assume (1) that at the interface of the droplet the solute concentration is constant and equal to \( c_s \), and (2) that the droplet shrinks sufficiently slowly, then this situation is mathematically equivalent to the buoyant flow around a hot sphere of constant temperature \( T \) and fixed radius \( R \). The second statement relies on a quasi-steady approximation, and is supported by the fact that, firstly, \( \tau/(R^2_0/D) \gg 1 \), reflecting that the time scale of the boundary layer effects is much smaller than the total dissolution time of the droplet. Secondly, due to dissolution, the droplet interface moves at a speed \( \dot{R} \ll 1 \) µm s\(^{-1}\), from which we can calculate the Péclet number \( Pe = \dot{RR}/D \ll 1 \), indicating that advective mass transport due to the motion of the droplet interface is much smaller than the diffusive mass transport. This, combined with the fact that the measured bulk velocities are \( >10 \) µm s\(^{-1}\) \( \gg \dot{R} \), allows us to safely regard the droplet as quasi-steady. In the context of thermal convection, the flow structure in both the boundary layer and plume are well known (see e.g. Fujii 1963; Bejan 1993). In this section we recapitulate the main findings in terms of the dissolution problem and compare them directly to our observations.
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\[ R(t) = \left( \frac{3V}{(2\pi)} \right)^{1/3} \] as a function of time for 1-pentanol droplets (a) and 1-heptanol droplets (b) dissolving in water. The inset in (a) illustrates how, for each droplet with initial radius \( R_0 \), the lifetime \( \tau \) is estimated to lie between the lower estimate \( \tau_l \) at the end of the experiment, and the upper estimate \( \tau_u \), found by extrapolation using (5.1) (green dashed line). The lifetime \( \tau \) is plotted as a function of \( R_0 \) in panels (c) and (d) for 1-pentanol and 1-heptanol measurements, respectively. The lines in panels (c) and (d) illustrate the \( \tau \sim R_0^2 \) and \( \tau \sim R_0^{5/4} \) relations, as expected for diffusion and convection, respectively. The vertical line in (d) indicates the transition \( Ra \) number \( Ra_t = 12 \), which marks the transition between convection (\( Ra > Ra_t \)) and diffusion (\( Ra < Ra_t \)). For 1-pentanol, \( Ra_t = 12 \) corresponds to \( R = 0.07 \) mm.

4.1. Dissolution rate

To study the droplet dissolution dynamics as a function of droplet liquid and size, individual droplets of varying initial volume and alcohol type were imaged throughout the dissolution process. Since the footprint radius \( R_{fp} \) shows steps due to the stick–jump mode dissolution (Dietrich et al. 2015; Lohse & Zhang 2015; Zhang et al. 2015), we define the equivalent radius \( R = (3V/(2\pi))^{1/3} \) to provide a continuously decreasing measure for the droplet size. Figure 6 shows \( R(t) \) for 1-pentanol (a) and 1-heptanol (b) droplets. From this, the mass loss rate \( \dot{m}_d \) was extracted and plotted as a function of \( R \) in figure 7 for all six alcohols. Note that, while \( \dot{m} < 0 \) for a shrinking droplet, we define the droplet mass loss rate as a positive amount, as it provides a more intuitive measure for the dissolution process. Using this, figure 7 shows that the measured mass loss rates for the various droplet sizes and alcohol types span two orders of magnitude. To find a universal description for the dissolution dynamics, we continue by defining dimensionless numbers, which take both the droplet size and liquid into account.

In (convective) heat exchange problems, the heat exchange is usually expressed in terms of the (dimensionless) Nusselt number, which is the ratio of the heat transfer...
Figure 7. (Colour online) Rate of mass loss as a function of $R$ for droplets of different alcohols. The mass loss rates are ordered as a function of alcohol solubility, with 1-pentanol and 1-octanol being the best and least soluble alcohols, respectively.

