1 Introduction

The static and dynamic wetting behaviour of simple and complex liquids on various surface types is highly relevant for many aspects of daily life, including cosmetics, cleaning and painting. It is also of great interest for many technical applications like, for example, printing, lubrication, and coating. Related phenomena have been studied for more than two hundred years.\(^1\) Many works published in the past decades consider (de)wetting on smooth homogeneous rigid solid surfaces,\(^4,5\) although more recently other substrate types have also attracted much attention. For instance, in many applications the substrate actually consists of a soft elastic solid,\(^6\) a viscous liquid,\(^7,8\) or a viscoelastic material.\(^9\) Lubricant-infused surfaces are another relevant example.\(^10,11\) In all these cases, the coupling of the dynamics of the liquid and of the underlying substrate is important. Such phenomena are of particular importance for micro- and mesoscale systems that are dominated by interface phenomena. The latter include adhesion and cohesion forces that, in the case of partially wetting liquids, cause droplet formation and also deform the substrate.

A particularly interesting situation arises for the wetting of soft elastic solid substrates.\(^6,12–14\) This situation is often referred to as “soft wetting”, and gives rise to many wetting phenomena that have no counterpart on a rigid solid. The most prominent feature of soft wetting is the appearance of a wetting ridge: an elastic deformation that is drawn out of the substrate by the vertical pulling force the liquid–gas interface exercises at the three-phase contact line.\(^15–18\) The ridge is responsible for intricate static and dynamic phenomena observed in soft wetting.\(^5\)

The fine-structure of the wetting ridge arises from an intricate balance of elastic and capillary forces, with recent literature identifying a central role for the surface tension of the solid.\(^16,19–21\) A central question is how the liquid angle, at equilibrium, is different from Young’s law for rigid substrates. It turns out that upon increasing the substrate softness, a double-transition is observed in the various angles between interfaces in the contact line region.\(^22\) Namely, when increasing softness from low values (i.e., starting with a rigid substrate), first a strong increase in the size of the wetting ridge is observed while the droplet shape remains approximately constant and the substrate surface in the inner region remains flat. Then, when the softness increases several orders of magnitude, a strong increase of the substrate depression under the droplet occurs a constant opening angle of the wetting ridge. The solid angle is governed by a balance of surface tensions,\(^19–21\) analogously to Neumann’s law known for floating lenses.
In comparison to liquids, however, the surface tension of solids brings in an additional complexity owing to its possible strain-dependence (i.e., the Shuttleworth effect).33-37 The wetting ridge further noticeably influences the dynamics of moving contact lines and droplet interactions. Depending on the substrate layer thickness and softness the wetting ridge may develop local depressions on both sides which counteract coarsening by droplet translation and can even result in an effective repelling force between them – also referred to as the “inverted Cheerios effect”.28 Another consequence is the so-called “viscoelastic braking”, describing the decreasing speed of a moving contact line with increasing softness of the substrate, due to increased dissipation within the latter. This effect was already investigated by Carré and Shanahan12,13 and Long et al.,29 followed up in recent years in ref. 30-32. New phenomena were discovered, such as intricate dependencies of sliding velocity on substrate thickness33 and stretching,34 as well as effects of free polymer chains that can be present in the substrate.35 Besides slowing down the contact line, the wetting ridge can also be the source of complex periodic stick-slip motion.9,31,36

From the modelling perspective, various approaches to soft wetting have been proposed. Most analytical results have been obtained using linear elasticity theory.16,17,19,21,22,29,31 In these approaches the surface tension forces are treated as tractions that are imposed as boundary conditions at the free surface. This approach has now been extended to large deformations, including the possibility of the Shuttleworth effect.27,37 Other studies employ an approach where the tractions are not imposed by hand, but computed via a Derjaguin (or disjoining) pressure – this is, a hybrid approach coupling elasticity to a thin-film static equation.39 Only recently, dynamic calculations coupling full hydrodynamic and nonlinear elasticity models have become available.40-45

Dynamic long-wave (or thin-film) models were developed in ref. 30 and 46-48. In particular, ref. 30, 46 and 47 present a model for the dynamics of a liquid drop on a viscoelastic layer while ref. 48 models the durotaxis of a liquid drop on a compliant Kirchhoff plate. Aspects of substrate elasticity also enter long-wave models for drops on polymer brushes and on growing layers of ice.50 These long-wave descriptions are further discussed after introducing our model in Section 2.1. Furthermore, some studies employ molecular dynamics simulations, exploring the elastocapillary interactions at the nanoscale.51,52

None of the reviewed modelling approaches lends itself easily to large-scale simulations of the coupled dynamics of elastic and liquid layer considering, e.g., the coarsening behaviour of the droplet ensembles on soft substrates. Also the incorporation of additional effects, e.g., the Shuttleworth effect or forces due to thermal or solutal Marangoni effects or condensation/evaporation, is a major effort. In the present work we present a simple qualitative model that captures main features of the coupled dynamics of elastic substrate and liquid drop or film. It is developed as a gradient dynamics model on a simplified free energy that in its simplest form captures compressional elasticity but can be extended to include effects such as those mentioned above. Such a model may then be employed in large-scale simulations similar to ref. 55 and 56 in the case of droplet ensembles on smooth rigid substrates.

The collective behaviour of such ensembles, e.g., their coarsening behaviour, is of large interest in practical contexts like, for instance, condensation or inkjet printing. Similar coarsening processes (Ostwald ripening) occur for emulsion droplets, quantum dots, or crystalline nanoparticles.59 In all cases, the mean cluster or drop size and their mean distance continuously grow following power laws. For simple nonvolatile liquids on horizontal homogeneous substrates coarsening is experimentally well studied60–63 and theoretically well understood through simulations and asymptotic considerations based on thin-film models.64-66 As explained more in detail below in Section 5 one distinguishes two coarsening modes – Ostwald ripening and migration – with the former often seen as dominant. However, for the dewetting process of a liquid film on a rigid substrate, ref. 65 and 68 find that, in contrast, there exist extended parameter regions where coarsening by migration dominates. We employ the developed simple model to investigate how substrate softness influences the dominant coarsening mode.

Here, the mesoscopic thin-film model is presented in the subsequent Section 2 where we also discuss how the quantities of the model are related to macroscopic interface tensions and, in consequence, how the classical Young and Neumann laws result as limiting cases. In Section 3 the model is employed to study the double transition of steady drops in its dependence on drop size, Section 4 investigates the spreading dynamics of a single drop in dependence of substrate softness for partially and completely wetting liquids, and Section 5 considers the coarsening behaviour of two drops and large drop ensembles, again in dependence of substrate softness. The various results are discussed in the context of literature results. Section 6 concludes and gives an outlook.

2 Modelling approach

2.1 Governing equations

The geometry of the problem is sketched in Fig. 1, where we define the liquid layer thickness $h(x,t)$, and the deviation of the elastic layer thickness from a uniform reference height $\xi(x,t)$. We employ a long-wave model consisting of a standard thin-film equation for $h(x,t)$ and a nonconserved Allen–Cahn-type relaxation dynamics for $\xi(x,t)$. In dimensional form we have

\[ \partial_t h = \nabla \cdot \left( \frac{h}{\eta} \nabla p_h \right), \]

\[ \partial_t \xi = -\frac{1}{\zeta} p_{\xi}. \]

Here, $p_h$ and $p_{\xi}$ are the pressures at the two interfaces that we will express in terms of functional derivatives. We further introduced the viscosity of the liquid $\eta$, and an “elastic friction constant” $\zeta$ that governs the relaxation of the elastic layer. We remark that the dynamics of the elastic layer is a spatially
This journal is © The Royal Society of Chemistry 2021 Soft Matter, 2021, 17, 10359–10375 | 10361

Fig. 1 The sketch shows the geometry of the considered two-layer system consisting of a liquid layer on an elastic layer. The description is based on the liquid layer thickness $h^\text{eq}$, $h$ and the vertical deviation $\xi^\text{eq}(x, t)$ of the elastic–liquid interface from a uniform reference height. The mean thicknesses of the layers are given by $h_0$ and $\xi_0$, respectively. The liquid–gas and the elastic–liquid interface have constant interface energies $\gamma_h$ and $\gamma_\ell$, respectively.

The system of governing equations is of gradient dynamics form, i.e., the pressures $p_h$ and $p_\ell$ are obtained as variations of an underlying free energy functional

$$
\mathcal{F}[h, \xi] = \mathcal{F}_\text{cap}[h, \xi] + \mathcal{F}_\text{vap}[h] + \mathcal{F}_\text{el}[\xi]
$$

with respect to $h$ and $\xi$, respectively. The energy eqn (2) features contributions due to capillarity of both interfaces, wettability of the liquid on the elastic layer, and elasticity. The first contribution consists of the interface energies in long-wave approximation, e.g., the area element $d^2s = \sqrt{1 + \nabla h^2}d^2x$ of the liquid–gas interface is approximated by $d^2s \approx \left(1 + \frac{1}{2}\nabla h^2\right)d^2x$.

