Computational Study of Surface Tension and Wall Adhesion Effects on an Oil Film Flow Underneath an Air Boundary Layer

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May 1998
Acknowledgments

The authors thank Sanford Davis and Murray Tobak of Ames for their mathematical advice and assistance; Lou Zechter of the Numerical Aerospace Simulation Facility for his computer expertise and resources; and U. Rist and the late H. Bestek of the Institut für Aero- und Gasdynamik for their support and encouragement.
CONTENTS

FIGURES

TABLES

1 INTRODUCTION

2 MODELING APPROACH

2.1 Considerations

2.1.1 Linear Solution and Lax-Wendroff Simulation

2.1.2 Dimensionless Numbers

2.2 Defining Equations

2.2.1 The Differential Navier-Stokes Equations in Conservation Form

2.2.2 Surface Tension

2.2.3 Wall Adhesion and Contact Angle

2.3 Computational Technique

2.3.1 VOF Method

2.3.2 CSF Model

2.3.3 Wall Adhesion in the CSF Model

2.3.4 Application of Tangential Surface Forces

2.3.5 Numerical Method

3 COMPUTATIONAL RESULTS

3.1 Geometry and Grid Dimensions

3.2 Comparison of the VSS Method and the XCSF Model

3.2.1 The “Standard” Case

3.2.2 The “High Sigma” Case

3.2.3 XCSF Vectors

3.2.4 Computational Conclusions

3.3 Comparison with Previous Studies; Surface Tension and Wall Adhesion Effects

3.3.1 Lax-Wendroff Results and the Linearized Equation

3.3.2 The Influence of Wall Adhesion

3.3.3 Surface Tension Effects
**FIGURES**

<table>
<thead>
<tr>
<th>FIGURE</th>
<th>DESCRIPTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Oil drop underneath an air boundary layer.</td>
</tr>
<tr>
<td>2</td>
<td>Molecular forces and surface tension.</td>
</tr>
<tr>
<td>3</td>
<td>Surface tension forces at the contact line.</td>
</tr>
<tr>
<td>4</td>
<td>SLIC aligns the surface in a cell with one of the cell coordinates (shaded regions represent reconstructed interfaces and numbers shown are volume fractions).</td>
</tr>
<tr>
<td>5</td>
<td>PLIC) allows a nonzero slope for the surface line (shaded regions represent reconstructed surfaces and numbers shown are volume fractions).</td>
</tr>
<tr>
<td>6</td>
<td>The piecewise constant/stair-step algorithm in RIPPLE allows the surface to &quot;stair step&quot; and align with more than one mesh coordinate within a given cell.</td>
</tr>
<tr>
<td>7</td>
<td>Cylindrical volume of integration used for the CSF model.</td>
</tr>
<tr>
<td>8</td>
<td>Characteristic step function ( c(\mathcal{I}) ). (Note: ( x ) is normal to the surface.)</td>
</tr>
<tr>
<td>9</td>
<td>Mollified function ( \bar{c}(\mathcal{I}) ).</td>
</tr>
<tr>
<td>10</td>
<td>( \nabla \bar{c}(\mathcal{I}) /[c] ) along the surface normal; ( F_{i+1}, F_{i+2} ) and ( F_{i+3} ) illustrate the discrete values of ( \bar{F}_v ) applied at grid points ( i + 1, i + 2, i + 3 ) in a numerical calculation.</td>
</tr>
<tr>
<td>11</td>
<td>Wall adhesion boundary condition in the CSF method; the unit vectors, ( \hat{n}_{bc} ), at the bottom vertices of cells adjacent to the wall are set normal to the equilibrium surface position given by ( \theta_s ).</td>
</tr>
<tr>
<td>12</td>
<td>The application of the shear stress boundary condition.</td>
</tr>
<tr>
<td>13</td>
<td>The local coordinate system (red) with the principal vectors ( \hat{e}_1 ) and ( \hat{e}_2 ) is rotated by ( \varphi ); ( \hat{n} ) is the surface unit normal vector pointing into the fluid.</td>
</tr>
<tr>
<td>14</td>
<td>Volume shear forces distributed on a grid.</td>
</tr>
<tr>
<td>15</td>
<td>Location of the variables in a grid cell.</td>
</tr>
<tr>
<td>16</td>
<td>Initial oil distribution and grid dimensions.</td>
</tr>
<tr>
<td>17</td>
<td>Drop shape evolution at 0.4 s, ( \sigma = 0.0208 ) N/m.</td>
</tr>
<tr>
<td>18</td>
<td>Drop shape evolution at 1.4 s, ( \sigma = 0.0208 ) N/m.</td>
</tr>
<tr>
<td>19</td>
<td>Drop shape evolution at 2.4 s, ( \sigma = 0.0208 ) N/m.</td>
</tr>
<tr>
<td>20</td>
<td>Surface tension volume forces, ( \sigma = 0.0208 ) N/m.</td>
</tr>
<tr>
<td>21</td>
<td>Drop shape evolution at 0.4 s; ( \sigma = 0.0624 ) N/m.</td>
</tr>
<tr>
<td>22</td>
<td>Drop shape evolution at 1.4 s, ( \sigma = 0.0624 ) N/m.</td>
</tr>
<tr>
<td>23</td>
<td>Drop shape evolution at 3.4 s, ( \sigma = 0.0624 ) N/m.</td>
</tr>
<tr>
<td>24</td>
<td>Velocity vectors XCSF computation, ( \sigma = 0.0624 ) N/m, ( t = 3.4 ) s.</td>
</tr>
</tbody>
</table>
25 Velocity vectors VSS computation, \( \sigma = 0.0624 \text{ N/m}, t = 3.4 \text{ s} \) ............................................. 23
26 Volume shear forces. ......................................................... 24
27 Lax-Wendroff and linearized equation, \( x = 1.5 \text{ mm} \) .................................................. 25
28 Drop evolution at 0, 0.4, 1.4, and 2.4 s. ................................................ 26
29 Drop evolution (wedge area), \( \sigma = 0.0208 \text{ N/m}, \theta_s = 3^\circ \) .................. 27
30 Drop shape evolution at 0.4 s. .................................................. 28
31 Drop shape evolution at 1.4 s. .................................................. 28
32 Drop shape evolution at 2.4 s. .................................................. 28
33 Surface tension volume forces. .................................................. 29
34 \( h(t) \) at \( x = 1.5 \text{ mm} \). .................................................. 29
35 The influence of grid resolution on \( h(t) \), \( x = 1.5 \text{ mm} \). ................................. 31
1 Values for $e_{\text{kin}}$ at $t = 2.4$ s ........................................ 21
2 Values for $f$ and $d_{\text{rel}}$ at $t = 2.4$ s ........................................ 30
1 INTRODUCTION

Half (or more) of a flight vehicle's drag arises from skin friction. It is therefore highly desirable to have accurate measurement tools that enable the measurement of skin friction distributions on surfaces. To date, various skin friction measurement methods have been developed. One is the fringe-imaging skin friction (FISF) technique, which was originally developed by Monson and Mateer (ref. 1) and has been used successfully in many different flows. With this technique, the flow pattern of an oil patch placed on the surface of the wind tunnel model is observed. To determine the skin friction magnitude, the thickness variation of the oil is measured using interferometry and related to the skin friction magnitude through lubrication theory.

In Squire's (ref. 2) theoretical work on the motion of a thin oil sheet under a steady boundary layer, a nonlinear partial differential equation that relates skin friction to the thickness variation of the oil was derived. Tanner and Blows' (ref. 3) integral approach yields a linear equation for oil layer thickness as a function of skin friction, time, oil viscosity, and position and was shown to be the linearized solution to a reduced form of Squire's equation. It forms the basis for the FISF technique.

Zilliac (ref. 4) performed a numerical simulation of Squire's reduced equation and proved the validity of the linearized solution. The numerical results are also in accordance with experimental data (ref. 4) although no comparisons are shown at an early time when surface tension effects are thought to be large.

The main assumptions, which were made to obtain the linearized equation used in the FISF technique, are that the oil flow is an incompressible, two-dimensional, slow viscous motion and that surface tension as well as wall adhesion effects can be neglected. The last two have been omitted entirely in previous studies.

In this paper a modified version of the computer program RIPPLE (ref. 5) is used to address surface tension and wall adhesion effects. RIPPLE computes finite difference solutions for incompressible, two-dimensional, laminar viscous flows with free surfaces. The volume-of-fluid (VOF) method (ref. 6) is used for surface-tracking and the continuum surface force (CSF) model (ref. 7) for inclusion of the surface tension effects. RIPPLE's capabilities have been shown in many examples; the collision of two water rods and the low-gravity jet-induced tank flow are two of them (ref. 5).

In the current study, the influence of surface tension and wall adhesion on the accuracy of the FISF technique is investigated. In particular, the validity of the linear relation between constant shear stress, downstream position, oil viscosity, and the height of the oil film in the presence of surface tension is questioned.
2 MODELING APPROACH

In the following we will give an overview of the flow problem studied in this work. Previous approaches, which will be used for comparison with the obtained computational results, are discussed briefly. The physical model (in particular the phenomena of surface tension and wall adhesion) including the defining equations is illustrated, and the computational technique with special focus on the VOF and CSF model is presented. The latter is extended for tangential tensile forces arising from skin friction.