rate and the rate for pure diffusion. The equivalent for solutal convection is the Sherwood number,

$$Sh \equiv \frac{\langle \dot{m}_d \rangle_A R}{D \Delta c}, \quad (4.1)$$

where $\langle \dot{m}_d \rangle_A$ is the actual (measured) mass transfer flux (rate per area), averaged over the droplet surface area $A$. This mass flux is compared to $D \Delta c / R$, the mass flux of pure (steady) diffusion from an equally sized spherical droplet (or a sessile droplet with $\theta = 90^\circ$). In the case of pure diffusion from our sessile droplets ($45^\circ < \theta < 75^\circ$), we expect to find a diffusion-limited Sherwood number $0.9 < Sh_d < 1.3$, the exact value of which depends on the droplet contact angle, as discussed in appendix B. For the case of laminar flow at high $Sc$ number, Bejan (1993) provides a complete and insightful derivation of the momentum equation, showing that the flow can be described using the Boussinesq approximation of the Navier–Stokes equation. This approximation assumes a slender boundary layer (i.e. $\delta_c \ll R$, which is justified by figures 3(c) and 3(d), which indeed show a thin boundary layer over the droplet), a constant pressure over the width of the boundary layer and a limited density difference. For high $Sc$ numbers, the buoyant force is balanced by viscosity and it can be shown that $\delta_c / R \sim Ra^{-1/4}$, independent of $Sc$ (Bejan 1993). Here $Ra$ is the Rayleigh number, which is the ratio of the buoyant force to the damping force,

$$Ra \equiv \frac{g \beta_c \Delta c R^3}{\nu D}. \quad (4.2)$$

Taking $\delta_c$ as the typical length scale over which diffusion takes place in the presence of convection, we find $\langle \dot{m}_d \rangle_A \sim D \Delta c / \delta_c$, so that

$$Sh \sim R / \delta_c \sim Ra^{1/4}, \quad (4.3)$$

again independent of $Sc$. If we recast the data from figure 7 in terms of the $Ra$ and $Sh$ numbers, all datasets from the six different alcohols collapse, as shown in figure 8. This figure also reveals that, for large $Ra$, the data follow the anticipated $Sh \sim Ra^{1/4}$ scaling, which is plotted as the dashed line. For small $Ra$ numbers, $Sh$ converges.
Figure 8. (Colour online) Sherwood number as a function of Rayleigh number. The plot shows the mean value and the spread for a total of 70 measurements on droplets with initial volumes $2 \text{ nl} \leq V_0 \leq 1200 \text{ nl}$. Since the $Ra$ number depends on both the droplet size and its material properties, high-$Ra$ droplets are easily made using large droplets of 1-pentanol, whereas low-$Ra$ droplets are best studied using small droplets of the poorly soluble 1-heptanol and 1-octanol. Equation (4.4) is plotted as the solid black line, using $Ra_t = 12.1$, $Sh_d = 1.2$ and $n = 1.0$.

to a plateau, as expected for diffusion. It is noteworthy that the $Sh(Ra)$ dependence from Enríquez et al. (2014), who studied the growth of CO$_2$ gas bubbles in slightly supersaturated water, is almost identical to our figure 8. This indicates that the flow structures around bubbles and droplets are very similar.

To better understand the transition between the convective and the diffusive behaviour, and to find the value of the transition $Ra$ number $Ra_t$, we fitted a crossover function of the form

$$Sh(Ra) = Sh_d \left[1 + \left(\frac{Ra}{Ra_t}\right)^n\right]^{1/(4n)}$$

(4.4)

to the data. Here $n$ is a fitting parameter that describes the sharpness of the transition. Equation (4.4) was fitted to the individual datasets of each alcohol, to obtain $Ra_t = 12.1 \pm 5.8$, $Sh_d = 1.2 \pm 0.2$ and $n = 1.0 \pm 0.5$. Equation (4.4) is plotted in figure 8, using the mean values. The fitted curve confirms that, for $Ra > Ra_t \approx 12$, the data follow the $Sh \sim Ra^{1/4}$ scaling.