Up to a constant, one then has

$$
\mathcal{F}_\text{cap}[h, \xi] = \int_\Omega \left[\frac{\gamma_h}{2}\nabla(\xi + h)^2 + \frac{\gamma_\ell}{2}\nabla\xi^2\right] d^2x
$$

where $\Omega$ is the considered domain, and $\gamma_h$ and $\gamma_\ell$ are the liquid-gas and the elastic–liquid interface energies per surface area, respectively. Partial wettability of the liquid on the elastic layer is incorporated into our mesoscopic model via

$$
\mathcal{F}_\text{vap}[h] = \int_\Omega f(h)d^2x
$$

where the local wetting energy $f(h) = \text{Derjaguin (or disjoining) pressure }$ is related to the Derjaguin (or disjoining) pressure $\Pi(h) = -\frac{1}{2}h\frac{\partial f}{\partial h}$. The coupling of an effective interface potential and elastic deformations was previously considered in ref. 39, 72 and 73. Here we use a simple form combining destabilising long-range van der Waals interactions and stabilising short-range adhesion forces, namely,

$$
f(h) = \frac{A_H}{2h^3} \left[\frac{2}{5} \left(\frac{h_0}{h}\right)^3 - 1\right].
$$

In this way, even macroscopically ‘dry’ spots are covered by a thin adsorption (or precursor) layer of thickness $h_0$ (a molecular scale). The Hamaker constant $A_H > 0$ is related to the (small) equilibrium contact angle $\theta_\text{eq}$ by $A_H = \frac{5}{3} \gamma_h h_0^2 \theta_\text{eq}^2$. As a result the effective elastic–gas interface energy is $\gamma_\text{sv} = \gamma_h + \gamma_\ell + f(h_0) = \gamma_h + \gamma_\ell - \frac{\gamma_h}{2} h_0^2/2$. In the limit of a rigid flat lower layer, this directly corresponds to the Young–Laplace law in long-wave approximation $\cos \theta_\text{eq} \approx 1 - \frac{\gamma_\text{sv}}{2 \rho g a^2}$.

Finally, we need to specify the elastic energy, which in linear elasticity theory can be expressed as a quadratic non-local functional of $\xi(x, t)$. Here we approximate the energy $\xi$ by a local form, as

$$
\mathcal{F}_\text{el} = \frac{k_\ell}{2} \int_\Omega \xi(x, t) \int_\Omega Q(x - x') \xi(x', t) d^2x' d^2x \approx \frac{k_\ell}{2} \int_\Omega \xi^2 d^2x.
$$

This gives a simple continuous spring potential $k_\ell x^2/2$, where $k_\ell$ corresponds to a spring constant per unit area. Such a free energy corresponds to the so-called Winkler foundation model, used also in problems of soft lubrication. In Appendix A we show how the above model can in principle be related to an elastic modulus $G$, see also ref. 78 for a detailed discussion.

Varying the functional (2) one obtains the pressures $p_h = \frac{\delta \mathcal{F}}{\delta h}$ and $p_\ell = \frac{\delta \mathcal{F}}{\delta \xi}$. After nondimensionalisation the governing eqn (1) become

$$
\frac{\partial h}{\partial t} = -\nabla \cdot \left(\frac{h^2}{2} \nabla(\Delta h + \xi + \Pi(h))\right),
$$

$$
\frac{\partial \xi}{\partial t} = \frac{1}{\tau} (\Delta h + \xi + \sigma \Delta \xi - \frac{\xi}{s}).
$$

where now all quantities are nondimensional. Here, the used rescaling for space and time is based on quantities of the liquid layer: vertical length scale $h_0$, horizontal length scale $L = \sqrt{3/5} h_0 / \theta_\text{eq}$, time scale $\tau_h = 27 h_0^2/25 \gamma_h \theta_\text{eq}^4$. Note that the employed long-wave approximation is valid if the ratio of the introduced vertical and horizontal length scales $s = h_0/L = \sqrt{3/5} \theta_\text{eq}$ is small. The dynamical system is governed by three dimensionless parameters that are all contained in the time-evolution equation for $\xi$:

$$
\frac{\gamma_\ell}{\gamma_h} \tau = \frac{5}{9} \frac{\gamma_h \theta_\text{eq}^2}{\eta} s = \frac{5}{3} \frac{\gamma_h \theta_\text{eq}^2}{k_\ell h_0^2}.
$$

The parameter $\sigma$ is the ratio of interfacial energies, $\tau$ reflects the ratio of viscous and viscoelastic dissipation, while $s$ is the dimensionless softness. Below we discuss how these dimensionless quantities can be interpreted in a macroscopic framework. Finally, eqn (7) contains the dimensionless Derjaguin pressure $\Pi(h) = h^6 - h^3$. In the scaled long-wave units, the equilibrium angle on a rigid substrate is $\sqrt{3/5}$.

We emphasize that all angles discussed below are long-wave angles, i.e., a long-wave angle $\hat{\theta}$ of $O(1)$ corresponds to a physical angle $\theta = o\hat{\theta}$ of $O(\epsilon)$. Further, we point out that for small physical angles $\hat{\theta}, h = \tan \theta \approx \theta$ implying that in long-wave scaling $\hat{\theta}, h$ directly holds. A diverging long-wave angle

\[\text{As given e.g., in ref. 29 in Fourier space.}\]
... modelled with a static long-wave description, i.e., the elastic layer dynamics is instantaneous and slaved to the dissipative liquid dynamics. In consequence, dissipation in the elastic layer is not considered. The long-wave model for a drop on a polymer brush in ref. 49 contains an elastic contribution resulting from entropic influences while ref. 50 describes a growing layer of ice under a droplet employing a periodic potential to capture microscale layering that resembles “periodic elasticity effects”. The latter two have a clear gradient dynamics structure similar to our model, however, in contrast to the present case they allow for transfer of material between the liquid drop and the substrate.

When determining steady states, \( \tilde{\theta}_h = \tilde{\theta}_c = 0 \) and one may integrate eqn (7) twice to obtain

\[ \begin{align*}
\frac{d}{dt} \Phi &= \int \frac{\delta \Phi}{\delta h} \frac{d}{dt} h + \frac{\delta \Phi}{\delta \xi} \frac{d}{dt} \xi d^2 x, \\
&= - \int h \left| \nabla^2 \frac{\delta \Phi}{\delta h} \right| d^2 x - \int \frac{1}{2} \left( \frac{\delta \Phi}{\delta \xi} \right)^2 d^2 x < 0
\end{align*} \tag{11} \]

where from (10) to (11) we use eqn (7) and (8) as well as integration by parts (in the first term). It is always negative and corresponds to the negative of the total dissipation \( \mathcal{D} \), i.e., \( \mathcal{D} = -d\mathcal{F}/dt \). A useful feature of eqn (11) is that it allows one to individually measure the contributions coming from the liquid and elastic layer, respectively, and to resolve them spatially using the dissipation density \( \Gamma(\bar{x}, t) \), i.e.,

\[ \mathcal{D} = \mathcal{D}_h + \mathcal{D}_c = \int_{\Omega} \Gamma_h d^2 x + \int_{\Omega} \Gamma_c d^2 x = \int_{\Omega} \Gamma d^2 x. \]

After having explained the gradient dynamics structure of our model for the coupled liquid and compressible substrate dynamics we briefly comment on related available long-wave models. The model in ref. 30, 46 and 47 describes such dynamics in the case of an incompressible viscoelastic material, however, its complexity does not allow to easily discern a gradient dynamics structure and underlying energy. Ref. 48 describes a drop durotactically sliding on a compliant Kirchhoff plate with position-dependent bending elasticity. However, the plate is...
to the microscopic thickness of the adsorption layer divided by the small equilibrium contact angle \( \theta_{eq} \).

A central feature of the macroscopic description is that it exhibits well-defined contact angles, as indicated in Fig. 2(a). Given that the drops are typically very large compared to the scale \( h_o \), these angles can also be observed in the mesoscopic model, at intermediate distances from the contact line (much larger than \( h_o \), much smaller than the drop size). If in addition we consider \( s \to \infty \), for which there is no elasticity and the substrate is a liquid layer, one can directly express the contact angles in the interfacial energies — using Neumann’s law.\( ^8 \) In the non-dimensional long-wave limit the Neumann conditions read

\[
\text{Vertical: } \theta_L - \sigma \theta_{SL} = [1 + \sigma + \tilde{c}^2 f(h_o)] \theta_{SV}, \quad (16a)
\]

\[
\text{Horizontal: } \theta_L^2 + \sigma \theta_{SL}^2 + 2f(h_o) = [1 + \sigma + \tilde{c}^2 f(h_o)] \theta_{SV}^2, \quad (16b)
\]

where all angles consistently correspond to long-wave angles, i.e. we dropped the tildes for these angles. In the absence of any elasticity, the liquid—liquid system approaches the shape of a liquid lens, i.e. as \( s \) increases the angle \( \theta_{SV} \) approaches zero. Neglecting this angle from the Neumann conditions (2), i.e. setting the right hand sides in eqn (2) to zero, the remaining angles may be expressed like

\[
\text{Vertical: } \theta_L = \sigma \theta_{SL}, \quad (17a)
\]

\[
\text{Horizontal: } \theta_{SL} = \sqrt{-2f(h_o)/(\sigma(1+\sigma))}. \quad (17b)
\]

These relations give a well-defined macroscopic meaning to the dimensionless parameter \( \sigma \) (ratio of surface energies) and the wetting energy of the adsorption layer \( f(h_o) \). We remind that in dimensionless units \( f(h_o) = -3/10 \), which governs the liquid angle for the rigid substrate (\( s = 0 \)), i.e., in long-wave scaling \( \theta_{eq} = \sqrt{-2f(h_o)} \). Indeed, we will verify below that the “liquid” and “rigid” limits are correctly recovered by the mesoscopic model.

