Investigation of Wetting Hydrodynamics Using Numerical Simulations

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I. INTRODUCTION

In both nature and in industrial processes, moving contact lines are ubiquitous, and they occur over a wide range of length scales—from microns for the movement of oil ganglia through porous rocks to meters in a variety of coating processes. A proper geometry-free characterization of the dynamic wetting process is therefore crucial if predictive models of these phenomena are to be developed. However, several complications arise when such a goal is attempted, largely because of the disparity between the submicroscopic length scales at which wetting occurs and the macroscopic length scales at which typical experimental observations are made. A further complication is that different local models yield the same dynamics at macroscopically observable levels. Therefore, in order to verify refined models, measurements must necessarily be made on a submicroscopic length scale over which this physics remains important. This is a formidable challenge, partly because rapid changes in interface curvature occur near the three-phase juncture, and sophisticated experimental techniques become necessary to produce adequate resolution of data in that region.

Challenges also exist from a continuum modeling perspective. For a pure liquid, the customary hydrodynamic analyses result in a multivalued velocity field at the dynamic contact line. If the liquid is Newtonian, a stress field is produced that is nonintegrable as the dynamic contact line is approached. One consequence of this stress divergence is an unbounded interface curvature at the dynamic contact line, resulting in the inability to specify a true dynamic contact angle. Because this angle serves as a boundary condition for the differential equation governing the shape of the liquid–fluid interface, an ill-posed problem results. This difficulty is often resolved by breaking the flow domain into two regions. An inner one, in the vicinity of the dynamic contact line, where slip is permitted (thereby removing the source of the double-valued velocity at the dynamic contact line), has dimensions characterized by a slip length $L_S$. An outer region, characterized by a much larger geometry-dependent macroscopic length scale $L_C$, utilizes the no-slip dynamic boundary at the solid surfaces. Singular perturbation solutions to the field equations in each region (ignoring inertial effects) are matched in the overlap region to produce a uniformly valid solution over the entire domain. In the double limit $Ca \rightarrow 0$, $\epsilon \rightarrow 0$, with $Ca \ln(\epsilon^{-1})$ of $O(1)$, where $Ca$ is the Capillary number $\mu U / \gamma$, $\mu$ is the liquid viscosity, and $\gamma$ is its surface tension, $U$ is a characteristic speed, and $\epsilon = L_S / L_C$, there is no overlap of the inner and outer regions, and a third, intermediate region must exist between the inner and outer regions of expansion. In cylindrical polar coordinates, the inner, intermediate, and outer regions are characterized by dimensionless lengths $r' / L_S$, $Ca \ln(r' / L_C)$ and $r' / L_C$, respectively. [Variables with a prime (') superscript are dimensional."

Analyses of the resulting boundary value problem to $O(1)$ yield a solution for the interface shape in the intermediate region of the form (assuming that the displaced fluid has zero viscosity)

$$\theta_{\text{intermediate}}(r') = g^{-1} \left[ g(\theta_E) + Ca \ln(r'/R) \right],$$

where

$$g(\theta) = \int_0^\theta \frac{\phi - \cos(\phi)\sin(\phi)}{2 \sin(\phi)} \, d\phi.$$

Here $\theta(r')$ is the angle between the moving solid and the tangent to the liquid free surface at a distance $r'$ from the...
dynamic contact line, the subscript indicating the value of this angle in the indexed region. In addition, \( \theta_R \) represents this angle at some distance \( R \) located within the intermediate region. As seen in Eq. (1), the slope of the interface within the intermediate region is independent of macroscopic geometry, that is, it does not depend upon the outer length scale \( L_C \). Although the exact magnitude of \( \theta_R \) depends upon the specific inner model, the form of Eq. (1) is independent of it.

If the geometry independence of Eq. (1) could be confirmed, that is, if for a certain \( \text{Ca} \), a single value of \( \theta_R \) can be used to correctly predict complete meniscus shapes for a range of outer length scales, then \( \theta_R \) could be used as a material boundary condition for the outer problem, replacing the true dynamic contact angle boundary condition. Cutting off the domain at \( r' = R \) and providing this slope characterization would permit a solution for all the field variables and the free-surface shape in the outer region without resorting to specifying any of the details of the fluid physics in the inner region (which are currently not well known). Clearly, this would represent a powerful advance in our ability to predict wetting behavior.