A transition exists around $Ra_t$, where the contribution of convective mass transport gradually decreases. When $Ra < Ra_t$, we obtain the diffusive limit $Sh \approx 1.2$, independent of $Ra$. A convective contribution to the evaporation of water droplets on mica has been claimed by Shahidzadeh-Bonn et al. (2006), for droplets with radii of approximately 1 mm, corresponding to $Ra = 10$. However, they did not directly measure or visualize a convective flow, and their finding was rebutted by Guéné, Poulard & Cazabat (2006), who studied the same system and excluded a convective contribution by flipping the system upside down. This hindered the development of a convective flow, and no difference in evaporation was reported whether the droplet was sessile or hanging, demonstrating that no convection developed around water droplets with radii up to 3 mm. The absence of convection around droplets with Rayleigh numbers well above the found value of $Ra_t$ can be explained by
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Shahidzadeh-Bonn et al.’s (2006) choice of the droplet diameter as characteristic droplet size, leading to an overestimation of the Rayleigh number. Water droplets fully wet the hydrophilic mica substrate, resulting in a large difference between the droplet lateral size and its height, and thus to only a seemingly high value of $Ra$, when taking the diameter. For the convective problem, it makes a great difference whether the solute-rich surface is parallel or perpendicular to the direction of gravity. This is highlighted by the fact that, in the case of a horizontal heated surface, there exists a critical value for the onset of convection, while this is not the case for a vertical heated surface, where a smooth transition is to be expected, as we find in the present case. We therefore believe that the length scale parallel to the direction of gravity is the more appropriate choice. As a final remark on this issue, we would like to encourage future work on droplet evaporation to be done in a humidity-controlled environment, as the density of air changes with humidity, which potentially could result in a different onset of convection and a different Rayleigh number.

4.2. Plume velocity

Similar to the concentration boundary layer around the droplet, the local velocity and structure of a convective plume are determined by a competition between buoyant and viscous stresses. For a thermal plume, this is described by the local thermal Rayleigh number $Ra(z)$, based on the local temperature difference between the centre of the plume and the surroundings (see e.g. Fujii 1963; Vázquez et al. 1996). The local solutal Rayleigh number can be written similarly, based on the local concentration difference $\Delta c(z)$,

$$Ra(z) \equiv \frac{g\beta c \Delta c(z) z^3}{vD}.$$  (4.5)

The plume can be linked to the dissolving droplet, as the transport of solutes inside the plume must equal the dissolution rate of the droplet,

$$\dot{m}_d \sim \delta_c^2 \Delta c v_p,$$  (4.6)

with $v_p$ the central velocity of the plume. Similarly to the previous derivation of the flow over the droplet interface, where again a slender boundary is assumed (as verified by figures 3(c) and 3(d)), $R$ is replaced by the vertical coordinate $z$, it can be shown (Fujii 1963; Bejan 1993) that for $Sc \gg 1$ the plume width $\delta_c$ behaves as

$$\delta_c \sim z (Ra(z))^{-1/4} \propto z^{1/2}.$$  (4.7)

and the plume velocity as

$$v_p \sim \frac{D}{z} (Ra(z))^{1/2} = \left(\frac{g\beta c \dot{m}_d}{v}\right)^{1/2}.$$  (4.8)

Substituting these relations into (4.6), we find the relation between $\Delta c$ and $\dot{m}_d$ as $\Delta c \sim \dot{m}_d/(Dz)$, so that (4.5) can also be written as

$$Ra(z) \sim \frac{g\beta c \dot{m}_d z^2}{D^2 v}.$$  (4.9)

Again by exploiting the analogy with the thermal case, one finds the following scaling behaviours for the width of the plume $\delta_c(z)$ and the central velocity $v_p$ of a solutal...
The plume Reynolds number as a function of $Ra$ number for three individual 1-pentanol droplets with $V_0 = 700$ nl (red), $V_0 = 550$ nl (green) and $V_0 = 140$ nl (blue). The expected $Re \sim Ra^{5/8}Sc^{-1}$ scaling is indeed found when $Ra \gg Ra_t$, but breaks down when $Ra$ approaches $Ra_t$. This marks the onset of the transition region (located around $Ra_t$), in which convective transport is gradually exceeded by diffusive transport. To obtain the fits to the two measurements, plotted in red and blue, an additional prefactor of 0.25 (red/green) and 0.1 (blue) was required.

