Self-propelled running droplets on solid substrates driven by chemical reactions

Karin John,1∗ Markus Bär,2† and Uwe Thiele1‡

1Max-Planck-Institut für Physik komplexer Systeme, Nöthnitzer Str. 38, D-01187 Dresden, Germany
2Physikalisch-Technische Bundesanstalt, Abbestr. 2–12, D-10587 Berlin, Germany

Abstract

We study chemically driven running droplets on a partially wetting solid substrate by means of coupled evolution equations for the thickness profile of the droplets and the density profile of an adsorbate layer. Two models are introduced corresponding to two qualitatively different types of experiments described in the literature. In both cases an adsorption or desorption reaction underneath the droplets induces a wettability gradient on the substrate and provides the driving force for droplet motion. The difference lies in the behavior of the substrate behind the droplet. In case I the substrate is irreversibly changed whereas in case II it recovers allowing for a periodic droplet movement (as long as the overall system stays far away from equilibrium). Both models allow for a non-saturated and a saturated regime of droplet movement depending on the ratio of the viscous and reactive time scales. In contrast to model I, model II allows for sitting drops at high reaction rate and zero diffusion along the substrate. The transition from running to sitting drops in model II occurs via a super- or subcritical drift-pitchfork bifurcation and may be strongly hysteretic implying a coexistence region of running and sitting drops.

PACS numbers: 68.15.+e, 47.20.Ky, 47.70.Fw 68.43.-h

∗Electronic address: john@pks.mpg.de
†Electronic address: markus.baer@ptb.de
‡Electronic address: thiele@pks.mpg.de, http:www.uwethiele.de
I. INTRODUCTION

The movement of droplets in external gradients fascinates scientists and layman alike at least since Newton’s description [1] of Hauksbee’s experiment with drops of orange oil that move between two non-parallel glass plates towards the point of smallest plate distance [2]. In another example a drop of liquid freely immersed in another liquid subject to a temperature gradient will move towards the higher temperature region due to Marangoni forces caused by surface tension gradients [3]. A drop sitting on a solid substrate also moves in a temperature [4] or wettability [5, 6, 7] gradient. Especially the Marangoni force is already used to manipulate droplets, for example in light induced drop movement [8]. Similar concepts of a directed movement of small amounts of soft matter in a given gradient are also realized in models of cell motility [9].

However, even more intricate are situations where matter spontaneously starts a directed movement in initially homogeneous settings. Small pieces of camphor that move on a liquid surface by emitting a surfactant have also been studied for centuries [10, 11, 12, 13]. More recently oil droplets containing volatile additives and interacting droplets of different volatile oils have been reported to move on solid surfaces due to the solutal Marangoni effect caused by evaporation/condensation [14]. Also intricate is the spontaneous movement of a juxtaposed pair of droplets with different wetting properties, so called bi-slugs, along a capillary tube [15]. This effect was already mentioned by Marangoni [16]. Another example are drops immersed in a second liquid. If the drops contain a soluble surfactant undergoing an isothermal chemical reaction at their surface the drops may start to move [17]. Apart from chemical reactions, drop movement can also be driven by surface phase transitions [18, 19]. The movement is possible because such active drops change their surrounding and produce a gradient that drives their motion.

![FIG. 1: Sketch of a right moving droplet driven by a self-produced wettability gradient.](image)

Recent experiments also found chemically driven running droplets on solid substrates [20, 21].
In these cases small droplets of solution are put on partially wettable substrates. A droplet changes the substrate by adsorption or desorption of a solute rendering the substrate underneath the droplet less wettable than the bare substrate. The radial symmetry still assured an equilibrium position that is, however, unstable. As a result, fluctuations break the symmetry and the drop starts to move in a self-sustained manner. In the course of its movement it changes the substrate and leaves a less wettable trail behind (see Fig. 1). We distinguish two types of experiments: In case I the substrate is changed in an irreversible way whereas in case II it recovers its initial state after the droplet has passed. The droplet does not rest again and follows a random trajectory until trapping itself between self-produced less wettable patches (type I) [21, 23]. Alternatively, a running drop may perform a periodic movement (type II) [24, 25]. The droplet often moves with nearly constant velocity for a rather long time (tens of minutes). However, in all cases the motion ceases when the overall system reaches thermal equilibrium. Next, we shortly recall the specific results of different sets of experiments, yielding running droplets.

In Ref. [21] droplets of \( n \)-alkanes (\( n \)-octane and \( n \)-dodecane) are used that contain 1,1,2,2-tetrahydroperfluorodecyltrichlorosilane (CF\(_3\)(CF\(_2\))\(_7\)(CH\(_2\))\(_2\)SiCl\(_3\)). The silane molecules form dense crafted monolayers on silicon or glass and render these surfaces hydrophobic. The droplet motion is stalled when no further hydrophilic surface is available. No limitation due to silane depletion inside the droplet is observed. For millimeter sized drops (smaller than the capillary length) droplet velocities between 1 mm/s and 10 cm/s are observed, depending on liquid viscosity, silane concentration and droplet size. The velocity increases with the silane concentration and the droplet size. The reaction rate is estimated to be \( 1100 \text{s}^{-1} \text{mol}^{-1} \). Typical reaction times are given as 0.01–0.2 s.

Another experiment is performed in a chemically different system [22, 23] using millimeter-sized droplets of a nonpolar solution of \( n \)-alkylamine (1 mM solution of C\(_6\)NH\(_2\) or C\(_{18}\)NH\(_2\)) in decahydro-naphthalene (\( \gamma = 41 \text{mJ/m}^2 \), \( \mu = 0.001 \text{Ns/m}^2 \)). Therein, silicon substrates with microprinted high-energy surfaces are employed, that expose a dense packing of carboxylic acid functionalities (CO\(_2\)H). The amines dissolved in the droplet adsorb at the substrate and produce a surface of lower energy exposing methyl groups. The effect of different adsorbates [22] and reaction kinetics [23] is investigated. The velocity decreases with increasing droplet size.

The only example of a type II experiment [24] features droplets of oil (5 mM iodine solution of nitrobenzene saturated with potassium iodide) on glass substrates immersed in aqueous solution of a cationic surfactant (1 mM stearyl trimethyl ammonium chloride). In this system the stearyl trimethyl ammonium ion (STA) absorbs at the glass substrate outside the oil droplet and renders
the glass lyophilic. When the oil droplet moves on top of the coating the STA desorbs into the oil. In this way a wettability contrast between front and back of the moving droplet is created and sustained. In contrast to the type I case the substrate recovers its lyophilic state soon after the droplet has passed, because `new’ STA absorbs at the glass. The aqueous phase can be seen as an infinite reservoir of adsorbent. Movement only finally stops when the oil droplet is saturated with STA.

Similar phenomena can be seen in metallurgic systems where droplets of liquid metals or alloys react with the metallic substrate, for instance, by alloying. The layer between the droplet and the substrate may be less wettable than the bare substrate resulting in the migration of reactive islands. This was studied for tin islands on copper surfaces that move and leave tracks of bronze behind [27]. Contrariwise, the layer can be more wettable than the bare substrate. This is typical for the related process of reactive spreading (also called reactive wetting) where a droplet of liquid on a (nearly) non-wettable surface starts to spread after forming a more wettable layer underneath [28, 29, 30, 31, 32, 33]. Variants are possible in which during spreading the substrate becomes less wettable in the center of the drop [34]. However, reactive spreading processes do normally not result in running droplets (but see the ’suddenly displaced’ droplets in Ref. [35]).

For type I experiments an implicit equation for the velocity \( v \) of the droplet was derived [36, 37] from a simple theoretical argument. Based on a balance of friction force and driving capillary forces one obtains
\[
v = C \tan \theta^* \left( 1 - \exp \left( -rL/v \right) \right),
\]
where \( r \) is the reaction rate, \( L \) the size of the droplet and \( C \) a constant. The dynamic contact angles at the advancing and receding ends of the droplet are then assumed to be identical \( \theta^* \), i.e. the droplet profile is approximated by a spherical cap with \( \theta^* \) given by
\[
\cos \theta^* = \left( \cos \theta^a_e + \cos \theta^r_e \right) / 2.
\]
The static contact angles at the advancing edge \( \theta^a_e \) and at the receding edge \( \theta^r_e > \theta^a_e \) are different due to the chemical gradient. The expression for the velocity is found from a first order reaction on the substrate that yields chemical concentrations \( \alpha_a = 0 \) and \( \alpha_r = 1 - \exp \left( -rL/v \right) \) at the respective ends of the droplet. The expression for the velocity predicts a monotone increase of the droplet velocity with the droplet length \( L \) and the reaction rate \( r \) in line with experimental observations [21]. The droplet velocity in the limiting case of a saturated chemical reaction was also given in [38]. However, the experiments of Ref. [23] show the opposite trend; the velocity decreases with increasing drop sizes and effective reaction rate. In the framework of Ref. [36] the decrease is explained as a result of the flattening of the drops by gravity. Related works discuss running droplets in a random medium [39] and the forced wetting of a plate immersed into a reactive fluid [40].
In this paper we propose and analyze dynamical models for self-propelled running droplets for both, type I and type II experiments. Our models consist of coupled evolution equations for the film thickness profile and the substrate coverage. Thereby the wettability of the substrate is modeled by a coverage dependent disjoining pressure. The different types of experiments are taken into account by incorporating different reaction terms. The models are capable of reproducing the different experimentally found regimes. A short account of a variant of the model for the type I experiments, i.e. for the case of a irreversibly changed substrate was recently presented [41]. There it was found that the dynamic contact angles at the advancing and receding ends of the droplet are not identical but rather resemble the corresponding static ones. Even for a strong driving force the deviation between dynamic and static angles is only about 10 percent. The bifurcation leading from sitting to running droplets for a finite reaction rate was identified as a drift-pitchfork bifurcation. Here we analyze the type I model in more detail and extend our analysis to the type II model.