2.1 Considerations

Figure 1 shows a schematic representation of the flow being studied. A line of oil is placed on a flat plate. A nontransient, turbulent air boundary layer forces the oil to spread in the downstream direction. The oil height is so small that it remains entirely in the viscous sublayer. The “idealized” drop shown in figure 1 is a hemispherical drop with the same fluid volume as the “initial drop” and it is solely used for the purpose of defining a suitable characteristic length. The initial drop is the oil distribution at time $t = 0$ for the RIPPLE calculations (and also for the Lax-Wendroff computations).

The linear equation used in the FISF technique, results of Zilliac’s numerical approach to computing the thinning rate of the oil film, and limited experimental data will be compared with RIPPLE’s results. The linear solution and Zilliac’s Lax-Wendroff–based simulations will be discussed briefly.

2.1.1 Linear Solution and Lax-Wendroff Simulation

Squire (ref. 2) performed a creeping-flow-conservation-of-mass analysis of an oil film flow and obtained an expression for the thinning rate of a three-dimensional oil patch. For a two-dimensional

Figure 1. Oil drop underneath an air boundary layer.
case, this equation becomes

\[
\frac{1}{\lambda} \frac{\partial h}{\partial t} = -\frac{\partial}{\partial x} \left[ \left( \frac{\partial u_{\text{air}}}{\partial y} \right)_{y=h} \frac{h^2}{2} - \frac{1}{\mu_{\text{air}}} \frac{h^3}{3} \frac{\partial p}{\partial x} \right]
\]

(1)

where \( \lambda \) is the ratio of the air viscosity and oil viscosity \( (\lambda = \mu_{\text{air}}/\mu_{\text{oil}}) \). Squire showed by order of magnitude considerations that the pressure gradient in the oil normal to the wall is negligible \( (\partial p/\partial y = 0) \), in other words: \( p(x)_{\text{air}} = p(x)_{\text{oil}} \). Hence the pressure gradient \( \partial p/\partial x \) and the velocity gradient \( (\partial u_{\text{air}}/\partial y)_{y=h} \) in equation (1) can be obtained directly from boundary-layer theory.

The boundary conditions at the air-oil interface \( (y = h) \) are:

(1) that the air and oil velocities are the same and

(2) that the shear stresses in the oil and the air are equal at the free surface.

Equation (1) can be further simplified by assuming that the pressure gradient term is small and by alignment of the coordinate system with the surface shear stress direction. This yields

\[
\frac{\partial h}{\partial t} = -\frac{1}{\mu_{\text{oil}}} \frac{\partial}{\partial x} \left( \frac{h^2}{2} \frac{\tau_{\text{wall}}}{\mu_{\text{oil}}} \right)
\]

(2)

with \( \tau_{\text{wall}} = (\partial u_{\text{air}}/\partial y)_{y=h} \). Equation (2) can be solved analytically resulting in the linear expression

\[
h = \frac{x_{\mu_{\text{oil}}}}{t_{\tau_{\text{wall}}}}
\]

(3)

In the FISF technique, \( h(t) \) is measured at a certain location \( x \) to obtain \( \tau_{\text{wall}} \), using equation (3). This linear solution of the partial differential equation (2) assumes an infinite initial oil thickness \( (h = \infty \text{ at } t = 0) \) and a constant shear stress \( \tau_{\text{wall}} \). (Note: in the above expression, \( x = 0 \) at the leading edge of the oil.)

Zilliac (ref. 4) applied the Lax-Wendroff algorithm to solve a finite difference form of equation (2). The linear equation and the Lax-Wendroff results were nearly identical at later times \( (t > 4 \text{ s}) \). However, at early times, a comparison cannot be made because of the infinite initial height condition required to obtain the linear solution.

### 2.1.2 Dimensionless Numbers

In flows where inertial, interfacial, gravitational, and viscous forces play a role, the following four dimensionless numbers are frequently defined: the Reynolds number, \( Re = ul/\nu \); the Bond number, \( B = \rho gl^2/\sigma \); the capillary (or Taylor-Saffman) number, \( Ca = u\mu/\sigma \); and the Weber number \( We = Re \cdot Ca \), where \( l \) is the characteristic length-scale and \( \sigma \) is the surface tension coefficient.

The Bond number can be interpreted as a ratio of gravity and surface tension. Thus a high Bond number indicates that gravitational forces have a much higher impact on the flow than interfacial forces.
Similarly, the capillary number indicates the importance of viscous forces relative to surface tension, while the Weber number compares inertial forces and surface tension.

Referring to figure 1, one can see that a characteristic length is hard to define because the geometry of the flow is not static (an initially defined characteristic length would change with time). However, a rough estimate of the four relevant dimensionless numbers can be made if the radius of an "idealized drop" is chosen as the characteristic length. Note that the velocity $u$ in the above dimensionless numbers refers to the velocity of the advancing contact line, which is in the current study approximately $u = 0.84 \text{ mm/s}$, and not to a free stream velocity normally used for obtaining the Reynolds numbers. The oil viscosity is given with $\nu_{\text{oil}} = 50 \text{ Cs}$. Given these assumptions, the Reynolds, Bond, capillary, and Weber numbers become:

$$Re = \frac{\text{inertial}}{\text{viscous}} = \frac{ul}{\nu} = 0.0063,$$

$$B = \frac{\text{gravitational}}{\text{interfacial}} = \frac{\rho gl^2}{\sigma} = 0.063,$$

$$Ca = \frac{\text{viscous}}{\text{interfacial}} = \frac{u\mu}{\sigma} = 0.0002,$$

$$We = \frac{\text{inertial}}{\text{interfacial}} = Re \cdot Ca = 1.27 \cdot 10^{-6}$$

respectively.

The values of $Ca$ and $We$ are well within the typical ranges for flows dominated by interfacial and viscous forces (ref. 8). This implies that surface tension forces may have a significant influence on the oil flow locally.

### 2.2 Defining Equations

#### 2.2.1 The Differential Navier-Stokes Equations in Conservation Form

The flow field in the oil film is governed by the Navier-Stokes equations for two-dimensional, incompressible, and viscous flows,

$$\nabla \cdot \mathbf{V} = 0$$

(4)

$$\frac{\partial \mathbf{V}}{\partial t} + \nabla \cdot (\mathbf{V} \mathbf{V}) = -\frac{1}{\rho} \nabla p + \frac{1}{\rho} \nabla \cdot \tau + \mathbf{g} + \frac{1}{\rho} \mathbf{F}_b$$

(5)

Here, $\mathbf{V} = V(x, y, t)$ is the velocity field, $\mathbf{g}$ is the gravity acceleration, $\mathbf{F}_b$ is a body force, $p$ is the pressure, $\tau$ is the viscous stress tensor for a Newtonian fluid, $\tau = \mu[\nabla \cdot \mathbf{V} + (\nabla \cdot \mathbf{V})^T]$, and $\rho$ is the oil density.

#### 2.2.2 Surface Tension

Within a fluid, the intermolecular forces of attraction are balanced in all directions. However, for the fluid molecules at or near the surface, the cohesive forces are unbalanced due to the discontinuity
of fluid properties at the interface (fig. 2). The unbalanced forces of intermolecular attraction at the surface cause the interface to behave like it were an elastic membrane under uniform tension. The magnitude of the tensile force per unit length of a line on the interface is called surface tension \( \sigma \). The value of \( \sigma \) depends on the fluids in contact and the temperature. In order to take the effects of surface tension into account, special boundary conditions must be applied to the Navier-Stokes equations at an interface. Assuming that the heat flux and mass transport through the surface are negligible, the boundary condition that must be satisfied at the boundary between the two fluids yields an equilibrium expression for all forces acting on both sides of a surface element at an instant in time,

\[
\hat{n} \left( p_{\text{fluid}} - \sigma \kappa \right) - \tau_{\text{fluid}} \cdot \hat{n} + \frac{\partial \sigma}{\partial s} = \hat{n} \left( p_{\text{vapor}} - \tau_{\text{vapor}} \cdot \hat{n} \right)
\]

where \( \hat{n} \) is the free surface unit normal vector directed into the fluid, \( \tau_{\text{fluid}} \) is the viscous stress tensor in the fluid, \( \tau_{\text{vapor}} \) is the viscous stress tensor in the vapor, and \( \kappa \) is the local curvature calculated from (ref. 9)

\[
\kappa = - (\nabla \cdot \hat{n})
\]

Notice that \( \partial \sigma / \partial s \) in equation (6) is the gradient of the surface tension coefficient along the surface.

Further, assuming that the stress tensor is continuous across the interface \( (\tau_{\text{fluid}} = \tau_{\text{vapor}} \text{ at the surface}) \) (ref. 10) and that \( \sigma \) can be regarded as constant \( (\partial \sigma / \partial s = 0) \), equation (6) reduces to

\[
p_{\text{fluid}} - p_{\text{vapor}} = \Delta p = \sigma \kappa
\]

The pressure jump \( \Delta p \) in the above expression has to be applied as a boundary condition of equation (5) at the fluid interface.