To test this scaling, we measure the maximum vertical velocity in the µPIV data at a height of 300 µm above the droplet, together with the size of the droplet, and use this to calculate $Re_p(Ra)$. The result of this analysis is shown in figure 9. For $Ra \gg Ra_t$, we find the anticipated $Re_p \sim Ra^{5/8}$. Note that this scaling breaks down already around $Ra \approx 400 \gg Ra_t$, reflecting the broad transitional regime also observed for the Sherwood number (figure 8) and as quantified by the value of the fitting parameter $n = 1$ in (4.4). For the measurements shown, there seems to be some dependence of

\[ Re_p \sim Ra^{5/8}Sc^{-1}. \]
the scaling prefactor on the initial size of the droplet. The smallest droplet displays a somewhat lower overall plume velocity than the two larger ones. We are not yet able to pinpoint the reason for this difference. One possible hypothesis might be that inertia of the bulk flow plays a role, and that the larger droplets create a stronger convection that persists throughout the dissolution. However, more work is required to confirm or rebut this hypothesis.

5. Dissolution time

The convective flow and the related increase in mass transport were described in the previous section. In this section we proceed by deriving an expression for the convective droplet dissolution rate and associated dissolution time $\tau_c$. However, we start by briefly introducing diffusive dissolution, which we will use later on.

An expression for the diffusive volume loss rate $\frac{dV}{dt}$ has been given by Popov (2005) in the context of evaporating sessile water droplets. Popov’s solution can be rewritten to find the rate of change of the droplet, expressed in terms of the previously introduced equivalent radius $R$,

$$\frac{dR}{dt} = -\frac{D\Delta c}{2\rho R} f(\theta) \left[ \frac{2}{2 - 3 \cos \theta + \cos^3 \theta} \right]^{1/3} \sin \theta,$$

where

$$f(\theta) = \frac{\sin(\theta)}{1 + \cos(\theta)} + 4 \int_0^\infty \frac{1 + \cosh(2\theta \epsilon)}{\sinh(2\pi \epsilon)} \tanh[(\pi - \theta) \epsilon] \mathrm{d}\epsilon$$

is a geometrical shape factor to describe the effect of the impenetrable substrate. Note that for simplicity we have neglected the intermittent contact line pinning which was observed in the experiments (Dietrich et al. 2015; Zhang et al. 2015), and (5.1) describes dissolution in the constant contact angle mode. Integration of (5.1) results in the dissolution time, with the associated diffusive time scale given by (1.1), i.e. in particular $\tau_d \propto R_0^2$.

We can perform a similar calculation for the convective mass exchange, again based on the cooling sphere analogy. An important difference between the cooling sphere and our dissolving droplet is that, in the latter case, the radius decreases in time. However, if the dissolution is slow, we can assume the process to be quasi-static and neglect this effect. We start by equating the rate of mass loss, $\dot{m}_d \sim -R^2 \rho (\frac{dR}{dt}) > 0$, to the rate at which mass is carried away in the convective plume, $\dot{m}_p = AD\Delta c Sh/R \sim AD\Delta c Ra^{1/4}/R$, with $A \propto R^2$ the area of the droplet–bulk interface. From this we obtain

$$\frac{dR}{dt} = -a \left( \frac{g\beta c \Delta c^5 D^3}{\nu \rho_0^4 R} \right)^{1/4},$$

with a prefactor $a$ of order 1. Separation of variables and integrating $R$ from $R=R_0$ till $R=0$, and time from $t=0$ till $t=\tau_c$, gives the dissolution time with the associated convective time scale $\tau_c$ with

$$\tau_c = \frac{4}{5a} \left( \frac{\nu \rho_0^4 R_0^5}{g\beta c \Delta c^5 D^3} \right)^{1/4}.$$