In section II the type I and type II dynamical models are presented. Results for the two models are discussed in section III and IV respectively. Stationary running droplets and sitting droplets are characterized in dependence of the control parameters reaction rate, droplet volume, diffusion constant of the adsorbate field and desorption rate of the adsorbate (for type II model only). The resulting families of solutions and their linear stability are used to derive phase diagrams that describe the existence regions for running and sitting droplets. To further elucidate the drift-pitchfork bifurcation that mediates the transition from sitting to running droplets also in the type II model we analyze the linear modes that destabilize sitting droplets close to the bifurcation. Before we conclude in section V we use numerical simulations to illustrate and analyze periodic droplet motion that is possible in the type II model. The simulations are compared with our continuation results and the type II experiment of Ref. [24].

II. MODEL

A. Evolution equations

The evolution of a thin liquid film on a horizontal smooth solid substrate is described by an equation for the thickness profile $h(x, y, t)$ derived from the Navier-Stokes equations using long-
wave or lubrication approximation [42]

\[
\frac{\partial t}{h} = -\nabla \cdot \left\{ \frac{h^3}{3 \eta} \nabla p \right\} .
\] (1)

The parameters \( \gamma \) and \( \eta \) are the surface tension and viscosity of the liquid, respectively. They define the viscous time scale \( \tau_v = \gamma L / \eta \), where \( L \) is a typical length for the system. The change in time of the film thickness profile equals the gradient of a flow that results as the product of a mobility and a pressure gradient. The mobility \( h^3 / 3 \eta \) corresponds to a parabolic velocity profile

\[
u(x, z) = \left( \frac{z^2}{2} - zh \right) \nabla p
\] (2)

within the film. The velocity component orthogonal to the substrate \( w \) can be obtained using the continuity equation

\[
\partial_x u + \partial_z w = 0 .
\] (3)

The pressure

\[
p = \gamma \Delta h + \Pi(h, \phi)
\] (4)

contains the curvature (or Laplace) pressure \(-\gamma \Delta h\) and the disjoining pressure \( \Pi(h, \phi) \). The latter comprises effective molecular interactions between the film surface and the substrate and accounts for the wetting properties of the substrate [43, 44, 45]. As discussed in detail below, here a mathematically simple function \( \Pi(h, \phi) \) is used that is common in the literature. In many situations the qualitative outcome only depends on very general characteristics of the disjoining pressure [46]. The used form allows for solutions of Eq. (1) that represent static (i.e. sitting) droplets with a finite mesoscopic equilibrium contact angle. The disjoining pressure is chosen such that the droplets coexist with an ultrathin precursor film.

The evolution of the density of the adsorbed layer on the substrate determines the wettability and is modeled by a reaction-diffusion equation for the dimensionless field \( 0 < \phi(x, y, t) < 1 \)

\[
\partial_t \phi = R(h, \phi) + d' \Delta \phi,
\] (5)

where the function \( R(h, \alpha) \) describes adsorption or desorption on the substrate. The second term allows for a diffusion of the chemical species along the substrate. For simplicity we assume that the adsorbate has the same diffusion constant on the bare and the droplet covered substrate. However, assuming different constants would not change the presented results qualitatively. Note that for type I experiments \( \phi \) directly corresponds to the substrate coverage. However, in a type II
experiment the droplet dissolves a more wettable coating. In this case the coverage corresponds to \(1 - \phi\). Also here one can use Eq. (5), however, the signs of adsorption and desorption term are switched. With this convention, in both cases the wettability decreases with increasing \(\phi\).

One can neglect the dynamics of the concentration field of the chemical in the bulk of the droplet by assuming a fast equilibration of the solute concentration within the moving droplet as compared to the reaction at the substrate. The fast equilibration is caused by diffusion and convective motion within the droplet whereas the latter is driven by the lateral movement of the droplet along the substrate. It corresponds to the limit of a small Damköhler number \(\text{Da} = r c_0 / (D/L)\) giving the ratio of reaction velocity at the substrate and diffusion velocity in the droplet \([47]\). The parameter \(r\) is a typical reaction rate, \(c_0\) stands for a typical concentration of the chemical species in the droplet and \(D\) is the diffusion constant in the droplet.

**B. Reaction term**

Corresponding to the two different sets of experiments that the model shall describe we use two different reaction terms. For type I the initial coverage of the substrate is zero (i.e. \(\phi = 0\)) and we only allow for adsorption underneath the droplet

\[
R_1(h, \phi) = r_{\text{in}} \xi(h) (1 - \phi).
\]  

(6)

The reaction saturates at \(\phi = 1\) because this is the maximal possible coverage. The function \(\xi(h)\) represents a (smooth) step function that approaches one and zero inside and outside the droplet, respectively. For type II the ‘rest’ state of the substrate without any droplet is the fully covered substrate (again corresponding to \(\phi = 0\), see above section II A) and we allow for desorption underneath the droplet and adsorption outside the droplet

\[
R_2(h, \phi) = r_{\text{in}} \xi(h) (1 - \phi) - r_{\text{out}} [1 - \xi(h)] \phi.
\]  

(7)

The time scales of the reactions at the substrate inside and outside the droplet are defined by the effective rate constants \(r_{\text{in}}\) and \(r_{\text{out}}\), respectively. Note that they have the dimension s\(^{-1}\). The function \(\xi(h)\) may be the step function

\[
\xi_1 = \Theta(h - h_c),
\]  

(8)

or the smooth function

\[
\xi_2 = \{\tanh[(h - h_c)/\Delta] + 1\}/2.
\]  

(9)
The value of $h_c$ is chosen slightly larger than the thickness of the precursor film and the maximal drop height is always $h_{\text{max}} \gg h_c$. A small value of $\Delta \ll h_{\text{max}}$ ensures that the switch between a predominant adsorption reaction and a predominant desorption reaction occurs over a small film thickness range. Note, that $\xi_2 \rightarrow \xi_1$ for $\Delta \rightarrow 0$. Changes in the details of the reaction term do not affect the results qualitatively.

C. Disjoining pressure

For the disjoining pressure we use throughout the present work

$$
\Pi(h) = \frac{2Sld_0^2}{h^3} + \left(1 - \frac{\phi}{g}\right)\frac{5Ssd_0^5}{h^6} \tag{10}
$$

where $d_0 = 0.158$ nm is the Born repulsion length that defines a lower cut-off for the film thickness. The parameter $S_l$ and $S_s$ are the long- and short-range components of the total spreading coefficient $S = S_l + S_s$ (for $\phi = 0$). We use $S_l < 0$ and $S_s > 0$ corresponding to a destabilizing long-range van der Waals and a stabilizing short-range interaction [48]. For $\phi = 0$ the pressure allows for drops sitting on a stable precursor film as can be seen studying the corresponding densities of the excess surface energy $f$, related to the disjoining pressures by $\Pi = -\partial_h f$.

The short-range interaction contains the influence of the coating and crucially influences the static contact angle [49]. To account for the varying wettability caused by the different substrate coverage we let the short-range part of the spreading coefficient $S_s$ depend linearly on the field $\phi(x, y, t)$. The signs are chosen in a way that for both types of experiments $g > 0$ assures that an increase in $\phi$ corresponds to a lower wettability, i.e. to a larger equilibrium contact angle $\theta_e$ given by $\cos \theta_e = S/\gamma + 1$ [49]. The constant $g$ relates the coverage to the wettability and therefore defines the magnitude of the possible wettability gradient. Note, that our Eq. (10) corresponds to the linear relation between $\cos \theta_e$ and $\phi$ assumed in Ref. [21, 23, 36].