### 2.2.3 Wall Adhesion and Contact Angle

When a drop of liquid is placed on a solid surface, the liquid spreads out under the influence of gravity. The rate of spreading depends on the relative sizes of intermolecular cohesive and adhesive forces. For example, if the fluid molecules possess a greater affinity for each other than for the solid
(i.e., cohesive forces are greater than adhesive forces) the drop tends to deform slightly but it does not "wet" the solid.

In the opposite case, where the adhesive force between the solid and the liquid is greater than the cohesive forces between the fluid molecules, the drop spreads out until it reaches a state of static equilibrium. At this point, the surface of the drop forms an angle of contact ($\theta_S$) with the solid surface (fig. 3) which can be measured experimentally (ref. 11). Young (ref. 11) assumed that the magnitude of the static contact angle depends on the surface tension coefficients of the three material boundaries ($\sigma_{\text{liquid/vapor}}$, $\sigma_{\text{liquid/solid}}$, and $\sigma_{\text{vapor/solid}}$). In order for the system to be in static equilibrium, the horizontal components must sum to zero at the contact line,

$$\sigma_{\text{vapor/solid}} - \sigma_{\text{liquid/solid}} - \sigma_{\text{liquid/vapor}} \cos \theta_S = 0$$

Although this equation ignores the force balance normal to the solid surface by assuming that the solid acts like a rigid body, it is a good approximation from the point of view of macroscopic capillary concepts. In the case of a moving contact line, the resulting dynamic contact angle is different from the static contact angle; it is at the very least a function of $\theta_S$ and the capillary number.

The general equations used to determine the static and dynamic contact angles are still a subject of research. Some success has been achieved using empirical methods but a general theory of dynamic contact lines has not been derived (ref. 11).

### 2.3 Computational Technique

RIPPLE is a computer program for simulating transient, two-dimensional, incompressible fluid flows governed by equations (4) and (5). Free surfaces are tracked using a VOF technique that was developed at the Los Alamos National Laboratory by Hirt and Nichols (ref. 6). Surface tension forces are incorporated applying the CSF model (ref. 9).

#### 2.3.1 VOF Method

The VOF technique for representing free surfaces assigns a characteristic value $F$ to each computational cell that specifies whether the cell is a fluid cell (cells completely filled with fluid), a surface cell (cells partially filled with fluid), or a void cell (empty cells). In two-fluid problems, such as the present one, the gas phase is not fully modeled by RIPPLE, and it is referred to as the "void region." In RIPPLE, $F = 1$ for a fluid cell, $F = 0$ for a void cell, and $0 < F < 1$ in cells containing the surface.

![Figure 3. Surface tension forces at the contact line.](image)
In these cells, $F$ represents the fractional volume of the cell occupied by fluid. The VOF function $F = F(\vec{r}, t)$ is a scalar field advected as a Lagrangian invariant in an Eulerian scheme,

$$\frac{dF}{dt} = \frac{\partial F}{\partial t} + u \frac{\partial F}{\partial x} + v \frac{\partial F}{\partial y} = 0 \tag{10}$$

where $F(\vec{r}, 0)$ is known. Equation (10) has to be solved in addition to the flow governing equations (4) and (5). The solution algorithm for equation (10) has to assure that the discontinuous property of $F$ (which is a fundamental premise in multi-fluid problems, where it is desirable to maintain a sharply defined interface) is preserved during the computation. Standard advection algorithms, which compute fluxes algebraically, tend to spread discontinuities owing to numerical diffusion. Even minimizing the diffusion with the objective of restricting the spreading to a width of two or three cells does not match the capabilities of VOF advection methods, which typically remain within one cell width.

In the past two decades, several VOF algorithms have been developed to advect volume fractions. Theses methods all have in common that they compute the variation of $F$ in a cell by reconstructing the surface from VOF data and computing the volume fluxes geometrically. The surface reconstruction method plays an important role and can have great effect on accuracy and robustness on the VOF method (refs. 12 and 13). The main difference between the various surface reconstruction algorithms is the assumed interface geometry within cells. Piecewise constant interface calculations, also called simple line interface calculation (SLIC), approximate the interface with a straight line aligned with one of the mesh coordinates (fig. 4) as opposed to piecewise linear interface calculations (PLIC), which allow a nonzero slope for the straight line (fig. 5).

One can see from figures 4 and 5 that the two VOF methods reconstruct different surface geometries from the same volume fraction data. The piecewise linear reconstruction algorithm yields a more accurate interface approximation which can lead to a smoother volume fraction distribution in translation and rotation problems. This can be important in numerical techniques where surface dynamics (i.e., surface tension) is included (like in RIPPLE) and depends on an accurate computation of surface normals, hence a "physically smooth" VOF distribution.

RIPPLE uses a piecewise constant/stair-step VOF advection, called the Hirt and Nichols (HN) method. This means the surface is allowed to align with more than one mesh coordinate within a

![Figure 4. SLIC aligns the surface in a cell with one of the cell coordinates (shaded regions represent reconstructed interfaces and numbers shown are volume fractions).](image-url)
A continuous surface line is not obtained

Figure 5. PLIC allows a nonzero slope for the surface line (shaded regions represent reconstructed surfaces and numbers shown are volume fractions).

Cell depending on volume fractions in neighboring cells and the local velocity (fig. 6). For a detailed discussion of this advection method see references 5–7.

For most flows, the HN method gives better results than a SLIC algorithm, but although it is more efficient, it is not as accurate as a PLIC VOF advection. In our computations, the surface remains smooth and does not undergo severe distortions, where the shortcomings of the HN advection, in comparison to the PLIC method, could be crucial. However, it is important to keep the type of surface reconstruction in mind when viewing results from RIPPLE computations, especially when surface smoothness, surface curvature, and resulting surface dynamics are viewed on a "microscopic" scale of a few cell widths.

The main advantage of the VOF technique over older surface tracking models like the marker and cell (MAC) method or the Lagrangian incompressible (LINC) technique, which use logically connected Lagrangian points instead of characteristic marker data, is that the VOF approach can track highly distorted and disconnected surfaces.

Reconstructed surface (stair-step)

Figure 6. The piecewise constant/stair-step algorithm in RIPPLE allows the surface to "stair step" and align with more than one mesh coordinate within a given cell.
2.3.2 CSF Model

The surface-tension-induced pressure jump $\Delta p$ given by equation (8) represents a boundary condition in the pressure field at the interface. In order to set the correct pressure at the free surface, one would have to compute the exact surface location and slope, which would require a time consuming reconstruction of a continuous surface. However, a fast and efficient algorithm for reconstructing a continuous surface line from VOF data has not been devised yet (ref. 7). An additional problem is that the interface line rarely coincides with grid points where boundary conditions such as the pressure jump can be easily applied. In addition to a continuous surface reconstruction, an interpolation scheme for setting the correct pressure in cells adjacent to surface cells would be required.

A very different approach to surface forces, in general, is given by the CSF model. It was developed by Brackbill, Kothe, and Zemach (ref. 9) and it makes very effective use of VOF data. In the following subsection, we will discuss the CSF method briefly to illustrate how it is derived and what the advantages are. For a detailed derivation and a proof of the CSF method for surface forces acting normal to the interface, see references 9 and 14.

In the CSF model, a surface tensile force per unit interfacial area (such as surface tension) is replaced by a localized volume force acting on fluid elements located in a thin transition region of a few cell widths. In the transition, the fluid properties are thought to change continuously from one fluid (or void) to another. This means surface forces are no longer applied as discrete boundary conditions at a discontinuity (the free surface) but as body forces $F_v$ acting everywhere in the transition region. Hence the exact surface location is no longer required and $F_v$ is applied at grid points located in the transition region.

The basic idea is that one can define a localized body force $F_v$, which volume integral in the limit of infinitesimally small transition-region-thickness $h$ is equal to the surface integral of the tensile force $F_s$.

$$\lim_{h \to 0} \int_{\delta V} F_v(\vec{x})dV = \int_{\delta S} F_s(\vec{x}_s)dS$$

(11)

where $\vec{x}_s$ is a position at the free surface. The surface integral is taken over a part of the free surface lying within the volume of integration $\delta V$. The orientation of $\delta V$ is such that its sides are perpendicular to the surface. Figure 7 shows an example using a cylindrical integration volume. In equation (11), $F_s$ is the classical surface tension force,

$$F_s(\vec{x}_s) = \sigma \kappa(\vec{x}_s)\hat{n}(\vec{x}_s)$$

(12)

where $\hat{n}(\vec{x}_s)$ is the unit vector normal to the surface. $F_s(\vec{x}_s)$ corresponds to the surface pressure $\Delta p$ acting on $S$: $|F_s(\vec{x}_s)| = \Delta p(\vec{x}_s)$.

