Finite element simulation of dynamic wetting flows as an interface formation process

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**Abstract**

A mathematically challenging model of dynamic wetting as a process of interface formation has been, for the first time, fully incorporated into a numerical code based on the finite element method and applied, as a test case, to the problem of capillary rise. The motivation for this work comes from the fact that, as discovered experimentally more than a decade ago, the key variable in dynamic wetting flows — the dynamic contact angle — depends not just on the velocity of the three-phase contact line but on the entire flow field/geometry. Hence, to describe this effect, it becomes necessary to use the mathematical model that has this dependence as its integral part. A new physical effect, termed the ‘hydrodynamic resist to dynamic wetting’, is discovered where the influence of the capillary’s radius on the dynamic contact angle, and hence on the global flow, is computed. The capabilities of the numerical framework are then demonstrated by comparing the results to experiments on the unsteady capillary rise, where excellent agreement is obtained. Practical recommendations on the spatial resolution required by the numerical scheme for a given set of non-dimensional similarity parameters are provided, and a comparison to asymptotic results available in limiting cases confirms that the code is converging to the correct solution. The appendix gives a user-friendly step-by-step guide specifying the entire implementation and allowing the reader to easily reproduce all presented results, including the benchmark calculations.

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1. Introduction

Reliable simulation of flows in which a liquid advances over a solid, known as dynamic wetting flows, is the key to the understanding of a whole host of natural phenomena and technological processes. In the technological context, the study of these flows has often been motivated by the need to optimize continuous coating processes that are routinely used to create thin liquid films on a product [1], for example, in the coating of optical fibres [2,3]. However, more recently, discrete coating, in particular inkjet printing of microdrops [4], has matured into a viable, and often preferable, alternative to traditional fabrication processes, e.g. in the additive manufacturing of 3D structures or the creation of P-OLED displays [5,6], and it is becoming a new driving force behind research into dynamic wetting phenomena. In most cases, such flows can be regarded as microfluidic phenomena, where a large surface-to-volume ratio brings in interfacial effects on the flow that are not observed at larger scales.

Obtaining accurate information about micro and nanofluidic flows experimentally is often difficult and usually costly so that, consequently, a desired alternative is to have a reliable theory describing the physics that is dominant for this class of...
flows and incorporate it into a flexible and robust computational tool which can quickly map the parameter space of interest to allow a specific process to be optimized. Such computational software could be validated against experiments at scales and geometries easily accessible to accurate measurement and then used to make predictions in processes inaccessible to experimental analysis.

The discovery that no solution exists for the moving contact-line problem in the framework of standard fluid mechanics [7,8] prompted a number of remedies to be proposed, which are summarized, for example, in [9, ch. 3]. Modelling approaches range from continuum theories through to mesoscopic approaches [10] right down to molecular dynamics simulations [11], see [12,13] for discussions of the various modelling approaches. Of these, most, based on continuum mechanics, are what we shall refer to as 'conventional' or 'slip' models, in which the no-slip condition on the solid surface is relaxed to allow a solution to exist, with the Navier-slip condition [14] being the most popular choice.

As a boundary condition on the free-surface shape at the contact line, one has to specify the contact angle formed between the free surface and solid. In conventional models, this angle is prescribed as a heuristic or empirical function of the contact-line speed and material parameters of the system, e.g. [15]. A variant of this approach is to claim that the actual contact angle is static, whilst the observed 'apparent' angle is produced as a result of viscous bending of the free surface near the contact line [16]. Such models provide predictions that adequately describe experiments at relatively large scales, often around the length scale of millimetres. It is well established that on such scales many of the proposed models work equally well and that finding significant deviations between their predictions, and hence ascertaining which best captures the key physical mechanisms of dynamic wetting, is practically impossible [17,18].

A physical phenomenon that gives an opportunity to distinguish between different models came to be known under a 'technological' name of the 'hydrodynamic assist of dynamic wetting' (henceforth 'hydrodynamic assist' or simply 'assist'). The essence of this effect, first observed in high accuracy experiments on the curtain coating process [19,20], is that for a given liquid spreading over a given solid at a fixed contact-line speed, the dynamic contact angle can still be manipulated by altering the flow field/geometry, for example, in the case of curtain coating, by changing the flow rate or the height from which the curtain falls. This effect has profound technological implications as it allows the process to be optimized by offsetting the increase of the contact angle with increasing contact-line speed by manipulating the flow conditions and hence postponing air entrainment [19].

The dependence of the dynamic contact angle on the flow field has also been reported in the imbibition of liquid into capillaries [21,22], in the spreading of impacted drops over solid substrates [23,24] and in the coating of fibres [3]. However, in many of these flows it is yet unclear whether hydrodynamic assist actually occurs, or whether the free surface bends significantly beneath the spatial resolution of the experiments, whereas for curtain coating the hope of attributing assist to the poor spatial resolution of experiments has been quashed by careful finite element simulations which show that the degree of free-surface bending under the reported resolution of the measurements is too small to account for the observed effect and that conventional models cannot in principle describe this important physical effect [25].

Currently, the only model known to be able to even qualitatively describe assist [26,27] is the model of dynamic wetting as an interface formation process, first introduced in [28] and discussed in detail in [9]. This model is based on the simple physical idea that dynamic wetting, as the very name suggests, is the process in which a fresh liquid–solid interface (a newly ‘wetted’ solid surface) forms. Qualitatively, the origin of the hydrodynamic assist is that the global flow influences the dynamics of the relaxation of the forming liquid–solid interface towards its equilibrium state and hence the value of this interface’s surface tension at the contact line, which, together with the surface tension on the free surface, determines the value of the dynamic contact angle. When there is a separation of scales between the interface formation process and the global flow, the ‘moving contact-line problem’ can be considered locally and asymptotic analysis provides a speed-angle relationship which is seen to describe experiments just as well as formulae proposed in other models [9]. However, in the situation where the scale of the global flow and that of the interface formation are no longer separated, the influence of the flow field on the dynamic contact angle will occur and hence no unique speed-angle relationship will be able to describe experiments. Then, the interface formation model becomes the only modelling tool, and, given that the processes of practical interest are free-surface flows under the influence of, at least, viscosity, capillarity and inertia, it is inevitable that, to describe such flows, one needs computer simulation, i.e. the development of accurate CFD codes, which, for the effect of hydrodynamic assist to be captured, have to incorporate the interface formation model.

Since the interface formation model came into wider circulation, there has been considerable interest, e.g. [29,23,24], in using it in its entirety as a practical tool for describing dynamic wetting phenomena, especially on the microscale. Review articles have also referred to the description of assist using this model as one of the main challenges in the field [12,13]. Although robust computational codes based on the conventional models already exist, e.g. [30,31], the extension of these codes to a numerical tool that incorporates the interface formation model is far from trivial. In particular, one has to solve numerically the Navier–Stokes equations describing the bulk flow subject to boundary conditions which are themselves partial differential equations along the interfaces and in their turn have to satisfy certain boundary conditions at contact lines confining the interfaces. These conditions determine the dynamic contact angle and hence influence the free-surface shape, which exerts its influence back on the bulk flow. Thus, the bulk flow, the distribution of the surface parameters along the interfaces and the dynamic contact angle that these distributions ‘negotiate’ become interdependent, with the dynamic contact angle being an outcome of the solution as opposed to conventional models where it is an input. This interdependency is, on the one hand, the physical essence of the experimentally observed effect of hydrodynamic assist to be described but, on
the other hand, it is this very interdependency that, coupled with the nonlinearity of the bulk equations and the flow geometry, is the reason why the model is difficult to implement robustly into a numerical code.

Some numerical progress has been made for the computationally less complex steady Stokes flows [27], but what is lacking is a step-by-step guide to the implementation of this model in the general case, with unsteady effects in the bulk and interfaces as well as nonlinearity of the bulk equations fully implemented. This would pave the way for incorporating the interface formation model into existing codes as well as developing new ones. Therefore, the first objective of the present paper is to address these issues and provide a digestible guide to the model’s implementation into CFD codes. Then, after giving benchmark computational results to verify this implementation, we consider a problem of imbibition into a capillary, compare the outcome with experiments and predict essentially new physical effects.

A starting point in the development of the aforementioned CFD code is to first develop an accurate computational approach for the simulation of dynamic wetting flows using the mathematically less complex conventional models and this was achieved in [32]. It was shown that many of the previous numerical results obtained for dynamic wetting processes are unreliable as they contain uncontrolled errors caused by not resolving all the scales in the conventional model, most notably the dynamics of slip and the curvature of the free surface near the contact line. Benchmark calculations in [32] for a range of mesh resolutions resolved previous misunderstandings about how to impose the dynamic contact angle and showed that implementing it using the usual finite element ideology, as opposed to ‘strong’ implementation of the contact angle, works most efficiently: it allows errors to be seen, and hence controlled, as the computed contact angle varies from its imposed value, instead of them being masked elsewhere in the code. Furthermore, in [33], we have shown that numerical artifacts which occur at obtuse contact angles are present in both commercial software and in academic codes where, misleadingly, they have previously been interpreted as physical effects. By comparing computational results to analytic near field asymptotic ones, we have shown that the previously obtained spikes in the distributions of pressure observed near the contact angle are completely spurious, and, to rectify this issue, a special method, based on removing the ‘hidden’ eigensolutions in the problem prior to computation, has been developed [33]. In [34], we showed that our code is capable of simulating unsteady high deformation flows by comparing to benchmark calculations published in the literature and performed by various research groups. In contrast, in [35] it has been shown that when commercial software is used to simulate relatively simple benchmark free surface flows, the solution, albeit converged, is not the correct one.

In this series of articles [32–34], our approach has been to carefully develop a robust CFD algorithm for the simulation of dynamic wetting flows. Thus far, this has been achieved for the conventional models, where we validated our results by performing mesh independence tests, comparing with asymptotic solutions in limiting cases and, where no analytic progress was possible, to previously obtained benchmark solutions published in the literature. In this article, we continue this series of papers and, for the first time, incorporate the interface formation model into our code in full. Notably, it will be shown that a code implementing the mathematically complex interface formation equations can easily recover the much simpler conventional models by setting certain parameters to zero. This allows the same code to be used to compare the predictions of dynamic wetting models for a range of flows and hence to determine, by a comparison with experiment, where the physics manifests itself in a significant way and where mathematically simpler conventional models are adequate. Here, we focus on dynamic wetting; the extension to other flows of interest where interface formation or disappearance also occurs is a straightforward procedure computationally, and it will be dealt with in forthcoming articles.

The layout of the present article is as follows. First, in Section 2, without limiting ourself to a particular flow configuration, the equations describing the dynamic wetting process are briefly recapitulated. Then, in Section 3, the finite element equations are derived for the dynamic wetting flow as an interface formation process. Local element matrices, and additional details about the finite element procedure are provided in the Appendix, which complements a ‘user-friendly’ step-by-step guide to finite element simulation given for this class of flows in [32] and allows one to reproduce the benchmark simulations in Section 4. These simulations are performed for the dynamic wetting flow through a capillary both in the case where the meniscus motion is steady and for the unsteady imbibition of a liquid into an initially dry capillary. Computations are checked for convergence both as the mesh is refined and towards asymptotic results in limiting cases. Having established the accuracy of our approach, in Section 5 new physical effects are discovered by considering the influence of capillary geometry on the dynamic wetting process. Finally, the computational tool’s ability to easily describe experimental data is shown in Section 6 and a number of advantages over current approaches, particularly in the initial stages of a meniscus’ motion into a capillary tube, are highlighted. Concluding remarks and areas for future research are discussed in Section 7.

2. Modelling dynamic wetting flows as an interface formation process

Consider the spreading of a Newtonian liquid, with constant density $\rho$ and viscosity $\mu$, over a chemically homogeneous smooth solid surface. The liquid is surrounded by a gas which, for simplicity, is assumed to be inviscid and dynamically passive, of a constant pressure $p_0$. Let the flow be characterized by scales for length $L$, velocity $U$, time $L/U$, pressure $\mu U^2/L$ and external body force $F_0$. In the dimensionless form, the continuity and momentum balance equations are then given by

$$\nabla \cdot \mathbf{u} = 0, \quad Re \left[ \frac{\partial \mathbf{u}}{\partial t} + \mathbf{u} \cdot \nabla \mathbf{u} \right] = \nabla \cdot \mathbf{P} + St \mathbf{F},$$

(1) where
is the stress tensor, $t$ is time, $\mathbf{u}$ and $p$ are the liquid’s velocity and pressure. $\mathbf{F}$ is the external force density and $\mathbf{I}$ is the metric tensor of the coordinate system. The non-dimensional parameters are the Reynolds number $Re = \rho U L / \mu$ and the Stokes number $St = \rho F_0 L^2 / (\mu U)$.

Boundary conditions to the bulk equations are required at the liquid–gas free surface $\mathbf{x} = \mathbf{x}_i(s_1, s_2, t)$, whose position is to be found as part of the solution, and at the liquid–solid interface $\mathbf{x} = \mathbf{x}_s(s_1, s_2, t)$, whose position is known, and at other bounding surfaces which will be specified by the problem of interest; here, $(s_1, s_2)$ are the coordinates that parameterize the surfaces. Boundary conditions along the free surface, the liquid–solid interface and the contact line at which they intersect are provided by the interface formation model [9], as follows.

### 2.1. The interface formation model

To represent the boundary conditions on an interface with a normal $\mathbf{n}$ in an invariant form, it is convenient to introduce a (symmetric) tensor $\mathbf{I} - \mathbf{n} \otimes \mathbf{n}$, which is essentially a metric tensor on the surface. If an arbitrary vector $\mathbf{a}$ is decomposed into a scalar normal component $a_n = \mathbf{a} \cdot \mathbf{n}$ and a vector tangential part $a_a$, so that $\mathbf{a} = a_n + a_a \mathbf{n}$, we can see that, because $\mathbf{n} \cdot (\mathbf{I} - \mathbf{n} \otimes \mathbf{n}) = \mathbf{0}$, the tensor $(\mathbf{I} - \mathbf{n} \otimes \mathbf{n})$ extracts the component of a vector which is tangential to the surface, i.e. $\mathbf{a} \cdot (\mathbf{I} - \mathbf{n} \otimes \mathbf{n}) = a_a$. Hereafter, $\mathbf{n}$ is the unit normal to a surface pointing into the liquid, and subscripts 1 and 2 refer to the free surface and solid surface, respectively.

The equations of interface formation on a liquid–gas free surface, which act as boundary conditions for the bulk Eq. (1), are given by

$$
\left( \frac{\partial \mathbf{x}_1}{\partial t} - \mathbf{v}_1 \right) \cdot \mathbf{n}_1 = 0, \tag{3}
$$

$$
Ca \mathbf{n}_1 \cdot \left[ \nabla \mathbf{u} + (\nabla \mathbf{u})^T \right] \cdot (\mathbf{I} - \mathbf{n}_1 \mathbf{n}_1) + \nabla \sigma_1 = \mathbf{0}, \tag{4}
$$

$$
Ca \left\{ p_s - p + \mathbf{n}_1 \cdot \left[ \nabla \mathbf{u} + (\nabla \mathbf{u})^T \right] \right\} = \sigma_1 \nabla \cdot \mathbf{n}_1. \tag{5}
$$

$$
\mathbf{v}_{1\parallel} - \mathbf{u}_1 = \frac{1 + 4\beta^2}{4\beta^2} \nabla \sigma_1, \tag{6}
$$

$$
(\mathbf{u} - \mathbf{v}_1) \cdot \mathbf{n}_1 = Q(\rho_1 - \rho_{1e}), \tag{7}
$$

$$
\epsilon \left[ \frac{\partial \rho_1}{\partial t} + \nabla \cdot (\rho_1 \mathbf{v}_1) \right] = -(\rho_1 - \rho_{1e}), \tag{8}
$$

$$
\sigma_1 = \lambda(1 - \rho_1), \tag{9}
$$

whilst at liquid–solid interfaces formed on a solid which moves with velocity $\mathbf{U}$, the equations of interface formation have the form

$$
(\mathbf{v}_2 - \mathbf{U}) \cdot \mathbf{n}_2 = 0, \tag{10}
$$

$$
Ca \mathbf{n}_2 \cdot \mathbf{P} \cdot (\mathbf{I} - \mathbf{n}_2 \mathbf{n}_2) + \frac{1}{2} \nabla \sigma_2 = \hat{\beta}(\mathbf{u}_1 - \mathbf{U}_1), \tag{11}
$$

$$
\mathbf{v}_{2\parallel} - \frac{1}{2}(\mathbf{u}_1 + \mathbf{U}_1) = \tilde{\alpha} \nabla \sigma_2, \tag{12}
$$

$$
(\mathbf{u} - \mathbf{v}_2) \cdot \mathbf{n}_2 = Q(\rho_2 - \rho_{2e}), \tag{13}
$$

$$
\epsilon \left[ \frac{\partial \rho_2}{\partial t} + \nabla \cdot (\rho_2 \mathbf{v}_2) \right] = -(\rho_2 - \rho_{2e}), \tag{14}
$$

$$
\sigma_2 = \lambda(1 - \rho_2). \tag{15}
$$

The differential term $\sigma_1 \nabla \cdot \mathbf{n}_1$, where $\sigma_1$ is the (dimensionless) surface tension on the free surface, describing the capillary pressure in the normal-stress Eq. (5) indicates that this equation requires its own boundary condition where the free surface terminates, i.e. at the contact line. There $\mathbf{n}_2$ is known and $\mathbf{n}_1$ must be specified by setting the dynamic contact angle $\theta_d$ at which the free surface meets the solid surface:
\[ \mathbf{n}_1 \cdot \mathbf{n}_2 = -\cos \theta_d. \]  \hspace{1cm} (16)

In the framework of continuum mechanics, the mass of the contact line (i.e., physically, the three-phase-interaction region), as well as the mass of the liquid–gas interface, is neglected, and hence one has that the forces acting on the contact line are in balance irrespective of the contact line’s acceleration. The tangential to the solid surface projection of the balance of forces acting on the contact line is given by Young’s equation [36]:

\[ \sigma_2 + \sigma_1 \cos \theta_d = \sigma_s, \]  \hspace{1cm} (17)

which introduces the contact angle via a physical law and where \( \sigma_s \) and \( \sigma_2 \) are the surface tensions of the liquid–solid interface and solid-gas interface, respectively, and the latter is henceforth assumed to be negligible \( \sigma_s \approx 0 \).