Two approaches can be taken to determine if Eq. (1) is valid. The first is to compare experimentally obtained meniscus shapes in the vicinity of the dynamic contact line, that is, in the intermediate region, with predictions from the perturbation analysis. For a given geometry and Capillary number, \( \theta_R \) can be chosen by minimizing the difference, in a least squares sense, between the experimental and analytically predicted meniscus shapes. This approach has been utilized by a number of different investigators.\(^4\)^\(^7\) The geometry-free nature of this expression for the shape of the interface in the intermediate region for \( \text{Ca} \leq 10^{-2} \) has been established.\(^6\) However, some systematic deviations were observed for \( \text{Ca} > 10^{-2} \), which could be attributable either to the difficulty in obtaining accurate meniscus shapes in the proximity of the dynamic contact line or to a possible failure of the \( O(1) \) asymptotic analysis in this Capillary number regime. Clearly, to resolve this issue, more accurate experimental meniscus shape measurements are required in the vicinity of the dynamic contact line. Such experiments, using rods entering liquid baths at various immersion angles (as a means of varying geometry) have now been completed.\(^7\)^\(^8\) Viscous deformation has been observed at distances of the order of a capillary length scale from the dynamic contact line at \( \text{Ca} \geq 10^{-2} \), an observation that has direct implications for the measurement of “apparent” dynamic contact angles in capillary tubes.\(^7\) Experiments conducted at \( \text{Ca} \sim 0.45 \) indicate that the breakdown of the model is caused by the inadequacy of the lowest-order perturbation solution in the geometry-free region, or because of contributions to the shape of the interface from the inner region.\(^8\)

Alternatively, numerical simulations of the boundary value problem can be used to obtain complete meniscus shapes, which can be compared with predictions from the perturbation analyses—this approach is used here. Because calculated meniscus shapes are not tainted by experimental errors, interface slopes accurate to a specified tolerance can be obtained over the whole domain, including the immediate vicinity of the dynamic contact line (at the expense of course, of computation time). Eliminating gravity in the model, the outer length scale becomes the container length, which can be changed easily. Thus, the geometry independence of Eq. (1) can be conveniently and rigorously tested. Furthermore, the simulation can be readily extended to high \( \text{Ca} \), where experimental measurements might become difficult, clearly identifying the range of validity of the perturbation analyses. These simulations can also be used to determine the limiting \( \text{Ca} \) value beyond which viscous effects start infiltrating the outer region, and static capillarity becomes inadequate for describing the outer meniscus shape.

The finite element\(^9\)^\(^\sim\)^\(^11\) and finite difference\(^12\)^\(^\sim\)^\(^13\) methods have been used for simulation of the steady motion of an interface between viscous liquids in a capillary tube. Comparisons of the simulations with experiments have been restricted to predictions of “apparent” contact angles. The goal of our work is to show that numerical simulations are a valid tool for understanding fluid physics in the vicinity of moving contact lines. Therefore, its scope is very different from previous numerical studies of wetting hydrodynamics.

II. THE MODEL

A prototype problem that captures the essential features of a dynamic wetting process is chosen for the numerical simulation, and is illustrated in Fig. 1. It is comprised of an incompressible liquid of infinite depth in a container of length \( L_C \), with the left sidewall moving into the liquid with constant velocity \( \mathbf{U} \). Gravity is ignored in the simulations, so
that \( L_C \) is the appropriate outer length scale. The upper boundary of the liquid, \( h'(x') \), is a free surface, so that its location is not known \textit{a priori}, but is obtained as part of the solution to the transport equations and applicable boundary conditions. A hypothetical lower boundary (shown by the dashed line, \( y' = 0 \)) restricts the computational domain. At this boundary, the flow is assumed unidirectional, with zero \textit{net} volumetric flow, and is sufficiently well removed from the free surface so that any further displacement has no impact on the shape of the free boundary. The volume/width of liquid within this computational domain is \( A \). The static contact angle is \( \theta_5 \).

For a single component Newtonian liquid, the field variables (velocities, pressures, free surface heights, and gas pressure) depend upon the Reynolds number, \( Re \), and the Capillary number, \( Ca \). Inertial effects will be ignored in the numerical simulations presented (\( Re = 0 \)), conforming to the assumptions used in the perturbation analyses.

The dimensionless steady-state governing equations and boundary conditions, including the slip model, are provided in Appendix A.