In order to find the body force $F_v(\vec{x})$ that satisfies equation (11), first a characteristic function $c(\vec{x})$ with the following properties is defined (fig. 8)

$$c(\vec{x}) = \begin{cases} 
    c_1 & \text{in fluid 1} \\
    c_2 & \text{in fluid 2} \\
    (c_1 + c_2)/2 & \text{at the interface}
\end{cases}$$

(13)
Figure 7. Cylindrical volume of integration used for the CSF model.

Figure 8. Characteristic step function \( c(\bar{x}) \). (Note: \( x \) is normal to the surface.)

From equation (13), one can see that \( c(\bar{x}) \) is a step function and that the interface is specified by the discrete value \( c(\bar{x}_s) = (c_1 + c_2)/2 = \langle c \rangle \). Hence the interface is an infinitesimally thin line.

In order to “create” a transition region between the two different fluids (marked by \( c_1 \) and \( c_2 \)), the interface is imaginary broadened to a small distance \( h \) by convolving \( c(\bar{x}) \) over \( h \) with a B-spline. This yields a mollified characteristic function \( \tilde{c}(\bar{x}) \) which varies smoothly from \( c_1 \) over \( \langle c \rangle \) to \( c_2 \), as shown in figure 9. The B-spline is defined so that \( \tilde{c}(\bar{x}) \) approaches \( c(\bar{x}) \) as the transition region thickness \( h \) goes to zero: \( \lim_{h \to 0} \tilde{c}(\bar{x}) = c(\bar{x}) \).

Figure 9. Mollified function \( \tilde{c}(\bar{x}) \).
With \( \tilde{c}(\vec{x}) \), a bell-shaped distribution function for the volume force \( \vec{F}_v \) is derived by taking the gradient of \( \tilde{c}(\vec{x}) \) and normalizing with \( [c] = c_2 - c_1 \). Figure 10 shows the distribution along a surface normal. The volume force \( \vec{F}_v \) is then formulated as

\[
\vec{F}_v(\vec{x}) = \sigma \kappa(\vec{x}) \frac{\nabla \tilde{c}(\vec{x})}{[c]}
\]  

(14)

To show that the expression on the right hand side of equation (14) satisfies the identity in the integral equation (11), one can apply the following relations.

Because \( \lim_{h \to 0} \tilde{c}(\vec{x}) = c(\vec{x}) \) (which is a step function) and the normalization of \( \tilde{c}(\vec{x}) \) with \( (c_2 - c_1) \),

\[
\lim_{h \to 0} \frac{\tilde{c}(\vec{x})}{[c]} = H(\hat{n}(\vec{x}_s) \cdot (\vec{x} - \vec{x}_s))
\]  

(15)

where \( H \) is the Heaviside unit function. With equation (15),

\[
\lim_{h \to 0} \int_{\delta V} \sigma \kappa(\vec{x}) \frac{\nabla \tilde{c}(\vec{x})}{[c]} dV = \int_{\delta V} \sigma \kappa(\vec{x}) \nabla H(\hat{n}(\vec{x}_s) \cdot (\vec{x} - \vec{x}_s)) dV
\]

Further,

\[
\int_{\delta V} \sigma \kappa(\vec{x}) \nabla H(\hat{n}(\vec{x}_s) \cdot (\vec{x} - \vec{x}_s)) dV = \int_{\delta V} \sigma \kappa(\vec{x}) \hat{n}(\vec{x}_s) \delta(\hat{n}(\vec{x}_s) \cdot (\vec{x} - \vec{x}_s)) dV
\]

where \( \delta(\vec{x}) \) is Dirac’s delta function, which is the derivative of the Heaviside unit function and has the property

\[
\int_{-\infty}^{\infty} \delta(x) g(x) d(x) = g(0)
\]

\( g(x) \) is any suitable continuous function and in our case, \( g(x) = \sigma \kappa(\vec{x}) \hat{n}(\vec{x}_s) \) and \( g(0) = \sigma \kappa(\vec{x}_s) \hat{n}(\vec{x}_s) \). Thus the delta function converts the volume integral into a surface integral,

\[
\int_{\delta V} \sigma \kappa(\vec{x}) \hat{n}(\vec{x}_s) \delta(\hat{n}(\vec{x}_s) \cdot (\vec{x} - \vec{x}_s)) dV = \int_{\delta S} \sigma \kappa(\vec{x}_s) \hat{n}(\vec{x}_s) dS
\]  

(16)

The integrand on the right hand side of equation (16) is the surface force defined in equation (12).

Figure 10. \( \nabla \tilde{c}(\vec{x})/[c] \) along the surface normal; \( F_{i+1}, F_{i+2} \) and \( F_{i+3} \) illustrate the discrete values of \( \vec{F}_v \) applied at grid points \( i + 1, i + 2, i + 3 \) in a numerical calculation.
Comments on $\vec{F}_v$ and $\kappa(\vec{x})$—It is interesting to note that $\vec{F}_v$ is not a function of the surface location $\vec{x}_s$; rather, it is a true volume force that depends on $\vec{x}$. The discrete boundary value problem is replaced with a continuous model. Since $\vec{c}(\vec{x})$ is constant in each fluid and changes only within the transition region, $\vec{F}_v$ becomes zero outside the interface region. $\vec{F}_v$ can be simply added to the momentum equation (5) setting $\vec{F}_b = \vec{F}_v$. In a numerical calculation, $\vec{F}_v$ is obtained at grid points that are located in the transition region (fig. 10).

Because nested surfaces of constant $\vec{c}$ in the transition region represent the interface in the CSF model, the normal vector of the interface is obtained by taking the gradient of the mollified characteristic function,

$$\vec{n}(\vec{x}) = \nabla \vec{c}(\vec{x})$$

Substituting the right hand side of equation (17) into equation (7) yields for the curvature

$$\kappa(\vec{x}) = -\nabla\left[ \frac{\nabla \vec{c}(\vec{x})}{|\nabla \vec{c}(\vec{x})|} \right]$$

In RIPPLE's implementation of the CSF model, the characteristic function $\vec{c}(\vec{x})$ is set equal to the VOF function $F(\vec{x})$,

$$\vec{c}(\vec{x}) = F(\vec{x}) = \begin{cases} 
  c_1 = 0 & \text{in the void,} \\
  > c_1, < c_2 & \text{in the interface region,} \\
  c_2 = 1 & \text{in the fluid} 
\end{cases}$$

since $F(\vec{x})$ offers all the properties postulated for $\vec{c}(\vec{x})$.

2.3.3 Wall Adhesion in the CSF Model

It is obvious from Young's equation (9) that if the fluid interface forms a contact angle $\theta$ with a rigid boundary that is different from the equilibrium contact angle $\theta_s$, a resulting force should be applied in order to restore the fluid to the equilibrium contact $\theta_s$. This force can easily be computed within the framework of the CSF technique by applying a wall adhesion boundary condition as follows.

The surface normal vectors at the wall are not computed from equation (17). Instead, the unit normal vectors are specified so that they form an angle with the wall normal that is equal to the equilibrium contact angle (fig. 11). This is done prior to evaluating the surface curvature (eq. (7)), and hence the volume force $\vec{F}_v$ in equation (14) in cells at the wall depends on the difference between $\theta$ and $\theta_s$. In figure 11, the case for $\theta > \theta_s$ is illustrated. One can see that the horizontal component of $\vec{F}_v$ is in accordance with Young's formula,

$$\sigma_{\text{vapor/solid}} - \sigma_{\text{liquid/solid}} - \sigma_{\text{liquid/vapor}} \cos \theta = \sigma_{\text{liquid/vapor}}(\cos \theta_s - \cos \theta) > 0$$

The assumption that the contact angle is constant is a physical approximation in the case of a moving contact line (refer to subsection 2.2.3). However, if the difference between the static and dynamic contact angle is small (i.e., if the static contact angle and the velocity of the advancing contact line is small, like in the current case), this assumption is likely to be a good approximation.
Equilibrium surface position (contact angle $\theta_s$)

Actual surface position (contact angle $\theta$)

Figure 11. Wall adhesion boundary condition in the CSF method; the unit vectors, $\hat{n}_{bc}$, at the bottom vertices of cells adjacent to the wall are set normal to the equilibrium surface position given by $\theta_s$.

2.3.4 Application of Tangential Surface Forces

The driving force of the flow studied in this work (i.e., the flow in the oil patch) is the shear force of the air boundary layer acting on the surface of the oil. Since in RIPPLE a solution of the flow governing equations is obtained only in the fluid and viscous effects at the surface are neglected, a method had to be developed that enabled the application of shear forces at the interface.

The Viscous Shear Stress Method-- It was first suggested to set the shear components of the viscous stress tensor in cells containing the surface and empty cells equal to that of the air boundary layer,

$$(\tau_{xy})_{oil} = (\tau_{xy})_{air} = \text{constant}, \forall \text{cells } 0 \leq \text{VOF} < 1 \quad (19)$$

Figure 12 shows how the shear stress of the air boundary layer is applied in the RIPPLE computation. This simple approach is a coarse approximation if the surface deviates from a straight line, has nonzero slope, or does not coincide with grid points. It was used to investigate RIPPLE's applicability to our case since the surface was thought to have a slope smaller than $10^\circ$ at all times. Its main advantage is the straightforward implementation of the boundary condition (eq. (19)) into the existing computer code. However, it will be shown later that the resulting flow field is nonphysical and the method was discarded.

