Self-Propelled Leidenfrost Droplets

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We report that liquids perform self-propelled motion when they are placed in contact with hot surfaces with asymmetric (ratchetlike) topology. The pumping effect is observed when the liquid is in the Leidenfrost regime (the film-boiling regime), for many liquids and over a wide temperature range. We propose that liquid motion is driven by a viscous force exerted by vapor flow between the solid and the liquid.

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The ability to control the dynamics of liquids is crucial in applications such as lab-on-a-chip technology, ink-jet printing, thermal management, spray cooling, and painting. Millimeter-scale liquid droplets on substrates can move spontaneously due to an imbalance of surface tension forces (the Marangoni effect) [1–3], caused by a chemical [4–10], thermal [4,11,12], or electrical [13] gradient. In these systems, hysteresis forces [3] due to the wetting contact between droplet and surface limit the droplet speed to a few mm/s, unless additional power is supplied [14]. The resulting displacement is usually a few cm or less, because a finite gradient along the direction of motion is needed. Here we report self-propelled Leidenfrost droplets, which are separated from the supporting solid by a lubricating vapor layer (also called the film-boiling regime). Droplet motion is driven by a thermal gradient perpendicular to the direction of motion and is controlled by a sawtooth shaped substrate topology, without the need for a gradient along the flow direction (Ref. [15], movie 1). Millimeter-sized droplets accelerate at rates of up to 1–2 m/s², can climb against inclines, and sustain speeds of order 5 cm/s over distances up to 1 m. We propose that droplet motion is due to a viscous force exerted by vapor flow between the droplet and the asymmetric surface.

Figure 1(a) shows a sequence of video images demonstrating droplet motion perpendicular to the thermal gradient that powers the liquid motion. A droplet of refrigerant R134a (1,1,1,2 tetrafluoroethane, boiling point \( T_b = -26.1 \, ^°C \)) was placed on a room-temperature, brass surface that was milled into a periodic, ratchetlike topology. The droplet is observed to move perpendicular to the ratchet features [in the positive \( x \) direction in Fig. 1(a)], reaching a terminal speed of several cm/s. The motion is sustained over the entire structured surface area, limited only by the evaporation time of the droplet (several tens of seconds). No external force or gradient (other than the vertical temperature difference between solid and liquid) is applied. We have observed this effect for all liquids we have investigated (including nitrogen, acetone, methanol, ethanol, water, and hexadecane, with boiling points ranging from \(-196 \, ^°C \) to \(+151 \, ^°C \)), provided the ratchet temperature is above the liquid’s Leidenfrost point \( T_L \) (about 20 °C for R134a on brass), where a vapor layer levitates the droplet. The effect is observed independent of the ratchet material and its precise geometry in the size ranges we have investigated [1 mm < \( s < 3 \) mm, 0.1 mm < \( d < 0.3 \) mm; see Fig. 1(a)].

Figure 2(a) shows the time evolution of the velocity of R134a droplets [radius \( r = 1.5 \) mm, comparable to the capillary length \( \kappa^{-1} = (\gamma/\rho g)^{1/2} = 1.1 \) mm, where \( \gamma = 15.5 \, \text{mN/m} \) is the liquid’s surface tension at \( T_b \), \( \rho = 1377 \, \text{kg/m}^3 \) is the mass density at \( T_b \), and \( g = 9.81 \, \text{m/s}^2 \) on a horizontally leveled brass ratchet]. Droplets were given an initial velocity \( v_0 \) before entering the ratchet (\( 5 \times 18 \) cm²). The droplet trajectory was recorded using a video camera (frame rate \( 30 \, \text{s} \)), digitized using the software VideoPoint (Lenox, MA), and the velocity’s \( x \)-component \( v_x \) was calculated. Droplets that...
initially move in the negative x direction in Fig. 1 ($v_{0x}<0$) turn around and accelerate until they reach a terminal velocity $v_i$ [see Fig. 2(a) and Ref. [15], movies 3 and 4]. The $v_i(t)$ data fit the equation

$$v_i(t) = \left( v_{0i} - \frac{a}{\beta/m} \right) e^{-t/(\beta/m)} + \frac{a}{\beta/m},$$

(1)

which, for a droplet of mass $m$, is the solution of the equation of motion $m(dv_i/dt) = -\beta v_i(t) + F$, with a velocity-dependent drag force ($-\beta v_i$), a positive, constant accelerating force $F = ma$, and resulting terminal velocity $v_i = a/(\beta/m)$. $v_{0i}$, $a$, and $\beta/m$ are fit parameters.