### III. NUMERICAL SOLUTION TECHNIQUE

Following a well-developed procedure, the Galerkin Finite Element technique is used to discretize the continuity and conservation of linear momentum equations, as well as the kinematic condition.\(^\text{14-16}\) The latter is distinguished in this scheme as the equation to be used for computing the free surface shape. The details leading to the appropriate weak forms of each of these equations are presented elsewhere.\(^\text{16}\)

The contact angle conditions replace the kinematic condition for calculating free surface heights at the contact lines. The unknown pressure in the inviscid gas phase, relative to a datum pressure at the lower right corner in the liquid is obtained using the volume constraint.

The liquid domain is subdivided into \( N_x \times N_y \) elements. Vertical spines originating at the lower hypothetical boundary form element borders, while the ends of these spines are used to represent the free surface. Element corner nodes are located proportionally along these spines. The velocity fields within each element are approximated by nine-node Lagrangian biquadratic basis functions, while the pressure field is approximated by four-node bilinear basis functions. The free surface location is expanded using one-dimensional quadratic basis functions. These choices have proven convergent for Newtonian flows, and provide \( C^0 \) continuity for all variables across interelement boundaries. As is customary in finite element practice, the basis functions are developed on a square parent element in an \((\eta, \xi)\) Cartesian coordinate system. This parent element is transformed onto the deformed quadrilateral element in the real domain through the use of isoparametric mapping. With this mapping, the free boundary coincides with \( \xi = +1 \) for each element bordering the surface. Isoparametric mapping also facilitates evaluation of the unit normal and tangent vectors.\(^\text{14}\) The residuals are calculated using the four-point tensor product Gaussian quadrature.

The discretization results in as many nonlinear algebraic equations as the number of unknowns (the unknowns consist of values of the two components of the velocity, the pressure, and free surface heights at appropriate finite-element nodes, and the gas pressure). This algebraic equation set is solved by Newton’s method. The \textit{linear} equation set to be solved at the \( n \)th Newton iteration is

\[
\overline{\text{corr}}_{\text{R}_1} = -\overline{R}_n \}
\]

where

\[
\overline{\text{corr}}_n \approx s_n + 1 - s_n \}
\]

The elements of vector \( \overline{R}_n \) are the residuals of the weak forms of the equations, the elements of vector \( s_n \) are the values of the unknowns at the \( n \)th Newton iteration, and the Jacobian matrix \( J_n = dR_n / ds_n \). The elements of \( J_n \) are obtained numerically using a one-sided forward difference scheme.\(^\text{15,16}\) At each Newton iteration, Eq. (3) is solved by frontal elimination.\(^\text{17}\) The Newton iterations are stopped when the \( L_2 \) norm and the \( L_\infty \) norm of \( \overline{R}_n \) are below \( 10^{-6} \). All the unknowns are updated at each Newton iteration, producing nearly quadratic convergence. Starting from initial guesses where only the known essential boundary conditions are specified, along with a static meniscus shape and zero values for all variables at all other nodes, convergence is achieved within nine iterations.

The convergence of numerically generated solutions with grid size has been carefully monitored, and details are provided below. This is especially critical in moving contact line problems, because large changes in velocity occur over dimensions comparable to the slip length, and an adequate tessellation must be provided to capture these features. Furthermore, roundoff errors associated with discretization of the transport equations and boundary conditions automatically produce slip, so that the force singularity referred to earlier is evident only when a plot of the force on the moving plate versus mesh size diverges as the mesh size is reduced.\(^\text{16}\) Therefore the mesh must be highly refined in the region around the dynamic contact line; we use a minimum of five nodes within a slip length.

The lower hypothetical boundary is displaced successively larger distances from the free boundary (through adjustment of the volume/width, \( A \)) until the \( L_2 \) norm and \( L_\infty \) norms of the slopes of the free surface from two successive values of \( A \) are \(<10^{-8}\) (note: because the piecewise quadratic expansion used for the free surface shape only guarantees \( C^0 \) continuity at interelement boundaries, slopes are calculated at Gauss points on the element surfaces). Our results indicate that this distance should be a minimum of \( 4L_C \). We have used \( A = 4L_C^2 \) in all the simulations.