Extension of the CSF for Tangential Surface Forces-- Applying the correct tangential forces at the free surface in a numerical scheme is analogous to applying normal forces, which was discussed in subsection 2.3.2. Thus a similar approach is used to overcome the restrictions of the viscous shear stress (VSS) method.

Since the boundary layer flow is parallel to the surface it is convenient to define a local coordinate system, the $x$- and $y$-axes of which lie in the surface tangential plane and the $z$-axis points in the outward normal direction. The two-dimensional case is shown in figure 13.
Figure 12. The application of the shear stress boundary condition.

Figure 13. The local coordinate system with the principal vectors $\hat{e}_1$ and $\hat{e}_2$ is rotated by $\varphi$; $\hat{n}$ is the surface unit normal vector pointing into the fluid.

The components of the boundary layer shear stress tensor $\boldsymbol{\tau}$ are given in the local coordinate system and can be transformed to the global coordinate system with the tensor transformation,

$$\tau_{ij} = c_{ik} c_{jl} \tau_{kl} \quad i, j, k, l = 1, 2, 3 \quad (20)$$

where $c_{ik}$ and $c_{jl}$ are the transformation coefficients, $c_{ik} = \cos(\angle \hat{e}_i, \hat{e}_k)$ and $c_{jl} = \cos(\angle \hat{e}_j, \hat{e}_l)$.

The force per unit interfacial area resulting from the shear stress tensor $\tau$ acting on a surface element is then given by

$$\bar{F}_\tau(\bar{x}_S) = -\tau(\bar{x}_S) \cdot \hat{n}(\bar{x}_S) \quad (21)$$

This tensile force can be converted to a localized volume force acting everywhere in the transition region utilizing the derivation used in subsection 2.3.2 to obtain the surface tension volume force. Setting $\bar{F}(\bar{x}_S) = \bar{F}_\tau(\bar{x}_S)$ in equation (11) yields for the volume shear force

$$\bar{F}_{\tau v}(\bar{x}) = -\tau(\bar{x}) \cdot \nabla \hat{e}(\bar{x}) \quad (22)$$

Figure 14 illustrates how the volume shear forces are distributed in the transition region when the surface is oriented horizontally.

All properties of the volume force $\bar{F}_v$ mentioned in subsection 2.3.2 apply, of course, also to $\bar{F}_{\tau v}$. Hence the body force $\bar{F}_b$ in the momentum equation (5) is calculated from $\bar{F}_b = \bar{F}_v + \bar{F}_{\tau v}$.  

14
Figure 14. Volume shear forces distributed on a grid.

2.3.5 Numerical Method

In RIPPLE, a finite difference representation of the flow governing equations (4) and (5) by central differences is solved. The location of the flow variables on the computational grid follows the MAC scheme which uses a staggered grid as illustrated in figure 15. In order to transcend the instability problems of time marching solutions for the incompressible Navier-Stokes equations, a two-step projection method (refs. 7 and 15) is applied.

A time discretization of the momentum equation (5) yields

$$\frac{\vec{\omega}^{n+1} - \vec{\omega}^n}{\Delta t} = -\nabla \cdot (\vec{\omega} \vec{\omega})^n + \frac{1}{\rho^n} \nabla p^{n+1} + \frac{1}{\rho^n} \nabla \cdot \vec{\tau}^n + \vec{g}^n + \frac{1}{\rho^n} \vec{F}_b^n$$

(23)

The superscripts $n$ and $n+1$ denote the variables taken at the old and new time step, respectively.

Figure 15. Location of the variables in a grid cell.
In the first step of RIPPLE's projection method, an intermediate velocity field \( \vec{v} \) is obtained from \( \vec{v}^n \) by dropping the implicit pressure term in equation (23),

\[
\frac{\vec{v} - \vec{v}^n}{\Delta t} = -\nabla \cdot (\vec{v} \vec{v})^n + \frac{1}{\rho^n} \nabla \cdot \vec{f}^n + \frac{1}{\rho^n} \vec{F}_b
\]  

Using the intermediate velocity field \( \vec{v} \), the second step becomes an expression for the new velocity field \( v^{n+1} \) and the new pressure field \( p^{n+1} \),

\[
\frac{\vec{v}^{n+1} - \vec{v}}{\Delta t} = -\frac{1}{\rho^n} \nabla p^{n+1} \tag{25}
\]

Multiplying each side of equation (25) with the Nabla operator and recalling the continuity condition, \( \nabla \cdot \vec{v}^{n+1} = 0 \), one gets a Poisson equation for the pressure,

\[
\nabla \cdot \left( \frac{1}{\rho^n} \nabla p^{n+1} \right) = \nabla \cdot \vec{v} \tag{26}
\]

The solenoidal velocity field \( \vec{v}^{n+1} \) can then be recovered from equation (25).

It should be noted that the momentum advection terms in equation (24) are estimated using the weakly monotonic, second-order upwind van Leer method. The solution of the implicitly discretized pressure Poisson equation (26) is obtained through an incomplete Cholesky conjugate gradient technique (ref. 16). A detailed presentation of the discretization methods used in RIPPLE is given in references 5 and 7.

The explicit treatment of the momentum and VOF advection, the viscous terms, and the body forces in equation (23) requires stability time step constraints. Hence for the advection terms a Courant condition,

\[
\Delta t < \min \left( C \left[ \frac{\Delta x}{\Delta |u|} \right] , C \left[ \frac{\Delta y}{\Delta |v|} \right] \right)
\]

is applied. The viscous time step constraint is evaluated as

\[
\Delta t < \frac{1}{3\nu} \left[ \frac{\left(\Delta x\right)^2 \left(\Delta y\right)^2}{\left(\Delta x\right)^2 + \left(\Delta y\right)^2} \right] \min \tag{27}
\]

The maximum allowable value for \( \Delta t \) is also restricted by a requirement which arises from the resolution of the propagation of capillary waves on the surface (ref. 7). Thus the time step constraint, resulting from the explicit treatment of surface tension forces in the CSF model for a uniform, rectangular grid, becomes

\[
\Delta t < \left[ \frac{\rho \left(\Delta x \Delta y\right)^{3/2}}{4\pi \sigma} \right]^{1/2} \min
\]

It will be shown in the next section that, for our case, the most limiting time step restriction was found to be the viscous time step constraint (eq. (27)).
3 COMPUTATIONAL RESULTS

This section is divided into three subsections. The first presents a brief discussion of initial oil shape and grid dimensions. Subsection 3.2 compares the previously discussed techniques for applying shear forces at the free surface, namely the VSS and the extended CSF (XCSF) model. In subsection 3.3, computational results of the XCSF, the Lax-Wendroff computation, and the linearized solution are analyzed and discussed, and the influence of surface tension on the oil flow is illustrated.

3.1 Geometry and Grid Dimensions

The initial oil shape is determined by the static contact angle, the fluid volume, and the gravity acceleration. Experiments were performed to provide the static contact angle and the dimensions of a typical oil drop used in the FISF technique (ref. 4). A contact angle of about $3^\circ$ was obtained. The measured geometric dimensions are shown in figure 16. The complete oil properties are listed in Appendix A.

Since RIPLE expects an analytical function for the initial fluid distribution (ref. 7), the initial oil shape is approximated with a circular arc as shown in figure 16. The resulting drop shape is in close accordance with the shape seen in the above mentioned experiments.

Various grid configurations were tested and the grid sketched in figure 16 was found to give the best compromise of resolution and computation time. Hence all computed cases, unless noted otherwise, are performed on a rectangular, uniform grid with a grid spacing of 0.025 mm and 0.01 mm in $x$ and $y$ directions, respectively. This yields a grid of 322 cells in width and 14 cells in height, including the ghost cells surrounding the actual physical domain.

Due to the very small cell widths ($\Delta x = 0.025$ mm, $\Delta y = 0.01$ mm) of the grid used, the most restricting time step constraint for all computations is the viscous time step constraint (eq. (27)). With the given grid spacing, it becomes $\Delta t = 5.747 \cdot 10^{-7}$ s.

![Figure 16. Initial oil distribution and grid dimensions.](image-url)
3.2 Comparison of the VSS Method and the XCSF Model

To compare the two methods for applying tangential surface forces, the flow discussed in subsection 2.1 is computed. Two different cases are presented in which the surface tension coefficient \( \sigma \) is varied. The first case is the "standard" case where \( \sigma = 0.0208 \) N/m was set (the \( \sigma \) measured by the oil manufacturer). In the second case, \( \sigma \) was set to a value three times higher. The value for the shear stress \( \tau = 2.394 \) N/m\(^2\), as measured using the FISF in a channel flow facility (ref. 4), was used.

When viewing the following figures, it is important to keep in mind that the \( y \)-axis is scaled by a factor of roughly 40 compared to the \( x \)-axis. Hence the oil layer is much shallower than suggested in figures 17–19 (and in figures 21–23 in subsection 3.2.2).