In Figs. 2(b)–2(d), we show results for droplets of R134a, ethanol ($T_b = 78.5^\circ C$, $\gamma = 18.2$ mN/m, $\rho = 727$ kg/m$^3$, $\kappa^{-1} = 1.6$ mm), and water ($\gamma = 58.9$ mN/m, $\rho = 957$ kg/m$^3$, $\kappa^{-1} = 2.5$ mm), respectively, as a function of superheat $\Delta T = (T_R - T_b)$, where $T_R$ is the bulk temperature of the ratcheted brass substrate. At relatively low superheat (within about 20–50$^\circ$C above the onset of self-propelled motion, regime L), we find the highest accelerations (up to 1–2 m/s$^2$), and the highest values for the drag parameter $\beta/m$. However, the results in regime L vary from droplet to droplet, and the velocity of individual droplets can fluctuate significantly [see inset of Fig. 2(a)]. At higher temperatures (regime H), we observe significantly lower $a$ and $\beta/m$, and much smaller fluctuations.

We speculate that in regime L droplets on a ratchet are not fully supported by a vapor layer, and nucleate boiling events introduce fluctuations. This interpretation is supported by our observation that in regime L the ratchet-cleaning method influences the droplet dynamics [Figs. 2(b) and 2(c)], and by the fact that a liquid’s Leidenfrost point is known to vary substantially with surface roughness and contamination [16,17]. In regime H, however, we find that surface contamination has little influence on $a$ and $\beta/m$, consistent with the liquid and substrate being fully separated by a vapor layer.

The orientation of the ratchet plane relative to gravity is not critical: droplets or slugs placed in a flat-bottomed, open channel (width 2–4 mm) with ratcheted, vertical side walls, accelerate against significant inclines, with the effect becoming stronger with decreasing channel width (Ref. [15], movie 5). This observation suggests that the role of gravity is limited to keeping the liquid in contact with the ratchet.

For ratchet periods $s = 1–2$ mm we do not observe motion for droplets with radius $r < 0.3s$. Droplets spanning multiple $s$ do accelerate (Ref. [15], movies 5 and 6), but puddles tend to break up into smaller droplets once $r$ exceeds several $\kappa^{-1}$. If confined into a channel, we observe acceleration of slugs tens of millimeters in length, interacting with many ratchet periods simultaneously (Ref. [15], movie 5).

To explain our observations, we propose the following model. As liquid evaporates at the bottom surface of the droplet, the pressure that levitates the droplet pushes out the vapor laterally. We propose that the ratchet surface partially rectifies this vapor flow, which exerts a net viscous force on the droplet. In the following, we calculate the magnitude of this force by estimating the pressure gradient underneath the droplet that drives the vapor flow. It is important to note that evaporation and vapor flow are powered by heat from the substrate. The droplets are thus essentially heat engines.

A droplet placed on a ratchet [see Fig. 3(a)] tends to curve concavely around the tops of the ridges (point $A$) while assuming a convex shape elsewhere. This variation in droplet curvature can be used to estimate the variation of the dynamic pressure along the vapor layer as explained in the following. The local difference between the droplet’s internal pressure $p_i$ (assumed constant along the bottom surface) and the pressure in the vapor film is given approximately by the Laplace pressure $\Delta p = \gamma/R$, where $R$ is the local radius of the curvature (assuming no curvature parallel to the ratchet ridges) [3]. A concave surface shape (near point $A$) corresponds to a curvature $R_A < 0$ and $p_A > p_i$, while the convex curvature at points $B_1$ and $B_2$ implies $R_B > 0$ and $p_B < p_i$, such that $p_A > p_B$. We therefore...
expect net vapor flow from point A to points $B_1$ and $B_2$. Flow from A to $B_2$ is expected to create a viscous force in forward direction, which we estimate below. In contrast, vapor flowing from A backwards is expected to escape sideways along the ratchet grooves (see Ref. [15], movie 6). Red (gray) arrows indicate expected thermocapillary flow. (c) A nucleate boiling event near the tape edge in Fig. 1(b) is expected to lead to vapor flow (black arrows) and to thermocapillary flow (red or gray arrows).

The force exerted by the vapor on the liquid between points A and $B_2$ has two components. First, a forward shear force due to Poiseuille vapor flow caused by the pressure differential $\Delta p = (p_A - p_B)$, using nonslip boundary conditions and a parallel-plate model, the horizontal component of this force is [19]

$$ F = 0.5A_{\text{eff}}h|dP/dx| \cos \theta, $$

where $A_{\text{eff}}$ is the total area over which this force contributes (depending on droplet size, multiple ratchet periods are involved), $h$ is the thickness of the vapor layer in this area, and $\theta$ is defined in Fig. 3(b). Second, if the droplet glides with $\nu_x$ relative to the substrate, there is a viscous drag force given by [19]

$$ -\beta \nu_x = - (\eta A_{\text{eff}}/h) \nu_x, $$

where $\eta$ is the vapor’s viscosity.