D. Dimensionless equations

We rewrite Eqs. (1) to (10) by introducing scales $\sqrt{3\gamma\eta/\kappa^2}$, $\sqrt{\gamma l/\kappa}$, and $l$ for $t$, $(x, y)$, and $h$, respectively. Then one obtains $2|S_l|d_0^2/l^3$ for the scaled spreading coefficient $\kappa$. As length scale $l$ we use the value of the film thickness where the local free energy $f(h)$ has its minimum, i.e., where $\Pi(h) = 0$. This gives $l = (5S_s/2|S_l|)^{1/3}d_0$ and implies that the ratio of the strength of
the two antagonistic effective molecular interactions is only an implicit parameter of the system. The length \( l \) also corresponds to the thickness of the precursor film for a sitting drop on the bare substrate.

Defining the dimensionless overall reaction rate \( r = 3 r_{in} \gamma \eta / l \kappa^2 \), diffusion constant \( d = 3 d' \eta / \kappa l^2 \), and ratio of reaction rates \( s = r_{in} / r_{out} \), we obtain from Eqs. (1) to (10) the dimensionless coupled evolution equations for the thickness profile \( h \) and the field \( \phi \):

\[
\partial_t h = -\nabla \left\{ h^3 \nabla \left[ \Delta h + \Pi(h, \phi) \right] \right\} \\
\partial_t \phi = r R(h, \phi) + d \Delta \phi
\]

(11) \hspace{1cm} (12)

with the disjoining pressure

\[
\Pi(h, \phi) = -\frac{1}{h^3} + \left(1 - \frac{\phi}{g}\right) \frac{1}{h^6}
\]

(13)

and the options for the reaction term

\[
R_1(h, \phi) = \xi(h) \left(1 - \phi\right) \\
R_2(h, \phi) = \xi(h) \left(1 - \phi\right) - s \left[1 - \xi(h)\right] \phi
\]

(14) \hspace{1cm} (15)

To give an impression of the influence of the coverage we give in Fig. 2 droplet profiles for different values of the coverage \( \phi \). Thereby, \( \phi \) is assumed to be independent of time and constant along the substrate. Increasing the ratio \( \phi/g \) clearly leads to increasing contact angles and decreasing droplet length.

In the following we explore the full non-linear behavior of the dynamical models presented above. The main interest lies in an exploration of the existence regions of qualitatively different solutions depending on control parameters like reaction rate \( r \), ratio of desorption and adsorption rates \( s \), and droplet volume \( V \). It was already shown that a type I model allows for three-dimensional running droplets that move with constant velocity \( v \) and droplet shape [41]. Here, we restrict our attention to two-dimensional drops to be able to explore a large part of the parameter space. To determine such droplets we therefore use Eqs. (11,12) replacing \( \nabla \) by \( \partial_x \). Then continuation techniques [50, 51, 52] are employed to calculate two-dimensional running droplets moving with constant speed. This is achieved by switching to the comoving frame \( x - vt \) and imposing appropriate boundary conditions. In the comoving frame running droplets correspond to steady solutions. Integration in time of the full system is also used in some cases. Details on all numerical techniques used can be found in the Appendix.
FIG. 2: Profiles of sitting droplets without reaction for different constant values of the wettability measure \( \phi/g \) as given in the legend. The coverage is assumed to be independent of time and constant along the substrate. The system size is \( L = 10000 \) and all droplets have the volume \( V \approx 3 \times 10^4 \).

III. RESULTS FOR MODEL I (ONLY ADSORPTION)

We start with model type I, where adsorption takes place only underneath the droplet, i.e. we combine the evolution equations for the film thickness profile (11) and the field of adsorbate coverage (12) with the reaction term (14) and the disjoining pressure (13).

A. Thickness and coverage profiles

First we focus on the characterization of the changing solution behavior in dependence of the overall reaction rate \( r \). Without diffusion along the substrate \( (d = 0) \) one finds unstable sitting droplets and stable running droplets for all \( r \). The droplets and the coverage profile move with constant velocity \( v \) and constant shape. We emphasize that this state corresponds to a subtle dynamical equilibrium given that coverage profile and droplet velocity depend on each other.

Fig. 3 shows for two different reaction rates profiles of moving droplets [(a) and (b)] where the streamlines correspond to contour lines of the stream function

\[
\psi(x, z) = \left( \frac{z^3}{2} - \frac{3z^2h}{2} \right) \partial_x p - vz,
\]  

(16)
and indicate the flow in the comoving coordinate system. Note that the velocity fields are obtained by \( (u, w) = (\partial_z \psi, -\partial_x \psi) \). Also shown are the corresponding profiles for the coverage \( \phi \) [(c) and (d)].

The two sets of profiles belong to two qualitatively different regimes that are prominently visible in the profiles of the coverage (corresponding to the results obtained for a different disjoining pressure in [41]).

\[ \begin{align*}
\text{FIG. 3: Profiles of running droplets in the comoving frame. Shown are (a,b) the droplet profiles } h \text{ and (c,d) the substrate coverage } \phi \text{ for two different reaction rates } r. \text{ Droplets on the left and right correspond to the non-saturated } (r = 10^{-5}) \text{ and saturated } (r = 0.001) \text{ regime and move with velocities } v \approx 0.006 \\
\text{and } v \approx 0.016, \text{ respectively. The streamlines plotted in (a) and (b) indicate the convective motion inside the droplets in the comoving frame and have the spacing } \Delta \psi = 0.025 \text{ and } 0.1, \text{ respectively. The direction of the movement is indicated by arrows. The remaining parameters are } g = 2, d = 0.001, L = 10000, \\
h = 3.8, h_c = 2.0, \Delta h = 0.2 \text{ and the droplet volume is } V \approx 3 \cdot 10^4.\end{align*} \]

B. Dependence on reaction rate

For low reaction rates [Fig 3(a) and (c)] the coverage starts to increase at the advancing contact zone and continues to increase up to the receding contact zone where it is still well below the saturation value of \( \phi = 1 \). We call this the non-saturated regime. For high reaction rates [Fig 3(b) and
the coverage starts to increase at the advancing contact zone as in the non-saturated regime. However, it increases much faster and reaches the saturation value $\phi = 1$ already underneath the droplet. At the receding contact zone it is always at the saturation value. We call this the saturated regime. The droplets in the two regimes behave qualitatively different when the reaction rate is changed as shown in Fig. 4.

![FIG. 4: Given is a characterization of stationary running droplets in dependence of the reaction rate $r$ (logarithmic scale) for different strength of the diffusion of the coverage $\phi$ along the substrate $d$. Shown are (a) the velocity $v$ for different diffusion constants $d$ (see legend); and (b) the advancing ($\theta_a$, dashed lines) and the receding ($\theta_r$, solid lines) dynamic contact angles for $d = 0.001$ (heavy lines) and $d = 0.1$ (thin lines). The arrows with the labels $r_{max}$, $r_{dp}$ and $r_{sn}$ in (a) indicate the locations for the maximum velocity, drift-pitchfork bifurcation and the saddle-node bifurcation, respectively, for $d = 0.1$. The remaining parameters are as in Fig. 3.]

For low reaction rates in the non-saturated regime the coverage does not reach the saturation value $\phi = 1$ at the rear of the droplet. This implies that the driving wettability gradient between front and rear of the droplet has not yet reached its maximally possible value. Therefore an increase in the reaction rate leads to a steeper increase in the spatial profile of the coverage underneath the droplet. In consequence a larger wettability gradient results implying a larger velocity. However, again it is subtle to determine the dynamic equilibrium. First, the higher velocity reduces the
contact time of the droplet with a given point of the substrate. Second, it also reduces the length of the droplet (see below). Both effects restrict the growth of the wettability gradient. The resulting increase of the velocity with increasing reaction rate can be seen in Fig. 4(a) for $r < 0.0001$. Note that it does only weakly depend on the diffusion along the substrate.

Fig. 4(b) shows the corresponding dynamical contact angles at the advancing and receding edges of the moving droplets. Below $r \approx 0.0001$ both angles increase, implying a decrease in the length of the droplet (that has constant volume). Because in the non-saturated regime the coverage in the contact zone at the advancing edge does effectively not depend on $r$, the increase in the advancing contact angle is caused by the larger velocity only. Seeing the full dynamic contact angle at a moving contact line as a superposition of a static (or equilibrium) part and a dynamic part, one can say that only the dynamic part of the advancing contact angle increases with $r$. The increase at the front is in line with the increase with velocity found for sliding drops on an incline (see for example Ref. [53]). However, at the receding edge the coverage in the contact zone depends strongly on $r$. Therefore the rather strong increase in the receding contact angle with increasing $r$ is caused by changes in both, the static and the dynamic part. For a larger $r$ the wettability at the back is smaller, i.e. the 'local equilibrium contact angle' increases. However, the dynamic receding contact angle normally decreases with increasing velocity [53]. This implies that the increase in the receding angle found here results from an increase of the static part only. It is even counteracted by a decrease of the dynamic part. Also in the contact angles, a relatively small $d$ has no visible influence in the non-saturated regime.