3.2.1 The "Standard" Case

In figures 17–19, the evolution of the drop shape computed by the XCSF and the VSS is shown at three different times. The initial shape as well as the applied shear stress, \( \tau = 2.394 \) N/m\(^2\), and the surface tension coefficient \( \sigma \) are the same for both computation methods. From figures 17–19, one can see that the drop shape can be divided into two main parts: the front part with the leading edge and the downstream part containing the trailing edge.

**The Leading Edge**—At early times, the leading edge of the drop is deformed to a wedge, with the leading edge angle that decreases and a wedge length that increases with time. In the wedge area, the XCSF results show a slightly smoother surface than the VSS computation, where the surface line tends to "stair step" (figs. 18 and 19).

It is interesting to see that the "step height" is of the same order of magnitude as \( \Delta y \). The "stair stepping" of the surface line is amplified when the interface approaches a horizontal orientation.

This difference in surface quality between the VSS and the XCSF method results from the refined application of surface shear stress forces in the XSCF. The VSS method does not take surface slope and surface sub-cell location into account as opposed to the XCSF method. In the VSS technique, the shear stress is set to \( \tau_{xy} = 2.394 \) N/m\(^2\) in cells containing the surface and empty cells (i.e., in cells where

![Figure 17. Drop shape evolution at 0.4 s, \( \sigma = 0.0208 \) N/m.](image-url)
Figure 18. Drop shape evolution at 1.4 s, $\sigma = 0.0208$ N/m.

Figure 19. Drop shape evolution at 2.4 s, $\sigma = 0.0208$ N/m.

$0 \leq \text{VOF} < 1)$. Since in RIPPLE the off-diagonal components of the viscous stress tensor are computed at cell vertices, the expression for the distance $d$ between the location of the surface boundary condition $\tau_{xy} = 2.394$ N/m$^2$ on the grid and the surface line becomes $0 < d < \Delta y$. This means that although the distance of the application point of the surface boundary condition to the actual surface within a cell is a transient function, the value of the VSS stress boundary condition is held constant. Furthermore, the application point is a noncontinuous function; it “jumps” by $\Delta y$ as soon as the fluid surface “swaps” into the next cell row.

Hence the stair-stepped character of the wedge computed by the VSS technique is thought not to be of physical origin and is therefore, obviously, an undesired behavior (with respect to the FISF technique). In the FISF, the thinning rate of the oil wedge is measured at a fixed $z$-location and it is assumed to be inversely related to time (see equation (3) and reference 4). However, the wedge thinning rate obtained from the stair-stepped surface line in the VSS computation does not show a linear behavior: each time a surface line step passes through the observation point, the height decreases more rapidly than in regions away from the step area.

The Trailing Edge—A significant difference in the two methods can be seen near the trailing edge of the drop, where the curvature (and hence surface tension) is higher than at the leading edge. At 0.4 s, the difference is still hardly visible, but figures 18 and 19 show that it becomes more pronounced.
at later times: the downstream contact line of the oil drop in the XCSF solution moves faster than the one computed using the VSS method.

Since the actual contact angle at the downstream edge of the drop is greater than the specified static contact angle, Young’s formula (eq. (9)) yields a force component that tries to advance the contact line downstream and therefore works in addition to the applied surface shear stress. It is surprising that the difference in the velocity of the advancing contact line is so high, since in both methods the contact angle is almost of the same value, and hence nearly the same accelerating force component acts on the trailing edge contact line.

The lower velocity of the downstream contact line in the VSS computation in comparison with the XCSF, in combination with similar thinning rates of the wedge in the front part of the drop, results in a greater maximum oil height and a more highly curved surface in the VSS result. It seems that the energy, which is introduced to the system through the external air flow, is to a great extent used to deform the drop rather than accelerate it. In the VSS method, more energy is converted into surface tension energy than into kinetic energy in comparison with the XCSF results.

Considering the units of $\mathbf{F}_v(\overline{\mathbf{x}})$ (surface tension volume force) as energy per unit volume [J/m$^3$] (eq. (14)), a comparison of the total amount of surface tension volume forces (the sum of surface tension volume forces in all cells) acting on the oil illustrates the difference in surface tension energy. One important point to recall is that the total amount of fluid specified at the beginning of the computation is the same in both techniques.

Figure 20 shows the comparison of $\mathbf{F}_v(\overline{\mathbf{x}})$ as a function of time. It can be seen that both functions start out at the same point, since the initial shape is the same for both computation methods. However, the total amount of surface tension volume forces, and hence surface tension energy, is clearly greater in the VSS case at all computed times. This leads to the conjecture that a greater retarding force must act on the fluid in the VSS method, which prevents the trailing edge of the oil from moving downstream at as high a rate as the XCSF drop and causes the high surface deformation.

![Figure 20. Surface tension volume forces, $\sigma = 0.0208$ N/m.](Image)
In order to verify that the amount of surface energy (i.e., surface tension volume force) due to the high surface deformation in the VSS technique leads to a lack of kinetic energy in the fluid, a kinetic energy balance was performed: the square of the magnitude of the velocity vector in each cell was multiplied with the corresponding value of the VOF function \( F \). (Note: \( F \) reflects the fractional volume, and hence fractional fluid mass, in a single cell.) These “scaled” velocity magnitudes were then summed over all \( N \) grid cells,

\[
e_{\text{kin}} = \sum_{i=1}^{N} F_i \cdot |\vec{v}_i|^2
\]

(28)

which yields the specific kinetic energy in the fluid. In table 1 the numerical values for \( e_{\text{kin}} \) are listed for \( t = 2.4 \) s; for other times, the difference in the value for \( e_{\text{kin}} \) is similar and will be not listed.

<table>
<thead>
<tr>
<th>Time</th>
<th>( e_{\text{kin}} ) VSS in ( \text{mm}^2/\text{s}^2 )</th>
<th>( e_{\text{kin}} ) XCSF in ( \text{mm}^2/\text{s}^2 )</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.4 s</td>
<td>1034.65</td>
<td>2696.53</td>
</tr>
</tbody>
</table>

### 3.2.2 The “High Sigma” Case

A lower contact line velocity yields a lower capillary number (\( Ca = u\mu/\sigma \)) which in turn means that interfacial forces play a more important role in the VSS technique. Hence the second case with \( \sigma = 0.0624 \) N/m (figs. 21–23) was computed to investigate the impact of surface tension on the velocity difference at the contact line and finally to determine which shear force method yields the more physical flow.

The results are basically the same as in the “standard” case described in subsection 3.2.1 with the difference being that the trailing edge contact line in the VSS computation does not move at all. This seems to be a paradox, since the contact angle is again the same in both computations, and hence the wall adhesion force should move the contact line of the oil drop downstream in the VSS computation as well. This is an illustration of the physical deficiencies of the VSS implementation.

![Figure 21. Drop shape evolution at 0.4 s; \( \sigma = 0.0624 \) N/m.](image)
In figures 24 and 25 the velocity profiles of the two techniques, in the area of greatest oil height, are shown. One can see that the velocity field of the oil flow in the XCSF computation looks very similar to a Couette flow, which was expected.

An unexpected result of the VSS method is that the oil in the vicinity of the surface flows in the same direction as the external air flow, but in the oil the fluid moves upstream against the air flow...
direction. This means that the oil flow circulates inside the drop instead of moving downstream with the air flow. The explanation for this unexpected flow behavior can be found through analyzing the VSS method by considering the action of the surface tension forces.

Setting the shear stress at the surface to a constant value (like in the VSS method) forces the fluid into a motion with velocity gradients. Or one can say that the boundary condition is satisfied when the flow field possesses velocity gradients at the surface that fulfill the expression

\[ \tau_{xy} = 2.394 \frac{N}{m^2} = \mu \left( \frac{\partial u}{\partial y} + \frac{\partial v}{\partial x} \right) S \approx \mu \left( \frac{\partial u}{\partial y} \right) S \]

Since the initial drop shape is the state of minimum surface tension energy (fig. 20), any deviation from this state induces surface forces that try to restore the initial surface position and hinder the acceleration of surface fluid elements. Hence the velocity gradient resulting from the shear stress boundary condition is established not only by accelerating fluid elements at the surface in the downstream direction, which was the desired effect, but also by moving fluid particles inside the flow in the upstream direction. This flow field satisfies all given boundary conditions but certainly does not reflect the physical facts as observed in the Fluid Mechanics Laboratory at Ames.

### 3.2.3 XCSF Vectors

Figure 26 shows the oil drop surface and the resulting volume shear force vectors in the region of highest curvature in the “standard” case. (It should be mentioned that the \( x \)- and \( y \)-axes are scaled equally in this plot.) The width of the transition region and the alignment of the XCSF vectors tangential to the surface can be seen very well and seem to be physically correct.