In Fig. 4(a) we show measured values of $a$ for water droplets as a function of their volume $V$. For droplets that are too small to cover about three full ratchet periods ($r < 1.5 \text{ s}$, where $s = 1.5 \text{ mm}$), there is considerable scatter in the data. We therefore focus on the range $V > 50 \mu l$ ($r > 2.3 \text{ mm}$) when we compare data and model in the following.

To calculate $a$ from Eq. (2), we estimate the fraction $\alpha = A_{\text{eff}}/A_c$ where $A_c(V) = (5/4)h^{1/2}V^{5/6}$ is the droplet’s volume-dependent contact area [20], by measuring the distance $l$ between points of type A and $B_2$ in high-resolution photographs of droplets with $r \approx 1 \text{ s}$ and find $\alpha \approx 0.6$ for water. The value of $h$ is typically 10–100 $\mu$m on flat surfaces [21,22], but varies with position on a ratchet surface [see Fig. 3(a)]. We determine $h$ from the measured values for $\beta/m$ using Eq. (3), neglecting contributions to the drag force from areas outside $A_{\text{eff}} = \alpha A_c(V)$. We find that $h$ varies weakly with droplet size for $V > 50 \mu l$ [see Fig. 4(b)] and use the averaged value in this range, $h = 10.2 \mu$m, for the further analysis. We determine $R_A$ and $R_B$ from high-resolution photographs to find pressure gradients $dP/dx = (p_A - p_B)/l$ of order $10^2 \text{ Pa/mm}$ [23], confirming that the hydrostatic variations in $p_i$ of the order of $\rho g d/s = 3 \text{ Pa/mm}$ are negligible.

The droplet acceleration $a = F/m$ of water droplets as a function of $V$ derived from Eq. (2) is plotted in Fig. 4(a), along with the observed data. The model quantitatively agrees with the data within the uncertainty of the calculation, which is mainly due to error in $\alpha$, $R_A$, $R_B$, and $h$. Note that the expression for $A_c(V)$ used here assumes circular droplets [20], while our droplets are typically
elongated in the $x$ direction. This may account for why the model deviates from the data for larger droplets ($r > 3 \text{ mm}$) which are more likely to elongate. For droplets with $x = r$, the droplet curvature changes with time in a complex manner, which may explain the scatter in the data for $a$ in this range.

The above model is relevant to regime H. We propose that in regime L nucleate boiling occurs near the ratchet ridges, explaining the occasionally observed high values of $B/m$ (hysteresis forces during wetting events introduce additional drag) and of $a$ (due to the additional bursts of vapor flow). Nucleate boiling may also contribute to the acceleration of a liquid nitrogen droplet across the edge of a piece of tape (VWR scientific tape) placed on a brass surface [see Fig. 1(b) and Ref. [15], movie 8]. We observe no acceleration when we use a brass step of equal height, and propose that vapor flow associated with a nucleate boiling event at the tape edge (which is expected to cool much faster than brass, promoting nucleate boiling) exerts a forward force on the droplet [see Fig. 3(c)].

Thermocapillary flow [3] along the droplet’s bottom surface may also be important. For instance, during a nucleate boiling event at a tape edge [Fig. 1(b)], the liquid surface closest to the wetting point is likely to be heated above $T_B$, inducing thermocapillary flow along the $x$ direction, away from the wetting point [Fig. 3(c)]. Because of the broken symmetry at the step, this may result in net pumping action, with the hysteresis forces at the wetting point providing a reaction force. A similar mechanism may be at work in periodic ratchets in regime L. In regime H, where frictional forces between liquid and solid are almost absent, momentum exchange between solid and liquid may nevertheless occur when the ratchet shape forces a change in the direction of thermocapillary flow [at point B in Fig. 3(b)]. We speculate that by relying on thermocapillary flow it may be possible to use a ratchetlike topology to propel droplets at $T_B < T_A$ if a ratchet with superhydrophobic coating is used to mitigate friction.

The ratchet effect [24] reported here may be used to construct pumps consisting of channels with ratchet-shaped inner walls. Such ratchet pumps could be powered by waste heat, making them attractive for use in millimeter-scale closed loop, two-phase cooling systems with no moving parts and no external power need, for example, for microprocessor cooling. It remains to be explored whether the pumping is sustained for ratchet dimensions below the millimeter scale, enabling microfluidic applications.

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