Therefore, for droplets dissolving in the convection-dominated regime, we expect a dissolution time $\tau_c \propto R_0^{5/4}$, with a material-dependent prefactor.
To test this scaling behaviour, we used the $R(t)$ curves in figure 6(a,b) and extracted the dissolution time from each droplet. Since the droplets could not be measured until complete dissolution, the actual value of $\tau$ had to be estimated. Therefore, we assumed that the last stage of the dissolution process was diffusion-limited and extrapolated the droplet evolution by integrating (5.1), using the smallest still measured droplet size of the experiment as initial value. This extrapolation (illustrated by the green dotted line in the inset of figure 6(a)) provides the upper bound of $\tau$, whereas the lower bound is given by the time at which the experiment was terminated. The values for $\tau$ thus obtained are plotted as a function of $R_0$ for 1-pentanol and 1-heptanol in figure 6(c,d), respectively. For larger droplets we indeed find $\tau \propto R_0^{5/4}$ as expected from (5.4), while for smaller 1-heptanol droplets $\tau \propto R_0^2$ is found, as expected for pure diffusion. The vertical line in figure 6(d) indicates $Ra = 12$, showing that the transition from diffusive to convective dissolution occurs around $Ra_t$, consistent with our findings in § 4.

Now that we have confirmed the $\tau_c \propto R_0^{5/4}$ behaviour for large droplets, we finally test whether (5.1) and (5.3) provide accurate descriptions of the dissolution dynamics in the diffusive and convective regimes, respectively. Moreover, we can test whether the transition between these regimes indeed occurs around $Ra_t = 12$, as found before. To this purpose, the curves in figure 6(a,b) are replotted in figure 10(a,b) as a function of the time $t - \tau$. Figure 10(c) and (d) provide a close-up of the final stage of dissolution, the outlined parts of panels (a) and (b), respectively. Along with the experimental data, we plotted the numerical integration of the convective dissolution model, (5.3), which is the upper black line in all panels. We also plotted the diffusive dissolution model, (5.1), represented by the lower black line in each panel. Figure 10(a–d) shows that, for $Ra \gg Ra_t$, (5.3) accurately captures the droplet dissolution process, reflecting convection-dominated dissolution. The value for the prefactor $a$ in (5.3) was adjusted for each alcohol to obtain a good fit in the convective regime. The values used are listed in table 2, along with $\theta$, for all alcohols. For $Ra \approx Ra_t$, the overlap between our convective model and the experiments becomes worse, consistent with the transition from convection-dominated to diffusion-dominated dissolution. The final stage of dissolution is best observed in figure 10(c,d). When $Ra < Ra_t$, the dissolution is purely diffusive, reflected by the good overlap between (5.1) and the measurements. The above findings confirm the applicability of our convective dissolution model when $Ra > Ra_t$, and validate $Ra_t \approx 12$ as the transitional $Ra$ number for the transition from convective to diffusive dissolution dynamics.

### 6. Conclusion

Sessile droplets of long-chain alcohols immersed in water are ideally suited to experimentally study the basic laws of mass transfer around small objects. By
choosing alcohols of different chain lengths, the alcohol’s solubility in water can be varied by almost two orders of magnitude, while its other properties remain practically the same. This large range of solubilities allowed us to vary the convective driving parameter, the Rayleigh number, by over six orders of magnitude, while keeping the droplets large enough to visualize their shrinkage and the flow around them.

Using a combination of μPIV and schlieren technique, we directly demonstrated that, above a transition Rayleigh number $Ra_t \approx 12$, a buoyant flow develops around a dissolving droplet, due to the density differences between the lighter alcohol–water mixture, as compared to the heavier clean water. By modelling the observed boundary layer structure at the droplet interface and in the plume, we derived a basic scaling relation $Sh \sim Ra^{1/4}$ for convective mass transport. Using this relation as a starting point, we derived expressions for the shrinkage rate of the droplet and the velocity of the plume. In the convective regime, these models are in good agreement with our data. However, once the droplet dissolves to a size close to $Ra_t$, diffusion gradually overtakes convective mass transport and the convective scaling relations break down.