The maximum wettability gradient, i.e. driving force, and therefore the maximum droplet velocity is reached for the bare substrate ($\phi = 0$) at the advancing edge and maximum coverage ($\phi = 1$) at the receding edge. The point of largest velocity [Fig. 4(a)] corresponds to the point of the largest difference between the receding and advancing contact angles [Fig. 4(b)]. We call the corresponding reaction rate $r_{max}$.

Next we discuss the behavior in the saturated regime ($r > 0.0001$ in Fig. 4). For larger reaction rates the coverage at the rear of the droplet remains at its saturation value. However, the driving force and in consequence the velocity decrease slightly with increasing $r$ (cf. also Ref. [41]). Note, that for small $d$ in Fig. 4(a) this is barely visible. The decrease from $r_{max}$ to $r = 0.1$ corresponds to about 2% of the maximal velocity.

As detailed next this slight decay is caused by the dynamics in the advancing contact zone. In the saturated regime the time scale of the reaction is short compared to the one of the droplet
movement. Therefore the coverage increases very steeply underneath the droplet leading to an
elevated coverage already in the advancing contact zone (see Fig. 3b). The resulting decrease of
the overall wettability gradient felt by the droplet causes the slight decrease of the velocity. In this
regime the slight increase of the advancing contact angle results from an increase of the static part
(because of the decreasing wettability) that overcomes a decrease of the dynamic part (decreasing
velocity). The slight increase of the receding angle comes from the dynamic part only.

We emphasize here that Fig. 4(b) clearly shows that the advancing contact angles are always
smaller than the receding ones. As was already shown in Ref. [41] also here the differences be-
tween the static and the dynamic contact angles at the front and the rear, respectively, are an order
of magnitude smaller than the difference between the two static (or the two dynamic) contact
angles.

C. Phase diagrams

1. Influence of Diffusion

Let us finally discuss the influence of diffusion along the substrate on the dependencies shown
in Fig. 4. The result without diffusion (not shown) coincides nearly perfectly with the shown
curves for $d = 0.001$. Only the decay in the saturated regime is slightly slower. We mention here
that the decay also depends slightly on the used cutoff $h_c$ (cf. Eq. (8) and Ref. [41]). Increasing the
diffusion has a rather small influence on the non-saturated regime but changes the saturated regime
even qualitatively. The velocity decreases with the reaction rate because the adsorbate produced
underneath the droplet close to the advancing edge diffuses onto the bare substrate ahead of the
moving droplet. This effectively reduces the wettability there, implying a slower movement. In
fact, running droplets cease to exist above a reaction rate $r_{dp}$ where the velocity drops to zero
again, corresponding to a supercritical bifurcation for large diffusion (see curve for $d = 1.0$).
The bifurcation structure can, however, be more involved. For a smaller $d = 0.1$ one finds that
the branch of running droplets joins the branch of sitting drops in a subcritical bifurcation at
$r_{dp}$, i.e. decreasing $v$ the running droplet branch first turns back in a saddle-node bifurcation at
$r_{sn}$ and then joins the sitting droplet branch at $r_{dp}$. We indicate the definition of $r_{sn}$ and $r_{dp}$ in
Fig. 4(a) using the curve for $d = 0.1$ as example. However, the subcritical part of the branch is
difficult to see. We refer the reader to the discussion of model II in section IV for more details.
FIG. 5: Existence of running and stationary droplets depending on the reaction rate $r$ and the diffusion constant $d$. The solid line indicates a drift-pitchfork bifurcation where a running droplet solution branches off a stationary droplet solution. The dotted line indicates a saddle-node bifurcation where two running droplet solutions merge. Shown is also the transition between non-saturated and saturated regime (dashed line). The remaining parameters are as in Fig.3.

Between $r_{dp}$ and $r_{sn}$ both the large velocity running droplet and the sitting droplet are linearly stable and correspond to metastable states. The branch emerging from the subcritical bifurcation is linearly unstable and corresponds to threshold solutions that separate the two stable solutions. The prominent features ($r_{max}$, $r_{dp}$ and $r_{sn}$) of the solution branches shown in Fig.4 can be used to determine phase diagrams by continuation techniques [52]. The phase diagrams show existence regions for the different types of solutions in the parameter space spanned, for instance, by the reaction rate and the diffusion constant (see Fig.5). To this end we followed the maximum of the $v(r)$ dependence at $r_{max}$, which marks the transition between the non-saturated and saturated regime for running droplets (dashed line in Fig.5). Continuation of the loci of the saddle-node bifurcation at $r_{sn}$ and the drift-pitchfork bifurcation at $r_{dp}$ gives the border of the existence region for stable running droplets (dotted line) and stable sitting droplets (solid line), respectively. The region between the two latter borders corresponds to a coexistence region where running and sitting droplets are metastable. Note, that unstable sitting drops do also exist everywhere in the
existence region of the running droplets. Such sitting droplets are steady solutions of the governing equations. However, because they are unstable, even infinitely small perturbations will grow with a characteristic rate. In consequence the droplets start to move and adopt the shape and speed of the stable running droplet solution.

2. **Influence of volume**

Practically, it is difficult to change the reaction rate over orders of magnitude. Experiments normally only cover a much smaller range \([21, 23]\). Nevertheless, the analysis of the \(v(r)\) dependence (Fig. 4) for droplets of fixed volume gives a very good characterization of the general system behavior. An experimentally important parameter is the size or volume of the droplet. The velocity may increase \([21]\) or decrease \([23]\) with increasing volume depending on the parameter regime \([41]\). We show in Fig. 6 the dependency of velocity on droplet volume (logarithmic scale) for a variety of reaction rates. Also there one can well distinguish a saturated and a non-saturated regime. In the non-saturated regime the droplet velocity increases with increasing size, whereas in the saturated regime it decreases. For small droplets one always finds a non-saturated regime whereas droplets of a very large size are always in a saturated regime (in Fig. 6 the latter is not yet reached for the curve with \(r = 10^{-5}\)). The explanation for this behavior is closely related to the one given for the \(v(r)\) dependence. For small droplets the reaction does not reach saturation until the rear of the droplet has passed. This implies that an increase in droplet size gives more time for the reaction leading to a larger value of \(\phi\) at the back and, in consequence, to a larger velocity.

For very large droplets the reaction has enough time to reach saturation \((\phi = 1)\) at the rear, i.e. an increase in size does not increase the driving wettability gradient. It may even decrease slightly due to an increase of \(\phi\) in the contact zone at the advancing edge (see above). However, a larger droplet has a larger viscous dissipation leading to a decreasing velocity with increasing size. Inspecting Fig. 6 shows that for \(r \geq 5 \cdot 10^{-4}\) and \(V = 30\,000\) (i.e. the volume used in Fig. 4) one is well in the saturated regime.

The volume \(V_{\text{max}}\) where the maximal velocity is obtained can be followed in the space spanned by droplet volume and reaction rate. The obtained existence region for non-saturated and saturated running droplets are shown in Fig. 7.

Having studied the type I model describing experiments where the passing droplet irreversibly changes the substrate we next report on results for our type II model where the substrate recovers.
FIG. 6: The transition between non-saturated and saturated regime is most clearly seen in the dependence of the velocity $v$ on drop size, i.e. droplet volume $V$ (logarithmic scale). Curves are plotted for different reaction rates $r$ (see legend). Remaining parameters are as in Fig. 3.

In contrast to type I this allows for the description of a periodic droplet movement as experimentally observed in Ref. [24].

IV. RESULTS FOR MODEL II (ADSORPTION AND DESORPTION)

This section is devoted to the type II model that extends and generalizes the model of type I discussed above. Model II accounts for experimental situations where both, the droplet and its surrounding medium are able to change the substrate by adsorption or desorption. In this way a moving droplet makes the substrate less wettable, but after the droplet has passed the substrate may relax to its initial state. Such systems are modeled by extending the reaction kinetics for the $\phi$-field (14) by an additional desorption term, i.e. we combine the evolution equations for the film thickness profile (11) and the adsorbate field (12) with the reaction term (15) and the disjoining pressure (13). We will refer to this set of equations as type II model. The reaction term (15) is chosen such that adsorption and desorption of $\phi$ take predominantly place underneath and outside the droplet, respectively.
FIG. 7: Existence of running and stationary droplets depending on the reaction rate $r$ and the droplet volume $V$ (in units of $10^4$) for two diffusion constants in the $\phi$-field $d = 0.001$ (a) and $d = 0.1$ (b). The solid and dotted lines indicate a drift-pitchfork and a saddle-node bifurcation, respectively. The dashed line indicates the transition between non-saturated and saturated regime. Remaining parameters are as in Fig. 3.