Eqs. (8) and (14) also require a boundary condition at the contact line where the two interfaces meet, and here we have the continuity of surface mass flux

\[ \rho^i_1 \left( \mathbf{v}^i_1 - \mathbf{U}_c \right) \cdot \mathbf{m}_1 + \rho^s_2 \left( \mathbf{v}^s_2 - \mathbf{U}_c \right) \cdot \mathbf{m}_2 = 0 \]  \hspace{1cm} (18)

where \( \mathbf{U}_c \) is the (dimensionless) velocity of the advancing contact line and \( \mathbf{m}_i \) is the inward unit vector tangential to surface \( i \) and normal to the contact line (see Fig. 1).

The interface formation model is described in detail in the monograph [9] and a series of preceding papers [e.g. 28,37] so that here only the main ideas are briefly recapitulated. The surface variables are in the ‘surface phase’, i.e. physically in a microscopic layer of liquid adjacent to the surface which is subject to intermolecular forces from two bulk phases. In the continuum limit, this microscopic layer becomes a mathematical surface of zero thickness with

\[ q \]

denoting its surface density (mass per unit area) and \( \mathbf{v}^i \) the velocity with which it is transported. The following non-dimensional parameters have been introduced \( \alpha = \sigma \beta / (UL) \), \( \beta = (\sigma \beta UL)/\sigma \), \( \epsilon = U \tau / L \), \( \lambda = \gamma \rho^i_0 / \sigma \) and \( Q = \rho^i_0 / \rho \) which are based on phenomenological material constants \( \alpha, \beta, \gamma, \tau, \alpha \) and \( \rho^i_0 \); in the simplest variant of the theory, the latter are assumed to be constant and take the same value on all interfaces; \( \sigma \) is the characteristic surface tension for which it is convenient to take the equilibrium surface tension on the liquid–gas interface.

Estimates for the material constants have been obtained by comparing the theory to experiments in dynamic wetting, e.g. in [38], but could just as easily have been taken from an entirely different process in which interface formation is key to describing the dynamics of the flow [39–42]. In other words, once obtained for a particular liquid in one set of experiments, the material constants determined can be used to describe all phenomena involving the same fluid in which interface formation dynamics ‘kicks in’.

The surface tension \( \sigma_i \) is considered as a dynamic quantity related to the surface density \( \rho^i \) via the equations of state in the ‘surface phase’ (9) and (15), which are taken here in their simplest linear form. Gradients in surface tension influence the flow, firstly, via the stress boundary conditions (4), (5) and (11), i.e. via the Marangoni effect, and, secondly, in the Darcy-type equations\(^1\) (6) and (12) by forcing the surface velocity to deviate from that generated in the surface phase by the outer flow. Mass exchange between the bulk and surface phases, caused by the possible deviation of the surface density from its equilibrium value \( \rho^i_0 \), is accounted for in the boundary conditions for the normal component of bulk velocity (7) and (13), and in the corresponding surface mass balance Eqs. (8) and (14).

One would expect a generalized set of boundary conditions to have the classical conditions as their limiting case. For the interface formation model this limiting case follows from the double limit \( \epsilon \to 0, \beta/Ca \to \infty \). When the limit \( \epsilon \to 0 \) is applied to (3)–(9), the liquid–gas interface equations are reduced to their classical form. Notably, if we apply \( \epsilon \to 0 \) to the liquid–solid equations (10)–(15), the conventional ‘slip’ model is obtained, that is the Navier-slip condition combined with impermeability.

\(^1\) The analogy with the Darcy equation is that the tangential surface velocity \( \mathbf{v}^i_1 \) is the average velocity of the interfacial layer and its deviation from that generated by the outer flow is proportional to the gradient of surface tension, which is the negative gradient of surface pressure as \( p^i = -\sigma \).
In this limit, the surfaces are in equilibrium so Young’s equation (17) gives that the dynamic contact angle is fixed as a constant at its equilibrium value \( \theta_d = \theta_e \). If we wish to go further with the conventional model approach and describe the dynamic contact angle as some function of the contact-line speed, then Young’s equation (17), which is essentially Newton’s second law projected on the tangent to the solid surface, has to be discarded in favour of this function, as the conventional models have no mechanism that would allow the surface tensions to be dynamic and hence make the contact angle, as observed in experiments, deviate from its equilibrium value while keeping the Young equation. Therefore, implementing the interface formation model into a numerical code allows one to test all conventional models of wetting proposed in the framework of continuum mechanics. By applying the limit \( \beta/\text{Ca} = \beta L/\mu \to \infty \) we recover the classical equations of no-slip and impermeability on the solid surface for which the moving contact-line problem is known to have no solution [7,8].

Despite the model’s complexity, in limiting cases, analytic progress on it can be made to obtain explicit relationships for the surface variables and, with some further assumptions, even a formula relating the contact-line speed to the dynamic contact angle. Such formulae are a useful test of our numerical solutions, and we will briefly recapitulate their outcome.

2.2. Asymptotic formulae in a limiting case

When the contact-line motion can be analyzed as a local problem, as opposed to cases where the interface formation and the bulk flow scales are not separated so that manipulating the global flow influences the relaxation process along the interfaces, asymptotic progress is possible. A full derivation of the results we use may be found in [9] and references therein; here we shall just outline the main assumptions and results.

If in the steady propagation of a liquid–gas free surface over a solid substrate in the Stokes regime (\( \text{Re} \ll 1 \)), the characteristic length scale of the interface formation process \( L = UL \) is small compared to the bulk length scale \( L \), we have that our non-dimensional parameter \( \varepsilon \ll 1 \). If in the limit \( \varepsilon \to 0 \) we also assume that the capillary number \( \text{Ca} \to 0 \), then, to leading order in \( \text{Ca} \), the normal-stress boundary condition (5) gives that the free surface near the contact line is planar, so that the problem may be considered locally in a wedge-shaped domain. Then, we can identify the following three asymptotic regions:

(a) The outer region, where, in a reference frame moving with the contact line, one has a flow in a wedge in the classical formulation, with a zero tangential-stress and a no-slip boundary, described in [43];
(b) The intermediate region with the characteristic (dimensionless) length scale \( \varepsilon \), where the surface-tension-relaxation process takes place and where, due to smallness of \( \text{Ca} \), to leading order the influence of the bulk flow on this process can be neglected;
(c) The inner region, with the characteristic length scale \( \varepsilon \text{Ca} \), through which the surface densities and velocities, to leading order, stay constant.

On the free surface, at leading order in the intermediate and inner regions, one has

\[
\rho_1^i = \rho_1 e, \quad v_1^i = u_f(\theta_d),
\]

where \( u_f(\theta_d) \) is the (dimensionless) radial velocity of the bulk flow in the far field on the liquid–gas interface given by [43]:

\[
u_f(\theta_d) = \frac{\sin \theta_d - \theta_d \cos \theta_d}{\sin \theta_d \cos \theta_d - \theta_d},
\] (20)

so that the surface mass flux into the contact line is \( -\rho_1^i u_f(\theta_d) \). Then, since the surface variables are constant through the inner region, the boundary conditions at the contact line (17), (18) can be applied to the distributions of the surface variables in the intermediate region. As a result, we have two first-order ODEs to solve for the distributions of \( \rho_2^i \) and \( v_2^i \) along the liquid–solid interface

\[
\frac{d\rho_2^i}{ds} = 4V^2(1 - v_2^i), \quad \frac{d(v_2^i v_2^i)}{ds} = - (\rho_2^i - \rho_2^e) \quad (s > 0)
\] (21)

where \( s = s/\varepsilon \) is the intermediate region’s variable and \( V^2 = \beta e/((1 + 4\varepsilon \beta)\lambda) \) is the non-dimensional contact-line speed, subject to (a) boundary condition (18), now taking the form \( \rho_2^i v_2^i = -\rho_2^e u_f(\theta_d) \) at \( s = 0 \), (b) the matching condition \( \rho_2^i \to \rho_2^e \) as \( s \to \infty \), and, as boundary condition (a) implicitly depends on the parameter \( \theta_d \), (c) Young’s equation (17), now taking the form \( \cos \theta_d = \lambda (\rho_2^i - 1) \). The equilibrium contact angle \( \theta_d \) is obviously related to \( \rho_2^e \) by \( \cos \theta_d = \lambda (\rho_2^e - 1) \), which can be used to replace \( \rho_2^i \) with \( \theta_d \).

The above problem is easily solved numerically; however, by taking an additional assumption \( \lambda \gg 1 \), one can obtain an analytic relation between the contact angle and the non-dimensional speed of the contact line \( V \):

\[
\cos \theta_e - \cos \theta_d = \frac{2V \cos \theta_e + (1 - \rho^e_1)^{-1}(1 + \rho^e_1 u_f(\theta_d))}{V + [V^2 + 1 + \cos \theta_e (1 - \rho^e_1)]^{1/2}},
\] (22)

where
\[ k = 2V(\rho_{2e}^s)^{-1}\left[\left(V^2 + \rho_{2e}^s\right)^{1/2} - V\right], \quad C = \frac{2V(\rho_{2e}^s + \rho_{2e}^I)\mu_l(\theta_d)}{\left(V^2 + \rho_{2e}^s\right)^{1/2} + V}. \]

The presence of \( \mu_l(\theta_d) \) in (22) highlights a connection between the flow in the outer asymptotic region and the value of the dynamic contact angle. When the contact line is ‘insulated’ from the global flow by the low-Reynolds-number region, the flow near the contact line is completely determined by the contact-line speed, and hence the mass flux into the contact line that ‘feeds’ the liquid–solid interface can be found from the local solution. This is the case considered in [28,44], where the theory shows excellent agreement of (22) with experiments. If the outer flow is to influence the mass flux into the contact line [27], this will affect the value of the contact angle.

The most important length scale \( L_d \) associated with the interface formation process is the characteristic distance over which the solid surface returns to equilibrium and the asymptotic result indicates that this is given, in non-dimensional units, by \( kL_d/\epsilon = 1 \) in (19) so that \( L_d = \epsilon\rho_{2e}^s\left[2V(\sqrt{V^2 + \rho_{2e}^s} - V)\right] \).

The expressions given above will be used in Section 4 below to compare our computations to in the situations where the underlying assumptions of the asymptotics are satisfied. Now, we shall consider the development of this computational algorithm for the general case without making any simplifying assumptions.

3. Finite element procedure

A finite element framework for the simulation of dynamic wetting flows using the conventional models of dynamic wetting was described in [32]. To handle the evolution of the free surface this framework uses an arbitrary Lagrangian Eulerian (ALE) scheme based on the method of spines, a computational approach which has been successfully applied to a range of coating flows over the past thirty years, e.g. in [45,46,25]. In [34], the framework was extended for the simulation of time-dependent free surface flows, with the code providing accurate solutions for the benchmark test case [47,48] of a freely oscillating liquid drop. This confirmed that the implementation of the viscous, inertial and capillarity effects is accurate, even when the mesh undergoes \( O(1) \) deformations.

Although the interface formation model introduces additional variables into the problem, these are confined to the surfaces of the domain so that this does not notably increase the computational cost of the scheme. Furthermore, even in the conventional models a small length scale associated with slip has to be resolved to obtain mesh-independent results (see [32]), and, as we shall see, to also capture the dynamics of interface formation does not significantly increase the computational burden. In this work, our aim was to implement the interface formation model into a computational scheme for the first time, so that we made no attempt to optimize the code. However, we have found that even for the highest resolution unsteady flows, it only take around a day to run on standard laptops. The issues related to optimization of the computations can be considered in future work where, in particular, the developed framework could be applied to fully three-dimensional phenomena.

What follows is the implementation of the interface formation equations into our framework. For a more detailed description of the basic components of the framework and a user-friendly step-by-step guide to implementing dynamic wetting flows into the finite element method, the reader is referred to [32] and in particular to the Appendix which makes it possible for the interested user to reproduce the results presented there. The Appendix in the present article provides additional details of the implementation and, in this sense, complements the Appendix in [32], allowing one to reproduce the results of Section 4 and 5.

3.1. Problem formulation in the ALE scheme

Consider how the equations of Section 2, written in Eulerian coordinates \( \mathbf{x} \), are formulated for an ALE system where the flow domain \( \chi = \chi(\mathbf{x}, t) \) deforms in time. This deformation must be accounted for in the temporal derivatives of variables whose position in the Eulerian system is evolutionary, in particular, in the Navier–Stokes equations where the material derivative \( D/Dt \) transforms as

\[ \frac{D\mathbf{u}}{Dt} = \frac{\partial\mathbf{u}}{\partial t} + \mathbf{u} \cdot \nabla \mathbf{u} = \frac{\partial\mathbf{u}}{\partial t} \bigg|_{\mathbf{x}} + (\mathbf{u} - \mathbf{c}) \cdot \nabla \mathbf{u}. \]  

Here, \( \mathbf{c}(\chi, t) = \frac{D\mathbf{x}}{Dt} \) is the velocity of the ALE coordinates with respect to the fixed reference frame [49,50]. It can be seen that, as should be expected, for \( \mathbf{c} = \mathbf{0} \) we have a Lagrangian scheme whereas for \( \mathbf{c} \neq \mathbf{0} \) the Eulerian system is recovered.