The algebraic grid generation scheme, using proportionally spaced spines, proves adequate for generating converged solutions for \( \epsilon \geq 10^{-4} \). The requirement that at least five nodes be placed within a distance of one slip length from the dynamic contact line implies a large disparity between element sizes in different parts of the computational domain. For \( \epsilon < 10^{-4} \), when the free surface nodal locations are perturbed in preparation for evaluation of the appropriate entry in the Jacobian matrix, the smallest elements bordering the free boundary get sufficiently distorted, so that the mapping to the isoparametric domain becomes singular. Furthermore,
variables approach constant values as the mesh is refined. We have used the 26 and 25, and 26. In going from (a) to (c), additional elements have been placed in the vicinity of the dynamic contact line. Here \( \text{Ca} = 10^{-3} \), \( \epsilon = 10^{-3} \), \( \theta_D = \theta_s = 45^\circ \), and \( A = 4L_C \).

the calculations presented here are restricted to surface shapes that are single valued in this simple representation using spines. Therefore this simple discretization scheme will certainly be inadequate at “low” or “high” contact angles, or where viscous deformation causes some part of the free surface to become tangent to a vertical spine. Alternate mesh generation schemes will then be required, but that remains outside the scope of this paper.

IV. RESULTS

As is customary in numerical simulations of transport problems, we demonstrate first that the numerical solution presented is not dependent upon the mesh size. Figure 2 illustrates the location and values of \( v_x \), \( v_{x_{\text{max}}}, v_{x_{\text{min}}} \), and \( v_y \) for successively refined meshes (23×23, 25×25, and 26×26—the last mesh is shown in Fig. 3). These variables approach constant values as the mesh is refined. We have used the 26×26 nonuniform mesh shown in Fig. 3 for the rest of our computations. This results in 6401 unknowns.

Since the location of the intermediate region is not known \textit{a priori}, the numerically generated solution is compared with a composite solution from the perturbation analysis. The outer solution to \( O(1) \), characterized by a constant curvature, is given by

\[
\theta_{\text{outer}}(r') = \omega_0 + 2 \sin^{-1} \left( \frac{g(\theta_R) + \text{Ca} \ln(L'/R)}{2} \right),
\]

where

\[
\omega_0 = g^{-1} \left[ g(\theta_R) + \text{Ca} \ln(L'/R) \right].
\]

The uniformly valid composite solution is obtained by summing Eqs. (1) and (5) and subtracting the common part \( \omega_0 \), and becomes

\[
\theta_{\text{composite}}(r') = g^{-1} \left[ g(\theta_R) + \text{Ca} \ln(L'/R) \right] + 2 \sin^{-1} \left( \frac{r'(\cos \omega_0 + \cos \theta_s) / 2L_C}{2} \right).
\]

For given inner model parameters \( \epsilon \) and \( \theta_D \), and Capillary number, the simulations have been run for a range of outer length scales \( 10L_S < L_C < 2000L_S \). Choosing \( R \) to be 10L_S (\( R \) must be a point within the intermediate region), the value of \( \theta_R \) for each \( L_C \) is obtained by a single parameter least square minimization of the difference between the numerically generated solution and Eq. (6) for \( r_1 < r' < r_2 \). The upper and lower bounds on \( r' \) are intended to limit the fitting to the intermediate region because \( \theta_R \) is an intermediate region parameter. In our simulations, \( r_1 = 5L_S \) and \( r_2 = 0.25L_C \) (we have varied \( r_2 \) by \( \pm 20\% \) and found no significant change in the value of \( \theta_R \)). The quality of the fit, using the \( L_2 \) norm of the numerical and composite solutions, is used to establish an upper bound for the Capillary number, above which this fitting procedure is no longer meaningful.

Computed meniscus shapes for a wide range of \( \text{Ca} \) and a fixed value of container length, \( L_C = 2000L_S \), are shown in Fig. 4. At low \( \text{Ca} \), the interface shape is dominated by surface tension, and is marked by near constant curvature. As \( \text{Ca} \) increases, viscous effects on the meniscus shape become more apparent—a rapid change in shape in the immediate vicinity of the moving contact line is followed by a region of low curvature. In order to obtain a more detailed understanding of the nature of these shapes, meniscus slopes are calculated at Gauss points along the free surface, and compared with the predictions from the perturbation analyses.