A. Thickness and coverage profiles

As above for model I, we use continuation techniques to calculate running droplet solutions of model II as solutions stationary in a comoving frame. Examples of the resulting droplet profiles along with the profiles of the $\phi$-field are shown for a very small diffusion $d$ in Figs. 8 and 9. Thereby, Fig. 8 displays the results for several values of the reaction rate $r$ for a fixed ratio $s$ of the desorption to adsorption rates. The drop profiles change only slightly but the $\phi$-profile undergoes prominent changes. As in model I there is practically no $\phi$-field in front of the droplet ($\phi = 0$). The field increases (i.e. the wettability decreases) underneath the droplet as in model I, but in contrast the $\phi$-field decreases behind the droplet due to the desorption. The concentration of the $\phi$-field at the rear of the droplet is increasing with $r$ until it reaches saturation. Because $r$ is the overall reaction rate also the desorption of $\phi$ behind the droplet becomes faster, i.e. the tail of the $\phi$-field becomes shorter.

Fig. 9 displays droplet and $\phi$ profiles for a fixed reaction rate $r$ but different values for the desorption to adsorption ratio $s$. The droplet shape and the form of the $\phi$-field underneath the droplet are almost independent of $s$ as one would expect. However the decay of the elevated $\phi$ value occurring behind the droplet is strongly influenced. For a small $s$, i.e. slow desorption
FIG. 8: Results of model II incorporating adsorption and desorption. Shown is the influence of the reaction rate $r$ (see legend) for (a) the profiles of moving droplets and (b) the corresponding profiles in the $\phi$-field. The drop profiles at different $r$ differ only slightly by visual inspection. The differences are more prominent in the $\phi$-profiles. Parameters are $s = 1$, $h_c = 2$, $\Delta h = 0.2$, $g = 2$, $d = 0.001$, $V \approx 30000$ ($L = 10000$, $\bar{h} = 3.8$).

As compared to adsorption, $\phi$ decays slowly approaching qualitatively the behavior of model I. Increasing $s$ reduces drastically the length of the $\phi$ tail behind the droplet. The length of the tail is very important when studying the periodic movement of a droplet on a finite stripe-like substrate (see below).

B. Phase diagrams

In the following the existence regions of running and sitting droplets are determined in their dependence on the control parameters reaction rate, desorption to adsorption ratio, diffusion constant and droplet volume. This is done along the lines explained in section III. Continuation gives branches of stationary solutions depending on one control parameter. On these branches special points that separate qualitatively different behavior are identified and followed in the space of parameters. The linear stability of the stationary solutions is also determined. Details on the used
FIG. 9: Shown is the influence of the ratio \( s \) of desorption and reaction (see legend) on (a) the profiles of moving droplets and (b) the corresponding profiles in the \( \phi \)-field for model II. The drop profiles at different \( s \) differ only slightly by visual inspection. The differences are prominent in the tail of the \( \phi \)-profile behind the droplet. \( r = 10^{-4} \) and the remaining parameters are as in Fig. 8.

Techniques can be found in the Appendix.

1. Influence of the desorption/adsorption ratio

Studying first the influence of \( r \) and \( s \) we present in Fig. 11 (a) and (b) branches of stationary solutions in dependence of \( r \) for different values of \( s \) characterized by their velocity and the resulting existence regions of running and sitting droplets in the \( r - s \) parameter plane, respectively. Beside the moving droplets there exist sitting droplets (steady states) for all values of \( r \) and \( s \). They are symmetrical with respect to the droplet maximum and may be stable or unstable. An example is shown in Fig. 10.

The curve for \( s = 1 \) in Fig. 11 (a) is used to better illustrate special values of the reaction rate introduced above in section 3. They mark the loci of the maximum of the \( v(r) \) dependence at \( r_{\text{max}} \), of the saddle-node bifurcation at \( r_{\text{sn}} \) and of the drift-pitchfork bifurcation at \( r_{\text{dp}} \). As before, continuation in \( r \) of these loci when changing \( s \) gives the phase diagram Fig. 11 (b). In particular one obtains the border between the non-saturated and saturated regime for running droplets.
FIG. 10: Shown are profiles of (a) the film thickness and (b) the φ-field for running and sitting droplets (see legend) for model II. The drop profiles differ only slightly by visual inspection. The differences are more prominent in the φ-profiles. $r = 10^{-4}$ and the remaining parameters are as in Fig. 8.

($r_{max}$), the border of the existence region for stable running droplets ($r_{sn}$), and the border of the existence region for stable sitting droplets ($r_{dp}$). In the small region where both, running and sitting droplets are stable, one finds metastability. There noise, for instance, in the form of substrate inhomogeneities may lead to intermittent droplet movement. The sitting droplets existing in the gray shaded area of Fig. 11(b) are all unstable.

Inspection of Fig. 11 reveals the rather strong influence of the desorption/adsorption ratio $s$ especially on the transition from the non-saturated to the saturated regime and the slowing down in the saturated regime. The extension in $r$ of the saturated regime is drastically reduced with increasing $s$. For decreasing $s$ the maximum of the $v(r)$ curve slowly transforms into a plateau and reaches for $s = 0$ the form typical for model I. Comparing the strong decrease of the velocity with increasing $r$ in the saturated regime [Fig. 11(a)] to the slight decrease found for $s = 0$ (model I, Fig. 4(a), curve for $d = 0.001$) poses the question why $s$ has such a strong influence.

The influence of $s$ parallels the influence of the diffusion $d$ in model I. There for large $d$ the adsorbate diffuses in front of the advancing droplet rendering the substrate there less wettable. Thereby it decreases the overall wettability gradient between front and rear of the droplet. Here,
FIG. 11: (a) Velocity of running droplets depending on the reaction rate $r$. The branch of running droplets emerges from zero reaction rate and undergoes a subcritical drift pitchfork bifurcation at finite $r > 0$. The reaction rate-velocity curve is shown for three different values of the ratio between desorption and adsorption $s$. The labels $r_{\text{max}}$, $r_{\text{dp}}$ and $r_{\text{sn}}$ indicate the locations for the maximum velocity, drift-pitchfork bifurcation and the saddle-node bifurcation, respectively, for $s = 1.0$. (b) Existence of running and sitting droplets depending on the reaction rate $r$ and the ratio between desorption and adsorption $s$. Shown is also the boundary between non-saturated and saturated regime (dashed line). The remaining parameters are as in Fig. 8.
for large $s$ the $\phi$-field is removed behind the rear of the droplet rendering the substrate there more wettable. That also implies a reduction of the overall wettability gradient.

2. The drift-pitchfork bifurcation

To elucidate the mechanism of the transition between moving and sitting drops we focus for a moment on the drift-pitchfork bifurcation at $r_{dp}$ that separates the metastable and the running droplet region. This type of bifurcation is well known from reaction-diffusion (see, for instance, [54, 55, 56, 57, 58] and references therein) and hydrodynamic systems (see, for instance, [59, 60, 61, 62] and references therein), where it mediates the transition between steady and travelling structures. In our system it breaks the reflectional symmetry of the sitting droplets leading to moving asymmetric droplets and accompanying travelling asymmetric adsorbate profiles. At the bifurcation a real eigenvalue switches sign, i.e. at $r_{dp}$ the velocity of the moving drops is zero. Beyond the bifurcation sitting droplets become unstable and start to move slowly. That makes it unlike a Hopf bifurcation (associated with the zero crossing of the real part of a pair of complex eigenvalues) where the travelling structure has a finite velocity at the bifurcation (for waves on flowing thin liquid films see the discussion in Ref. [63]). The bifurcation may be subcritical or supercritical (see Figs. 4-11 and 12) and in consequence the bifurcating branch corresponds to unstable or stable moving drops. Approaching the bifurcation their respective velocity goes to zero as $\sqrt{|r - r_{dp}|}$ providing an unambiguous signature of the drift-pitchfork bifurcation.

The basic mechanism of the drift-pitchfork bifurcation is connected to the behaviour of the neutral (or Goldstone) mode related to the translational symmetry of the system. This mode with eigenvalue zero is obtained by analyzing the linear stability of the stationary solutions. In general, each continuous symmetry is related to such a neutral mode. In a sense, the neutral modes are the modes that are 'closest' to zero. This implies that a perturbation or modulation of these modes in an additional degree of freedom (if existing) gives modes that are probable candidates to cross zero and become instability modes. For instance, the transversal (fingering) instability of a liquid front results from a transversal modulation of the longitudinal translational neutral mode [64]. Also one of the two coarsening modes of two liquid droplets is the combination of translational neutral modes of the individual droplets directed in opposite directions [65].