Before considering temporal derivatives occurring in the interface formation equations, it is convenient to introduce the surface gradient \( \nabla^s \), which is the projection of the usual gradient operator \( \nabla \) onto the surface \( \nabla^s = (\mathbf{I} - \mathbf{n} \mathbf{n}^T) \cdot \nabla \). An arbitrary surface vector \( \mathbf{a}^s \) is written in terms of components normal and tangential to the surface as \( \mathbf{a}^s = a_n^s \mathbf{n} + a_t^s \mathbf{a}^T \mathbf{n} \), where \( a_n^s = \mathbf{a}^T \cdot \mathbf{n} \), so that for its divergence one has \( \nabla^s \cdot \mathbf{a}^s = \nabla^s \cdot \mathbf{a}^T + a_n^s \mathbf{n} \cdot \nabla^s \mathbf{n}. \) In particular, as described in [51], points on the surfaces move with the normal surface velocity \( v_n \), i.e. according to the kinematic Eq. (3), and an arbitrary tangential component \( c^s_T \) which depends on the choice of mesh design. Then, on a given surface
\[ \mathbf{c}^i = \mathbf{c}^i_j + \mathbf{v}^i_j \mathbf{n}, \quad \mathbf{c}^i_j = \frac{\partial \mathbf{x}}{\partial t} \cdot (I - \mathbf{n} \mathbf{n}). \]

Therefore, in the ALE framework the left-hand side of the surface mass balance Eqs. (8) and (14), become

\[ \frac{\partial \rho_j^s}{\partial t} + \nabla \cdot \left( \rho_j^s \mathbf{v}^s_j \right) = \frac{\partial \rho_j^s}{\partial t} \bigg|_{\mathbf{x}^e} - \mathbf{c}^s_j \cdot \nabla \rho_j^s + \nabla \cdot \left( \rho_j^s \mathbf{v}^s_j \right), \]

where \( \gamma = 1.2 \) refers to the liquid–gas and liquid–solid interface, respectively, and \( \mathbf{x}^e_j = \mathbf{x}^e_j(\mathbf{x}, t) \) are the corresponding coordinates. Then, (8) and (14) take the form

\[ \epsilon \left[ \frac{\partial \rho_j^s}{\partial t} - \mathbf{c}^s_j \cdot \nabla \rho_j^s + \nabla \cdot \left( \rho_j^s \mathbf{v}^s_j \right) + \rho_j^s \mathbf{v}^s_m \nabla \mathbf{n} \cdot \mathbf{c}^s_j \right] + \rho_j^s - \rho_j^{s_e} = 0, \quad (\gamma = 1.2), \]

where we have used that \( \mathbf{n}_j \cdot \nabla \rho_j^s = 0 \). Eq. (24) can be rearranged to obtain

\[ \epsilon \left[ \frac{\partial \rho_j^s}{\partial t} + \rho_j^s \nabla \mathbf{c}^s_j + \nabla \cdot \left( \rho_j^s \mathbf{v}^s_j \right) \left( \mathbf{v}^s_j - \mathbf{c}^s_j \right) \right] + \rho_j^s - \rho_j^{s_e} = 0, \quad (\gamma = 1.2). \]

In the limiting case, where the surface moves only normal to itself, so that \( \mathbf{c}^s_j = \mathbf{0} \), the usual Eulerian equations are recovered:

\[ \epsilon \left[ \frac{\partial \rho_j^s}{\partial t} + \nabla \cdot \left( \rho_j^s \mathbf{v}^s_j \right) + \rho_j^s \mathbf{v}^s_m \nabla \mathbf{n} \cdot \mathbf{c}^s_j \right] + \rho_j^s - \rho_j^{s_e} = \epsilon \left[ \frac{\partial \rho_j^s}{\partial t} + \nabla \cdot \left( \rho_j^s \mathbf{v}^s_j \right) \right] + \rho_j^s - \rho_j^{s_e} = 0, \quad (\gamma = 1.2), \]

whilst if the surface moves in a Lagrangian way, \( \mathbf{c}^s_j = \mathbf{v}^s_j \), then the term under the divergence becomes identically zero, and we have

\[ \epsilon \left( \frac{\partial \rho_j^s}{\partial t} + \rho_j^s \nabla \mathbf{v}^s_j + \rho_j^s \mathbf{v}^s_m \nabla \mathbf{n} \cdot \mathbf{v}^s_j \right) + \rho_j^s - \rho_j^{s_e} = \epsilon \left( \frac{\partial \rho_j^s}{\partial t} + \rho_j^s \nabla \cdot \mathbf{v}^s_j \right) + \rho_j^s - \rho_j^{s_e} = 0, \quad (\gamma = 1.2). \]

Here, as should be expected, there is no term representing the convection of surface density by the surface velocity, i.e. a term of the form \( \mathbf{v}^s \cdot \nabla \rho_j^s \) does not appear.

Having reformulated the equations for the ALE system and introduced surface operators, we can now derive the appropriate FEM residuals.

3.2. Forming the finite element residuals

The defining feature of the FEM is that the computational domain \( V \) is tessellated into a finite number of non-overlapping elements, each containing a fixed number of nodes at which the functions’ values are determined. Between these nodes the functions are approximated using interpolating functions whose functional dependence on position is chosen. In what follows, \( N_p \) is the total number of nodes in \( V \) at which the pressure is determined, \( N_n \) is the number of nodes at which the velocity components are to be found, \( N_s \) is the number of nodes on the free surface \( A_1 \), \( N_2 \) the number of nodes on the solid surface \( A_2 \), and \( N_t \) the number of nodes along the contact line.

The procedure of generating the finite element equations is well known and a detailed explanation of how this is achieved for dynamic wetting flows described by the conventional model is given in [32], so that here we just give the main details. Functions are approximated as a linear combination of interpolating functions each weighted by the corresponding nodal value. In particular, we use mixed interpolation so that the Ladyzhenskaya–Babuška–Brezzi [52] condition is satisfied\(^2\) with linear basis functions \( \psi_j \) to represent pressure and quadratic ones \( \phi_j \) for velocity:

\[ p = \sum_{j=1}^{N_p} p_j \psi_j(\mathbf{x}), \quad \mathbf{u} = \sum_{j=1}^{N_n} \mathbf{u}_j \phi_j(\mathbf{x}), \quad (\mathbf{x} \in V). \]

In the Galerkin finite element method, the basis functions which are used to discretize the functions of the problem are also used as weighting functions to create the weak form of the problem, see [Section 3 of 32] for specific details. Note that here we use Roman letters for the indices (i, j, etc) to refer to the nodal values and approximating functions spanning the whole domain (globally); in the Appendix, where all the numerical details are given, these indices will be used alongside italicized ones (i, j, etc), which will refer to local, element-based, values and functions.

Surface variables are also approximated quadratically with basis functions on surface \( A_1 \) (\( \gamma = 1.2 \)) denoted by \( \phi_j(\mathbf{s}_1, \mathbf{s}_2) \). So, all of the interface formation variables, represented by an arbitrary surface variable \( a_j^s \), as well as the shape of the free surface \( \mathbf{x}_1 \), are approximated as

\(^2\) Equal-order methods may also be used with stabilization, e.g. [53,54], but these issues lie beyond the scope of this paper.
To determine the free surface shape, i.e. the nodal values \( x_{1,j} \), a function \( h = h(s_1, s_2) \) is found as part of the solution at every free-surface node, so that \( h_j = h(s_1, s_2) \) for \( j = 1, \ldots, N_1 \), which points in a direction linearly independent from both \( s_1 \) and \( s_2 \) at each node, i.e. 'out' of the free surface. For example, in the simplest case of a Cartesian coordinate system one could have the free surface at \((x, y, z) = (s_1, s_2, h(s_1, s_2))\), so that \( h \) is the height above the \((x,y)\)-plane, or, in a two-dimensional example, one may have a polar coordinate system with the free surface given by \((\theta, r) = (s_1, h(s_1))\), in which case \( h \) is the distance of the free surface from the origin for every \( \theta \).

The basis functions used to approximated the variables are now used to derive the weak form of the problem, i.e. the finite element equations. From (1), the continuity of mass (incompressibility of the fluid) residuals \( R^c_i \) are

\[
R^c_i = \int_V \psi_i \nabla \cdot \mathbf{u} \; dV \quad (i = 1, \ldots, N_p).
\]

After projecting the momentum Eq. (1) onto the unit basis vectors of the coordinate system \( \mathbf{e}_x \; (x = 1, 2, 3) \) and using (23), our momentum residuals \( R^{M,x}_i \) take the form

\[
R^{M,x}_i = \int_V \left[ \psi_i \mathbf{e}_x \cdot \left[ \nabla \cdot \left( \frac{\partial \mathbf{u}}{\partial t} + (\mathbf{u} - \mathbf{c}) \cdot \nabla \mathbf{u} \right) - \nabla \cdot \mathbf{P} - St \mathbf{F} \right] \right] dV \quad (i = 1, \ldots, N_u).
\]

Integrating by parts and using the divergence theorem, as shown in [32], one can rewrite (27) in terms of volume and surface contributions:

\[
\begin{align*}
R^{M,x}_i & = \left( R^{M,x}_i \right)_V + \left( R^{M,x}_i \right)_A \quad (i = 1, \ldots, N_u), \\
\left( R^{M,x}_i \right)_V & = \int_V \left[ \psi_i \mathbf{e}_x \cdot \left[ \nabla \cdot \left( \frac{\partial \mathbf{u}}{\partial t} + (\mathbf{u} - \mathbf{c}) \cdot \nabla \mathbf{u} \right) - \nabla \cdot \mathbf{P} - St \mathbf{F} \right] \right] dV, \\
\left( R^{M,x}_i \right)_A & = \int_A \left[ \psi_i \mathbf{e}_x \cdot \mathbf{n} \right] \mathbf{n} \cdot dA.
\end{align*}
\]

In (28), only when node \( i \) is on the surface \( A \) will \( \psi_i \) be non-zero, i.e. it is nodes on the surface of \( V \) which contribute to the momentum residuals via (28). This term allows us to incorporate stress boundary conditions naturally, i.e. by adding them as contributions to the momentum equations at the surfaces, which is a well known advantage of the finite element method over other numerical approaches where special boundary approximations have to be constructed.

To incorporate our free-surface stress boundary conditions into (28), Eqs. (4) and (5) are rewritten into the computationally favourable form

\[
Ca \left( \mathbf{n}_i \cdot \mathbf{P} + \mathbf{n}_i \cdot \mathbf{F} \right) + \nabla^\top \cdot [\mathbf{\sigma}_i (\mathbf{I} - \mathbf{n}_i \mathbf{n}_i)] = \mathbf{0},
\]

where \( \mathbf{\sigma}_i (\mathbf{I} - \mathbf{n}_i \mathbf{n}_i) \) is the surface stress, playing the same role on the surface as \( \mathbf{P} \) does in the bulk. Then (28) can be rewritten on the free surface as

\[
\int_{A_1} \mathbf{e}_x \cdot \mathbf{n} \; dA_1 = -\frac{1}{Ca} \int_{A_1} \phi_1 \mathbf{e}_x \cdot \left[ \nabla^\top \cdot [\mathbf{\sigma}_1 (\mathbf{I} - \mathbf{n}_1 \mathbf{n}_1)] + Cap_s \mathbf{n}_1 \right] dA_1.
\]

It was initially suggested by Ruschak [55], and generalized for three-dimensional problems in [56,57], that, by using the surface divergence theorem, one could lower the highest derivatives and thus both reduce the constraints on the differentiability of the interpolating functions which are used to approximate the free surface shape and give a natural way to impose boundary conditions on the shape of the surface where it meets other boundaries. This is achieved by, first, using the product rule

\[
\int_{A_1} \mathbf{e}_x \cdot \mathbf{n} \; dA_1 = -\frac{1}{Ca} \int_{A_1} \left\{ \nabla^\top \cdot (\phi_1 \mathbf{e}_x \cdot (\mathbf{I} - \mathbf{n}_1 \mathbf{n}_1)) - \mathbf{\sigma}_1 \nabla^\top \cdot (\phi_1 \mathbf{e}_x) + Cap_s \phi_1 \mathbf{e}_x \cdot (\mathbf{n}_1) \right\} dA_1,
\]

and then using the surface divergence theorem [58, p. 224], which for a surface vector \( \mathbf{a}^s \) with no normal component, \( \mathbf{a}^s = \mathbf{a}_i^s \), is given by

\[
\int_A \mathbf{n}^\top \cdot \mathbf{a}_i^s \; dA = -\int_C \mathbf{a}_i^s \cdot \mathbf{m} \; dC,
\]

where the unit vector \( \mathbf{m} \) is the inwardly facing normal to the contour \( C \) that confines \( A \) (Fig. 1), with \( \mathbf{a}_i^s = \phi_1 \mathbf{e}_x \cdot (\mathbf{I} - \mathbf{n}_1 \mathbf{n}_1) \), so that

\[
\int_{A_1} \mathbf{e}_x \cdot \mathbf{n} \; dA_1 = \frac{1}{Ca} \int_{A_1} [\mathbf{\sigma}_1 \nabla^\top \cdot (\phi_1 \mathbf{e}_x) - Cap_s \phi_1 \mathbf{e}_x \cdot (\mathbf{n}_1)] dA_1 + \frac{1}{Ca} \int_{C_1} \phi_1 \mathbf{e}_x \cdot \mathbf{m} \; dC_1.
\]
Thus, on the free surface, the term (28) in the momentum residual is now replaced by a different surface contribution and a line contribution

\[
(R_i^{M,x})_{A_1} = \frac{1}{Ca} \int_{A_1} [\sigma_1 \nabla \cdot (\phi_1 \mathbf{e}_2) - C_{ap} \phi_1 \phi_2 (\mathbf{e}_2 \cdot \mathbf{n}_1)] \, dA_1, \quad (R_i^{M,x})_{C_1} = \frac{1}{Ca} \int_{C_1} \phi_1 \sigma_1 \mathbf{e}_2 \cdot \mathbf{m}_1 \, dC_1.
\]  

The same procedure of integrating by parts and using the divergence theorem has been used on both the surface stress term \( \nabla \cdot [\sigma_1 (1 - \mathbf{n}_i \mathbf{n}_1)] \) and the bulk stress term \( \nabla \cdot \mathbf{P} \). In both cases, this has created contributions from the boundary of that term’s domain, i.e. the confining contour and surface, respectively. Consequently, the momentum residual now contains a cascade of scales

\[
R_i^{M,x} = (R_i^{M,x})_A + (R_i^{M,x})_C.
\]  

which represent, respectively, the volume, surface and line contributions. In particular, part of the contour \( C_{i_1} \) which bounds the free surface is the contact line \( C_{i_1} \) where the free surface meets the solid. Other boundaries to the free surface further away from the contact line, for example an axis or plane of symmetry, are treated in the same way but are not to be formulated until specific problems are considered in Section 4.

At the contact line, it is useful to rearrange the term in the integrand of (30) by representing the vector \( \mathbf{m}_1 \) in terms of a linear combination of its components parallel to \( \mathbf{n}_2 \) and \( \mathbf{m}_2 \) (Fig. 1):

\[
\mathbf{m}_1 = (\mathbf{m}_1 \cdot \mathbf{m}_2) \mathbf{m}_2 + (\mathbf{m}_1 \cdot \mathbf{n}_2) \mathbf{n}_2.
\]

This identity can be used to make the contribution to (31) coming from the contact line dependent on the known vector \( \mathbf{n}_2 \), the vector \( \mathbf{m}_2 \), varying along the contact line but independent of the free-surface shape, and \( \theta_\alpha \), defined by (16) and determined by Young’s Eq. (17), so that

\[
(R_i^{M,x})_{C_i} = \frac{1}{Ca} \int_{C_i} \sigma_1 \phi_1 \mathbf{e}_2 \cdot (\mathbf{m}_2 \cos \theta_\alpha + \mathbf{n}_2 \sin \theta_\alpha) \, dC_i \quad (i = 1, \ldots, N_c).
\]

If the contact line’s tangent vector is \( \mathbf{t}_e \), then \( \mathbf{m}_2 = \pm \mathbf{t}_e \times \mathbf{n}_2 \), with the sign chosen to ensure the inward facing vector \( \mathbf{m}_2 \) is selected. Thus, Eq. (16), which defines the contact angle, can be applied in a natural way, that is without needing to drop another equation in order to make room for an equation that would fix the shape of the free surface at the contact line. The contact angle itself is determined from Young’s Eq. (17), which, when put in residual form as an integral over the contact line contour, gives

\[
R_i^l = \int_{C_i} \phi_1 \sigma_1 \cos \theta_\alpha + \sigma_2 \, dC_i \quad (i = 1, \ldots, N_c).
\]

At the liquid–solid interface the approach developed in [32] is used. Instead of dropping the momentum equation normal to the solid to impose a Dirichlet condition on the normal velocity (13), we use it to determine the normal stress acting on the liquid–solid interface which allows the contact line, where boundary conditions of different type meet, to be treated in a manner consistent with standard FEM ideology. Specifically, we introduce a new unknown \( \Lambda \) on the liquid–solid interface which is defined by the equation

\[
\Lambda = \mathbf{n}_2 \cdot \mathbf{P} \cdot \mathbf{n}_2.
\]  

(32)

It should be pointed out that this particular implementation simplifies the finite element procedure independently of the dynamic wetting model chosen and is particularly useful when considering a surface non-aligned with a coordinate axis. In this case, the procedure of rotating the momentum equations to align with the coordinate axes [59] is cumbersome whereas our approach is independent of both the free surface and the solid’s shape, that is the curved nature of a surface is as easy to handle as, say, a planar surface aligned with coordinate axes.

On the liquid–solid interface the contribution to the momentum equations from the stress on the surface, which contains contributions from both the generalized Navier condition (11) for the tangential stress and (32) for the normal stress gives

\[
(R_i^{M,x})_{A_2} = \int_{A_2} \phi_{2i} \left[ \Lambda (\mathbf{e}_2 \cdot \mathbf{n}_2) + \frac{\beta}{Ca} \mathbf{e}_2 \cdot (\mathbf{u}_i - \mathbf{U}_i) - \frac{1}{2Ca} \mathbf{e}_2 \cdot \nabla \sigma_2 \right] \, dA_2 \quad (i = 1, \ldots, N_2).
\]

where \( \phi_{2i} \) is a interpolating function for the liquid–solid interface corresponding to the ith node.