Finally, let us comment on the final dimensionless number \( \tau \), which expresses a ratio of timescales. The substrate eqn (8) contains a stiffness \( k_c \) and dissipative constant \( \zeta \). As is common for the Kelvin–Voigt model, a timescale is obtained by taking the ratio \( \zeta/k_c \). Exciting the solid at slower timescales gives a nearly elastic response, while at faster timescales the layer is highly dissipative. We thus introduce the substrate’s viscoelastic timescale \( \tau_\text{v} \), defined as

\[
\tau_\text{v} = \frac{\zeta}{k_c} \Rightarrow \tilde{\tau} = \frac{\tau_\text{v}}{\tau_b} = \frac{\tau_{ec}^2}{\sigma}. \quad (18)
\]

In the second step we expressed the ratio \( \tau_b/\tau_\text{v} \), comparing the viscoelastic timescale to that of the viscous fluid. If one chose to scale the model on the substrate properties, most notably using \( \tau_{ec} \) and \( \tau_\text{v} \), eqn (8) would take the form

\[
\frac{\partial h}{\tau_\text{v}} = \frac{1}{\zeta} \left[ \frac{\tilde{c}^2}{\sigma} \Delta(h + \zeta) + \tilde{c}^2 \Delta h - \zeta \right]. \quad (19)
\]

In what follows, we stick to the original formulation, using \( \tau, \sigma, s \) as dimensionless numbers, obtained when scaling the variables on the fluid properties. For a physical interpretation of the results, and in order to compare our model to existing theory and experiments, we will make use of the connections (15) and (18).

### 2.3 Numerical approach

In the following we employ the developed model in a number of static and dynamic situations using two numerical techniques, namely the continuation of steady states and direct time simulations. For both methods the one-dimensional spatial domain \( \Omega \) of size \( D \) is discretized into a normally nonuniform adaptive grid with periodic or Neumann boundary conditions.

Steady solutions are obtained by solving the second order eqn (12) and (13) by pseud-arclength continuation \(^{80-82} \) employing the packages auto-\texttt{0p} \(^{83} \) and PDE2path.\(^{84} \) They are frequently used for thin-film descriptions of layers and drops of simple or complex liquids on solid or liquid substrates.\(^{7,71,85,86} \)

Continuation is started at an analytically or numerically known steady state, e.g., a flat film with a small harmonic modulation of a wavelength given by a linear analysis as critical value for a surface instability. Then the steady state is followed in parameter space using, e.g., softness as control parameter, and fixing the drop volume \( \text{via} \) a Lagrange multiplier (here \( \mu \)). An individual continuation step combines a tangent predictor and Newton correction steps.\(^{82,87} \)

For the direct time simulations of eqn (7) and (8) the FEM-based software package omph-lib\(^{88} \) is employed. An adaptive time stepping is used based on a backward differentiation method of order 2 (BDF2) from which the next state is obtained \text{via} a Newton procedure. The efficient adaptive time stepping and mesh refinement routines allow for a treatment of even very large systems. Fig. 3 illustrates that even quite sharp corners like the tip of a wetting ridge are properly resolved.

In the following sections we employ the model to study the double transition of steady drops (Section 3), the spreading dynamics of a single drop (Section 4) and the drop coarsening behaviour (Section 5). The results are compared to findings in the literature to evaluate the validity of the developed model.

### 3 The double transition for static drops

First, we investigate a static situation, namely, the dependence of the properties of sitting drops on the substrate softness \( s \). As described in Section 2.3 this is done by employing numerical continuation to track solutions to the nondimensional steady eqn (12) and (13). Typical drop and substrate profiles are given in Fig. 4(a), showing how the equilibrium morphology changes as the substrate changes from very rigid to very soft. Here we explore the essential features, and compare the results to existing literature.

Of particular interest are the angles sketched in the inset of Fig. 4(b), namely, \( \theta_L \), the angle the liquid–gas interface takes with the horizontal and \( \theta_{SL} + \theta_{SV} \), the combined angles with the...
horizontal on both sides of the wetting ridge. Note, that the left axis of Fig. 4(b) and 5 gives \(-\theta_{\text{SL}} + \theta_{\text{SV}}\), as this is the long-wave equivalent to \(\pi - \theta_s\) such that the plots visually resemble Fig. 3 of ref. 22 (cf. Fig. 2 for angle definitions). Both, \(\theta_s\) and \(\theta_{\text{SL}} + \theta_{\text{SV}}\), are expected to smoothly change from fulfilling Young’s law for a rigid substrate \((s \to 0)\) to fulfilling Neumann’s law in the limiting case of a liquid substrate \((s \to \infty)\). The change of both angles with softness is given in Fig. 4(b) with the scale for \(\theta_L \ [\theta_{\text{SL}} + \theta_{\text{SV}}]\) given on the right [left] hand side axis. As in the full elastic theory for drops with contact angle \(\pi/2\) in ref. 22 two distinguished separate transitions occur. Starting in the rigid limit and increasing the softness, first, one observes the formation of wetting ridges. As the overall shape of the drop itself barely changes, in this first transition \(\theta_L\) remains approximately constant while \(\theta_{\text{SL}} + \theta_{\text{SV}}\) strongly increases (in Fig. 4(b) at about \(s = 1 \times 10^3\)). In parallel, the wetting ridge grows as shown in Fig. 4(c). Increasing the softness further, the substrate below the drop is compressed owing to the Laplace pressure imposed by the drop. In consequence, the drop gradually sinks into the elastic layer accompanied by an inwards rotation of the wedge structure in the three-phase contact region. This rotation marks the second transition (in Fig. 4(b) at about \(s = 10^7\)...) now \(\theta_L\) strongly decreases while \(\theta_{\text{SL}} + \theta_{\text{SV}}\) remains approximately constant. The second transition is also reflected as a maximum in \(\theta_{\text{SL}} + \theta_{\text{SV}}\) given on the right [left] hand side axis. In the full elastic theory for drops with contact angle \(\pi/2\) in ref. 22 two distinguished separate transitions occur. Starting in the rigid limit and increasing the softness, first, one observes the formation of wetting ridges. As the overall shape of the drop itself barely changes, in this first transition \(\theta_L\) remains approximately constant while \(\theta_{\text{SL}} + \theta_{\text{SV}}\) strongly increases (in Fig. 4(b) at about \(s = 1 \times 10^3\)). In parallel, the wetting ridge grows as shown in Fig. 4(c). Increasing the softness further, the substrate below the drop is compressed owing to the Laplace pressure imposed by the drop. In consequence, the drop gradually sinks into the elastic layer accompanied by an inwards rotation of the wedge structure in the three-phase contact region. This rotation marks the second transition (in Fig. 4(b) at about \(s = 10^7\)...) now \(\theta_L\) strongly decreases while \(\theta_{\text{SL}} + \theta_{\text{SV}}\) remains approximately constant. The second transition is also reflected as a maximum in \(\theta_{\text{SL}} + \theta_{\text{SV}}\) given on the right [left] hand side axis. In the full elastic theory for drops with contact angle \(\pi/2\) in ref. 22 two distinguished separate transitions occur. Starting in the rigid limit and increasing the softness, first, one observes the formation of wetting ridges. As the overall shape of the drop itself barely changes, in this first transition \(\theta_L\) remains approximately constant while \(\theta_{\text{SL}} + \theta_{\text{SV}}\) strongly increases (in Fig. 4(b) at about \(s = 1 \times 10^3\)). In parallel, the wetting ridge grows as shown in Fig. 4(c). Increasing the softness further, the substrate below the drop is compressed owing to the Laplace pressure imposed by the drop. In consequence, the drop gradually sinks into the elastic layer accompanied by an inwards rotation of the wedge structure in the three-phase contact region. This rotation marks the second transition (in Fig. 4(b) at about \(s = 10^7\)...) now \(\theta_L\) strongly decreases while \(\theta_{\text{SL}} + \theta_{\text{SV}}\) remains approximately constant. The second transition is also reflected as a maximum in \(\theta_{\text{SL}} + \theta_{\text{SV}}\) given on the right [left] hand side axis. In the full elastic theory for drops with contact angle \(\pi/2\) in ref. 22 two distinguished separate transitions occur. Starting in the rigid limit and increasing the softness, first, one observes the formation of wetting ridges. As the overall shape of the drop itself barely changes, in this first transition \(\theta_L\) remains approximately constant while \(\theta_{\text{SL}} + \theta_{\text{SV}}\) strongly increases (in Fig. 4(b) at about \(s = 1 \times 10^3\)). In parallel, the wetting ridge grows as shown in Fig. 4(c). Increasing the softness further, the substrate below the drop is compressed owing to the Laplace pressure imposed by the drop. In consequence, the drop gradually sinks into the elastic layer accompanied by an inwards rotation of the wedge structure in the three-phase contact region. This rotation marks the second transition (in Fig. 4(b) at about \(s = 10^7\)...
down the spreading of the drop. Here we verify whether this viscoelastic braking is correctly accounted for in our model. Beyond that, we compare the case of a drop of partially wetting liquid, i.e., employing the wetting energy (5) as in Section 3, and the case of completely wetting liquid. The latter is realized by using a positive sign for the long-range contribution to the wetting energy (5) resulting in a strictly repulsive disjoining pressure. As described in Section 2.3 the spreading is analysed by direct time simulation of eqn (7) and (8). Note that here we make use of the reflection symmetry w.r.t. the maximum of the drop, and only simulate one half of the drop using Neumann boundary conditions.