Here, the drift instability is associated with a mode representing a relative shift between the translational modes of the height and the $\phi$ profiles. Right at the bifurcation this mode corresponds
exactly to the translational neutral mode. Although elsewhere one can still identify the translational
mode when only looking at the sub-mode for the height profile or the one for the $\phi$ profile, looking
at the complete mode one realizes that the relative weight of the two sub-modes is shifted in favor
of one of them. This corresponds to the introduction of a relative shift between the two fields. The
relative shift breaks the overall reflection symmetry and leads to the movement of the drops.

3. **Influence of diffusion**

Next we discuss the influence of $r$ and $d$ presenting in Fig. [12] dependencies of droplet velocities
on reaction rate for different diffusion constants and the resulting phase diagram in the $r - d$ plane
for a fixed desorption/adsorption ratio $s$. One notes first that the strength of diffusion has nearly no
influence on the non-saturated regime and the transition value $r_{\text{max}}$ [the dashed line in Fig. [12](b) is
practically vertical]. Note that Fig. [12] focuses on relatively small diffusion. For large diffusion the
droplets will also stop as shown for model I in Fig. [5]. However, the saturated regime does depend
on $d$ quantitatively as well as qualitatively. For fixed $r$, with increasing $d$ the stable running
droplets become slower. In parallel its existence range in $r$ shrinks slowly. As in model I this
behavior is mainly caused by the diffusion of the $\phi$-field in front of the advancing droplet. There
it increases the coverage thereby reducing the overall wettability gradient, i.e. slowing down the
droplets. The qualitative change concerns the character of the drift-pitchfork bifurcation. With
increasing $d$ it becomes less subcritical and at a critical $d_c$ it becomes supercritical, i.e. there
exists no coexistence region for sitting and running drops any more. For small diffusion $d < d_c$
the existence range of stable sitting drops shrinks with increasing $d$ whereas for larger diffusion
$d > d_c$ it grows.

4. **Influence of volume**

Finally, in Fig. [13] we present the phase diagram for the dependence on reaction rate and droplet
volume for fixed desorption/adsorption ratio $s$ and diffusion constant $d$. The locations of the
saddle-node, the drift-pitchfork bifurcation and the velocity maximum are all shifted slightly to-
wards smaller $r$ when increasing the droplet volume. The range in $r$ of the non-saturated regime
shrinks slightly, but the range of the saturated regime and the metastable region remain practically
constant, they are only shifted towards smaller $r$.
FIG. 12: (a) Velocity of running droplets depending on the reaction rate $r$. The branch of running droplets emerges from zero reaction rate and undergoes a subcritical drift pitchfork bifurcation at finite $r > 0$. The reaction rate-velocity curve is shown for three different values of the diffusion constant in the $\phi$-field $d$.

(b) Existence of running and stationary droplets depending on the reaction rate $r$ and the diffusion constant of the $\phi$-field $d$. Shown is also the transition between non-saturated and saturated regime (dashed line). Remaining parameters are as in Fig.8.
FIG. 13: Existence of running and stationary droplets depending on the reaction rate $r$ and the droplet volume $V$ (in units of $10^4$). Shown is also the transition between non-saturated and saturated regime (dashed line). Remaining parameters are as in Fig. 8.

Obviously the two contact regions and the substrate outside the droplet are not affected by a change in the droplet volume. However, an increase in droplet size increases the viscous forces and therefore stalls the droplet movement at smaller reaction rates.

C. One-dimensional numerical simulations

Finally, we employ numerical simulations of running droplets and show that the type II model is able to describe different experimentally found modes of periodic droplet movement [24, 25]. To perform the simulations we use the routine ‘d02cjc’ provided by the NAG library [66]. It is based on a variable order, variable step size Adam’s method. The simulations either utilize periodic boundary conditions (to model droplets on a ring-like track [24]) or boundary conditions that mimic non-wettable borders of an otherwise wettable channel (to model droplets on finite stripe-like tracks) [24, 25]. Simulations are started from steady droplet solutions which develop in the absence of a chemical field, i.e. imposing $\phi = 0$. Then the droplet movement is initiated by breaking the symmetry of the $\phi$-field by imposing a small gradient and starting the adsorption/desorption reaction. After a short initial transient the running droplets follow periodic trajectories that do not
depend on details of the initial symmetry breaking. For the periodic boundary conditions the initial solution was one stationary droplet, whereas in the case of non-wettable boundaries the initial solution were two stationary droplets.

In the case of periodic boundary conditions the droplets move with constant speed and shape after an initial phase. Fig. 14(a) shows space-time plots of the evolution of the film thickness for different reaction rates $r$. The droplet velocity increases with the reaction rate as expected from our continuation results. Fig. 14(b) shows a comparison of the droplet velocities obtained in the simulations and by continuation. The values for the simulations are estimated after the droplets have reached a constant speed and shape. One finds that both velocities match fairly well. However, for higher reaction rates the simulations slightly overestimate the velocities compared to the continuation results. This results from the lower and equidistant discretization used in the simulation in time.

Experiments with droplets in a finite wettable channel found 'regular rhythmic motion' [24] or different types of 'shuttling motion' along with slowing and stopping behavior [25]. The here performed simulations show a smooth transition between the different types of periodic movements depending on our control parameters.

In general, the droplets in a wettable channel with non-wettable walls move periodically between the two walls as shown for different reaction rates $r$ in the space-time plots in Fig. 15(a). The initial solution of two stationary droplets quickly coarses upon starting the chemical reactions. The prevailing droplet oscillates between the channel walls with a frequency that depends on $r$.

The droplet movement can be classified into two identical but antisymmetric halfcycles (i.e. with different signs in the velocity, and profiles that are related by reflection). Fig. 15(b) shows the droplet velocity depending on time during one halfcycle exemplary for two reaction rates $r$. We find that each half cycle typically contains three phases, distinguishable by their different velocities. After meeting the non-wettable boundary (phase I) the droplet velocity is very low or even zero in the case of very small reaction rates $r$. In this phase, the $\phi$-field, which has been produced by the passing droplet has first to decrease, until the droplet can return on its own path. This phase of very small velocity is followed by a short phase of a very high velocity (phase II) until the droplet returns to a medium velocity (phase III), that is kept until it meets the opposite wall and the next half cycle begins. The subphases can also be very well distinguished in Fig. 16, were we show hidden line plots of the film thickness profiles (top) and coverage profiles (bottom) for one period of droplet movement. One can clearly see that in phase I (velocity close to zero),
FIG. 14: Simulations of running droplets on a one-dimensional circular stripe (periodic boundary conditions). Shown are (a) space time plots of the film thickness for different reaction rates \( r \) (indicated on the right) and (b) a comparison of drop velocities obtained by numerical simulation (\( \bigcirc \)) and continuation (solid line) for different values of the reaction rate \( r \). Remaining parameters are \( g = 2, s = 1, d = 0.001, L = 100, \bar{h} = 1.5, h_c = 2 \) and \( \Delta h = 0.2 \). Dark (light) colors correspond to small (large) film thickness.

After the droplet has encountered the non-wettable wall, the concentration in the \( \phi \)-field is very high and drops steeply in phase II when the droplet starts moving again. Similar subphases of droplet motion have also been observed in Ref. [25]. In the context of that work the continuous transition from Fig. 15 (a) between \( r = 0.002 \) (top) and \( r = 0.0001 \) (bottom) can be seen as their transition between 'shuttling motion' and 'intermittent shuttling motion'.

A simple bead-spring model based on a mechanical analogy is used in Ref. [25]. It models the
FIG. 15: Simulations of running droplets on a one-dimensional finite stripe bounded by non-wettable borders. Shown are (a) space time plots of the film thickness for different reaction rates $r$ (indicated on the right) and (b) the velocity of the center of mass of the film thickness depending on time examplary for $r = 0.001$ (top) and $r = 0.0001$ (bottom). The thin dotted line in (b) at $v = 0$ is meant to guide the eye. Remaining parameters are as in Fig.14. Dark (light) colors correspond to small (large) film thickness.

different experimentally observed regimes varying the wettability of the walls. Although it well captures the overall behaviour it can not resolve the more hydrodynamic aspects of the motion like the flow field inside the running droplets or the dynamical contact angles.
FIG. 16: Hidden-line space-time plots showing one halfcycle of the periodic movement of a running droplet on a one-dimensional stripe bounded by non-wettable borders. Shown are the film thickness (top) and the $\phi$-field (bottom). $r = 0.0005$ and remaining parameters are as in Fig.[13]

V. CONCLUSIONS

In the present work we have developed and analysed two models for chemically-driven self-propelled running droplets on solid substrates. Such moving droplets were described for several
experimental systems using different liquids, substrates and reacting substances [20, 21, 22, 23, 24, 25]. The movement of the droplets is driven by a self-produced wettability gradient that is perpetuated with the droplet itself by means of a desorption or adsorption reaction underneath the droplet.