In addition to the boundary conditions involving stress on each interface, we have an additional equation involving the velocity normal to each interface. On the liquid–gas free surface this is the kinematic condition (3) whose residuals \( R_i^k \) are given by

\[
R_i^k = \int_{h_1} \phi_{1i} \left[ \left( \frac{\partial \mathbf{r}_1}{\partial t} - \mathbf{v}_1 \right) \cdot \mathbf{n}_1 \right] \, dA_1 \quad (i = 1, \ldots, N_1),
\]

(34)

\footnote{Labelled \( \iota \) in [32].}
whilst on the liquid–solid interface we have a condition of impermeability of the solid (10) with residuals \( R_i^s \) given by
\[
R_i^s = \int_{A_i} \phi_{2,i} (v_i^s - U) \cdot n_s dA_i \quad (i = 1, \ldots, N_2).
\]

In keeping with the framework presented in [32], all momentum equations are applied at both the liquid–gas free surface and at the liquid–solid interface and hence, once the two boundaries meet at the contact line, the contact line conditions can be implemented naturally, without dropping any of the equations there. A crude, but useful, interpretation is to think of the momentum equations as determining the bulk velocities, the kinematic condition on the free surface as determining the unknowns that specify its position, and the impermeability condition, which is the geometric constraint of the prescribed shape, as determining the ‘extra’ unknown \( \Lambda \), i.e. the normal stress.

Thus far, the equations are assumed to determine the bulk velocity, the shape of the free surface and the normal stress on the liquid–solid interface (\( \Lambda \)). In addition to these equations we have (6)–(9) on the free surface and (12)–(15) on the liquid–solid interface which determine the surface velocity \( v^s \), surface density \( \rho^s \) and surface tension \( \sigma \). In particular, we can think of the Darcy-type Eqs. (6) and (12) as determining the surface velocity tangential to the surface \( v^s \), which in a fully three-dimensional flow will have two components in the \( e_i \) and \( e_n \) direction, where \( e_i \) is a basis vector in the direction of increasing \( s_i, v = 1.2 \). The residuals \( R_i^{e_i,v} \) and \( R_i^{e_n,v} \) are
\[
R_i^{e_i,v} = \int_{A_i} \phi_{1,i} e_i \cdot \left( v_{1,i} - u_i - \frac{1 + 4\beta}{4\beta} \nabla^s \sigma_1 \right) dA_i \quad (i = 1, \ldots, N_1),
\]
\[
R_i^{e_n,v} = \int_{A_i} \phi_{2,i} e_n \cdot \left( v_{2,i} - \frac{1}{2} \left( u_i + U_i \right) - 2\nabla^s \sigma_2 \right) dA_i \quad (i = 1, \ldots, N_2).
\]

Eqs. (7) and (13) can be thought of as determining the normal component of the surface velocity on the surface \( A_i (\gamma = 1.2) \) and the residuals \( R_i^{e_n,n} \) from these equations take the same form on each interface:
\[
R_i^{e_n,n} = \int_{A_i} \phi_{1,i} \left[ (u - v_i^s) \cdot n_i - \frac{\rho_i - \rho_i^s}{\gamma} \right] dA_i \quad (i = 1, \ldots, N_1; \quad \gamma = 1.2).
\]

The corresponding residuals \( R_i^{e_i,n} \) from Eqs. (8) and (14) to determine the evolution of the surface density are
\[
R_i^{e_i,n} = \int_{A_i} \phi_{1,i} \left\{ \left( \frac{\partial \rho_i}{\partial t} + \nabla^s \cdot \left[ \rho_i^s \left( v_i^s - c_i^s \right) \right] \right) + \rho_i^s \nabla^s \cdot c_i^s + \rho_i^s \left( v_i^s \cdot n_i \right) n_i \right\} dA_i \quad (i = 1, \ldots, N_1; \quad \gamma = 1.2).
\]

Using the standard FEM ideology, which will give us a method for applying boundary conditions on the surface flux \( \rho^s v_i^s \) where the surface meets a boundary, we integrate the divergence term in (39) by parts to obtain
\[
R_i^{e_i} = \int_{A_i} \left\{ -\rho_i^s \left( v_i^s - c_i^s \right) \cdot n_i \phi_{1,i} + \phi_{1,i} \left[ \left( \frac{\partial \rho_i}{\partial t} + \rho_i^s \nabla^s \cdot c_i^s + \rho_i^s \left( v_i^s \cdot n_i \right) n_i \right) + \rho_i^s - \rho_i^s \right] \right\} dA_i \quad (i = 1, \ldots, N_1; \quad \gamma = 1.2).
\]

Then, using the surface divergence theorem (29) with \( a_i^s = \phi_{1,i} \rho^s \left( v_i^s - c_i^s \right) \) we have a contribution from both the surface and the contour bounding that surface, so that
\[
R_i^{e_i} = \left( R_i^{e_i} \right)_{A_i} + \left( R_i^{e_i} \right)_{C_i},
\]
\[
\left( R_i^{e_i} \right)_{A_i} = -\int_{A_i} \phi_{1,i} \left\{ \frac{\partial \rho_i}{\partial t} + \rho_i^s \nabla^s \cdot c_i^s + \rho_i^s \left( v_i^s \cdot n_i \right) n_i \right\} dA_i \quad (i = 1, \ldots, N_1; \quad \gamma = 1.2),
\]
\[
\left( R_i^{e_i} \right)_{C_i} = -\int_{C_i} \phi_{1,i} \rho_i^s \left( v_i^s - c_i^s \right) \cdot n_i dC_i \quad (i = 1, \ldots, N_1; \quad \gamma = 1.2).
\]

Consequently, we are able to apply any boundary conditions on surface mass flux by replacing the contour contribution (43) with the appropriate value. In particular, at the contact line contour \( C_{d} \) we have the surface mass flux continuity condition (18). This condition could potentially be applied by using it to replace either the contour contribution to the free surface or the liquid–solid interface equations, and to determine which way is correct we study the structure of the interface formation equations. Specifically, an asymptotic approach to the dynamic wetting problem for small \( C a \) and \( \epsilon \) recapitulated in Section 2.2 shows that it is the flux of surface mass into the contact line from the free surface, which depends on the global flow via the velocity in the far field on the free surface \( u_f (\phi_e) \) that determines the relaxation process on the solid surface. Therefore, in a numerical implementation of this problem one should allow the free surface equations to determine the flux into
the contact line and then use the surface mass continuity condition (18) on the liquid–solid side of the contact line to take this flux as the surface mass supply into the liquid–solid interface.

Therefore, on the free surface we have

$$
\left( R_{s1}^{\rho} \right)_{cl} = - \int_{cl1} \epsilon \phi_{11} \rho_{1} \left( v_{1} - c_{1} \right) \cdot m_{1} \, dC_{cl},
$$

whilst on the liquid–solid interface, using (18) to rewrite the flux $\rho_{2} \left( v_{2} - c_{2} \right) \cdot m_{2}$ into this interface in terms of the flux into the contact line from the free surface, we have

$$
\left( R_{s2}^{\rho} \right)_{cl} = \int_{cl1} \epsilon \phi_{21} \rho_{1} \left( v_{1} - c_{1} \right) \cdot m_{1} \, dC_{cl}.
$$

Then, the flux into the liquid–solid interface is given in terms of the flux that goes into the contact line from the free surface.

The final residuals from the surface equations, $R_{s}^{\rho}$, are obtained from the surface equation of state (9) and (15), which express the surface tension as an algebraic function of the surface density:

$$
R_{s}^{\rho} = \int_{A_{s}} \phi_{i} \left( \sigma_{i} - z \left( 1 - \rho_{i}^{*} \right) \right) \, dA_{i} \quad (i = 1, \ldots, N_{f}; \, \gamma = 1, 2).
$$

In this section, we have derived the finite element residuals required to solve dynamic wetting flows using the interface formation model. In particular we have the following coupling of unknowns

$$
R_{i}^{M}, \, R_{i}^{C}, \, R_{i}^{K}, \, R_{i}^{\rho}, \, R_{i}^{\sigma}, \, R_{i}^{\theta}, \, R_{i}^{\sigma}, \, R_{i}^{\beta}, \, R_{i}^{\gamma}, \quad (\alpha = 1, 2, 3; \, \gamma = 1, 2; \, \nu = 1, 2),
$$

each of which has its corresponding residual

$$
\left( R_{i}^{M}, \, R_{i}^{C}, \, R_{i}^{K}, \, R_{i}^{\rho}, \, R_{i}^{\sigma}, \, R_{i}^{\theta}, \, R_{i}^{\sigma}, \, R_{i}^{\beta}, \, R_{i}^{\gamma} \right) \quad (\alpha = 1, 2, 3; \, \gamma = 1, 2; \, \nu = 1, 2).
$$

The subscripts i above will have different limits that are the same in both the variable and its corresponding residual, i.e. the same number of equations and unknowns has been assured.

4. Validation of the code and some benchmark calculations for Problem A

Computations are performed for cases which are axisymmetric or two-dimensional in simple geometries so that calculations may be easily reproduced, thus giving benchmark results for future investigators. In what follows, we consider a meniscus rising against gravity through a cylindrical capillary of radius R. The computational domain is a region in the $(r, z)$-plane, and the free surface is parameterized by arclength s. Motion is assumed to be axisymmetric about the $z$-axis which runs vertically through the centre of the capillary with the radial axis perpendicular and located at the base of the capillary (Fig. 2). First, we consider the steady propagation of a meniscus through a capillary (hereafter Problem A) to check convergence of our code as the mesh is refined and to compare our computations to the asymptotic results summarized in Section 2.2. Then, we will examine Problem A from the view point of analyzing the influence of the radius R of the capillary.
on the interface formation dynamics. Finally, the unsteady imbibition of a liquid into a capillary will be considered (hereafter Problem B) and the results compared to published experimental data for this type of flow.

In the \((r, z)\)-plane where the solution is computed, in addition to the equations formulated in Section 2, on the axis of symmetry for both Problem A and B we have the symmetry conditions in the form of impermeability and zero tangential stress

\[
\mathbf{u} \cdot \mathbf{n} = 0, \quad \mathbf{n} \cdot \mathbf{P} \cdot \mathbf{t} = 0, \quad (r = 0, 0 < z < z_0),
\]

where \(z_0\) is the a-priori unknown apex height. Additionally, at the apex we have the conditions of (a) smoothness of the free-surface shape and (b) the absence of a surface mass source or sink:

\[
\mathbf{n} \cdot \mathbf{e}_r = 0, \quad \rho_1^s \mathbf{v}_l^r \cdot \mathbf{e}_r = 0, \quad (r = 0, z = z_0).
\]

Before doing the calculations we need to consider estimates for the model’s parameters for the two problems to be studied, leaving free the radius of the capillary as a parameter, whose influence will be examined in Section 5.

4.1. Typical parameter regime

To obtain estimates for our parameters, consider the flow of a water-glycerol mixture through a capillary of radius \(R\) at speed \(U = 0.01\) m s\(^{-1}\). At 60% glycerol this gives fluid properties of \(\rho \approx 10^3\) kg m\(^{-3}\), \(\mu \approx 10^{-2}\) kg m\(^{-1}\) s\(^{-1}\) and \(\sigma \approx 7 \times 10^{-2}\) N m\(^{-1}\) [38]. On the molecular scale, the interface is a layer of finite thickness \(\ell\) and, as discussed in [38], the generalized Navier equation and Darcy-type equation are analogous to what one would have for the averaged quantities for flow in a channel of width \(\ell\). Using this analogy, one would have \(\beta \sim \mu/\ell\) and \(\alpha \sim \ell/\mu\), so that, taking the coefficients of proportionality as unity as a first approximation, and estimating \(\rho_\ell^s \approx 0.6\) (so that \(\lambda = 2.5\)), one has from dynamic wetting experiments [38] that the relaxation time scales as \(\tau = \tau_\mu\) with \(\tau_\mu\) extracted from the data as \(\tau_\mu \approx 7 \times 10^{-3}\) kg m\(^{-1}\) s\(^{-2}\). The equilibrium contact angle is taken as \(\theta_c = 30^\circ\), so that \(\rho_{\ell e}^s = 1.35\). Using these estimates, we have the following parameters of what we will regard as the base state. The parameters independent of the length scale are

\[
Ca = 10^{-2}, \quad Q = 0.1, \quad \rho_1^s = 0.6, \quad \rho_{2e} = 1.35, \quad \lambda = 2.5, \quad \theta_c = 30^\circ,
\]

and those dependent on the length scale are

\[
Re = 10^{-2} (\mu m)^{-1} R, \quad St = 10^{-5} (\mu m)^{2} R^{2}, \quad \bar{x} = 10^{-1} (\mu m)^{R^{-1}}, \quad \bar{\beta} = 10 (\mu m)^{R^{-1}}, \quad \epsilon = 10^{-2} (\mu m)^{R^{-1}}.
\]

As capillary sizes of interest can range from the millimetre scale \(R = 10^2\) mm, relevant, for example, for applications in microgravity [60], right down to a few tens of nanometres \(R = 10^{-2}\) mm [22], the flow can be dominated by different physical factors. In particular, in large capillaries bulk inertia becomes important as \(Re > 1\) and the surface tension relaxation effect is localized \((\epsilon \ll 1)\), being important only close to the contact line, whilst for nanoscale flows inertia becomes negligible \((Re \ll 1\) and the surface tension relaxation length becomes comparable to the capillary’s width \(\epsilon \approx 1\).

Now, we will present benchmark calculations for the steady propagation of a meniscus through a capillary (Problem A) and then consider the time-dependent rise of a meniscus into a capillary (Problem B), which will be compared to experiments.

4.1.1. Problem A: steady propagation of a meniscus through a capillary

For the steady propagation of a meniscus, the ‘base’ of the domain \((z = 0)\) is sufficiently far (quantified below) from the meniscus so that the base can be considered as a ‘far field’. In the far field, the flow is fully developed and the surface variables take their equilibrium values.

The velocity distribution across the capillary base is a Poiseuille profile adapted to allow for slip at the liquid–solid interface that follows from the Navier-slip condition and an additional flux of mass out of the domain via the liquid–solid interface which occurs when \(Q\) is non-zero:

\[
u = 0, \quad w = -1 + \frac{1 + \bar{q}}{1/2 + (2 + \bar{q})/\bar{\beta}/Ca}(1 + 2/(\bar{\beta}/Ca) - r^2), \quad \bar{q} = 2\epsilon Q \rho_{2e}.
\]

where \(u\) and \(w\) are the radial and axial velocity components, respectively. The derivation of this condition is given in the Appendix. Notably, if \(\bar{q} = 0\) and \(\bar{\beta}/Ca \to \infty\), then \(w = 1 - 2r^2\) which is the usual Poiseuille profile satisfying \(dw/dr = 0\) at \(r = 0, w = -1 \) at \(r = 1\) and \(\int_0^1 wdr = 0\).

Taking the radius of the capillary as \(R = 100\) mm, the parameters dependent on the capillary width become \(Re = 1.5 St = 10^{-1}, \bar{x} = 10^{-3}, \bar{\beta} = 10 \) and \(\epsilon = 10^{-4}\). In this case, \(\epsilon = 10^{-4}\) is so small that the distance of the far field is determined by the need for the bulk flow to settle as opposed to for the surface phase to relax to its equilibrium state. It is known that putting the contact line a distance of ten radii away from the ‘far field’ is more than enough for this to be satisfied [61,32].

The streamlines for this flow are shown in Fig. 3. It can be seen that the motion of the solid with respect to the meniscus drags fluid away from the contact line and that this in turn leads to a flux of fluid into the contact-line zone from the region near the free surface. In Fig. 4, one can see that, as predicted by experiments, the motion of the liquid in the vicinity of the
contact-line is ‘rolling’, with the liquid–solid interface formed by adsorbing fluid from the bulk. The Poiseuille profile, indicated by parallel streamlines in Fig. 3, is established relatively quickly, so that the truncated far field is certainly far enough away from the contact-line region to not influence the results. The mesh is graded, with small elements near the contact line and larger ones further away, so that the essential physics of wetting, occurring on a smaller length scale than the bulk flow, is fully resolved whilst the problem remains computationally tractable. The details of the construction of this mesh are given in [32], where it is shown how the bipolar coordinate system can be utilized to provide circular spines near the contact line and straight ones further away to match with the domain’s shape.

4.1.2. Mesh independence tests
The mesh is constructed such that the number of elements in the body of the capillary and those near the meniscus can be separated. This allows us to study the convergence of the surface equations to a solution without being concerned by the bulk dynamics. The number of nodes along the surfaces can be chosen as well as how fast the element size is increased as one moves away from the contact line. Here, we fix the increase in element size as 10%, and check that the solution converges as the mesh is refined, i.e. as we increase the resolution near the contact line.