4.1 Partial wetting
The initial profile is a drop of parabolic shape on a flat substrate with an initial contact angle that is three times larger than the equilibrium contact angle resulting from Young’s law, i.e., \( \theta_{\text{ini}} = 3 \theta_{\text{eq}} = 3 \sqrt{3}/5 \). The initial drop height is \( h_{\text{max}} = 1000 \), i.e., three orders of magnitude larger than the adsorption layer height. A typical time evolution at intermediate softness \( s = 10^5 \) is shown in Fig. 7. Given that the initial contact angle is larger than its equilibrium value, the disjoining pressure will drive a flow of liquid towards the contact line. This induces a contact line motion towards equilibrium, as visible in Fig. 7(a). The corresponding evolution of the wetting ridge is indicated in Fig. 7(b). At very early time, a ridge is growing out of the initially flat substrate and subsequently moves along with the contact line. This behavior closely resembles that observed in experiments. The corresponding dissipations in the liquid and in the substrate are reported, respectively, in Fig. 7(c) and (d). For both, the largest dissipation happens near the contact line. For this set of parameters, it turns out that the dissipation peak is larger in magnitude inside the solid, showing that the substrate dissipation dominates over liquid dissipation. This is the viscoelastic braking effect that we quantify in detail below.

A common way to characterize dynamical wetting is via the dependence of the (dynamic) liquid angle \( \theta_L \) on contact line velocity \( v \). To extract this information from our simulations we trace the estimated macroscopic contact line position \( x_C \), and its velocity \( v = dx_C/dt \) during the spreading process. There are different ways of defining \( x_C \) in the mesoscopic model. Here, we use

\[ 4 \quad \frac{d\theta_L}{dt} = \frac{(h_{\text{ini}} - h_{\text{eq}})}{2} \]

4 Drop spreading

We now investigate the dynamical process of droplet spreading. Of particular interest is the effect of viscoelastic braking, i.e., the decrease of contact line speed during spreading with increasing substrate softness. Any motion of the contact line will displace the wetting ridge, which induces a time-dependent deformation. Owing to the viscoelasticity of the solid, this offers an additional source of dissipation that slows down the spreading of the drop. Here we verify whether this viscoelastic braking is correctly accounted for in our model. Beyond that, we compare the case of a drop of partially wetting liquid, i.e., employing the wetting energy (5) as in Section 3, and the case of completely wetting liquid. The latter is realized by using a positive sign for the long-range contribution to the wetting energy (5) resulting in a strictly repulsive disjoining pressure. As described in Section 2.3 the spreading is analysed by direct time simulation of eqn (7) and (8). Note that here we make use of the reflection symmetry w.r.t. the maximum of the drop, and only simulate one half of the drop using Neumann boundary conditions.

![Fig. 5](image1)

Fig. 5 Shown is the dependence of the angles \( \theta_L \) (l.h.s. axis, blue curves) and \( -\theta_{SV} + \theta_{SL} \) (r.h.s. axis, red curves) on substrate softness \( s \) for steady drops of different volumes \( V \) as given in the legend. The remaining parameters are as in Fig. 4. As the volume increases, the two distinguished transitions become increasingly separated and for the angles the approach of the Young–Laplace (YL) and Neumann (N) law in the rigid and liquid limit, respectively, improves. The respective analytically predicted values are marked by blue (for \( \theta_L \)) and red (for \( \theta_{SV} + \theta_{SL} \)) arrows and thin horizontal lines.

![Fig. 6](image2)

Fig. 6 The softness values where the two distinguished transitions occur are shown in dependence of the drop volume [cf. Fig. 5]. The transition values are determined as the values where the respective angle is \( \theta = (\theta_{\text{max}} - \theta_{\text{ini}})/2 \).

Common approximations for \( x_C \) are: (i) the use of the position of steepest slope of the liquid–gas interface \( \theta_L \) is obtained as slope of the corresponding tangent, (ii) fitting a parabola to the drop apex and evaluate \( x_C \) and \( \theta_L \) at its intersection with the reference flat liquid–elastic interface at \( z = 0 \), (iii) the use of the intersection of the tangent from (i) and \( z = 0 \), and (iv) using the position of the maximum of the wetting ridge. Our tests show that method (i) appears too often to shift \( x_C \) into the droplet by a non-negligible amount. Further, we find method (ii) to be unreliable as it entirely ignores the local geometry of the three-phase contact region, a problem partly shared by method (iii). Finally, method (iv) is problematic when the contact line surfs on the wetting ridge, i.e., when the wetting ridge falls behind. Beyond that, method (iv) requires a fitting routine to prevent errors resulting from the discretization. Ref. 30 uses method (iii) to predict certain subtle differences in the \( x_C \) dependence of the spreading of partially and completely wetting liquids on softness. Here, we only consider the dependence of dynamic contact angle on contact line speed \( dx_C/dt \) and method (iii) gives robust results.
we determine $x_c$ as the intersection point of the tangent to the liquid–gas interface at the position of its steepest slope with the reference flat liquid–elastic interface at $z = 0$.

Fig. 8 presents the resulting dynamic liquid contact angles, for different softness (varying $s$) and different degrees of substrate viscoelasticity (varying $\tau$). Specifically, we report the difference in angle with respect to its equilibrium value, $\Delta \theta_L = \theta_L - \theta_{eq}$, as a function of the contact line velocity $v = dx_c/dt$. In all cases there is an extended regime of proportionality $\Delta \theta_L \sim v$ where the change in contact angle is linear with the contact line velocity. Such a linear dependence is indeed expected here. The liquid dynamics is governed by Stokes flow, for which the dynamic contact angle in partial wetting is linear with velocity.\(^5,^{89,90}\) Similarly, a Kelvin–Voigt solid (as in the present case), will give a linear dependence of the dynamic contact angle.\(^31\) We remark that in experiments, the exponent is much lower than unity, with typical values around 0.5. This is because the dynamic contact angle inherits the scaling of the loss modulus – for the Kelvin–Voigt model this is linear in frequency but in crosslinked polymer networks typically exhibit smaller exponents.\(^6\)

However, we still need to disentangle the relative roles of liquid and solid dissipation. Fig. 8(a) corresponds to a very small value of $\tau$, so that the solid very quickly adapts to any changes in the liquid. We observe that the spreading dynamics is completely independent of the softness $s$ in this case. This implies that the substrate has no influence on the spreading, in spite of the presence of a wetting ridge. Indeed, at very small $\tau$ the solid’s response is nearly instantaneous, which corresponds to a perfectly elastic limit where there is no dissipation inside the substrate. By contrast, for larger values of $\tau$ the softness plays an important role, as seen in Fig. 8(b). Softer substrates (larger $s$) give a larger departure from the equilibrium angle. This indeed is a signature of the viscoelastic braking effect: for a given value of $\theta_L - \theta_{eq}$, the contact line velocity is lower on softer substrates. To verify whether the substrate dissipation indeed dominates over liquid dissipation, we try to collapse the same data using a characteristic velocity of the substrate. Following ref. 6 and 31, this velocity is given by the ratio $\ell_{el}/\tau$, which in the present scaling implies a dependence on softness as $\sim 1/\sqrt{s}$. Fig. 8(c) therefore shows the dynamic contact angle plotted against the rescaled dimensionless substrate velocity $v \sqrt{s}$: the data now collapses, confirming that the spreading dynamics is indeed completely governed by the substrate’s viscoelasticity.

**4.2 Complete wetting**

The case of a completely wetting fluid can be analogously treated, by using a positive instead of the negative sign in the wetting energy (5). A corresponding typical time evolution is displayed in Fig. 9, for the same initial conditions as for the partially wetting case. At early times, the dynamics is very similar to that of the partially wetting case. The liquid starts to spread and a ridge is pulled out of the initially flat solid, however, the ridge remains quite small. Owing to the complete wetting, however, the drop does not spread toward a shape with finite equilibrium angle but, instead, the contact line continues to spread. During this process the wetting ridge gradually decays, cf. Fig. 9(b), in contrast to the growth observed under partial wetting conditions. We attribute this trend to the continuous decrease of the contact angles reached during spreading, which diminishes the upward pulling force of surface tension. In consequence, the dissipation in the substrate decays more rapidly than the dissipation in the fluid, as seen in Fig. 9(c) and (d).

Fig. 10 shows the dynamic contact angle $\theta_L$ against the spreading velocity $v$ for various softness values $s$. The chosen
ripening, or Ostwald ripening.65,68,93,94

A corresponding time evolution can be seen in HeSH2021-SupplMat-Video03-fig09.mp4 of the ESI.†

value of the relaxation $\tau = 1$ exhibits a strong viscoelastic braking in the partially wetting conditions – the complete wetting case, by contrast, shows no significant dependence on the substrate softness $s$. This reflects the observations in Fig. 9, where only a comparatively small and soon decaying wetting ridge is found. The observed relation for the dynamic angle is consistent with the scaling $L \sim v^{1/3}$, which is the classical Tanner–Cox–Voinov relation, for viscous spreading under complete wetting conditions. We thus conclude that in complete wetting, the softness of the substrate does not influence the spreading dynamics.

5 Droplet coarsening

When an ensemble of drops coarsens the average drop size and distance increase in time while their number decreases. This may occur at overall fixed liquid volume or involve condensation (and evaporation). In any case, the underlying elementary process is the coarsening of two (and sometimes three) drops into one. For such a coarsening step, there exist two basic coarsening modes.