An additional comment should be made at this point about the computation of \( \vec{F}_{\tau v}(\vec{z}) \). In RIPPLE, equation (22) is implemented with an additional term,

\[ \vec{F}_{\tau v}(\vec{z}) = -\sigma(\vec{z}) \nabla \vec{F}(\vec{z}) \cdot 2\vec{F}(\vec{z}) \]  \hspace{1cm} (29)
The multiplication by $2F(x)$ is permissible, since it does not affect the validity of

$$\lim_{h \to 0} \int_{\delta V} \bar{r}_s(x) dV = \int_{\delta S} \bar{r}_s(x) dS$$

which is the equivalent expression to equation (11) for the volume shear force. This is obvious because through the definition of the mollified function (i.e., $F(x)$ in Ripple), the limit expression (eq. (15)) becomes

$$\lim_{h \to 0} [F(x) \cdot 2F(x)] = \frac{F_1 + F_2}{2} \lim_{h \to 0} F(x) = H(n(x_s) \cdot (x - x_s))$$

The additional factor in equation (29) biases the volume force more into the fluid than suggested in figure 14. Figure 26 shows how the shear force vectors are more pronounced in the part of the transition region that lies inside the fluid.

The biasing yields the advantage that accelerations due to the shear forces are proportional to $\nabla F$ rather than $F$ itself. $F_{\nu T}$ becomes then a true body force. (A similar procedure for interfacial volume forces due to surface tension is discussed in references 7 and 9.)

### 3.2.4 Computational Conclusions

In an effort to extend Ripple's capabilities to model surface shear forces, two different approaches were implemented and evaluated.

One is the VSS method, which simply sets the viscous shear stress in surface cells to the desired value. The second approach, which is a more refined stress application technique, is an extension of the CSF model that was derived and implemented.

Both methods were tested with the "standard case," which reflects the physical situation of the flow channel facility where the 3D-FISF technique was developed (ref. 4).

It was found that the XCSF model gives more physical results. In particular, the circulating velocity field inside the drop computed by the VSS method is thought to be unphysical. The velocity of the contact line at the trailing edge of the oil, resulting from the VSS computations, does not agree with the theory of advancing contact lines.
Thus the investigation of surface tension and wall adhesion effects on the FISF technique in the following subsection is carried out using the XCSF model.

3.3 Comparison with Previous Studies; Surface Tension and Wall Adhesion Effects

In this subsection, we will first present $h(t)$ graphs obtained from the Lax-Wendroff algorithm and the linearized equation (3), which is used in the FISF technique.

Subsequently, the impact of wall adhesion is investigated through the variation of the static contact angle. The effects of surface tension are illustrated through computational results obtained using different values of the surface tension coefficient.

3.3.1 Lax-Wendroff Results and the Linearized Equation

Figure 27 shows the $h(t)$ plots of the linear solution and Lax-Wendroff computation at a distance $x = 1.5$ mm from the leading edge. One can see that the plots lie on top of each other after about 10 s. At this time, the fluid height is approximately $3 \cdot 10^{-3}$ mm. In order to resolve the flow in an oil drop of comparable height using RIPPLE, a much finer grid than described in subsection 3.1 would be necessary. Since the limiting time step in all RIPPLE computations was determined by the viscous time step constraint, the maximum allowable time step decreases by the power of 2 with a grid refinement (eq. (27)). In addition, the number of grid points would increase quadratically.

Thus a grid that resolves the oil flow in a $3 \cdot 10^{-3}$ mm high drop would yield a tremendous increase of computation time. Currently, the CPU time for a typical computation of the drop evolution up to 2.4 s amounts to 108 CPU hours on an SGI R10000 machine.

![Figure 27. Lax-Wendroff and linearized equation, $x = 1.5$ mm.](image)
Since surface tension forces are localized body forces, it is assumed that surface tension effects are limited to regions of high curvature. From the drop evolution figures in the preceding section, one can see that the area of high curvature is located near the trailing edge, which moves downstream—away from the region of interest as far as the FISF measurement technique is concerned. Hence the effects of surface tension on \( h(t) \) at \( x = 1.5 \text{ mm} \) are expected to decrease with time and the time frame of 2.4 s is thought to be sufficient for the investigations.

An additional note must be made about the “high sigma” computation (\( \sigma = 0.0624 \text{ N/m} \)). Although the resulting oil height at \( x = 1.5 \text{ mm} \) from the leading edge is lower than one cell width, the oil flow was computed up to 3.4 s. This was done in order to compare the global behavior of the oil drop flows computed with the XCSF and VSS methods only (refer to subsection 3.2). The behavior of the flow in the wedge area, and in particular \( h(t) \) for \( x = 1.5 \text{ mm} \), is not expected to be resolved sufficiently at times greater than approximately 2.4 s. Hence it is not referenced in the following discussion.

### 3.3.2 The Influence of Wall Adhesion

In order to determine the effect of wall adhesion on the oil flow, two cases were computed with RIPPLE, where the contact angle was set to \( \theta_s = 0.1^\circ \) and \( \theta_s = 6^\circ \), respectively. (\( \sigma \) was specified to 0.0208 N/m in both cases.)

The relatively low velocity of the contact line suggests that the dynamic contact angle does not differ significantly from the static contact angle. Thus the given values for \( \theta_s \) are thought to be extrema, which are well suited to answer the question of the influence of wall adhesion on the drop motion, and hence on the FISF technique.

Figure 28 shows the results of the two cases. The computed drop shapes are almost identical at all plotted times. In particular, the surface lines coincide in the wedge area, which is the interesting area of the oil drop as far as the FISF technique. Even in the vicinity of the two contact lines, no significant differences are visible.

This result is not surprising, since it follows from Young’s formula (eq. (3)) that the resulting horizontal force acting on the contact line in the case of nonequilibrium is proportional to \( \cos \theta_s \), which varies between 0.999 and 0.995 for the specified angles. This means that the difference between the contact line forces in the two cases is marginal and has no significant effect on the oil flow.

![Figure 28. Drop evolution at 0, 0.4, 1.4, and 2.4 s.](image-url)
Two conclusions can be drawn from the results shown in figure 28: (1) wall adhesion forces do not affect the oil drop motion for a wide range of $\theta$, and (2) the simplification made in RIPPLE, that the contact angle at the moving contact line is constant and equal to the dynamic one, is a valid approximation for the flow problem studied in this work.

3.3.3 Surface Tension Effects

**RIPPLE Compared with Lax-Wendroff**—Figure 29 displays the drop evolution in the wedge area obtained from the Lax-Wendroff computation and RIPPLE. The spatial restriction to the front part of the drop arises from assumptions implicit in the implementation of the Lax-Wendroff algorithm. It should be noted that the $x$-direction grid spacing used for these two computations is the same.

One can see that the oil quickly forms a wedge in both computations. The slope of the surface line computed with the Lax-Wendroff technique decreases faster at early times than the one obtained from RIPPLE. This behavior is not surprising, since surface tension was identified to be a retarding force that tries to establish the initial shape and is omitted entirely in the Lax-Wendroff calculation.

**Variation of $\sigma$ in RIPPLE**—The retarding effect of surface tension on the oil motion is also visible in figures 30–32, where the drop evolutions from RIPPLE computations for $\sigma = 0.0208$ N/m and $\sigma = 0.0624$ N/m are shown.

The trailing edge moves slower in the downstream direction and the surface line is less curved for the case with $\sigma = 0.0624$ N/m. The point where the drop shape deviates from the wedge form is located more upstream than in the “standard” case.

In figure 33, surface tension volume forces are plotted as functions of time for the two cases. It is interesting to see that although the surface line is less curved in the “high sigma” case, more energy per unit volume is used for the surface deformation. This explains the slower downstream motion of the oil drop in the high sigma case: since the flow energy introduced to the oil through the external air flow is the same at $t = 0$ in both computations, the higher surface tension energy causes a reduction of kinetic energy.

![Figure 29. Drop evolution (wedge area), $\sigma = 0.0208$ N/m, $\theta_s = 3^\circ$.](image-url)
It looks like surface tension not only affects the surface line curvature but also the overall fluid flow. This seems to be in contradiction with the statement made previously that surface tension forces are localized body forces with local effect on the fluid flow.

Looking at the global drop shape, all computations show a similar flow behavior: the back part of the drop surface is more or less curved while the front part becomes a wedge with a straight surface line. After a certain period of time, all computed drops show quite a uniform downstream motion, which means that the thinning rate of the wedge, $h(t)$ (which is measured in the FISF technique), does not differ fundamentally when surface tension is included in the calculation. It is thought that once the uniform state of motion is reached, the change of $h(t)$ by including surface tension can be extrapolated in time in order to estimate the total effect of surface tension on the FISF technique. This is presented in the following subsection.
The Influence on h(t) and the FISF Technique—The influence of surface tension on the wedge thinning rate is illustrated in figure 34, where the RIPPLE results for $\sigma = 0.0208 \text{N/m}$ and $\sigma = 0.0624 \text{N/m}$ as well as the Lax-Wendroff results are included.

The h(t) graphs lie on top of each other until about 0.2 s. After this time, the surface deformation is high enough so that the resulting surface tension force restrains the drop motion, and hence retards the evolution of the wedge. One can see that increasing $\sigma$ in the RIPPLE computations yields a greater wedge height at the same location and time.