Figures 5(a)–5(d) show the numerical solution, the best fit composite solution, as well as the outer and intermediate solutions for \( L_C = 2000L_S \) and a range of \( \text{Ca} \). As illustrated in Figs. 5(a) and 5(b), the fit over the full container length is excellent for \( \text{Ca} = 10^{-2} \). Figure 5(c) reveals that at \( \text{Ca} = 10^{-1} \), the fit within the intermediate region is good, while the numerically generated meniscus shape shows a lower curvature.
in the outer region than that predicted from the analysis. The leading-order perturbation analysis appears to adequately describe the strong viscous effects in the proximity of the contact line, but viscous effects, not accounted for in the leading-order solution, are starting to infiltrate the outer region. At \( \text{Ca} = 3.5 \times 10^{-1} \), there is a complete breakdown of the fit between the numerically generated and the leading-order perturbation solution over the entire domain, as shown in Fig. 5(d). At this \( \text{Ca} \), higher-order corrections to both the intermediate and outer solutions must be included in order to adequately describe meniscus shapes.

The quality of the fit for the numerical and best fit composite solutions (fitted in the intermediate region) for both the intermediate and outer regions at each \( \text{Ca} \), is assessed by calculating the \( L_2 \) norms, using

\[
L_2 \text{ norm} = \sqrt{\sum_{j=1}^{m} (\theta_j \text{ numerical} - \theta_j \text{ composite})^2/m},
\]

where \( m \) is the number of data points used in each region. The results are shown in Fig. 6. For \( \text{Ca} \leq 10^{-2} \), the difference
between the numerical and composite solutions remain small for the intermediate as well as the outer regions. For $10^{-2} < \text{Ca} \leq 10^{-1}$, a good match is maintained in the intermediate, while the fit deteriorates in the outer region. For $\text{Ca} > 10^{-1}$, the fit becomes poor for both regions.

The infiltration of viscous effects into the outer region is further illustrated in Fig. 7, where the pressure in the inviscid gas phase $P_g$, relative to a datum pressure in the liquid at the lower right corner of the domain, is plotted as a function of Capillary Number for various outer length scales $L_C$. The geometry independence of $\theta_R$ is maintained up to $\text{Ca} \sim 10^{-1}$. No real significance can be attached to the apparent geometry dependence for $\text{Ca} > 10^{-1}$, because the leading-order perturbation solution becomes inadequate for this range of Ca.

![FIG. 6. Variation of the pressure in the inviscid surrounding phase (—), relative to an arbitrary datum pressure of 100 at the lower right corner of the domain, with Capillary number. The low Ca asymptote, shown as the inclined dashed line, is given by $P_g \sim \text{Ca}^{-1}(\cos \theta_D + \cos \theta_S)$, while the high Ca asymptote, shown as the horizontal dashed line, is given by $P_g \sim -6\alpha L_C^2 r$. The small constant difference between the simulated and asymptotic results at high Ca is the end effect caused by the presence of the bounding liquid surface. Both asymptotes are independent of $L_C$. The calculated $P_g$ values are also independent of $L_C$, indicating that these results are independent of macroscopic geometry. Viscous effects infiltrate into the outer region for $\text{Ca} > 10^{-1}$.](image1)

![FIG. 7. Variation of the pressure in the inviscid surrounding phase (—), relative to an arbitrary datum pressure of 100 at the lower right corner of the domain, with Capillary number. The low Ca asymptote, shown as the inclined dashed line, is given by $P_g \sim \text{Ca}^{-1}(\cos \theta_D + \cos \theta_S)$, while the high Ca asymptote, shown as the horizontal dashed line, is given by $P_g \sim -6\alpha L_C^2 r$. The small constant difference between the simulated and asymptotic results at high Ca is the end effect caused by the presence of the bounding liquid surface. Both asymptotes are independent of $L_C$. The calculated $P_g$ values are also independent of $L_C$, indicating that these results are independent of macroscopic geometry. Viscous effects infiltrate into the outer region for $\text{Ca} > 10^{-1}$.](image2)

![FIG. 8. Variation of the angle $\theta_R$ as a function of Capillary number for various outer length scales $L_C$. The geometry independence of $\theta_R$ is maintained up to $\text{Ca} \sim 10^{-1}$. No real significance can be attached to the apparent geometry dependence for $\text{Ca} > 10^{-1}$, because the leading-order perturbation solution becomes inadequate for this range of Ca.](image3)
inner model. In order to test this directly, we have completed a set of simulations for $L_C = 200L_S$ and 2000$L_S$ using an entirely different slip behavior,\(^\text{18}\) where the liquid velocity along the plate is given by $v_y = -U[(h(0)-y)/\varepsilon]^2/[(1 + [h(0)-y]/\varepsilon)^2]$. Up to $C_a = 0.1$, the values of $\theta_0$ using this model are geometry independent, and are within $\pm 1^\circ$ of those using the exponential slip model with the same $\varepsilon$.