In [32], it was shown that the dynamic contact angle \( \theta_d \) imposed into the weak formulation can differ from the computed one \( \theta_c \), i.e. the one obtained from the computed free-surface shape, if the mesh is under resolved near the contact line. Rather than imposing the angle in the strong form, and moving the errors to other less observable parts of the numerical scheme, as has often been the case in previous works, it was demonstrated that the difference \( |\theta_d - \theta_c| \) provides a simple error-indicator for the scheme. When considering the conventional model, it was shown\(^4\) that to resolve both the dynamics of slip and the free-surface shape near the contact line, so that for the computed angle one has, say, \( |\theta_d - \theta_c| < 0.1 \), the smallest element size

\(^4\) Note that in this previous publication the non-dimensionalization is slightly different with \( \gamma = \beta L/\mu \) whereas here it is \( \gamma/Ca = \beta L/\mu \).
Consider 13 meshes with smallest elements ranging over five orders of magnitude from $l_{\text{min}} = 5 \times 10^{-3}$ down to $l_{\text{min}} = 5 \times 10^{-8}$. Coarser meshes were unable to provide a solution as the computed angle $\theta_c$ increased past 180°, despite the imposed angle $\theta_d$ being less than 60°, in the same way as observed in [32]. The change in the free-surface shape during the mesh refinement procedure is shown in Fig. 5, where one can see just how bad the approximation is on our most coarse mesh, represented by curve 1. It can be seen that convergence appears to take place in two stages, so that decreasing the smallest element from 3 \times 10^{-4} (curve 2) to 2 \times 10^{-5} (curve 3) has little influence on the free-surface shape, but, for the reason explained below, these shapes are yet far from the converged free-surface shape (curve 5) which is obtained only when the mesh elements decrease much further.

The two-stage convergence can be explained by examining the aforementioned angles as the mesh is refined, as shown in Fig. 6. One can see that at around $l_{\text{min}} = 10^{-5}$ the computed angle $\theta_c$ becomes indistinguishable from the dynamic angle $\theta_d$ that goes into the weak formulation, but at this stage $\theta_d$, which is a part of the solution, is yet to converge to its final value. This corresponds to curves 2 and 3 in Fig. 5. More specifically, we see that $|\theta_d - \theta_c| < 0.1°$ at $l_{\text{min}} = 7 \times 10^{-6}$, close to $10^{-5}$ predicted from [32], i.e. the mesh is already sufficiently resolved for the computed angle $\theta_c$ to be close to the angle $\theta_d$ that features in the mathematics of the problem and goes into the weak formulation. However, in our problem, the angle $\theta_d$ is itself a variable and the deviation of $\theta_d$ from its final value $\Theta_d = 60.5°$ is still large $|\theta_d - \Theta_d| = 3.8°$. The convergence towards a final solution is then dominated by the requirement to resolve the interface formation dynamics. We have $|\theta_d - \Theta_d| < 0.1°$ at $l_{\text{min}} = 4 \times 10^{-7}$ which again falls close to our estimate and shows the need to resolve the length scale $\epsilon \text{Ca} = 10^{-6}$. The two-stage convergence in this problem is what one should expect given that there are two small length scales associated with the model we are using. Once these have been resolved, there is no reason to expect any further interesting convergence behaviour.

The information obtained suggests a new estimate which builds upon that given in [32], that for errors in the angle of the order of 0.1°, we require

$$l_{\text{min}} \leq \min(5 \times 10^{-2} \beta^{-1}, \text{Ca} \beta^{-1}, 10^{-1} \epsilon \text{Ca}),$$

which, respectively, ensure that the code resolves the free surface near the contact line (particularly at high Ca), the length scale on which slip occurs and the dynamics of interface formation.

A frequently used way of simplifying the problem of modelling the flow through a capillary and of easily interpreting theoretical and experimental results for this flow is to approximate the free-surface shape as a spherical cap [62,63]. Then, the free-surface shape is fully specified by the difference between the contact-line height and apex height $h = z_c - z_a$. Usually, the spherical-cap approximation is characterized by the so-called ‘apparent’ contact angle $\theta_a$, which is the angle that a spherical cap fitted through the apex and the contact line makes with the solid:

$$\theta_a = \min \left\{ \theta_{c15}, \theta_{c28}, \theta_{c08} \right\}.$$
A spherical cap is the free-surface shape obtained in the limit of $Ca \to 0$ and by computing $\theta_a$ we will also have a measure of the deviation from this shape due to viscous bending of the free surface.

In Fig. 5 the spherical cap approximation for the converged solution is plotted (dashed line). Deviations from the actual free-surface shape (curve 5) can be observed and the apparent angle is considerably larger, by 11°, than the dynamic contact angle whose asymptotic value for this set of parameters is 102°.

Three data sets are considered, with $(Ca, \epsilon) = 1 : (5 \times 10^{-2}, 5 \times 10^{-4}), 2 : (10^{-3}, 2.5 \times 10^{-2}), 3 : (10^{-3}, 5 \times 10^{-4})$, so that we can ascertain the influence of decreasing either $\epsilon$ or $Ca$ by a factor of 50. The simplest ‘integral’ measure of convergence is the dynamic contact angle whose asymptotic value for this set of parameters is 102.1°. The obtained solutions for $\theta_a$ for $1 – 3$ are 104.1°, 107.7° and 102° so that the deviations from the asymptotic result are 2°, 5.6° and 0.1°, respectively. Therefore, we have the convergence of the numerically obtained dynamic contact angle to the asymptotic one, with an indication how this convergence is controlled by $\epsilon$ and $Ca$.

To examine the issue of convergence in more detail, consider the distributions of the surface parameters and the bulk velocity along the interfaces. Deviations of the surface density on the free surface $\rho_1^s$ from its equilibrium value $\rho_{1e}$, and the distributions of the tangential components of the surface velocity $v_{1t}^s$ and the bulk velocity tangential to the free surface $u_{1t}$, are shown in Fig. 7. As one can see, the magnitude of the variation in $\rho_1^s$ along the free surface is governed by $Ca$, but it is $\epsilon$ that determines how far the ‘level’ of $\rho_1^s$ is from its equilibrium value. As $\epsilon \to 0, Ca \to 0$, we have that $\rho_1^s \to \rho_{1e}$.

In agreement with the asymptotic results, the surface velocity $v_{1t}^s$ coincides with $u_{1t}$ in the intermediate region and the two velocities begin to deviate from each other at around $s/\epsilon \sim Ca$, i.e. in the inner region, where, as expected, $v_{1t}^s$ remains constant whereas $u_{1t}$ decreases as the contact line is approached. This pattern is clearly seen for data set 3, where both the surface and bulk velocity attain the value of $u_j = -0.68$ in the outer region, $s/\epsilon \gg 1$, then, as the intermediate asymptotic region is entered, $v_{1t}^s$ and $u_{1t}$ remain at this level through this region $Ca \ll s/\epsilon \ll 1$, and as the inner region is reached, the surface velocity $v_{1t}^s$ remains constant up to the contact line, whereas $u_{1t}$ varies. Since $\rho_1^s$ is also constant through the inner region, it is the surface mass flux of the intermediate region that goes into the contact line and hence determine the mass flux that feeds the liquid–solid interface. Notably, as one could already see in Fig. 4, for $Q \neq 0$ the contact line is not a stagnation point for the bulk flow, so that $u_{1t} \neq 0$ at the contact line.
As one can see, the computed curves converge to the asymptotic distributions of Section 2.2 as \( \epsilon \) is smaller than the one assumed to be asymptotically small, e.g. for \( \epsilon = 10^{-6} \). Now, we can do the reverse and use the code to ascertain the accuracy with which the boundary-value problem for the ODEs (21) gives the dynamic contact angle and the accuracy of the formula (22) obtained under the additional assumption \( \lambda \gg 1 \) when both are applied in a range of parameters where the assumptions of the asymptotics may not be satisfied. In general, over the considered range the asymptotics gives a remarkably good approximation to its equilibrium value at the contact line, and it takes a finite distance for the interface to fully form. At the contact line, the surface velocity tangential to the solid is lower than its equilibrium value and it increases to reach the solid’s velocity on the same characteristic length scale.

The relaxation of the surface variables along the liquid–solid interface to their equilibrium far-field values is shown in Fig. 8. As one can see, the computed curves converge to the asymptotic distributions of Section 2.2 as \( \epsilon, \Ca \to 0 \) with data set 3 being practically indistinguishable from the asymptotic curves (dashed lines). In all cases the surface density is below its equilibrium value at the contact line, and it takes a finite distance for the interface to fully form. At the contact line, the surface velocity tangential to the solid is lower than its equilibrium value and it increases to reach the solid’s velocity on the same characteristic length scale.

The convergence of our computational results to the asymptotic ones in the limit considered has now been confirmed. In Table 1 a range of values of \( \epsilon, \Ca \) are considered with the parameters as above except for \( \tilde{\beta} = \tilde{\lambda}^{-1} = 10^3 \), which is now fixed, so that \( V^2 \) varies. In general, over the considered range the asymptotics gives a remarkably good approximation to the dynamic contact angle. The formula (22) also gives a very reasonable approximation which improves considerably for smaller values of \( \epsilon \). This good agreement is obtained despite the fact that \( V^2 \) is smaller than the value of \( \Ca \) in certain cases, i.e. a parameter considered finite in the asymptotics is smaller than the one assumed to be asymptotically small, e.g. for \( \epsilon = 10^{-4} \) we have \( V^2 = 8 \times 10^{-2} \), and yet for \( \Ca = 10^{-1} \) the computed angle is still only 1.6° away from the asymptotic value.

The aforementioned benchmarking has been achieved without special consideration of the pressure field, which, as is known for this class of flows, can exhibit singularities, both inherent in the mathematical model and spurious, arising as...
numerical artifacts. Specifically, very close to the contact line, for both the conventional models as well as the interface formation model, for acute contact angles computations exhibit logarithmically singular behaviour of pressure, whilst for obtuse angles spurious pressure modes dominate [33]. Known methods, which have been previously implemented into our numerical codes, exist for both capturing the logarithmic singularity using knowledge of its asymptotic structure, see [64,25], as well as for identifying and removing spurious pressure modes, see [33]. However, the flow field and contact angle remain insensitive to the precise way in which the pressure is handled near the contact line, so that benchmark results have been obtained without the need for these special methods. It is interesting to note that, in spite of progress in [33], the precise cause of spurious pressure behaviour near the contact line is unclear and a robust method for removing singular modes in computations of this class of flows, which are straightforward to identify and eliminate in analytic work, remains an open problem.

Thus, it has been shown that our computational scheme, which incorporates the interface formation model, is mesh-independent once all the length scales are resolved and converges to the asymptotic results in the limiting case. The computational results reported so far can be used as benchmarks for those who are interested in putting the interface formation model into a numerical code. It is useful to know that the asymptotic result gives a good prediction of the key feature of the wetting process, the value of the dynamic contact angle, with the formula (22) giving a reasonable approximation, well outside its strict limits of applicability. Now, we will use the developed numerical tool to make novel predictions on the size-dependence of the dynamic wetting flow through a capillary.

5. Size dependence of flow characteristics in Problem A

Now, consider how the flow characteristics of steady capillary rise (Problem A) depend on the radius of the capillary. We will vary the radii over three orders of magnitude, from the \( R = 100 \, \mu m \) considered previously down to \( R = 0.1 \, \mu m \). In all cases, the far field is placed at a sufficient distance from the contact line, ensuring that the liquid–solid interface is in equilibrium there and that moving it further away does not alter the dynamics of the interfacial relaxation process.

Conventional theories, where the dynamic contact angle is an input, give that, because the material properties of the system remain unchanged and the meniscus is travelling at the same speed, the dynamic contact angle must be the same in all of these cases, i.e. independent of \( R \). In contrast, in the interface formation model, where the dynamic contact angle is an output determined by the relaxation process along the interfaces, i.e. it is part of the solution, changes to the flow field near the contact line due to changes in \( R \) can alter the angle.

In Fig. 9, the bulk and surface velocities tangential to the free surface are plotted against \( s/\varepsilon = 10^3 \, (\mu m)^{-1} \, R \). In the geometry of a capillary, the apex is a stagnation point in the flow; it slows down the flow in its vicinity so that, as one can see in the Figure, the smaller \( R \) becomes, the lower the bulk velocity is along the free surface. As the contact line is

| \( \varepsilon = 10^{-2} \) | \( \varepsilon = 10^{-3} \) | \( \varepsilon = 10^{-4} \) |
|-----------------|-----------------|-----------------|
| \( s/\varepsilon \) | \( s/\varepsilon \) | \( s/\varepsilon \) |
| \( \rho_j \) | \( \rho_j \) | \( \rho_j \) |
| \( u_j \) | \( u_j \) | \( u_j \) |
| \( v_j \) | \( v_j \) | \( v_j \) |

Fig. 9. Distribution of surface variables along the free surface in capillaries of sizes 1: \( R = 100 \, \mu m \), 2: \( R = 10 \, \mu m \), 3: \( R = 1 \, \mu m \), 4: \( R = 0.1 \, \mu m \), against the scaled distance from the contact line \( s/\varepsilon \). Left: surface density \( \rho_j \). Right: velocities tangential to the free surface from the bulk \( u_j \) and in the interface \( \rho_j \).
approached, the surface velocity follows the bulk velocity until \( s \approx 10\epsilon \) and then the bulk velocity decreases whilst the surface velocity remains approximately constant. Thus, when the capillary is relatively large (\( R \gg 10\mu m \)), as for curves 1, 2, there is enough space for the bulk velocity and surface velocity to increase from zero at the apex to a 'far field' value, which is not far from \( u_f(\theta_d) \) given by the asymptotic solution. Then it is this value that the surface velocity keeps up to the contact line. However, as the radius of the capillary decreases and the apex gets closer to the contact line, a 'far field' no longer exists and the velocities do not have enough room to accelerate to reach the same 'far field' value, and, as a result, the surface velocity at the contact line becomes much reduced. As can be seen from Fig. 9, the surface density on the free surface is less affected, remaining close to its equilibrium value even for the smallest capillary and consequently it is the surface velocity which determines the surface mass flux into the contact line. For smaller capillaries this flux is significantly reduced.

Fig. 10. Distribution of surface variables along the solid surface in capillaries of sizes 1: \( R = 100 \mu m \), 2: \( R = 10 \mu m \), 3: \( R = 1 \mu m \), 4: \( R = 0.1 \mu m \), against the scaled distance from the contact line \( s/\epsilon \). Left: surface density \( \rho_s \). Right: velocities tangential to the solid surface from the bulk \( u_2 \) and in the interface \( v_2 \).

Continuity of surface mass flux across the contact line means that, as \( R \) goes down, the flux into the liquid–solid interface is also reduced, as can be seen in Fig. 10, where, as is the case for the free surface, changes in the surface velocity are more pronounced than those in surface density. The general trend for both of these surface variables along the liquid–solid interface is the same exponential profile, with the value at the contact line differing in each case. Notably, as the capillary size decreases, the influence of slip on the velocity at the liquid–solid interface becomes more pronounced and this accounts for curve 4 of the surface velocity deviating from the other curves for large \( s/\epsilon \).

The value of the dynamic contact angle is 'negotiated' (via Young’s equation) by the surface tensions at the contact line, and, although the changes in the surface density, and hence surface tension, are less pronounced than those in surface velocity, it can be seen in Fig. 11 that the former still lead to appreciable changes in the dynamic contact angle as the capillary radius is reduced. There is a general trend upwards which is consistent with previous studies [9] showing that, if the forming liquid–solid interface is 'starved' of surface mass flux coming from the free surface, this results in higher dynamic contact angles. In Fig. 11, the lower dashed-line corresponds to the asymptotic value, whilst the upper dashed-line is the value

Fig. 11. Dynamic contact angle \( \theta_d \) as the capillary radius \( R \) is varied from 100 nm up to 100 \( \mu m \). The lower dashed line is the asymptotic value using \( u_f(\theta_d) \) obtained from (20) whilst the upper one is obtained using \( u_f = 0 \).
for the angle which would be obtained if the velocity in the far field is taken as $u_r = 0$ in the asymptotic formula (22). Our curve is wedged between these two potentially limiting values, and it will be the subject of further work to investigate whether this simple estimate is actually a bound on the variation of the dynamic contact angle in this geometry or whether additional factors come into play in other parameter regimes.