Two types of experimental systems were reported on: (I) adsorption underneath the droplet decreases the wettability of the substrate irreversibly [21, 22, 23]; and (II) desorption underneath the droplet removes a more wettable coating that is recovered behind the droplet through an adsorption from a surrounding medium [24, 25]. We have described the droplet dynamics for both types of experimental systems using coupled evolution equations for the profiles of the film thickness and the adsorbate coverage. The equations can be derived from the Navier-Stokes equations using a long-wave or lubrication approximation [42] and assuming a small Damköhler number. The two types of experiments have been mapped onto two types of models. In the type I model a wettability-decreasing reaction takes place underneath the droplet. In the type II model this mechanism is extended by additionally introducing a wettability-increasing reaction that takes place at the substrate outside the droplet.

The wettability of the substrate enters both models through a disjoining pressure supplementing the Laplace pressure in the thin film equation. We have chosen here a disjoining pressure consisting of a long-range destabilizing part $\sim h^{-3}$ and a short-range stabilizing part $\sim h^{-6}$ used, for instance, in Ref. [67] to study coarsening in dewetting. The long-range part corresponds to van der Waals interaction and is not influenced by the adsorbate. All the influence of the adsorbate goes into the short-range part that in the simplest case selected here depends linearly on the adsorbate coverage.

Using continuation techniques and numerical simulations we have analyzed the solution behaviour of both models. Thereby we have focused on stationary running and sitting droplets in two dimensions. Both models display a transition from a non-saturated to a saturated regime with increasing reaction rate. The transition is also obtained when increasing the droplet volume. In the non-saturated regime an increase in the reaction rate leads to a larger wettability gradient implying a larger droplet velocity. In the saturated regime an increase in the reaction rate does not increase the coverage at the rear of the droplet, i.e. it does neither lead to a larger wettability gradient nor to a larger droplet velocity. However, it has turned out that the driving force and in consequence the velocity are decreasing slightly with increasing reaction rate. This effect is due to a rise in the adsorbat concentration in the advancing contact zone. A similar behavior occurs when the
droplet volume increases. There, however, in the saturated regime the velocity clearly decreases with increasing volume because the constant driving force (wettability gradient) is counteracted by an increasing viscous friction. The latter dependencies found for the non-saturated and the saturated regime correspond very well to experimental results of Ref. [21] (Fig. 5) and Ref. [23] [Fig. 7 (a)], respectively. To our knowledge there exist, however, no experimental results for a physico-chemical system that show the qualitative change between the two regimes in dependence of the droplet velocity on its volume. With the combination of materials used in Ref. [23] this should be possible because their Fig. 6 (a) shows the transition from the non-saturated to saturated regime for increasing solute concentration within the droplet. This corresponds directly to the dependence on the reaction rate shown here in Fig. 4.

Allowing for diffusion of the adsorbate along the substrate does not affect the results in the non-saturated regime too much. However, it leads to a stronger decrease of the velocity in the saturated regime because adsorbate is transported to the substrate in front of the running droplet thereby decreasing the overall wettability gradient. This may even lead to a transition towards sitting droplets. This transition occurs either continuously through a supercritical drift-pitchfork bifurcation or discontinuously through a saddle-node bifurcation. In the latter case a metastable parameter region exist where running and sitting drops can coexist.

We have determined the existence regions of running droplets depending on the reaction rate, the diffusion constant in the surface coating, the droplet volume and the desorption/adsorption ratio (only for model II). Both models show a strong dependence of the existence regions of sitting and running droplets on the diffusion constant and only a week dependence on the droplet volume. The strong dependence on the diffusion constant results mainly from the diffusion of the $\phi$-field in front of the advancing droplet, thereby greatly affecting the driving force and the droplet velocity. The type II model also shows a strong dependence on the desorption/adsorption ratio, which is due to a decreased substrate coverage in close proximity behind the droplet, effectively decreasing the driving force. The primary effect of the desorption reaction is the possibility for the droplet to return on its own path, which we have illustrated by numerical simulations of droplets moving on a wettable stripe with non-wettable borders in an oscillatory manner, which has also been observed experimentally [24, 25]. Finally, we have shown that different modes of periodic movement called 'shuttling motion' and 'intermittent shuttling motion' in Ref. [25] are covered by the presented model.

The analysis of both models has shown that for the chemically-driven running droplets the ad-
vancing contact angle is always smaller than the receding one. As was already shown in Ref. [41] also here the differences between the static and the dynamic contact angles at the front and the rear are one order of magnitude smaller than the difference between the two dynamic contact angles. This challenges the assumption of equal dynamic contact angles at the front and the rear that was used in Ref. [36, 37] to develop a simple description of self-propelled running droplets. A simple quantitative theory should instead be based on the assumption that the respective dynamic contact angles equal the different static contact angles at the front and rear.

The proposed type I and type II model based on thin film or lubrication theory can very well reproduce the main features of droplet motion that have been observed experimentally. However, they fail to reproduce the reported damped oscillations in the droplet shape overlaying the continuous droplet movement [25]. These oscillations could be on the one hand the result of a weakly inhomogeneous surface, since the authors of Ref. [25] themselves suggested that they have no full control of the experimental surface properties. On the other hand the oscillations could be a sign of a Hopf bifurcation along the solution branch of running droplets. We are well aware that the models presented in this paper are minimal models that, however, reproduce qualitatively most aspects of the behaviour of chemically self-propelled droplets. Refinements of the presented theory could include the viscous motion of the ambient medium as present in the type II experiments. This extension can still be based on the lubrication approximation, for instance, along the lines of the two-layer systems studied in Refs. [65, 68]. A second important extension could cover the case of a higher Damköhler number. Such a model has to include a description of the transport of the solute within the droplet.

APPENDIX A: NUMERICAL TECHNIQUES

1. Continuation

Sitting and running droplets are steady solutions in the laboratory and comoving frame, respectively. They can be followed in parameter space using numerical continuation techniques [50, 51], for instance, employing the continuation software AUTO97 [52]. The following section highlights some technical details of the employed techniques.

The basic idea behind continuation is that unknown solutions of an algebraic system for a certain set of control parameters are obtained by iterative techniques from known solutions nearby
in parameter space. Differential equations of the form

\[ u'(x) = f(u(x), p) \quad \text{with} \quad f, u \in \mathbb{R}^n \]  

subject to initial, boundary and integral constraints are discretized in space and then the resulting algebraic system is solved iteratively. Here the dash indicates the first derivative with respect to \( x \) and \( p \) denotes the set of control parameters. The presence of boundary conditions and/or integral conditions requires the presence of free parameters which are determined simultaneously and are part of the solution to the differential equation. The package AUTO97 is limited to the continuation of ordinary differential equations (ODE’s), thus it can only be used to compute droplet solutions in two dimensions. As an example we consider the continuation of stationary running droplets, steady in a comoving frame. After transforming Eqs. (11) and (12) into the comoving frame with velocity \( v \) and integrating the resulting time-independent thin-film equation we have the system of ODE’s

\[
\begin{align*}
    h_1' &= h_2 \\
    h_2' &= h_3 \\
    h_3' &= \frac{\mu - vh_1}{h_3^3} - \Pi_x(h_1, \phi_1) \\
    \phi_1' &= \phi_2 \\
    \phi_2' &= -\frac{1}{d}(R(h_1, \phi_1) + v\phi_2)
\end{align*}
\]

where \( \mu \) is an integration constant. It has the physical meaning of a mean flow in the comoving frame. \( h_1, h_2, h_3, \phi_1 \) and \( \phi_2 \) denote \( h, \partial_x h, \partial_{xx} h, \phi \) and \( \partial_x \phi \), respectively. The dashed quantities denote first derivatives with respect to \( x \). The system is flux conservative, thus we need to specify the integral condition

\[ 0 = \frac{1}{L} \int_0^L h_1 \, dx - \bar{h} \]  

where \( L \) is the system length and \( \bar{h} \) is the mean film thickness. Furthermore, for a complete description of the system we introduce periodic boundary conditions for the film thickness

\[
\begin{align*}
    h_1(0) &= h_1(L) \\
    h_2(0) &= h_2(L) \\
    h_3(0) &= h_3(L)
\end{align*}
\]
and the mixed boundary conditions

\[ 0 = \xi(h_1(0))\lambda_1 + \{\xi(h_1(0)) + s[1 - \xi(h_1(0))]\} \times (\phi_2(0) - \phi_1(0))\lambda_1 \]

(A11)

\[ 0 = \xi(h_1(0))\lambda_2 + \{\xi(h_1(L)) + s[1 - \xi(h_1(L))]\} \times (\phi_2(L) - \phi_1(L))\lambda_2 \]

(A12)

for the \( \phi \)-field (see appendix \[A2\]). Since translation of a running droplet solution in the \( x \)-direction also yields a valid solution one needs to introduce a pinning condition in the form of an additional boundary or integral condition, which will not be specified further in here.