This is, on the one hand, the mass transfer mode of coarsening: transport of material occurs from one droplet to the other one(s) while the centers of mass of the drops do not move. The mass flux is always directed from the small drop to the larger one (guaranteeing coarsening) due to the difference in Laplace pressure in the two coarsening drops. The process ends when the smaller drop has vanished and all the mass is contained in the remaining drop. There are different transport channels for the mass transfer mode: transport may occur via the vapour phase (for volatile liquids) or through an ultrathin adsorption layer (for nonvolatile, partially wetting liquids). This coarsening mode is in the literature referred to as volume or mass transfer mode, (drop) collapse mode, diffusion-controlled ripening, or Ostwald ripening.65,68,93,94

On the other hand, there is the translation mode of coarsening: the entire drops migrate toward each other, until their contact lines touch and the drops fast coalesce. This mode is also referred to as collision, coalescence, or migration mode.65,68,93,94 Sketches of mass transfer and translation modes can be found in ref. 65 and their relation to the translation symmetry mode for single fronts or contact lines in a homogeneous system is discussed in Section 3.4 of ref. 95. The stabilization of both coarsening modes by substrate heterogeneities gives rise to intricate bifurcation behaviour.96

The two coarsening modes can contribute to the coarsening process in a mixed way and can nearly not be distinguished in the resulting scaling laws describing the increase of mean drop volume and drop distance for coarsening large drop ensembles. For the most common cubic mobility function (resulting from no-slip boundary conditions at the solid substrate) the scaling laws for the two modes only differ by a logarithmic factor.65 Ref. 68 shows for drops on a one-dimensional solid substrate that beyond a certain threshold with increasing mean drop size and decreasing mean drop distance the translation mode becomes more prevalent until it finally dominates (cf. Fig. 15 of ref. 68 and also ref. 98 and 99). Below the threshold, only the mass transfer mode is found.

Here we briefly show that the developed model is well suited to study drop coarsening on soft substrates. In particular, the following two sections discuss the influence of softness on the individual coarsening process of two drops and on the collective coarsening behaviour of a large ensemble, respectively.

5.1 Case of two drops

Here, we employ the developed model to investigate the dynamics of the individual coarsening step for two neighboring drops and discuss how the dominant coarsening mode depends on substrate softness. The process is initiated with two parabolic drops of identical volume $V$ that are placed at a distance $L_0$ (between the contact lines) on an initially flat but soft substrate. Their contact angle corresponds to the equilibrium value in the corresponding rigid limit. The resulting profile is perturbed by
some small initial noise to slightly break the initial reflection symmetry about the center line \(x = 0\). Typical time evolutions for selected softnesses \(s\) at fixed \(V = 100\) and \(L_0 = 50\) are given in Fig. 11 as space–time plots. In all cases, initially, up to about \(t = O(10^3)\) the substrate–liquid interface relaxes, i.e., the drops sink into the soft substrate (not shown). The amount of sinking, i.e., the size of the indentation in the substrate, and the size and height of the developing wetting ridge all depend on softness (cf. Section 3). Then, the coarsening process sets in on a much larger time scale. Depending on softness, it is usually terminated at \(t = 10^5\ldots10^7\).

Inspecting Fig. 11, we find that the coarsening process qualitatively changes with softness in a nonmonotonic manner. In panels (a) and (d), respectively corresponding to the rigid and soft limits, the droplets exchange volume with minor or without lateral translation (volume mode). By contrast, at intermediate softness panels (b) and (c) exhibit a clear lateral motion of the drops towards one another (translation mode). This nonmonotonic change in coarsening behaviour with \(s\) is reminiscent of the nonmonotonic change of the wetting ridge height with softness, observed for static drops in Fig. 4(c). This suggests that the translation regime is mediated by elastic deformation, as a form of the Cheerios effect.28

To quantify the relative importance of volume and translation mode, we introduce an asymmetry measure for the considered two-drop configuration, namely,

\[
A(t) = \frac{1}{2V} \int_D \left| h(x, t) - h_0 \right| dx. \tag{20}
\]

which corresponds to the center of mass of the drop volume above the adsorption layer \((h_a = 1)\). A coarsening process exclusively involving the translation mode that symmetrically moves the drops towards each other would keep \(A = 0\) as the center of mass remains at \(x = 0\). By contrast, an exclusive transfer of liquid from one to the other drop via the volume transfer mode would result in a maximal increase to \(A = A_{\text{max}}\) as all the volume is finally centered about the initial position of the remaining drop. If both coarsening modes contribute the resulting value of \(A\) lies between the two extremes.

For the time evolutions in Fig. 11, the corresponding measures \(A(t)\) are given in Fig. 12(a) allowing us to quantitatively discuss the changes of the coarsening mechanism with increasing softness: for (nearly) rigid substrates \([s = 10^{-1}, \text{Fig. 11(a)}]\) the volume transfer mode dominates, as one drop shrinks and vanishes while the other one grows. The asymmetry measure \(A(t)\) monotonically increases from zero and reaches a final value of \(A_{\text{fin}} \approx 22 < A_{\text{max}} \approx 40\) [Fig. 12(a)]. It is interesting to note that in parallel to the volume transfer both drops slightly migrate into the same direction, pointing from the growing to the shrinking drop. Even after the small drop vanishes the remaining drop briefly keeps traveling in that direction. This effect is well known for drops on rigid substrates, see e.g. ref. 68 and 93, and is related to liquid motion in the adsorption layer.

Increasing the softness to \(s = 10\), volume transfer remains dominant. The amount of migration into the same direction decreases resulting in an increase of \(A_{\text{fin}}\) to about 30. The increase can be well appreciated in Fig. 12(b) where the final asymmetry measure \(A_{\text{fin}}\) is shown as a function of softness \(s\). Reaching the regime of intermediate elasticity, at about \(s = 10^2\) [Fig. 11(b)] the behaviour abruptly changes and coarsening becomes entirely dominated by the translation mode with \(A_{\text{fin}} \approx 0\), as seen from the dotted line in Fig. 12(a). When further increasing \(s\) by about one magnitude, the volume mode starts to become important again: here, translation is still dominant during the early stages of the process, but volume

Fig. 11 Shown are space–time plots for the coarsening of two drops of partially wetting liquid into a single drop. The softness increases from (a) \(s = 10^{-1}\) in the rigid limit, via (b) \(s = 10^{0}\) and (c) \(s = 10^{1}\) (intermediate elasticity) to (d) \(s = 10^{3}\) (liquid limit). With increasing softness the dominant coarsening mode changes as discussed in the main text. The volume of each drop is \(V = 100\), the initial distance of their “inner” contact lines is \(L_0 = 50\) and the domain size is \(D = 400\). The remaining parameters are as in Fig. 7. The corresponding time evolutions can be seen in (a) HeSH2021-SupplMat-Vide04-fg11.mp4, (b) HeSH2021-SupplMat-Vide05-fg11.mp4, (c) HeSH2021-SupplMat-Vide06-fg11.mp4 and (d) HeSH2021-SupplMat-Vide07-fg11.mp4 of the ESI.†

Fig. 12 (a) Shown is the temporal change in the asymmetry measure \(A(t)\) [eqn (20)] for two-drop coarsening processes at softness values given in the legend (including the values of Fig. 11). The maximal possible value for the given drop volume and distance corresponds to the absolute value of the initial drop position \(A_{\text{max}} = |h_a| = 40\). Panel (b) gives the intricate dependence of the final value \(A_{\text{fin}}\) on softness \(s\). It shows that the volume mode is dominant apart from a finite range of intermediate softness \((s \approx 10^{1}\ldots10^{3})\) where the translation mode dominates. See main text for further discussion.
transfer becomes relevant in the final phase [see $s = 10^3$, Fig. 11(c), $A_{\text{fin}} \approx 10$]. Then, in approaching the liquid limit at larger $s$, the translation mode loses all its impact and the volume mode again dominates coarsening. In contrast to the rigid limit, now no drop migration occurs [$s = 10^4$, Fig. 11(d)], and $A_{\text{fin}}$ approaches $A_{\text{max}}$. The described intricate change from volume transfer to translation mode and back can best be appreciated in Fig. 12(b). Independently, of the non-monotonic change in mode dominance, Fig. 12(a) clearly indicates coarsening slows down with increasing softness.

To gain a deeper understanding of the transition between modes, we perform a linear stability analysis for the alternative constellation of a periodic drop array. Namely, two identical drops of volume $V$ are placed in a domain of finite size $D = 200$ at respective positions $x = \pm D/4$. The corresponding unstable periodic steady two-drop state and its linear stability are continued (using PDE2path). Starting at $s = 10^{-3}$ in the rigid limit, the softness is increased up to $s = 10^3$ in the liquid limit. The resulting real eigenvalues corresponding to volume transfer mode (the changes in film height profile are antisymmetric w.r.t. the center line between drops) and translation mode (symmetric changes in film height profile) of coarsening are presented in Fig. 13 for three selected $V$. The blue curves indicate the volume transfer mode, while red curves correspond to the translation mode. The volume mode is strongest in the rigid limit and decreases nearly monotonically with increasing softness (up to the small dip after the strong step-like decrease at intermediate softness $s = 10^3 \ldots 10^5$). It further slightly decreases overall with increasing drop volume, a change which is more pronounced in the rigid limit.

By contrast, the translation mode is nearly absent apart from intermediate softness where it develops a peak that becomes more pronounced with increasing volume. In consequence, the translation mode eventually dominates the volume transfer mode for sufficiently large drops at intermediate $s$ in accordance with our time simulations in Fig. 12(b). The dominance of the translation mode in the range of intermediate softness approximately coincides with the intermediate regime for the “double transition” of static drops discussed in Section 3, compare in particular, Fig. 6 at $V = 100$ to 1000 with Fig. 12(b) and 13. It is in this regime that a prominent wetting ridge occurs (cf. Fig. 4(c)), through which the droplets interact via substrate deformations. Hence, we indeed conclude that the dominant translation mode is a direct consequence of the Cheerios effect.

This is further confirmed by stability calculations as in Fig. 13 for other domain sizes $D$ (not shown). We find that the critical drop volume $V_c$ where the translation mode first becomes dominant at some critical softness $s_c$ increases with increasing $D$. The corresponding $s_c$ always nearly coincides with the left edge of the $s$-range where the second transition occurs in Section 3. This agrees with the observed proportionality $\sqrt{V_c} \sim \sqrt{s_c} \sim \xi_c \sim A_c$ where $A_c$ is the critical distance of the contact lines of the two drops.