6. Problem B: unsteady imbibition into a capillary

Next, the time-dependent imbibition of a liquid into an initially dry capillary is considered, i.e. the case in which a capillary comes into contact with a reservoir of liquid leading to the fluid being sucked into the capillary until the meniscus reaches Jurin’s height, at which gravitational forces balance the capillary ones that drive the flow. The capillary is aligned vertical with respect to the gravitational field so that the body force $\mathbf{F} = -\mathbf{g}$. It is assumed that the base of the capillary is maintained at a constant pressure, the atmospheric one, and that the flow at the inlet is parallel to the walls of the capillary so that

$$u = 0, \quad p = p_s, \quad (0 < r < 1, \ z = 0).$$

As an initial condition, we assume that a small amount of liquid is in the capillary, of height one-tenth of the radius of the capillary $h = 0.1$, that this liquid is at rest, and that the meniscus has not had time to adjust to its surroundings so that the interface is flat with $z_1 = h$ for $0 < r_1 < 1$. The interface formation variables on the free surface are taken to be the equilibrium values, so that $\sigma_1 = 1$, and the liquid–solid interface is assumed not to be formed yet so that $\rho^2 = \rho^0$. This corresponds to $\sigma_2 = 0$ and hence is consistent with our assumption of a flat interface as the Young Eq. (17) gives $\theta_0 = 90^\circ$. Thus, as initial conditions we have

$$u = 0, \quad \rho_1 = \rho_{1e}, \quad \rho_2 = \rho_{2(0)}, \quad h = 0.1.$$

As a benchmark test case, parameters are obtained from the experiments in [65]. In particular, we compare our simulations to experimental data in Fig. 6, 8 of [65], where the imbibition of a silicon oil of density $\rho = 9.8 \times 10^3$ kg m$^{-3}$, viscosity $\mu = 12.25$ kg m$^{-1}$ s$^{-1}$ and a surface tension with air of $\sigma = 2.13 \times 10^{-2}$ kg s$^{-2}$ into capillaries with perfectly wettable walls, $\theta_0 = 0^\circ$, was studied. The two capillary radii studied were $R = 0.036$ cm and $R = 0.074$ cm. Rather than fitting the interface formation parameters for this liquid–solid combination, we use, as a first approximation, estimates for the model’s parameters considered previously. Using these parameters and taking as a characteristic velocity $U = \sigma/\mu = 1.74 \times 10^{-3}$ ms$^{-1}$, so that $Ca = 1$, we have for $R = 0.036$ cm that

$$Re = 5 \times 10^3, \quad St = 5.8 \times 10^2, \quad \epsilon = 4.1 \times 10^{-4}, \quad \bar{a} = 5.6 \times 10^{-6}, \quad \bar{b} = 1.8 \times 10^5, \quad Q = 1.3 \times 10^{-2}, \quad \rho_{1e} = 0.6, \quad \rho_{2e} = 1.4, \quad \lambda = 2.5, \quad \theta_e = 0^\circ, \quad (51)$$

whilst for $R = 0.074$ cm

$$Re = 1 \times 10^4, \quad St = 2.5 \times 10^{-1}, \quad \epsilon = 2 \times 10^{-4}, \quad \bar{a} = 2.7 \times 10^{-6}, \quad \bar{b} = 3.7 \times 10^5,$$

and the remaining parameters are the same.

A relevant reference case for comparison is the so-called Lucas–Washburn approximation [62], often used to analyze experiments [66] on the unsteady imbibition into capillary tubes. In this approximation, the Navier–Stokes equations are volume averaged to obtain the meniscus height $h$ as a function of time under the assumptions that (a) Poiseuille flow occurs throughout the entire capillary, (b) that the driving force for the flow is the difference $p_s - p_e$ in pressure between the pressure at the inlet $p_s$ and the pressure at the meniscus $p_e$, (c) the meniscus is a spherical cap at all times so that $p_s - p_e = 2\sigma \cos \theta_e / R$ and, in the original form of the theory, (d) the contact angle is assumed to be equal to its equilibrium value the entire time, so that, in our case, $\theta_e \equiv 0^\circ$ and hence $p_s - p_e = 2\sigma / R$. When there is no gravitational field, the meniscus continues to propagate indefinitely into the capillary, being slowed by the gradually increasing viscous drag as the distance between the meniscus and the inlet increases. If gravity is taken into account, then the meniscus will eventually reach its equilibrium Jurin height $h_e$, calculated from balancing the driving force $p_s - p_e$ with the body force on the fluid $\rho g h_e$, i.e. $h_e = 2\sigma / (\rho R g)$. Inertial effects can also be easily included, resulting in a nonlinear ODE to be solved for the meniscus height which in dimensional form is given by

$$\rho \left[ hh'' + (h')^2 \right] = -\frac{8\mu}{R^2} hh' + \frac{2\sigma}{R} - \rho g h,$$

where the prime denotes differentiation with respect to time and, to compare with our calculations, the initial conditions at $t = 0$ are $h = 0.1R, h' = 0$. The terms on the left hand side of (52) are inertial forces, the first term on the right hand side represents viscous forces, the second one is the capillary driving force and the last term is the body force of gravity. There are papers that have been concerned with improving the basic Lucas–Washburn model [65,17,11], but here we just use the original model for comparison as a reference point.

As the Lucas–Washburn model predicts that the meniscus is always a spherical cap with, in the case of perfect wetting, $\theta_e = 0^\circ$, we take the predicted averaged height $h$, and then fit the spherical cap to it ensuring mass conservation which results
in the apex at \( z_a = h - 2R/3 \) and the contact line at \( z_c = h + R/3 \). This means that at the instant when the simulation starts, the interface jumps from its initially flat shape to a spherical cap which extends both above and below the initial interface, i.e. partly back into the reservoir. This is a known unphysical feature of the model \cite{66}, and forces it to predict infinite speeds as \( t \to 0 \).

In Fig. 12, the predictions of the two models are compared for each capillary. The Lucas–Washburn model always predicts a much faster uptake of the liquid, whilst in our simulation the meniscus is slowed as it takes time for the dynamic contact angle to decrease, and hence the capillary pressure to increase, from an initially flat interface, so that no unphysical jumps in variables are observed in the initial stages. Notably, at the end of the time periods considered, the Lucas–Washburn result predicts that the menisci are close to their Jurin height, \( h_\text{J} = 1.23 \text{ cm} \) and \( h_\text{J} = 0.6 \text{ cm} \), respectively, whilst the results from our simulation are such a distance away from \( h_\text{J} \) that a comparison of the results with experimental ones should be able to easily distinguish between the two models’ predictions.

The results from our simulations are compared to both the experimental results of \cite{65} as well as to the Lucas–Washburn curve in Fig. 13, where two sets of data have been shown in the second graph to illustrate the repeatability of the experimental results. Considering that no parameters have been fitted, our computations are seen to approximate the data exceptionally well and are vastly superior to the Lucas–Washburn curve which hugely overpredicts the speed at which the capillary uptakes the fluid. The inadequacy of the Lucas–Washburn equation has been recognized before and attributed to the need to account for a variable dynamic contact angle \cite{65}. In a forthcoming article, our code will be used to assess how all the models proposed for imbibition into a capillary perform, establish the individual effects contributing to the meniscus’ behaviour and, since none of the models account for the effect of the capillary size on the contact angle, ascertain bounds of applicability of the models previously proposed. Here, we are satisfied with the current result which demonstrates the code’s ability to describe experimental data with ease.

In Fig. 14, the relationship between capillary number based on the contact line speed and contact angle is presented. Notably, as the two curves are for the same fluid, it is only the variation of the contact-line speed that causes the variation of the capillary number, so that the latter can be viewed just as the dimensionless contact-line speed. Surprisingly, the curves for the two different capillary sizes are graphically almost indistinguishable. One may naively think that this is an argument in favour of using a speed-angle formula; however, as one can see from Fig. 14, the velocity dependence of the contact angle is multivalued so that it is not a function. For example, at \( C_a = 0.03 \) the contact angle \( \theta_a = 45.7^\circ \) or \( 84.2^\circ \). Furthermore, the upper branch of the curve corresponds to a decrease in the contact angle as the contact-line speed increases. The upper branch of the curve is far away from the asymptotic line (dashed line) computed from \( (22) \) whilst the lower one is very close to it (the asymptotic value at \( C_a = 0.03 \) is \( \theta_a = 46.4^\circ \)).

The reason for this double-valuedness of the dynamic contact angle versus the contact-line speed plot can be explained very simply. The liquid started moving from rest and at \( t = 0 \) the contact angle had the value of \( \theta_a = 90^\circ \). Therefore, if the model does not produce unphysical singularities, with dynamic characteristics experiencing instant jumps, then both the contact-line speed and the contact angle have to evolve continuously from their initial values, in our case, to 0 and \( 90^\circ \), respectively. Hence there appears a branch in the speed-angle plot stemming from the initial state and leading towards the quasi-steady state. This manifestation in the speed-angle relationship of the initial conditions rules out in principle the possibility of treating the dynamic contact angle as a prescribed input into the model. It is important to emphasize that this is a general argument based on the necessity not to have unphysical singularities in the mathematical model of a physical phenomenon, and the plot produced by using the interface formation model is merely an illustration that a model regarding the dynamic contact angle as part of the solution satisfies the above requirement in a natural way.

Fig. 12. Snapshots of the free-surface shape taken at constant time intervals as predicted by the Lucas–Washburn model (left of each figure) compared to our simulation (right of each figure), not to scale. Left: Capillary of radius \( R = 0.036 \text{ cm} \) with snapshots at intervals of 100 s for the first 2500 s. Right: Capillary of radius \( R = 0.074 \text{ cm} \) with snapshots at intervals of 50 s for the first 600 s.
Following the initial period, where the dynamic contact angle evolves from its initial value and the contact-line speed increases, the flow ‘settles’ into the ‘regular’ quasi-steady pattern. There the velocity-dependence of the contact angle falls onto the one given by the asymptotics outlined in Section 2.2. It is important to point out here that, although the radii of the capillaries considered in this section are large enough for the influence of $R$ on the flow to be negligible, the unsteadiness of the process makes it necessary to use the interface formation model in its entirety and not just the asymptotic result for the speed-angle relationship that follows from the model in the limiting case. The latter becomes applicable at the later stages of the imbibition, whereas at the onset both (22) and any of the velocity-dependencies of the contact angle used in the conventional models will make the contact-line speed jump from zero straight up to a large finite value.

7. Concluding remarks

The necessity to develop a robust scheme of incorporating the model of dynamic wetting as an interface formation process into a numerical code is dictated by the need to describe the experimentally observed effect of the dynamic contact angle being dependent on the entire flow field/geometry. This effect becomes more pronounced as the size of the system goes down and has far-reaching implications for a host of emerging technologies.

In this paper, such a scheme is described and the resulting numerical code is thoroughly validated. The exposition is purposefully laid out in a ‘digestible’ manner, with specific details given in the Appendix, allowing an interested reader to reproduce the code and compare the results with the benchmark calculations provided and, together with our previous publication [32], giving the first guide for simulating fluid flows with forming interfaces. A particular emphasis is put on the estimates for the required mesh resolution and the physics associated with the smallest length scales for different
regimes. These estimates as well as the benchmark calculations remain valid independently of the particular numerical scheme used.

The considered test problem of a steady imbibition into a capillary highlights a unique feature of the interface formation model, namely that the dynamic contact angle is an output, part of the solution, as opposed to being a prescribed function of the contact-line speed and other parameters. For this test problem, our computations show that, as the radius $R$ of the capillary is reduced, the velocity-dependence of the dynamic contact angle becomes influenced by $R$, with the dynamic contact angle for the same contact-line speed being higher for smaller radii. This is an essentially new physical effect predicted using the interface formation model. Given that, when the influence of the flow field/geometry was first discovered experimentally [19], the term ‘hydrodynamic assist of dynamic wetting’ was coined to emphasize the possibility of lowering the dynamic contact angle by manipulating the flow field, here we may term the newly discovered phenomenon as ‘hydrodynamic resist to dynamic wetting’. The focus of this paper is on numerical implementation of the interface formation model and so we postpone the full exploration of the effect of ‘hydrodynamic resist’ to a future publication in which our framework will be compared to experiments, e.g. [21], where resist-like behaviour has been observed but is yet to be theoretically described.

The problem of unsteady imbibition into a capillary demonstrates the above unique feature of the interface formation model from a different perspective. This problem shows that the velocity-dependence of the dynamic contact angle bears the influence of the initial conditions even in the situation where otherwise the effect of the flow field/geometry would be negligible. In particular, in the unsteady flow the dynamic contact angle can decrease with the increasing contact-line speed. Qualitatively, this possibility follows from a general argument about the continuity of variation of the flow characteristics necessary to avoid unphysicality, and the interface formation model provides a quantitative illustration of how this unphysicality is removed.

With regard to the discovered effect, it is worth pointing out that the familiar to ‘practitioners’ of dynamic wetting angle-vs-speed curves showing an increase of the dynamic contact angle with the contact-line speed come from experiments with steady flows and, with some variants, a few experiments with unsteady flows with low temporal resolution, with the characteristic time scales much larger than that of the surface tension relaxation. By contrast, the effect described in this work is essentially unsteady and requires special experimental techniques to be captured. In this regard, electrical measurements that are being utilized to probe the coalescence of two liquid drops, e.g. [67], have demonstrated an ability to probe spatio-temporal scales inaccessible to optical measurements, and they have already led to the discovery of new flow regimes. If these methods could be adapted to consider dynamic wetting phenomena, where optical measurement suggest the existence of new regimes at short time scales [68], then the effects discovered here could be experimentally verified.

The validation of our numerical code against experimental data, demonstrating excellent agreement without the need to fit any parameters, opens up two lines of inquiry. First, it would be interesting to consider how scaling down of the system towards submicron and nanoscales affects the flow and, by comparing the results with experimental data, identify the scales where specific ‘nanoeffects’ become important. Secondly, the code can be used to validate different simplified models for the capillary imbibition by identifying the regions of parameters where different effects play the dominant role.

What has been shown is only a small part of the capabilities of the interface formation model fully incorporated into a numerical framework. In [34], the dynamics of freely oscillating liquid drops and of drops impacting and spreading on chemically patterned surface is considered, demonstrating the framework’s capability to simulate flows in which changes in free-surface shape are extreme. The ability of the interface formation model to incorporate additional physical effects is shown in [42,69], where a viscous flow over a solid surface whose wettability varies has been described, with the variations in wettability affecting the flow even in the absence of any meniscus. Further influences on the wetting dynamics, such as those associated with the solid substrate’s properties, e.g. its porosity [70], or thermal and electromagnetic effects, can be added as required, safe in the knowledge that the underlying framework’s accuracy has been completely confirmed.

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Appendix A

In [32], a user-friendly step-by-step guide to the finite element simulation of dynamic wetting flows using the conventional models is given. Many of the aspects dealt with in that paper remain unchanged when we introduce the interface formation model into this framework, and these are not repeated here. In particular, we do not repeat: the re-writing of tensorial expressions in terms of a particular coordinate system, the mesh design which utilizes the bipolar coordinate system, the numerical scheme used.
system, the mapping of integrals onto a master element, the calculation of the corresponding integrals using Gaussian quadrature or the solution procedure based on the Newton–Raphson method. Here, the main change is that we are now solving many more equations along the boundaries of the domain, and, consequently, the element-level residuals change from those in [32] and are thus listed in full. Therefore, in parallel with [32], this guide enables one to easily reproduce all the results presented in the paper. For a more detailed exposition about using the FEM to solve fluid flows the reader is referred to [71,72], whilst specific information on free surface flows, including alternative mesh designs, can be found in [73–76,57]. We have provided information of our own specific implementation of the framework described in the main body of the paper. Many alternatives exist (e.g. different element types) which may produce equally accurate results but here we present what we have used and tested, which for a ‘practitioner’ gives a ready-to-use algorithm.

Two-dimensional and three-dimensional axisymmetric flows are considered simultaneously by using a variable n which takes the values n = 0 in the former case and n = 1 in the latter. Two-dimensional flow occurs in a Cartesian coordinate system (r, z) whilst axisymmetric flow is in a cylindrical polar coordinate system (r, z, ), where r is now the radial coordinate and  is the azimuthal angular coordinate around the z-axis about which symmetry is assumed (e.g. Fig. 2). For both coordinate systems, the governing equations are solved in the (r, z)-plane. In this coordinate system, the surface velocity is \( \mathbf{v}^t = \nu^t \mathbf{t} + \nu^s \mathbf{n} \), and \( \mathbf{v}^s = \nu^s \mathbf{t} \) as, in the n = 1 case, \( \mathbf{v}^s \cdot \mathbf{e}_s = 0 \). Then, notably, the divergence of the tangential component of a surface vector is given by

\[
\nabla^s \cdot \mathbf{a}^s = \mathbf{t} \cdot \frac{\partial (a^s_i \mathbf{t})}{\partial s} + \frac{na^s_i}{r} = \frac{\partial a^s_i}{\partial s} + \frac{na^s_i}{r}
\]

where we have used that \( \mathbf{t} \cdot \mathbf{a}^s = 0 \) and \( a^s_i = a^s \cdot \mathbf{e}_s \).