The geometry independence of $\theta_0$ for $C_a \leq 0.1$ implies that it can be used to replace the true dynamic contact angle as a material boundary condition when modeling fluid flow problems containing dynamic contact lines within this range of $C_a$. Essentially, the troublesome region containing the dynamic contact line can be eliminated, and the modeling can proceed on a truncated domain, to which geometry-free boundary conditions along with the usual hydrodynamic conditions can be applied.

V. CONCLUSIONS

Numerical simulations of a model problem containing a dynamic contact line have been utilized to establish the geometry-independent nature of the slope of the free surface in the immediate vicinity of the moving contactline for $C_a \leq 10^{-1}$. Within this region, local viscous effects have a strong impact on the shape of the free surface, so that the macroscopic geometry does not play a role. The slope of the interface at any point within this region can be used as a material boundary condition for simulation of dynamic wetting problems. For $10^{-2} < C_a < 10^{-1}$, viscous effects infiltrate into the outer region, significantly diminishing the quality of fit between the numerically computed and lowest-order perturbation solutions in this region. For $C_a > 10^{-1}$, higher-order corrections to the perturbation solutions appear necessary for both the intermediate and outer regions to adequately described meniscus shapes.

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APPENDIX: DIMENSIONLESS GOVERNING EQUATIONS

All lengths are scaled with $L_C$, velocities with the plate speed $U$, and the pressure and viscous stresses with $\mu U/L_C$. All variables presented here are dimensionless.

**Continuity:**

\[
\nabla \cdot v = 0, \nonumber
\]

(A1)

Here $v = v_x e_x + v_y e_y$, where $e_x$ and $e_y$ are unit vectors in the $x$ and $y$ directions.

**Conservation of momentum (creeping flow, no gravity):**

\[
-\nabla P + \nabla \cdot \tau = 0. \nonumber
\]

(A2)

\[
\tau = \nabla v + (\nabla v)^T
\]

is the viscous stress tensor.

**1. Boundary conditions**

No slip and no penetration of liquid on all stationary solid walls,

\[
v = 0. \nonumber
\]

(A3)

At the free surface $y = h(x)$,

\[
\mathbf{n} \cdot \mathbf{T} = C_a^{-1} 2Hn - n\mathbf{P}_g, \nonumber
\]

(A4)

\[
\mathbf{n} \cdot v = 0. \nonumber
\]

(A5)

Equations (A4) and (A5) represent the stress balance and kinematic conditions, respectively. Here $\mathbf{T} = -P\mathbf{I} + \tau$, $\mathbf{I}$ is the identity tensor, $\mathbf{n}$ is the unit outward pointing normal from the free liquid surface, $H$ is the local mean curvature of the interface $2H = -\nabla H \cdot \mathbf{n}$, where $\nabla$ is the surface divergence operator $(\mathbf{I} - \mathbf{n}\mathbf{n}) \cdot \nabla$, and $P_g$ is the unknown pressure in the surrounding inviscid phase, relative to an arbitrarily chosen datum pressure at the lower right-hand corner of the domain.

**At the contact lines:**

\[
\mathbf{n}_{\text{solid}} \cdot \mathbf{n} = \cos \theta. \nonumber
\]

(A6)

At the dynamic contact line, the microscopic dynamic contact angle is $\theta = \theta_0$ and at the static contact line, $\theta = \theta_S$.

**Slip model:** The liquid speed along the moving plate is

\[
v_y = -1 + \exp \left( -[h(0)-y]/\varepsilon \right), \nonumber
\]

(A7)

where $h(x)$ is the dimensionless height of the free surface and $\varepsilon = L_S/L_C$.

**At the hypothetical lower height, $y = 0$,**

\[
v_x = 0, \nonumber
\]

(A8)

\[
v_y = -v_y 0 \left( 1 - x \right) \left( 1 - 3x \right). \nonumber
\]

(A9)

Here $v_{y0}$ is the dimensionless liquid speed at $(x,y) = (0,0)$, and is defined using the slip model (A7). Note that the imposition of Eqs. (A8) and (A9) is equivalent to using the boundary condition $\partial v_y/\partial y = 0$ at that location.

The mass of liquid within the computational domain must be conserved. For a liquid of constant density, this reduces to the volume constraint,

\[
\int_0^1 h(x) dx = A/L_C, \nonumber
\]

(A10)

where $A$ is the liquid volume/width within the computational domain.

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