The observed convection and associated increase in mass transport confirms earlier work on growing bubbles in supersaturated water (Enríquez et al. 2014) and evaporating droplets (Kelly-Zion et al. 2013a), indicating that it is a universal phenomenon. It should be noted that the considerations in this work assume that the plume is transported away from the substrate, against gravity. In the opposite case

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**Figure 10.** (Colour online) Equivalent radius $R = (3V/(2\pi))^{1/3}$ as a function of time $t−\tau$ to complete dissolution for 1-pentanol droplets (a) and 1-heptanol droplets (b). (c,d) Zooms of the outlined parts of panels (a) and (b), respectively. The upper and lower black curves in each panel represent the integration of (5.3) and (5.1), respectively. When $Ra > Ra_t = 12$, (5.3) accurately describes the dissolution dynamics. Around $Ra_t$, which is indicated by the horizontal line in each panel, a transition exists, in which convection is progressively overtaken by diffusion. In the final stage ($Ra < Ra_t$), the dissolution is well described by diffusion, i.e. (5.1). The colour coding for the individual measurements is the same as in figure 6.
in which the surface is placed upside down, different effects such as the formation of a stable stratification layer near the surface will also play an important role. Also we expect that the prefactor $a$ found in our experiments will be affected by parameters such as the geometry of the droplet, confinements around the droplet and the inclination of the substrate.

The value for $Ra_t$ presented in this work provides an important indication to determine the dominant transport mechanism in droplet dissolution. In conjunction with this, the convective dissolution model allows for more accurate predictions of droplet dissolution times. The demonstrated predictability of the dissolution behaviour of single sessile droplets on a horizontal substrate invites the testing of the derived scaling relations and fitting parameters in more complicated situations. For example, the applicability of the derived scaling relations and measured value for $Ra_t$ could be tested in the context of bubble growth or droplet evaporation. Moreover, interesting changes in the flow profile can be expected, for example, when the orientation or the wettability of the substrate is changed. Other possible research directions include placing multiple droplets close together, to study their interaction, or making the droplets so large ($R > \lambda_c$) that they form a puddle.

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Supplementary movies

Supplementary movies are available at http://dx.doi.org/10.1017/jfm.2016.158.

Appendix A. Micro-particle image velocimetry data

The flow structure around the dissolving droplet has been measured using $\mu$PIV, utilizing the procedure and set-up described above. In figure 11, the time evolution of the velocity field around a dissolving 1-pentanol droplet ($V_0 = 700$ nl) is shown. Shortly after deposition of the droplet (figure 11a), the data reveal the presence of a toroidal vortex above the droplet, and a strong narrow plume originating from the droplet apex. At 600 s into the dissolution process, shown in figure 11(b), the centre of the vortex is observed to have moved away from the droplet, causing the plume to slow down and broaden. Figure 11(c) shows the droplet at 2000 s after deposition: the droplet has shrunk considerably; however, a small plume is still visible, along with a small mean flow, from right to left. Close to the end of the experiment ($t = 2800$ s, figure 11d), the droplet has almost disappeared, as has the plume. A small right-to-left mean flow is still observable.

To show that the plume is caused by solutal convection, and not simply by the presence of a spherical object at the interface, the experiment is repeated using an equally sized droplet of 1-decanol. This alcohol has negligible solubility, $c_s = 0.037$ g l$^{-1}$ (Kinoshita et al. 1958). For this droplet, $Ra \approx 10$, which means that solutal convection should be absent. The velocity field around this 1-decanol droplet is shown in figure 12. From this figure, it is clear that the presence of the droplet does not cause the formation of a plume. A slight mean flow is present, which can be seen to flow around the droplet. Although not visible in figure 12, it should be noted
that, in this particular experiment, tracer particles adhered to the droplet interface, something that did not happen in all other experiments where soluble droplets were studied.
Hypothetically, the convection could be induced by surface tension gradients as well, developing over the droplet–water interface. These gradients would cause Marangoni convection over the alcohol–water interface (Kostarev, Zuev & Viviani 2004). If this were the case, the plume would always have the same shape and orientation with respect to the droplet and substrate, regardless of the direction of gravity. To check whether this is the case, the substrate with a 1-pentanol droplet in place is mounted vertically in the centre of the tank. The resulting flow, shown in figure 13, is found to be mainly parallel to the substrate. Liquid is replenished by inflow from the side and bottom, creating a large convection roll. The fact that the plume orients in a direction opposite to gravity, regardless of the orientation of the substrate, confirms that the convection is buoyancy-driven.