A.1. Finite element procedure

The finite element method presented here is not restricted to the capillary geometry and our only assumption about the free surface is that in the (r, z)-plane, i.e. the computational domain, it is a line parameterized by arclength s with position defined by a function of one variable \( h = h(s) \) which depends on the particular mesh design. In other words, to determine the free-surface shape we must determine the value of \( h \) at every node on the free surface \( i = 1, \ldots, N_f \).

The domain is tessellated into \( e = 1, \ldots, E \), non-overlapping subdomains of area \( E_e \), the elements, which in our case will be two-dimensional curved-sided triangles whose positions are defined by the spines of the mesh and hence by the values of free surface unknowns \( h \). The boundary of the computational domain is composed of one-dimensional elements of arclength \( s_e \) formed from the sides of the bulk elements which are adjacent to the boundary. We refer to element-level quantities as local and those defined over the entire domain, which were used in Section 3, as global. Each element contains a set number of local nodes, and it is the set of all local nodes which form the global nodes referred to in Section 3. We have used Roman letters (i, j) for global quantities and italicized letters (i, j, k, l) for local ones. In the FEM, one must store a function \( f \) which relates each local node \( i \) in each element \( e \) to its global node number \( i \) so that \( i = I(e, i) \).

The global functions and residuals are constructed in a piecewise manner from local quantities, by ensuring that, as required in the FEM, the interpolating function associated with a given node is constructed to be zero outside the elements to which that node belongs. This allows us to derive expressions for the residuals in an arbitrary element which, after summing up contributions from every element in the domain, will form a set of algebraic equations whose coefficients are integrals to be calculated by Gaussian integration as shown in [32].

To avoid having to construct each local interpolating function in an element-specific manner, they are constructed on a master element with coordinates \( (\xi, \eta) \), so that \( \phi_i = \phi_i(\xi, \eta), \psi_j = \psi_j(\xi, \eta) \). Mixed interpolation is achieved, to ensure the Ladyzhenskaya–Babuška–Brezzi [52] condition is satisfied, by using the V6P3 Taylor–Hood triangular element which approximates velocity by means of bi-quadratic local interpolating functions \( \phi_i(\xi, \eta) \) \((i = 1, \ldots, 6)\) and pressure using bi-linear ones \( \psi_i(\xi, \eta) \) \((i = 1, \ldots, 3)\), with explicit expressions given in [32]. Over an element \( e \) the following functional forms are used

\[
(u, w) = \sum_{j=1}^{6} (u_j, w_j) \phi_j(\xi, \eta), \quad p = \sum_{j=1}^{3} p_j \psi_j(\xi, \eta), \quad (r, z) = \sum_{j=1}^{6} (r_j, z_j) \phi_j(\xi, \eta).
\]

By ensuring that elements whose sides form the free surface always have local nodes 2, 6, 3 associated with them and that those on the solid surface are always \( j = 1, 5, 2 \), we can define surface interpolating functions \( \phi_{1,5}(\eta) = \phi_1(\xi = -1, \eta) \) and \( \phi_{2,6}(\eta) = \phi_5(\xi = 1, \eta) \). Then, a surface variable \( a^s \) and the free-surface shape \( (r_1(h), z_1(h)) \), which is determined from the free surface unknowns \( h \), are approximated on the surfaces quadratically as

\[
a^s_i = \sum_{j=2,6,3} a^s_{ij} \phi_{1,5}(\xi), \quad (r_1, z_1) = \sum_{j=2,6,3} (r_{1j}, z_{1j}) \phi_{1,5}(\xi), \quad a^s_2 = \sum_{j=1,5,2} a^s_{2j} \phi_{2,6}(\eta).
\]

The global residuals (the \( R_e \)‘s) presented in Section 3, which involve integrals over the entire domain, are formed by summing up local residuals (the \( R_e \)‘s), denoted with a subscript \( e \) to indicate which element the local residual is calculated in, obtained by integrating over each of the \( N_e \) elements in the global domain. Let \( N_e \) be the set of numbers of those elements that form part of the free surface and \( N_f \) be the numbers of elements forming a part of the liquid–solid interface. Then, the continuity of
mass, momentum equations,\(^5\) kinematic equation on the free surface and impermeability equation at the liquid–solid interface are:

\[
R^c_i = \sum_{e=1}^{N_e} \sum_{i=1}^{3} R^c_{e,ij}, \quad R^{M,X}_i = \sum_{e=1}^{N_e} \sum_{i=1}^{6} R^{M,X}_{e,ij}, \quad R^k = \sum_{e=1}^{N_e} \sum_{j=1}^{N_{2,e}} R^k_{e,j}, \quad R^l_i = \sum_{e=1}^{N_e} \sum_{j=1}^{N_{2,e}} R^l_{e,j},
\]

where the constraint under the summation symbols ensures that, through \(i = l(e,i)\), the local residuals are assigned to the correct global ones and, via \(e \in N\), that the surface equations are only evaluated when an element is adjacent to the boundary of the domain. In addition, we now have residuals from the interface formation equations given by

\[
R^{\nu}_{i} = \sum_{e=1}^{N_e} \sum_{i=1}^{2} R^{\nu}_{e,ij}, \quad R^{\nu}_{i} = \sum_{e=1}^{N_e} \sum_{i=1}^{2.6} R^{\nu}_{e,ij}, \quad R^{\nu}_{i} = \sum_{e=1}^{N_e} \sum_{i=1}^{2.6} R^{\nu}_{e,ij}, \quad R^{\nu}_{i} = \sum_{e=1}^{N_e} \sum_{i=1}^{2.6} R^{\nu}_{e,ij},
\]

and the residual from Young’s equation \(R^y\) which is only applied at the contact line node.

Taking the equations of Section 3 over an arbitrary element \(e\), with area \(A_e\) and boundary \(s_e\), we now derive the local residuals required in (54) and (55).

### A.2. Element-level residuals

For an arbitrary element \(e\), with local nodes \(i = 1, \ldots, 6\), contributions from the bulk of the element \(E_e\) to the momentum residuals are

\[
R^{M,T}_{e,ij} = Re \left[ M_{ij} \frac{du_j}{dt} + A_{ij} \left( u_j - \frac{dr_j}{dt} \right) \right] + K^{ij}_{y} u_j + K^{12}_{y} w_j + C^{1}_{ik}\nu k,
\]

\[
R^{M,Z}_{e,ij} = Re \left[ M_{ij} \frac{dw_j}{dt} + A_{ij} \left( w_j - \frac{dz_j}{dt} \right) \right] + K^{ij}_{y} u_j + K^{12}_{y} w_j + C^{2}_{ik}\nu k + St \ G_i, \quad j = 1, \ldots, 6 \quad k = 1, \ldots, 3,
\]

where summation over repeated indices is henceforth assumed. The \(M\) terms are the mass matrices, the \(A\) terms are from the nonlinear convective terms, the \(K\) terms are associated with viscous forces, the \(C\) terms are with pressure forces and \(G\) terms correspond to the body force acting on the liquid due to gravity. Temporal derivatives will be considered in Section A.3.

For \(i = 1, \ldots, 3\) we have the incompressibility residuals

\[
R^c_{e,j} = C^1_{i\nu} u_j + C^2_{i\nu} w_j \quad j = 1, \ldots, 6.
\]

If an element \(e\) forms a part of the free surface, \(e \in N_t\), then for \(i = 2, 6, 3\), i.e. the free surface nodes, there are additional contributions to the momentum equations from capillary stress terms, so that

\[
R^{M,T}_{e,ij} = R^{M,T}_{e,ij} + \frac{1}{Ca} F^i_{j} \sigma_{ij} - I_{n}^{j} \nu_{p_{e},j} \quad j = 2, 6, 3,
\]

\[
R^{M,Z}_{e,ij} = R^{M,Z}_{e,ij} + \frac{1}{Ca} F^j_{i} \sigma_{ij} - I_{n}^{j} \nu_{p_{e},j} \quad j = 2, 6, 3,
\]

where \(p_{e} \) is the nodal value of the gas pressure, which in this work has been assumed to remain constant. If an element \(e\) forms the part of the free surface, \(e \in N_{t,2}\), adjacent to the contact line, so that local node \(i = 2\) is the contact-line node, then there is an additional contribution

\[
R^{M,T}_{e,2} = R^{M,T}_{e,2} + \frac{1}{Ca} T^1 \sigma_{1,2},
\]

\[
R^{M,Z}_{e,2} = R^{M,Z}_{e,2} + \frac{1}{Ca} T^2 \sigma_{1,2},
\]

which is where the contact angle is imposed into the weak formulation. The contact angle is determined from the Young equation, now applied only at the contact–line node, whose residual is given by

\[
R^y = \sigma_{1,2} \cos \theta_{d} + \sigma_{2,1,2}.
\]

Additionally, for \(i = 2, 6, 3\) one has the kinematic equation

\[
R^{k}_{e,j} = I_{n}^{j} \frac{dr_{j}}{dt} + I_{n}^{j} \frac{d\nu_{j}}{dt} - l_{j} \nu_{n_{j}} \quad j = 2, 6, 3.
\]

\(^5\) Note the typo in [32] where the limit for \(i\) in \(R^c_{e,1}\) and \(R^{M,X}_{e,1}\) should be 3 and 6, respectively.
If an element $e$ forms a part of the liquid–solid interface, $e \in \mathcal{N}_2$, then for $i = 1, 5, 2$ there are additional contributions to the momentum equations of

$$R_{e1}^{M} = R_{e1}^{M, r} + \frac{\hat{\beta}}{C_1} \left( N_{ij}^{1} (u_j - U_j) + N_{ij}^{2} (w_j - W_j) \right) + \frac{1}{2Ca} \left( S_{ij}^{1} \sigma_{ij, r} \right) + \frac{1}{2Ca} \left( S_{ij}^{2} \sigma_{ij, r} \right) \quad j = 1, 5, 2,
$$

$$R_{e2}^{M} = R_{e2}^{M, r} + \frac{\hat{\beta}}{C_1} \left( N_{ij}^{1} (u_j - U_j) + N_{ij}^{2} (w_j - W_j) \right) + \frac{1}{2Ca} \left( S_{ij}^{1} \sigma_{ij, r} \right) + \frac{1}{2Ca} \left( S_{ij}^{2} \sigma_{ij, r} \right) \quad j = 1, 5, 2,
$$

(60)

where the $N$ terms come from the slip terms in the generalized Navier condition (33), whilst the $S$ terms are associated with gradients in surface tension in (33) and the terms containing $\Lambda$ are from the normal stress on the interface. The substrate speed$^6$ has been decomposed into scalar components as $\mathbf{U} = \mathbf{U}_e + \mathbf{W}_e$. Additionally, for $i = 1, 5, 2$ one has the impermeability Eq. (46) in the form

$$R_{e1}^l = I_{ij} \nu_{ij} - \left( I_{ij} U_j + I_{ij}^2 W_i \right) \quad j = 1, 5, 2.
$$

Moving on to the equations of the interface formation model, which do not appear in [32], the Darcy-type Equations (36) and (37) give

$$R_{e1}^{v_{ij}} = I_{ij} \nu_{ij} - \left( I_{ij} U_j + I_{ij}^2 W_i \right) - \frac{1 + 4 \frac{\beta}{\beta}}{4 \frac{\beta}{\beta}} D_0 \sigma_{ij}, \quad i, j = 2, 6, 3,
$$

$$R_{e2}^{v_{ij}} = I_{ij} \nu_{ij} - \frac{1}{2} \left[ I_{ij}^2 (u_j + U_j) + I_{ij}^2 (w_j + W_j) \right] - 2D_0 \sigma_{ij}, \quad i, j = 1, 5, 2.
$$

The surface velocity normal to each interface $\gamma = 1, 2$ is determined from

$$R_{e1}^{v_{ij}} = \left( I_{ij} U_j + I_{ij}^2 W_i \right) - I_{ij} v_{ij} - Q_{ij} \left( \rho^s_{ij} - \rho^e_{ij} \right) \quad \gamma = 1: i, j = 2, 6, 3, \quad \gamma = 2: i, j = 1, 5, 2
$$

where we have allowed for the extension to problems in which $\rho^e$ varies spatially.

The terms appearing in the surface mass balance Eq. (41) are more complicated due to the presence in this equation of temporal derivatives and the divergence of surface vectors. First, recall that on each surface $\gamma = 1, 2$ the finite element surface mass balance equation is

$$R_{e}^{c_{ij}} = \left( R_{e1}^{c_{ij}} \right)_{h_1} + \left( R_{e2}^{c_{ij}} \right)_{c_1}
$$

\begin{align*}
\left( R_{e1}^{c_{ij}} \right)_{h_1} &= \int_{h_1} \left[ -\epsilon \rho^c_{ij} \left( \mathbf{v}_{ij} - \mathbf{c}_{ij} \right) \cdot \nabla \phi_{ij, l} + \epsilon \phi_{ij, l} \left( \frac{\partial \rho^c_{ij}}{\partial t} + \rho^c_{ij} \nabla \cdot \mathbf{c}_{ij} + \rho^c_{ij} \mathbf{v}_{ij} \cdot \mathbf{n} \right) \cdot \nabla \mathbf{c}_{ij} + \phi_{ij, l} \left( \rho^e_{ij} - \rho^c_{ij} \right) \right] dA_{ij} \\
\left( R_{e2}^{c_{ij}} \right)_{c_1} &= -\int_{c_1} \epsilon \phi_{ij, l} \rho^c_{ij} \left( \mathbf{v}_{ij} - \mathbf{c}_{ij} \right) \cdot \mathbf{m} dC_{ij} \quad (i = 1, \ldots, N_\gamma).
\end{align*}

(61)

Before deriving the individual terms, it proves convenient to introduce the component of the surface’s mesh velocity tangential to that surface as a new variable which we label $c_{ij}^t$. This is discretized in the usual way on each surface $\gamma = 1, 2$

$$c_{ij}^t = \sum_{j} c_{ij}^t \phi_{ij, l} \quad \gamma = 1: j = 2, 6, 3, \quad \gamma = 2: j = 1, 5, 2
$$

This new variable is in the $(r, z)$-plane, so that $c_{ij}^t = c_{ij}^t \cdot \mathbf{t}$ and is thus defined by the finite element equation

$$R_{e1}^{c_{ij}} = I_{ij} \frac{dc_{ij}^t}{dt} - \left( I_{ij} \frac{dr_{ij}}{dt} + I_{ij} \frac{dz_{ij}}{dt} \right) \quad \gamma = 1: i, j = 2, 6, 3, \quad \gamma = 2: i, j = 1, 5, 2
$$

The first of the three nonlinear terms in (61) is

$$\int_{s_{\gamma e}} \frac{\partial \phi_{ij, l}}{\partial s} \rho^c_{ij} \left( \mathbf{v}_{ij} - c_{ij}^t \right) r^a ds_{\gamma e} = \rho^c_{ij} \left( \mathbf{v}_{ij, l} - c_{ij}^t \right) \int_{s_{\gamma e}} \frac{\partial \phi_{ij, l}}{\partial s} \phi_{ij, k} r^n ds_{\gamma e} = p_{ij} \rho^c_{ij} \left( \mathbf{v}_{ij, l} - c_{ij}^t \right) = P_{ij} \left( \mathbf{v}_{ij, l} - c_{ij}^t \right) \rho^c_{ij, l}.
$$

(62)

The second one is

$$\int_{s_{\gamma e}} \phi_{ij, l} \rho^c_{ij} \nabla \cdot \mathbf{c}_{ij}^t r^a ds_{\gamma e} = \rho^c_{ij} \int_{s_{\gamma e}} \phi_{ij, l} \phi_{ij, l} \frac{\partial \phi_{ij, k}}{\partial s} r^n ds_{\gamma e} + \rho^c_{ij} \frac{dr_{ij}}{dt} \int_{s_{\gamma e}} \phi_{ij, l} \phi_{ij, k} \phi_{ij, k} ds_{\gamma e} = w_{ij} \rho^c_{ij} \frac{dc_{ij}^t}{dt} + \frac{nw_{ij} \rho^c_{ij}}{dt} = W_{ij} (c_{ij}^t, r_{ij}) \rho^c_{ij}.
$$

(63)

$^6$ Which should also have been present in [32].
The third term gives
\[ \int_{s_{\gamma}} \phi_{ij} \rho_i^* v_{in} \nabla v^* \cdot n \, r^n \, ds_{\gamma} = \rho_{ij}^* v_{m,k}^* \int_{s_{\gamma}} \phi_{ij} \phi_{j,k} k \tau d s_{\gamma} = \gamma_{ij} \rho_{ij}^* v_{m,k} = \gamma_{ij} (v_{m,k}^* \rho_{ij}^*), \]
where \( \kappa = t_{\gamma} \frac{\partial n_{\gamma}}{\partial x} + t_{\gamma z} \frac{\partial n_{\gamma}}{\partial z} + n(n_{\gamma}^2 / r) \) is the curvature.