### 5.2 Case of drop arrays

Finally, we show that the developed model is also capable of describing the coarsening dynamics of large drop ensembles. Starting with a flat film of height $h_0 = 3$ on a rather large domain ($D = 5000$), a small added noise initiates the early spinodal dewetting process. It produces a large ensemble of about 100–200 droplets which then undergo coarsening.

Fig. 14 gives an exemplary space–time plot showing part of the domain and process for an ensemble of initially $\approx 200$ drops on a substrate of intermediate softness ($s = 10^3$). The subsequent Fig. 15 shows space–time plots of the positions of all drop maxima for the same case of intermediate softness and, additionally, for the rigid and the liquid limit at $s = 10^{-1}$ and $s = 10^4$, respectively. The corresponding dependencies of drop number on time are presented in Fig. 16.

Inspecting Fig. 15(b) (also cf. the magnification in Fig. 14) shows that at intermediate softness nearly all coarsening events at early times occur by drop translation, i.e., two lines meet and...
latter, are, however, much weaker than the two-drop modes. Once coarsening starts at the end of the plateau, the drop number continuously decreases. Due to the discrete measure of the number of drops, individual steps in the curves are often visible and become prominent at late times when few drops remain. Qualitatively, the substrate softness does not seem to have an effect on long-time coarsening as the three curves are very similar. Empirically, a power law fit \( N(t) \sim t^{-2} \) gives approximately identical scaling exponents for all values of \( s \) studied in Fig. 16. Based on the present low statistics one might even be tempted to distinguish two power laws. One in the range from \( t \approx 10^4 \) to \( t \approx 10^6 \) with \( x \approx 0.275 \) and one in the range from \( t \approx 10^6 \) to \( t \approx 10^8 \) with \( x \approx 0.4 \).

Finally, we briefly discuss how the various contributions to the total energy depend on time. Fig. 17(a) gives the mean interface energy density \( \mathcal{F}_{\text{cap}} + \mathcal{F}_{\text{wet}} / D \) encompassing wetting and capillary contributions [see eqn (3) and (4)]. It monotonically decreases with time and decreases with increasing substrate softness. Fig. 17(b) gives the mean elastic energy density \( \mathcal{F}_{\text{el}} / D \) [eqn (6)]. It is negligible in the rigid limit, small and nonmonotonic in the liquid limit, and monotonically decreases during the coarsening process (i.e., after the plateau) at intermediate softness. Even at intermediate softness, where elasticity is most prominent, the elastic energy is much smaller than the interface energy. This indicates that coarsening is dominated by interface effects; this might offer an explanation why the coarsening exponent is not sensitive to the substrate softness.

6 Summary and outlook

We have developed a qualitative long-wave model that captures main features of the statics and dynamics of a liquid drop on an elastic substrate. The transparency and simplicity of the model rests in its gradient dynamics structure and the underlying free

Fig. 15 Space–time plots of the positions of all drop maxima on a logarithmic time scale for (a) the rigid limit at \( s = 10^{-1} \) (fewer initial drops and coarsening mainly via volume transfer, slight migrations are visible), (b) intermediate softness at \( s = 10^{-2} \) (more initial drops, translation mode first dominates before the volume transfer mode prevails at larger drop size and distance), and (c) the soft limit at \( s = 10^4 \) (further increased initial drop number, dominating volume mode, no visible drop migration). The domain size is \( D = 5000 \), i.e., only a part is shown. The remaining parameters are as in Fig. 7.

Fig. 16 Dependence of drop number on time on a doubly logarithmic scale for softness \( s \) as given in the legend. The initial stage of drop formation (dewetting) is finished at about \( t = 10^3 \) (indicated by the vertical line). Coarsening sets in shortly after when the respective plateaus end. Empirically, one may distinguish two coarsening regimes with slightly different scaling laws as indicated by the short solid straight lines.

** Note that during the dewetting phase (not shown in Fig. 16) our counting routine gives the number of maxima in the film profile, i.e., a mean wave number.
energy that reflects capillarity, wettability and compressional elasticity. The model has first been employed to investigate the double transition that occurs in the equilibrium contact angles when increasing the softness from an ideally rigid substrate towards a very soft, i.e., liquid substrate.\(^{22}\) Both, the dependency on softness and the scaling of the two transitions with drop volume given by our model, closely resemble the behaviour observed in ref. 22 with a full elasticity model for drops of contact angle \(\pi/2\) on thick incompressible elastic layers. In particular, the scaling of the typical softnesses where the two transitions occur with drop volume and the scalings of the wetting ridge height with softness in the different regimes agree very well. This very promising qualitative agreement should in the future be quantified by numerical models based on fully nonlinear elastic theory, in the limit of small contact angles; this would enable to determine effective parameters for the present long-wave model in terms of experimental parameters. Alternatively, one can attempt to include a more realistic elastic energy into the long-wave model by adopting the more intricate Green’s functions described in the Appendix. Of specific interest would be to consider how the dissipation in the substrate is dominant and increases with softness, i.e. that viscoelastic braking is present.\(^{12,13}\) Beyond that, the simplicity of our model allowed us to also investigate the case of completely wetting liquids. We find that viscoelastic braking is present in the completely wetting case as well, in contradiction to ref. 30. However, this effect is rather small and such is the shift between the curves for dynamic contact angles (Fig. 10) due to the less distinctive wetting ridge.

Third, we have considered the coarsening dynamics of a pair of drops and of large ensembles of drops. In the case of two drops we have found that the dominant coarsening mode nonmonotonically changes when increasing the substrate softness. In both, the rigid and the soft limits, the volume transfer mode of coarsening (mass transfer mode, collapse mode, diffusion-controlled ripening, or Ostwald ripening) dominates while at intermediate softness the translation mode of coarsening (collision, coalescence, migration mode) dominates for sufficiently large drops. As such, the model recovers the “inverted Cheerios effect”, describing interaction of liquid drops mediated by elastic deformation of the solid substrate\(^{29}\) (in contrast to the Cheerios effect describing interaction of solid particles on a liquid substrate). In the rigid limit the volume transfer mode is accompanied by migration of both drops into the direction of the smaller one in accordance with literature.\(^{68,93}\) Although, the nonmonotonic change in mode dominance has mainly been investigated by time simulations, it has furthermore be confirmed through a linear stability analysis for a closely related unstable steady two-drop constellation.

Based on the results on two-drop coarsening, finally, we have considered large drop ensembles and found that the change in dominance of the coarsening mechanisms can be found there as well. However, the strength of the translation mode decreases faster with increasing separation of the drops than the strength of the volume mode does. Hence, the translation dominated regime ends at a critical value as the drop number decreases and the mean distance between the drops becomes too large. As the initial drop contribution emerges from a dewetting film the number of drops at the beginning of the coarsening process is not controlled. Indeed the initial drop number was found to increase with increasing softness, which might be reasoned by the reduced width of drops that (partially) sink into the substrate. It will be interesting to apply the model in the future to investigate the coarsening of large ensembles of drops on a two-dimensional substrate.

Beside of direct extensions of the presently studied examples, the presented model can be directly applied to a number of different situations, e.g., to study the sliding of individual drops under lateral driving, e.g., on an incline. Further, the model lends itself easily to the investigation of moving contact lines on elastic substrates, e.g., in a Landau–Levich geometry.\(^{101,102}\) This allows
one to investigate the appearance of stick-slip cycles as empirically observed in ref. 9, 36 and 103. In its present form the model can be employed as an element of more complex models to study the interaction of complex fluids with soft substrates. For instance, it may be combined with thin-film models of active media, or biofilms to investigate the motion of active drops on soft substrates and the growth of biofilms and cell aggregates on such substrates.

Finally, we point out that the presented qualitative model may be expanded in a number of ways. One can explicitly introduce the effect of finite thickness on sliding, which also enables one to study phenomena such as durotaxis, and to systematically account for other rheologies. Furthermore, it will be interesting to see how the approach can be expanded to incorporate a description of the Shuttleworth effect, to account for substrate stretching, bending elasticity, or to systematically account for other rheologies and incompressibility of the elastic layer. The latter should then allow one to capture drop-drop repulsion and attraction via the inverted Cheerios effect not only the attractive case as the present model.

The data that support the findings of this study are openly available.

Conflicts of interest

There are no conflicts to declare.

Appendix A: elastic energy per area in the framework of linear elasticity

In this Appendix we discuss how elastic energy functionals can in principle be derived from the theory of linear elasticity. The central objects in elasticity theory are the displacement vectors \( \mathbf{u} \) and the Green’s function \( G \). When varying the displacement at the free surface, one performs a mechanical work that is equal to the stored elastic energy. Hence, the energy can in principle be expressed as a functional of \( \mathbf{u} \) at the free surface. In the case of linear elasticity, this energy must be of quadratic form,

\[
\mathcal{F}_d = \frac{1}{2} G \int d^2x' d^3x' \mathbf{u}(x) \cdot K(x - x') \cdot \mathbf{u}(x').
\]  

In this expression \( G \) is the elastic (shear) modulus while \( K \) is a Green’s function for an elastic layer that we assume to be homogeneous.