Appendix B. Derivation of the Sherwood number in the diffusion limited case

The Sherwood number has been defined in (4.1). In the diffusion-limited case, the mass loss rate $\dot{m}$ can be calculated from the droplet properties and its size. For a spherical droplet with radius $R$, floating in an infinite bulk, the steady-state mass loss is (Epstein & Plesset 1950)

$$\frac{dm}{dt} = -4\pi RD\Delta c,$$  \hspace{1cm} (B 1)

resulting in a Sherwood number of $Sh_d = 1$.

For a sessile droplet, the presence of a substrate changes the dissolution, and a suitable correction factor has to be used (Popov 2005):

$$\frac{dm}{dt} = -\pi R_p D\Delta cf(\theta),$$  \hspace{1cm} (B 2)

with

$$f(\theta) = \frac{\sin(\theta)}{1 + \cos(\theta)} + 4 \int_0^\infty \frac{1 + \cosh(2\theta \epsilon)}{\sinh(2\pi \epsilon)} \tanh[(\pi - \theta)\epsilon] \, d\epsilon$$  \hspace{1cm} (B 3)
and $R_{fp}$ the footprint diameter of the droplet. Using goniometry, both $R_{fp}$ and the droplet surface area $A$ can be expressed in terms of the volume (and thus the equivalent radius $R = (3V/(2\pi))^{1/3}$) and the contact angle. From this, the Sherwood number for a sessile droplet with contact angle $\theta$, dissolving purely via diffusion, is found to be

$$\text{Sh}_d = f(\theta) \sin \theta \left(1 + \tan^2 \frac{\theta}{2}\right) \left(\frac{2}{2 - 3\cos \theta + \cos^3 \theta}\right)^{1/3}, \quad (B\,4)$$

which indeed depends only on the droplet contact angle. Note that we defined the mass loss rate as a positive quantity, and dropped the minus sign from (B 2), resulting in $\text{Sh}_d > 0$. By solving $f(\theta)$ numerically, $\text{Sh}_d(\theta)$ is plotted as the black line in figure 14. When $\theta = 90^\circ$, the droplet has the shape of a hemisphere, and $\text{Sh}_d$ is equal to that of a free sphere, $\text{Sh}_d = 1$. When $\theta > 90^\circ$, the mass transport is reduced (as compared to that from a free sphere) and hence $\text{Sh}_d < 1$.

When $\theta < 90^\circ$, $\text{Sh}_d$ decreases towards zero, which is an implication of the choice of our characteristic length scale: for practical reasons, the equivalent radius $R$ is chosen as the characteristic length scale. In the extreme case of dissolution from a flat disk, $\theta \to 0$, $V \to 0$, and thus $R \to 0$, resulting in $\text{Sh}_d = 0$. This does not provide a proper physical representation of the actual system, as it would result in zero mass exchange in the case of complete wetting.

Then an alternative characteristic length scale is the footprint radius $R_{fp}$, in which case $\text{Sh}_d$ is given by the red curve in figure 14. By using $R_{fp}$ as the characteristic length scale, $\text{Sh}_d$ for evaporation from a flat disk with radius $R_{fp}$, can be calculated exactly: $f(\theta = 0) = 4/\pi$ (Stauber et al. 2014), which gives $\text{Sh}_d = 4/\pi$. However, the drawback of using $R_{fp}$ as the length scale appears for $\theta > 90^\circ$. Especially when $\theta \to 180^\circ$, $R_{fp} \to 0$, resulting once again in $\text{Sh}_d \to 0$.

So what experimental value for $\text{Sh}_d$ is expected? The Sherwood number scales mass exchange with respect to a diffusive, free and spherical droplet. Hence $\text{Sh}_d = 1$ when
θ = 90°. Owing to the substrate, the mass loss from a surface droplet with θ < 90° is larger as compared to the mass exchange from the same segment of a free and spherical droplet (Hu & Larson 2002). The opposite is true when θ > 90° (Stauber, Wilson & Duffy 2015a). Still, $Sh_d$ is always of order 1 for practical droplets (10° < θ < 160°) and independent of droplet size. The detailed dependence, as proposed in figure 14, could be the subject of future work, where small droplets (ensuring $Ra < 10$) dissolve on substrates of various wettability.

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