Finally, this gives the surface mass balance equation as
\[ \left( K_{ij}^T \right)_{\gamma} = \epsilon \left[ \frac{d \rho_{ij}^*}{dt} - p_{ij} \rho_{ij}^* + W_{ij} \rho_{ij} + Y_{ij} \rho_{ij}^* \right] + I_0 \left( \rho_{ij}^* - \rho_{ij} \right) \]
Whenever the interface reaches the contact line, where now \( m_1 = t_1 \) and \( r = r_c \), we have an additional contribution on both the free surface and solid surface sides as:
\[ \left( R_{ij} \right)_{C_{\gamma}} = -\epsilon \left[ \rho_{ij}^* (v_{ij}\hat{t}_{i,j} - c_{\gamma}^i v_{ij} \hat{t}_{i}) r^n \right] \]
Lastly, the equations of state (46) are given by
\[ R_{ij}^o = I_1 \sigma_{ij} - \lambda (I_1 - I_0 \rho_{ij}^*) \]
where \( I_1 = \sum I_{ij} \).

A.2.1. Element-level matrices

It should be pointed out that, in contrast to [32], in this paper non-dimensional parameters are not contained in the element-level matrices below as they have already been factored out.

The mass matrix is
\[ M_{ij} = \int_{E_i} \phi_i \phi_j r^n \, dE_e. \]
and temporal derivatives are handled in Section A.3.

The viscous terms are
\[ K_{ij}^{11} = 2k_{j11} + k_{j22} + 2nk_j, \quad K_{ij}^{12} = k_{j21}, \quad K_{ij}^{21} = k_{j12}, \quad K_{ij}^{22} = k_{j11} + 2k_{j22}, \]
where
\[ k_{ijkl} = \int_{E_i} \frac{\partial \phi_i}{\partial y_k} \frac{\partial \phi_j}{\partial y_l} r^n \, dE_e, \quad k'_j = \int_{E_i} \phi_i \phi_j r^n \, dE_e, \]
with \( y_1 = r, \ y_2 = z \). Nonlinear convective terms are
\[ A_{ij}(u, w) = a_{ijk1} u_k + a_{ijk2} w_k \quad k = 1, \ldots, 6, \]
where
\[ a_{ijkl} = \int_{E_i} \phi_i \phi_k \frac{\partial \phi_j}{\partial y_l} r^n \, dE_e. \]
The C matrices represent pressure terms whilst their transpose is required in the continuity of mass equation:
\[ C_{ij}^1 = -\int_{E_i} \phi_i \left( \frac{\partial \phi_j}{\partial r} + \frac{n \phi_j}{r} \right) r^n \, dE_e, \quad C_{ij}^2 = -\int_{E_i} \phi_i \frac{\partial \phi_j}{\partial z} r^n \, dE_e. \]
The gravitational force contributes through the G term and is given by
\[ G_i = \int_{E_i} \phi_i r^n \, dE_e. \]
On a free surface (one-dimensional) element \( s_{1e} \), the capillary stress terms\(^7\) are
\[ F_{ij}^1 = \int_{s_{1e}} \left( t_{11} \frac{\partial \phi_{11}}{\partial s} + \frac{n \phi_{11}}{r} \right) \phi_{11} r^n \, ds_{1e}, \quad F_{ij}^2 = \int_{s_{1e}} t_{12} \frac{\partial \phi_{12}}{\partial s} \phi_{11} r^n \, ds_{1e}. \]
The components of the vectors tangential \( t = (t_r, t_z) \) and normal \( n = (n_r, n_z) \) to surface \( \gamma = 1, 2 \) are found in the usual way, see [32].

At the contact line \( (r_c, z_c) \) contributions will occur at local node \( i = 2 \) of the first free surface element so that
\[ T^1 = \left[ (t_{2r} \cos \theta_2 + n_{2r} \sin \theta_2) r^n \right], \quad T^2 = \left[ (t_{2z} \cos \theta_2 + n_{2z} \sin \theta_2) r^n \right]. \]

\(^7\) Note that there was a typo in [32].
On a (one-dimensional) element $S_{2e}$ on the liquid–solid interface, the tangential stress terms from the Navier–slip boundary condition for $k,l = 1,2$ are

$$\mathbf{N}_{ij} = \int_{S_{2e}} \phi_{2i}\phi_{2j}t_{2k}t_{2l}r^m \, ds_{2e},$$

with $t_{21} = t_{22}$ and $t_{22} = t_{22}$. The terms associated with gradients in surface tension are given by

$$S_{ij}^{1} = \int_{S_{2e}} \phi_{2i}\frac{\partial \phi_{2j}}{\partial S} t_{21} r^n \, ds_{2e}, \quad S_{ij}^{2} = \int_{S_{2e}} \phi_{2i}\frac{\partial \phi_{2j}}{\partial S} t_{22} r^n \, ds_{2e}.$$  

On a surface $\gamma = 1.2$ the following other expressions have been used

$$I_{ij}^{1} = \int_{S_{2e}} \phi_{1i}\phi_{1j}r^n \, ds_{\gamma e}, \quad I_{ij}^{2} = \int_{S_{2e}} \phi_{1i}\phi_{1j}n \, ds_{\gamma e},$$

$$l_{ij}^{1} = \int_{S_{2e}} \phi_{1i}\phi_{1j}t_{1l} r^n \, ds_{\gamma e}, \quad l_{ij}^{2} = \int_{S_{2e}} \phi_{1i}\phi_{1j}t_{1j} r^n \, ds_{\gamma e},$$

$$D_{ij} = \int_{S_{2e}} \phi_{1i}\frac{\partial \phi_{1j}}{\partial S} r^n \, ds_{\gamma e}, \quad P_{ij} = p_{ijk} \left( v_{t,k} - c_{t,k} \right),$$

where

$$p_{ijk} = \int_{S_{2e}} \frac{\partial \phi_{1i}}{\partial S} \phi_{1j}\phi_{1k} r^n \, ds_{\gamma e}.$$

$$S_{ijk}^{1} = \int_{S_{2e}} \frac{\partial \phi_{1i}}{\partial S} \phi_{1j}\phi_{1k}t_{1l} r^n \, ds_{\gamma e}, \quad S_{ijk}^{2} = \int_{S_{2e}} \frac{\partial \phi_{1i}}{\partial S} \phi_{1j}\phi_{1k}t_{1j} r^n \, ds_{\gamma e},$$

$$W_{ij} = w_{ijk}c_{t,k} + nw_{ijk} \frac{dt_{z,k}}{dr},$$

where

$$w_{ijk} = p_{ijk}, \quad w'_{ijk} = \int_{S_{2e}} \phi_{1i}\phi_{1j}\phi_{1k} ds_{\gamma e},$$

$$Y_{ij} = y_{ijk} v_{t,n,k},$$

where

$$y_{ijk} = \int_{S_{2e}} \phi_{1i}\phi_{1j}\phi_{1k} \frac{\partial r^n}{\partial S} \, ds_{\gamma e}.$$

A.3. Temporal discretization

The result of our spatial discretization is a system of non-linear differential algebraic equations (DAEs) of index 2 ([72]). However, it is well known that the same methods that apply to ODEs can be used for DAEs [77], and for a review of the methods available we refer the reader to [72].

The second-order backward differentiation formula (BDF2), which has been applied successfully to similar problems [46], is implemented into our scheme. Below, the method is applied to a scalar equation $\dot{y} = f(y,t)$, with the extension to derivatives in the Navier–Stokes and interface formation equations being a straightforward task. For a time $(n + 1)$ with step $\Delta t$, the method applied to the scalar equation gives

$$\frac{y_{n+1} - y_n}{\Delta t} = \frac{1}{3} y_n - \frac{2}{3} y_{n-1} + \frac{2}{3} \dot{y}_{n+1},$$

where the subscript indicates the time step at which a variable is evaluated and $dy/dt = \dot{y}$. Alternatively, it may be written as

$$3y_{n+1} - 4y_n + y_{n-1} = 2\Delta t \dot{y}_{n+1},$$

which is the second-order accurate one-sided Taylor series expansion of $y$ at $t_{n+1}$.
Often, during a physical process there will be different stages that are characterized by different time scales. It is important that our temporal discretization takes these different scales into account so that the largest possible time step, that enforces a certain accuracy, is chosen automatically. This is achieved by choosing a ‘step’ so that the local truncation error \( d_n = y(t_{n+1}) - y(t_n) \), where \( y(t_{n+1}) \) is the exact solution, is maintained below a certain tolerance. By using an explicit second-order Adams–Bashforth method (AB2) \[71\] to predict the solution \( y^p_{n+1} \), and then comparing the difference between the actual solution and the predicted one, after solving the non-linear equations we are able to deduce this error. For the equation, \( y = f(y, t) \), having obtained the solution \( y_{n+1} \) we follow the analysis of \[71\] to obtain a new time step. The BDF2 for variable step size gives

\[
\frac{y_{n+1} - y_n}{\Delta t_n} = \frac{\Delta t_n}{2 \Delta t_n + \Delta t_{n-1}} \left( \frac{y_n - y_{n-1}}{\Delta t_n} + \frac{\Delta t_n + \Delta t_{n-1}}{2 \Delta t_n + \Delta t_{n-1}} \frac{y_n - y_{n-1}}{\Delta t_{n-1}} \right),
\]

(64)

while the predictor gave

\[ y^p_{n+1} = y_n + \left( 1 + \frac{\Delta t_n}{\Delta t_{n-1}} \right) \frac{\Delta t_n}{2 \Delta t_n + \Delta t_{n-1}} \left( \frac{y_n - y_{n-1}}{\Delta t_{n-1}} \right)^2. \]

The local truncation error of (64) is given by

\[
d_n = \frac{(\Delta t_n + \Delta t_{n-1})^2}{\Delta t_n(2 \Delta t_n + \Delta t_{n-1})} \frac{\Delta t_n^2}{6} + O(\Delta t_n^4),
\]

(65)

whilst the predictor’s error is

\[
y^p_{n+1} - y(t_{n+1}) = - \left( 1 + \frac{\Delta t_n}{\Delta t_{n-1}} \right) \frac{\Delta t_n}{2 \Delta t_n + \Delta t_{n-1}} \left( \frac{y_n - y_{n-1}}{\Delta t_{n-1}} \right)^2 + O(\Delta t_n^4). \]

(66)

The exact solution’s contribution \( y(t_{n+1}) \), which, of course, in general will not be known, is eliminated from (65) and (66) to give

\[
d_n = \frac{(1 + \Delta t_{n-1}/\Delta t_n)^2}{1 + 3(\Delta t_{n-1}/\Delta t_n) + 4(\Delta t_{n-1}/\Delta t_n)^2 + 2(\Delta t_{n-1}/\Delta t_n)^3} (y_{n+1} - y^p_{n+1}),
\]

so that the error is linearly proportional to the difference between the predicted and actual solution. This error estimate is then used to compute the next step size

\[
\Delta t_{n+1} = \Delta t_n \epsilon/ \|d_n\|^{1/3}, \quad \|d_n\|^2 = d_n^T d_n/(N\epsilon_{\max}^2).
\]

Here, \( N \) is the total number of nodes, \( \epsilon_{\max} \) is an estimate of the maximum value of \( y \) in the domain and \( \epsilon \) is the relative error tolerance parameter. Then the error of the approximate solution is bounded by \( \|d_{n+1}\| \leq \epsilon \epsilon_{\max} \). This allows us to choose the largest possible time step whilst ensuring that the error of the temporal integration remains below a chosen tolerance.

The variable step method outlined above can be extended for use with the Navier–Stokes and interface formation equations and the reader is referred to [72, p. 797], for details. In this book a number of ‘rules of thumb’ are suggested based on the ratio of the new and previous step sizes, the so-called Delta T scale factor, \( \text{DTSF} = \frac{\Delta t_{n+1}}{\Delta t_n} \), and these can be adapted to the problem of interest as required.

### A.4. Far-field velocity profile for the steady propagation of a meniscus through a capillary

To ensure mass is conserved in our domain for this steady problem in which an adsorption-desorption process will occur along the interfaces, we must derive appropriate conditions at the base of the capillary, i.e. in the truncated far field. Assuming the problem is steady, so that \( \epsilon = \frac{\partial}{\partial r} = 0 \), noting that there is no surface mass flux through the axis of symmetry and that there is no sink or source of mass at the contact line, we have

\[
0 = \int_V \nabla \cdot \mathbf{u} \, dV = \int_A \mathbf{u} \cdot \mathbf{n} \, dA = \int_A \mathbf{u} \cdot \mathbf{n} \, dA_{\text{ff}} + Q \int_{A_1} (\rho^1 \dot{v}_1 - \rho^1 v_1) \, dA_1 + Q \int_{A_2} (\rho^2 \dot{v}_2 - \rho^2 v_2) \, dA_2
\]

\[
= \int_{s_f} w \, dA_{\text{ff}} - \epsilon Q \left[ \int_{s_1} \nabla \cdot \left( \rho^1 v_{1|_1} \right) \, dA_1 + \int_{s_2} \nabla \cdot \left( \rho^2 v_{2|_2} \right) \, dA_2 \right]
\]

\[
= (2\pi)^2 \left( \int_{s_f} w \, r \, dr - \epsilon Q \left( -\rho^1 v_{1|_1} \cdot \mathbf{m}_1 \, r_{|_i} - \rho^1 v_{1|_1} \cdot \mathbf{m}_1 \, r_{|_f} - \rho^2 v_{2|_2} \cdot \mathbf{m}_2 \, r_{|_f} - \rho^2 v_{2|_2} \cdot \mathbf{m}_2 \, r_{|_i} \right) \right)
\]

\[
= (2\pi)^2 \left( \int_{s_f} w \, r \, dr + \epsilon Q \rho^2 v_{2|} \big|_{s_f} \right)
\]

with subscripts \( \text{ff} \) referring to far field variables.
Given that \( w = w(r) \) in the far-field, from the equations of motion we see that
\[
\frac{dp}{dz} = \frac{1}{r^2} \left( r^2 \frac{dw}{dr} \right).
\]
and, after integrating once, with \( \frac{dp}{dz} = G \), this gives
\[
\frac{Gr}{n+1} + Ar^{-n} = \frac{dw}{dr}.
\]
Noting that the surface velocity in the far field is \( \frac{dw}{dr} = 0 \) at \( r = 0 \), so that \( A = 0 \), we integrate again to find
\[
w = \frac{Gr^2}{2(n+1)} + B.
\]
Applying the Navier condition \( \frac{dw}{dr} = (\beta/\sigma)(w - W) \) at \( r = 1 \), assuming that in the far field there are no gradients in surface tension, we have that
\[
-\frac{G}{n+1} = \frac{\beta}{\sigma} \left( \frac{G}{2(n+1)} + B - W \right),
\]
so that
\[
B = W - \frac{G}{2(n+1)} \left[ 1 + \frac{2}{\beta/\sigma} \right],
\]
and, finally
\[
w = W + \frac{G}{2(n+1)} \left[ r^2 - 1 - \frac{2}{(\beta/\sigma)} \right].
\]
Now, we calculate the pressure gradient, \( G \), required to maintain a steady flux in and out of the domain for a given \( W \) by calculating
\[
\int_{r=0}^{1} wr^mdr = \left[ \frac{W}{n+1} + \frac{G}{2(n+1)} \left( \frac{r^{n+1}}{n+3} - \frac{r^{n+1}}{n+1} - \frac{2r^{n+1}}{(\beta/\sigma)(n+1)} \right) \right]_{r=0}^{1}
\]
\[
= \frac{W}{n+1} + \frac{G}{2(n+3)(n+1)^2} \left[ n+1 - 3 - \frac{2(n+3)}{(\beta/\sigma)} \right].
\]
Noting that the surface velocity in the far field is \( v_0 = (1/2)(W + W) \), where \( q = \epsilon q_0 \), we have that
\[
0 = \frac{W}{n+1} \left[ \frac{G}{(n+3)(n+1)^2} \left( 1 + \frac{3}{(\beta/\sigma)} \right) + qW - \frac{Gq}{2(n+1)(\beta/\sigma)} \right]
\]
so that
\[
G = \frac{2W(n+1)(1+q)}{2/(n+3) + (2+q)/(\beta/\sigma)}
\]
with \( \bar{q} = q(n+1) \) and therefore finally
\[
w = W + \frac{W(1+\bar{q})}{2/(n+3) + (2+q)/(\beta/\sigma)} (r^2 - 1 - 2/(\beta/\sigma)),
\]
where, in Section 4, we had \( n = 1 \) and \( W = -1 \).

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