We will now restrict ourselves to the case where the interface is shear free, and use that for small displacements \( u_z(x) \approx h(x) \). Then, the elastic energy can be inferred from the interface shape

\[
\mathcal{F}_d = \frac{1}{2} G \int d^2x \int d^2x' K(x - x') h(x) h(x'),
\]  

where now \( K \) is a scalar Green’s function, relating normal displacement to normal stress. Explicit calculation of the Green’s function requires solving the bulk-elastic problem inside the layer. In case the problem is invariant along \( y \), the Green’s function is known analytically in the form of a Fourier transform:

\[
\tilde{K}(q)^{-1} = \frac{(1 - \nu)}{q} \frac{\sinh(2qd) - 2qd}{\cosh(2qd) + 2(qd)^2 + 5 - 12\nu + 8\nu^2}.
\]  

Here it is given as an inverse of the Green’s function, which maps the traction to the displacement (while \( \tilde{K}(q) \) maps the displacement to the traction). It is of interest to consider some limits. In the short-wave limit \( (qd \gg 1) \), we find

\[
\tilde{K}(q)^{-1} = \frac{1 - \nu}{|q|}.
\]  

In real-space this inverse Green’s function is a logarithm (i.e. the surface displacement \( h(x) \) due to a point force is logarithmic on a half-space). In the long-wave limit \( (qd \ll 1) \), the limiting behavior depends on the Poisson ratio. For \( \nu \neq 1/2 \), we find

\[
\tilde{K}(q)^{-1} = \frac{d(1 - 2\nu)}{2(1 - \nu)}.
\]  

which is independent of \( q \). This implies that the real-space Green’s function is a \( \delta \) function, i.e.

\[
K(x) = \frac{2(1 - \nu)}{d(1 - 2\nu)} \delta(x).
\]  

We notice from this expression that there is a problem at \( \nu = 1/2 \), so that an incompressible layer cannot deform under long-wave tractions. This property is a consequence of \( \tilde{K}(q = 0)^{-1} = 0 \). Therefore, the long-wave expansion for incompressible media needs to go to the next order, to yield

\[
\tilde{K}(q)^{-1} = \frac{d^2}{3\nu},
\]  

which in real space implies

\[
h(x) = \frac{d^3}{3G} \frac{\partial^2 p}{\partial x^2}.
\]  

Note the resemblance with the viscous thin-film response \( \frac{1}{3} \frac{\partial^2 p}{\partial x^2} \): the Green’s function in elasticity plays the role of the mobility in viscous fluids.

We thus conclude that the assumption of a Green’s function that is a \( \delta \) function is rigorous in a specific limit: the long-wave limit \( qd \ll 1 \) for compressible layers (\( \nu \neq 1/2 \)). In that case, the elastic energy reads

\[
\mathcal{F}_d = \frac{1}{2} G \int d^2x h(x)^2,
\]  

which is identical to eqn (6) in the main text. In this specific limit, we recover the connection

\[
\kappa_v = \frac{2G(1 - \nu)}{d(1 - 2\nu)}.
\]  

Notice once again that the formulation does not apply for incompressible layers (\( \nu = 1/2 \)). Another implicit assumption here is that we ignored any horizontal displacements that can be caused by shear stress induced by the fluid. In the
considered long-wave limit, however, one can argue that shear stress is asymptotically small compared to the pressure.

Acknowledgements
UT and JS acknowledges support by the Deutsche Forschungsgemeinschaft (DFG) via Grants TH781/12 and SN145/1-1, respectively, within SPP 2171. We are grateful to Stefan Karpitschka and Johannes Kemper for discussions and calculations, respectively, within SPP 2171. We are grateful to Stefan Hartmann and E. H. van Brummelen for discussions with Linda Cummings, Lou Kondic and Simon Hartmann are acknowledged.