As mentioned earlier, the number of boundary conditions (NBCD) and integral conditions (NINT) imposes a constraint on the number of parameters (NPAR), that have to be varied during the continuation process. Specifically NPAR=NBCD+NINT-NDIM+1, where NDIM is the dimensionality of the system of ODE’s. This condition leaves us here with three free parameters, which are the principal continuation parameter (e.g. the reaction rate \( r \)), the mean flow \( \mu \) and the droplet velocity \( v \). AUTO97 uses the method of Orthogonal Collocation for discretizing solutions, where the solution is approximated by piecewise polynomials with 2–7 collocation points per mesh interval. The mesh is adaptive as to equidistribute the discretization error. Having specified the ODE system in standard form with boundary and integral conditions AUTO97 then tries to find stationary solutions to the discretized system, by using a combination of Newton and Chord iterative methods. Once the solution has converged AUTO97 proceeds along the solution branch by a small step in the parameter space defined by the free continuation parameters and restarts the iteration.

The challenge usually is, to provide AUTO97 with a nonuniform starting solution for the continuation. For our purpose it is sufficient to start the continuation close to the point of the primary bifurcation point, where the stable uniform solution of the ODE system \([A2]–[A6]\) or a system similar to \([A2]–[A6]\) with periodic boundary conditions undergoes a Hopf-bifurcation. In the vicinity of the bifurcation one can determine analytically small-amplitude sinusoidal stationary traveling waves. By selectively using reaction, boundary or integral conditions as primary continuation parameter one finally computes fully nonlinear solutions for the film thickness and the coverage.

AUTO97 is not only able to follow solution branches but can also detect bifurcations, like saddle-node bifurcation or branching points, and can then follow these bifurcations in parameter space.
2. Boundary conditions

The use of periodic boundary conditions does not rule out interactions between droplets in consecutive periods either via the film thickness or the \( \phi \)-field. Interactions through the \( \phi \)-field arise if the desorption of the coating is slow compared to the drop movement. One way to avoid this problem is to use very large periods, such that the \( \phi \)-field has enough time to recover. This approach is successful in suppressing interactions but is very costly from a computational point of view. An alternative approach is to use other than periodic boundary conditions for the \( \phi \)-field. The following paragraphs illustrate this approach.

We generally consider a running droplet steady in a comoving frame, that has its maximum approximately in the center of the computational domain of length \( L \). We assume that the film thickness profile obeys periodic boundary conditions \( h_b \equiv h(0) = h(L) \) and \( \partial_z h(0) = \partial_z h(L) \ll 1 \). The latter restriction ensures that the minimal film thickness is close to equilibrium and the period is large compared to the droplet.

a. Model I

In model I we assume that there is no chemical reaction taking place outside the droplet. Therefore, outside the computational domain for \( x \leq 0 \) and \( x \geq L \) the following ordinary differential equation holds

\[
0 = d\phi_{xx} + v\phi_x, 
\]

which has the general solution

\[
\phi(x) = c + c' e^{-\frac{x}{\bar{z}}} 
\]

with \( c \) and \( c' \) being yet undetermined constants. For simplicity we assume that \( v > 0 \), i.e. the droplet is moving to the right. In front of the droplet it is assumed that \( \phi \to 0 \) as \( x \to \infty \). Furthermore we assume that the \( \phi \)-field adopts a finite value behind the droplet such that \( \phi \to \phi_\infty \) as \( x \to -\infty \). First we consider the boundary at \( x = 0 \) with the boundary value for the \( \phi \)-field \( \phi(0) = \phi_0 \) and the first derivative of the \( \phi \)-field \( \phi_x(0) = \phi_{x0} \). Since \( \phi_\infty \) is a finite value we find \( c' = 0 \), which leads to the Neumann boundary condition

\[
\phi_{x0} = 0. 
\]
Next we consider the boundary at \( x = L \) with the boundary value for the \( \phi \)-field \( \phi(L) = \phi_L \) and the first derivative of the \( \phi \)-field \( \phi_x(L) = \phi_{xL} \). Integrating (A13) we find

\[
0 = d\phi_x + v\phi + c'' ,
\]

where \( c'' \) is an integration constant. Since \( \phi \to 0 \) as \( x \to \infty \) and also \( \phi_x \to 0 \) as \( x \to \infty \) we find \( c'' = 0 \), which leads us to the mixed boundary condition

\[
0 = \phi_L v + \phi_{xL} .
\]

b. Model II

We assume for the \( h \)-field outside the computational domain with \( x \leq 0 \) or \( x \geq L \) \( h(x) = h(0) = h(L) = h_b = \text{const.} \). Then the concentration profile for the \( \phi \)-field for \( x \leq 0 \) or \( x \geq L \) can be computed analytically by solving a linear 2\(^{nd}\) order ordinary differential equation in \( x \) of the form

\[
0 = R_2(h_b, \phi) + d\phi_{xx} + v\phi_x ,
\]

which has the general solution

\[
\phi(x) = c + c'e^{\lambda_1 x} + c''e^{\lambda_2 x} ,
\]

where

\[
c = \frac{\xi(h_b)}{\xi(h_b) + s[1 - \xi(h_b)]} ,
\]

\[
\lambda_{1,2} = -\frac{v}{2d} \pm \sqrt{\frac{v^2}{4d^2} + \frac{r}{d}\{\xi(h_b) + s[1 - \xi(h_b)]}\}}
\]

and \( c' \) and \( c'' \) are yet undetermined constants. For \( x \to \pm \infty \) \( \phi \) adopts a small but finite value \( \phi_\infty \).

First we consider the boundary at \( x = 0 \). This leaves us with \( c'' = 0 \). Using the boundary values for the \( \phi \)-field \( \phi(0) = \phi_0 \) and the first derivative of the \( \phi \)-field \( \phi_x(0) = \phi_{x0} \) at the boundary \( x = 0 \) the following system of equations holds

\[
0 = \phi_0 - c - c' \quad (A22)
\]

\[
0 = \phi_{x0} - c'\lambda_1 . \quad (A23)
\]

Solving for \( c' \) one finds the following mixed boundary condition

\[
0 = \xi(h_b)\lambda_1 + \{\xi(h_b) + s[1 - \xi(h_b)]\} (\phi_{x0} - \phi_0\lambda_1) .
\]

(A24)
The same procedure can be performed at the boundary \( x = L \) with \( \phi(L) = \phi_L \) and \( \phi_x(L) = \phi_{xL} \) yielding the condition

\[
0 = \xi(h_b)\lambda_2 + \{\xi(h_b) + s [1 - \xi(h_b)]\} (\phi_{xL} - \phi_L \lambda_2) 
\] (A25)

In the continuation algorithm the two periodic boundary conditions for the \( \phi \)-field were substituted by the two mixed boundary conditions obtained above.

Acknowledgments

We thank E. Knobloch for his comments on the drift-pitchfork bifurcation, and acknowledge support through the EU RTN "Unifying principles in non-equilibrium pattern formation" (Contract MRTN-CT-2004–005728).

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Note, that the disjoining pressure used in Ref. [41] was
\[ \Pi(h) = \frac{2S_a d_0^3}{h^3} + \frac{S_p}{h} \left( 1 + \frac{\phi}{g} \right) \exp \left[ \frac{d_0 - h}{l} \right] \]
where for \( \phi = 0 \), \( S_a \) and \( S_p \) are the apolar and polar components of the total spreading coefficient \( S = S_a + S_p \), respectively, and \( l \) is a correlation length [49]. One usually describes the choice \( S_a > 0 \) and \( S_p < 0 \) as a combination of a stabilizing long-range van der Waals and a destabilizing short-range polar interaction. The apparent contradiction of qualitative similar results for model I for different verbal descriptions and combinations of signs used here and in Ref. [41] results from a subtle feature of the combination of exponential and power law. Combining a term \( \sim 1/h^3 \) and one \( \sim \exp(-h) \) leads for a proper choice of parameters to a dominance of \( 1/h^3 \) for large and very small \( h \). The exponential only dominates for intermediate thicknesses (see Ref. [69] for a related phase diagram). This implies that above verbal description only covers part of the feature of the disjoining pressure. On the contrary, the combination of two power laws used here clearly attributes the long range and short range forces to the terms \( h^{-3} \) and \( h^{-6} \), respectively. We therefore believe, that the chosen disjoining pressure more accurately represents the physical situation.

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\[ \Pi(h) = \frac{2S_a d_0^3}{h^3} + \frac{S_p}{h} \left( 1 + \frac{\phi}{g} \right) \exp \left[ \frac{d_0 - h}{l} \right] \]
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