References
1 P.-G. de Gennes, F. Brochard-Wyart and D. Quéré, Capillarity and wetting phenomena: Drops, bubbles, pearls, waves, Springer, New York, 2004.
2 V. M. Starov, M. G. Velarde and C. J. Radke, Wetting and spreading dynamics, Taylor and Francis, Boca Raton, 2007, vol. 138, p. 544.
3 E. Bormashenko, Physics of Wetting: Phenomena and Applications of Fluids on Surfaces, De Gruyter, Berlin/Boston, 2017.
4 P.-G. de Gennes, Rev. Mod. Phys., 1985, 57, 827–863.
5 D. Bonn, J. Eggers, J. de Inekeu, J. Meunier and E. Rolley, Rev. Mod. Phys., 2009, 81, 739–805.
6 B. Andreotti and J. H. Snoeijer, Annu. Rev. Fluid Mech., 2020, 52, 285–308.
7 A. Pototsky, M. Bestenhorn, D. Merkt and U. Thiele, J. Chem. Phys., 2005, 122, 224711.
8 S. Bommer, F. Cartellieri, S. Jachalski, D. Peschka, R. Seemann and B. Wagner, Eur. Phys. J. E: Soft Matter Biol. Phys., 2013, 36, 87.
9 T. Kajiyama, A. Daerr, T. Narita, L. Royon, F. Lequeux and L. Limat, Soft Matter, 2013, 9, 454–461.
10 F. Schellenberger, J. Xie, N. Encinas, A. Hardy, M. Klapper, P. Papadopoulos, H. J. Butt and D. Vollmer, Soft Matter, 2015, 11, 7617–7626.
11 A. Keiser, L. Keiser, C. Clanet and D. Quere, Soft Matter, 2017, 13, 6981–6987.
12 A. Carré, J. C. Gastel and M. E. R. Shanahan, Nature, 1996, 379, 432–434.
13 A. Carré and M. E. R. Shanahan, Langmuir, 2001, 17, 2982–2985.
14 R. W. Style, A. Jagota, C. Y. Hui and E. R. Dufresne, Annu. Rev. Condens. Matter Phys., 2017, 8, 99–118.
15 C. W. Extrand and Y. Kumagai, J. Colloid Interface Sci., 1996, 184, 191–200.
16 E. R. Jerison, Y. Xu, L. A. Wilen and E. R. Dufresne, Phys. Rev. Lett., 2011, 106, 186103.
17 J. B. Bostwick, M. Shearer and K. E. Daniels, Soft Matter, 2014, 10, 7361–7369.
18 S. J. Park, B. M. Weon, J. S. Lee, J. Lee, J. Kim and J. H. Je, Nat. Commun., 2014, 5, 4369.
19 R. W. Style and E. R. Dufresne, Soft Matter, 2012, 8, 7177–7184.
20 A. Marchand, S. Das, J. H. Snoeijer and B. Andreotti, Phys. Rev. Lett., 2012, 108, 094301.
21 L. Limat, Eur. Phys. J. E: Soft Matter Biol. Phys., 2012, 35, 134.
22 L. A. Lubbers, J. H. Weijis, L. Botto, S. Das, B. Andreotti and J. H. Snoeijer, J. Fluid Mech., 2014, 747, R1.
23 B. Andreotti and J. H. Snoeijer, Europhys. Lett., 2016, 113, 66001.
24 Q. Xu, K. E. Jensen, R. Boltyanskiy, R. Sarfati, R. W. Style and E. R. Dufresne, Nat. Commun., 2017, 8, 555.
25 Q. Xu, R. W. Style and E. R. Dufresne, Soft Matter, 2018, 14, 916–920.
26 R. D. Schulman, M. Trejo, T. Salez, E. Raphael and K. Dalnoki-Veress, Nat. Commun., 2018, 9, 982.
27 A. Pandey, B. Andreotti, S. Karpitschka, G. J. van Zwieten, E. H. van Brummelen and J. H. Snoeijer, Phys. Rev. X, 2020, 10, 031067.
28 S. Karpitschka, A. Pandey, L. A. Lubbers, J. H. Weijis, L. Botto, S. Das, B. Andreotti and J. H. Snoeijer, Proc. Natl. Acad. Sci. U. S. A., 2016, 113, 7403–7407.
29 D. Long, A. Ajdari and L. Leibler, Langmuir, 1996, 12, 5221–5230.
30 V. Charlatatos and S. Kumar, Soft Matter, 2020, 16, 8284–8298.
31 S. Karpitschka, S. Das, M. van Gorcum, H. Perrin, B. Andreotti and J. H. Snoeijer, Nat. Commun., 2015, 6, 7891.
32 M. van Gorcum, S. Karpitschka, B. Andreotti and J. H. Snoeijer, Soft Matter, 2020, 16, 1306–1322.
33 M. H. Zhao, J. Dervaux, T. Narita, F. Lequeux, L. Limat and M. Roche, Proc. Natl. Acad. Sci. U. S. A., 2018, 115, 1748–1753.
34 K. Smith-Mannschott, Q. Xu, S. Heyden, N. Bain, J. H. Snoeijer, E. R. Dufresne and R. W. Style, Phys. Rev. Lett., 2021, 126, 158004.
35 A. Hourlier-Fargette, A. Antkowiak, A. Chateauoinois and S. Neukirch, Soft Matter, 2017, 13, 3484–3491.
36 M. van Gorcum, B. Andreotti, J. H. Snoeijer and S. Karpitschka, Phys. Rev. Lett., 2018, 121, 208003.
37 H. B. Wu, Z. Z. Liu, A. Jagota and C. Y. Hui, Soft Matter, 2018, 14, 1847–1855.
38 V. M. Starov and M. G. Velarde, J. Phys. Condens. Matter, 2009, 21, 464121.
39 L. R. White, J. Colloid Interface Sci., 2003, 258, 82–96.
40 S. Aland and D. Mokbel, Int. J. Numer. Methods Eng., 2021, 122, 903–918.
41 S. Aland and P. Auerbach, Int. J. Numer. Methods Eng., 2021, 122, 4114–4128.
42 M. S. Roudbari and E. H. van Brummelen, Math. Models Meth. Appl. Sci., 2019, 29, 995–1036.
43 E. H. van Brummelen, M. S. Roudbari, G. Simsek and K. G. van der Zee, in Fluid-Structure Interaction, ed. S. Frei, B. Holm, T. Richter, T. Wick and H. Yang, De Gruyter, Berlin, 2017, pp. 283–328.
44 J. Bueno, Y. Bazilevs, R. Juanes and H. Gomez, Extreme Mech. Lett., 2017, 13, 10–16.
45 J. Bueno, Y. Bazilevs, R. Juanes and H. Gomez, Soft Matter, 2018, 14, 1417–1426.
46 O. K. Matar, V. Gkanis and S. Kumar, J. Colloid Interface Sci., 2005, 286, 319–332.
47 M. Gielok, M. Lopes, E. Bonaccorso and T. Gambaryan-Roisman, Colloids Surf., A, 2017, 521, 13–21.
48 H. Gomez and M. Velay-Lizancos, Eur. Phys. J.: Spec. Top., 2020, 229, 265–273.
49 U. Thiele and S. Hartmann, Eur. Phys. J.: Spec. Top., 2020, 229, 1819–1832.
50 D. N. Sibley, P. Llombart, E. G. Noya, A. J. Archer and L. G. MacDowell, Nat. Commun., 2021, 12, 239.
51 J. H. Weij, B. Andreotti and J. H. Snoeijer, Soft Matter, 2013, 9, 8494–8503.
52 H. Y. Liang, Z. Cao, Z. L. Wang and A. V. Dobrynin, Langmuir, 2018, 34, 7497–7502.
53 U. Thiele, A. J. Archer and L. M. Pismen, Phys. Rev. Fluids, 2016, 1, 083903.
54 U. Thiele, Colloids Surf., A, 2018, 553, 487–495.
55 M. Wilczek, W. Tewes, S. Engelknemper, S. V. Gurevich and U. Thiele, Phys. Rev. Lett., 2017, 119, 204501.
56 S. Engelknemper and U. Thiele, Europhys. Lett., 2019, 127, 54002.
57 P. Taylor, Adv. Colloid Interface Sci., 1998, 75, 107–163.
58 R. D. Vengrovich, Y. V. Gudyma and S. V. Yarema, Semiconductors, 2001, 35, 1378–1382.
59 T. Kinkel, J. Nordmann, A. N. Raj and M. Haase, Nanoscale, 2014, 6, 14523–14530.
60 C. Andrieu, D. A. Beysens, V. S. Nikolayev and Y. Pomeau, J. Fluid Mech., 2003, 453, 427–438.
61 R. Limary and P. F. Green, Langmuir, 2003, 19, 2419–2424.
62 J. Blaschke, T. Lapp, B. Hof and J. Vollmer, Phys. Rev. Lett., 2012, 109, 068701.
63 D. Baratian, R. Dey, H. Hoek, D. van den Ende and F. Mugele, Phys. Rev. Lett., 2018, 120, 214502.
64 M. B. Gratton and T. P. Witelski, Physica D, 2009, 238, 2380–2394.
65 G. Klasner, F. Otto, T. Rump and D. Slepecev, Eur. J. Appl. Math., 2009, 20, 1–67.
66 G. Kitavtsev, Eur. J. Appl. Math., 2014, 25, 83–115.
67 T. P. Witelski, AIMs Math., 2020, 5, 4229–4259.
68 K. B. Glaser and T. P. Witelski, Phys. D, 2005, 209, 80–104.
69 A. Oron, S. H. Davis and S. G. Bankoff, Rev. Mod. Phys., 1997, 69, 931–980.
70 R. V. Craster and O. K. Matar, Rev. Mod. Phys., 2009, 81, 1131–1198.
71 U. Thiele, J. Phys.: Condens. Matter, 2010, 22, 084019.
72 S. Das, A. Marchand, B. Andreotti and J. H. Snoeijer, Phys. Fluids, 2011, 23, 072006.
73 M. Napiorkowski, L. Schimmele and S. Dietrich, Europhys. Lett., 2020, 129, 16002.
74 M. N. Popescu, G. Oshanin, S. Dietrich and A. M. Cazabat, J. Phys.: Condens. Matter, 2012, 24, 243102.
75 K. L. Johnson, Contact Mechanics, Cambridge University Press, Cambridge, 1987.
76 J. M. Skotheim and L. Mahadevan, Phys. Rev. Lett., 2004, 92, 245509.
77 T. Salez and L. Mahadevan, J. Fluid Mech., 2015, 779, 181–196.
78 T. G. J. Chandler and D. Vella, Proc. R. Soc. A, 2020, 476, 20200551.
79 R. Masurel, M. Roché, L. Limat, I. Ionescu and J. Dervalux, Phys. Rev. Lett., 2019, 122, 248004.
80 Numerical Continuation Methods for Dynamical Systems, ed. B. Krauskopf, H. M. Osinga and J. Galan-Vioque, Springer, Dordrecht, 2007.
81 H. A. Dijkstra, F. W. Wubs, A. K. Cliffe, E. Doedel, I. F. Dragomirescu, B. Eckhardt, A. Y. Gelfgat, A. Hazel, V. Lucarini, A. G. Salinger, E. T. Phipps, J. Sanchez-Umbria, H. Schutteelaars, L. S. Tuckerman and U. Thiele, Commun. Comput. Phys., 2014, 15, 1–45.
82 S. Engelknemper, S. Gurevich, H. Uecker, D. Wetzel and U. Thiele, in Computational Modeling of Bifurcations and Instabilities in Fluid Mechanics, ed. A. Gelfgat, Springer, 2019, pp. 459–501.
83 E. Doedel, H. B. Keller and J. P. Kernevez, Int. J. Bifurcat. Chaos, 1991, 1, 745–772.
84 H. Uecker, D. Wetzel and J. D. M. Rademacher, Numer. Math.-Theory Methods Appl., 2014, 7, 58–106.
85 S. Engelknemper, M. Wilczek, S. V. Gurevich and U. Thiele, Phys. Rev. Fluids, 2016, 1, 073901.
86 S. Trinschek, F. Stegemerten, K. John and U. Thiele, Phys. Rev. E, 2020, 101, 062802.
87 U. Thiele, Zeno, 2021, DOI: 10.5281/zenodo.4544848.
88 M. Heil and A. L. Hazel, in Fluid-Structure Interaction: Modelling, Simulation, Optimisation, ed. H.-J. Bungartz and M. Schäfer, Springer, Berlin, Heidelberg, 2006, pp. 19–49.
89 E. B. Dussan, Ann. Rev. Fluid Mech., 1979, 11, 371–400.
90 U. Thiele, M. G. Velarde, K. Neuffer, M. Bestehorn and Y. Pomeau, Phys. Rev. E, 2001, 64, 061601.
91 L. H. Tanner, J. Phys. D, 1979, 12, 1473–1484.
92 D. Fritter, C. M. Knobler and D. A. Beysens, Phys. Rev. A, 1991, 43, 2858–2869.
93 L. M. Pismen and Y. Pomeau, Phys. Fluids, 2004, 16, 2604–2612.
94 S. B. Dai, Nonlinearity, 2010, 23, 325–340.
95 U. Thiele, in Thin Films of Soft Matter, ed. S. Kalliadasis and U. Thiele, Springer Vienna, Vienna, 2007, pp. 25–93.
96 U. Thiele, L. Brusch, M. Bestehorn and M. Bär, Eur. Phys. J. E: Soft Matter Biol. Phys., 2003, 11, 255–271.
97 A. Münch, B. Wagner and T. P. Witelski, J. Eng. Math., 2005, 53, 359–383.
98 K. B. Glaser and T. P. Witelski, Phys. Rev. E, 2003, 67, 016302.
99 K. B. Glaser, SIAM J. Appl. Math., 2008, 69, 473–493.
100 U. Thiele, M. G. Velarde and K. Neuffer, Phys. Rev. Lett., 2001, 87, 016104.
101 J. H. Snoeijer, B. Andreotti, G. Delon and M. Fermigier, J. Fluid Mech., 2007, 579, 63–83.
102 M. Galvano, D. Tseluiko, H. Lopez and U. Thiele, Phys. Rev. Lett., 2014, 112, 137803.
103 T. Kajiy, P. Brunet, L. Royon, A. Daerr, M. Receveur and L. Limat, Soft Matter, 2014, 10, 8888–8895.
104 A. Loisy, J. Eggers and T. B. Liverpool, Soft Matter, 2020, 16, 3106–3124.

105 S. Trinschek, K. John, S. Lecuyer and U. Thiele, Phys. Rev. Lett., 2017, 119, 078003.

106 J. Lober, F. Ziebert and I. S. Aranson, Soft Matter, 2014, 10, 1365–1373.

107 S. Douezan, J. Dumond and F. Brochard-Wyart, Soft Matter, 2012, 8, 4578–4583.

108 A. Cont, T. Rossy, Z. Al-Mayyah and A. Persat, eLife, 2020, 9, e56533.

109 R. W. Style, Y. Che, S. J. Park, B. M. Weon, J. H. Je, C. Hyland, G. K. German, M. P. Power, L. A. Wilen, J. S. Wetland and E. R. Dufresne, Proc. Natl. Acad. Sci. U. S. A., 2013, 110, 12541–12544.

110 C. Henkel, J. Snoeijer and U. Thiele, Data Supplement for “Gradient-dynamics model for liquid drops on elastic substrates”, 2021, DOI: 10.5281/zenodo.5607074.

111 M. Hannah, Q. J. Mech. Appl. Math., 1951, 4, 94–105.

112 M. H. Essink, A. Pandey, S. Karpitskha, C. H. Venner and J. H. Snoeijer, J. Fluid Mech., 2021, 915, A49.