At about the 0.6 s point, the various solutions for h(t) approach straight lines in the logarithmic scaled plot of figure 34. After this time, RIPPLE's h(t) graphs remain parallel to the Lax-Wendroff
graph, which approaches the linear equation,

\[ h(t) = \frac{\mu_{\text{oil}} x}{\tau_{\text{wall}} t} = \frac{\text{constant}}{t} \]

as shown in figure 27.

Functions of this form (having different y-values but the same slope in a logarithmic scale) differ only by a constant factor. Thus an analytic expression for \( h(t) \) obtained from RIPPLE can be derived when “scaling” the Lax-Wendroff solution with an appropriate factor \( f \).

\[ h(t)_{\text{RIPPLE}} = f \cdot h(t)_{\text{Lax-Wendroff}} = f \cdot \frac{\text{constant}}{t}, \quad \forall t \geq 1.5 \text{ s} \]

With this equation, the relative height difference between RIPPLE’s results (including surface tension) and the Lax-Wendroff computation,

\[ d_{\text{rel}} = \frac{h(t)_{\text{RIPPLE}} - h(t)_{\text{Lax-Wendroff}}}{h(t)_{\text{RIPPLE}}} = \frac{f - 1}{f} \]

becomes a constant expression rather than a function of time.

To determine \( f \), the heights at the latest possible time are used, in order to be as close as possible to the point where the Lax-Wendroff computation coincides with the linear expression (fig. 27). Since the computation with \( \sigma = 0.0624 \text{ N/m} \) is a numerical experiment not reflecting the physical situation, \( f \) was calculated only for \( \sigma = 0.0208 \text{ N/m} \). The results are summarized in table 2:

<table>
<thead>
<tr>
<th>Time</th>
<th>( h_{\text{Lax-Wendroff}} )</th>
<th>( h_{\text{RIPPLE}} )</th>
<th>( f )</th>
<th>( d_{\text{rel}} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.4 s</td>
<td>( 1.07511 \cdot 10^{-2} \text{ mm} )</td>
<td>( 1.15685 \cdot 10^{-2} \text{ mm} )</td>
<td>1.0707</td>
<td>7.07%</td>
</tr>
</tbody>
</table>

Comments on \( f \) - RIPPLE’s results show that the specified shear stress value yields a 7% smaller oil thinning rate when surface tension is included than predicted by the Lax-Wendroff computation and the linearized solution. In order to obtain the same oil thinning rate with RIPPLE, one would have to specify a 7% higher shear stress value. This means that the shear stress level measured with the FISF technique (which uses the linear expression to relate shear stress to oil height) is underpredicted by 7%. However, the value for \( f \) listed in table 2 is based on data that are taken at a very early time (2.4 s) compared with the typical time in the FISF technique (e.g., 100 s). In addition, it was assumed that the linear oil thinning behavior in RIPPLE after 1.5 s persists, which is not a certainty since the oil flow is governed by nonlinear equations. There is limited evidence measured in the laboratory that indicates that the linear oil thinning behavior does not persist at later times. If this proves to be the case then the actual underprediction of \( \tau_{\text{wall}} \) may be much less. Thus 7% underprediction for \( \tau_{\text{wall}} \) by the FISF is a very rough estimate, which requires more investigation. An additional uncertainty arises from the fact that in RIPPLE only the oil flow is modeled and the effect of the surrounding air flow is reduced to the
assumption that shear forces at the interface are the only way the air flow acts on the oil. Any further interaction between the oil flow and the air boundary layer (e.g., pressure gradients) is omitted. Thus the value for $f$ should be interpreted as a more qualitative statement rather than an absolute "correction factor" for the FISF technique.

It is also expected that $f$, and hence the influence of surface tension, is less for larger volumes of oil. By enlarging the oil drop, surface line curvature decreases while the area where shear forces are applied increases. This means that surface tension forces are weakened and the viscous forces at the interface are amplified.

### 3.3.4 Influence of Grid Resolution

The purpose of the last subsection of this section is to verify the adequacy of the chosen grid resolution. For this purpose, the previously described "standard" case was computed on a grid, where $\Delta x = 0.05 \text{ mm}$ and $\Delta y = 0.02 \text{ mm}$. This corresponds to a doubling of the grid spacing in both directions.

In figure 35, $h(t)$ is plotted for the two different grids. The two graphs lie on top of each other until the oil height in the coarse grid case is significantly less than one cell height. At that time (i.e., approximately 2 s), the oil height begins to decrease more rapidly in the computation with the coarse grid. The fact that the oil thinning rate in the case with the doubled grid spacing is almost the same as on the finer grid (until about 2 s) shows that the grid in the previously discussed computations is fine enough for our purposes.

![Figure 35. The influence of grid resolution on $h(t)$, $x = 1.5 \text{ mm}$.](image-url)
4 CONCLUSIONS AND RECOMMENDATIONS

(1) A computational study was performed to investigate the effects of surface tension and wall adhesion on the accuracy of the FISF technique for the measurement of skin friction.

The computer code RIPPLE, which solves the two-dimensional, incompressible, transient Navier-Stokes equations for free surface flows, was used to compute the flow of an oil line placed on a flat plate in an air boundary layer. RIPPLE was modified in order to model shear forces acting on the free surface. In this context, two different approaches for applying shear stresses at the interface were developed and discussed. One is the viscous shear stress model, where the value for the skin friction measured with the FISF technique is set explicitly in grid cells containing the interface. The second approach is an extension of the continuum surface force model (CSF) for tangential forces, where skin friction is converted into a localized body force and simply added to the momentum equations.

The two models were compared and it was shown that the extended continuum surface force model is ideally suited for the application of tangential surface forces (i.e., shear forces). It was used for the subsequent study of surface tension and wall adhesion effects on the oil drop motion and hence the accuracy of the FISF technique.

(2) The surface tension coefficient $\sigma$ was varied to illustrate the effect on the oil drop motion and deformation. Different static contact angles were specified to investigate the influence of wall adhesion. In all case studies, special focus was made on the thinning rate of the leading edge of the oil drop, since this is the drop area where the FISF technique is applied. In addition, RIPPLE's result for the oil thinning rate was compared with previous studies discussed in reference 4.

(3) It was demonstrated that the variation of the contact angle does not affect the flow of the oil drop, and hence wall adhesion has no impact on the accuracy of the FISF technique.

The variation of the surface tension coefficient and the comparison with Zilliac's work (ref. 4) showed that surface tension has a retarding effect on the oil thinning rate, which may lead to an underprediction of the skin friction measured with the FISF technique. This effect can be reduced by enlarging the drop volume, which yields a lower surface curvature and a higher sum of driving tangential shear forces at the interface.

(4) Finally, some recommendations for improving the results presented in this paper and reducing the uncertainty of the FISF accuracy estimation are now made in terms of future work:

- Establish an implicit treatment of the viscous term in the momentum equation in order to overcome the restricting viscous time step constraint and to be able to compute further in time on a even finer grid.
- Extend RIPPLE to two fluids in order to model the entire physics involved in the flow.
- Include a turbulence model for modeling the turbulent air boundary layer.
- Extend RIPPLE to three dimensions to validate the 3D-FISF technique.

Some of these items, such as the capability of modeling two fluids, are currently under development.
APPENDIX A

OIL PROPERTIES

A.1 Dow Corning Silicone Fluid

<table>
<thead>
<tr>
<th>Name</th>
<th>200 Fluid</th>
</tr>
</thead>
<tbody>
<tr>
<td>Manufacturer</td>
<td>Dow Corning Corporation, USA</td>
</tr>
<tr>
<td>Density at 25°C</td>
<td>9600 kg/m³</td>
</tr>
<tr>
<td>Surface tension at 25°C</td>
<td>0.0208 N/m</td>
</tr>
<tr>
<td>Kinematic viscosity at 25°C</td>
<td>50 Cs</td>
</tr>
</tbody>
</table>

A.2 Description

The 50 Cs 200 Fluid from Dow Corning is a medium viscosity poly-dimethyl-sil-oxane polymer. It has the following typical chemical composition:

\[(\text{CH}_3)_3\text{SiO}[\text{SiO(CH}_3)_2]^n\text{Si(CH}_3)_3\]
APPENDIX B

RIPTTLE

The original version of RIPPLE is available through the National Energy Software Center, 9700 South Cass Avenue, Argonne, IL 60439.

B.1 A Typical RIPPLE Input Deck

Oil drop
$numparam
conserve=.false., alpha=2.0,
twfin=2.6, prtdt=500.2, pltdt=0.2,
delt=0.0000001, dtmax=0.1,
restart=-1.0,
dmpdt=0.2,
kb=2,kt=2,kl=2,kr=2,
autot=1.0D0,
sym=.true.,
gfncrn=.true.,
$end
$fldparam
icyl=0, isurf10=1,
gy=-9799.3,
cangler=3.0,cangle1=3.0,
cangle2=3.0,cangleb=3.0,
xnu=50.0, rho=0.000096, sigma=20.8,
psat=101300.0,
$end
$mesh
nkx=1, xl=0.0,0.0, xc=4.0,
nx1=160, nxr=160, dxmn=0.025,
nky=1, yi=0.0,0.12, yc=0.01,
nyl=1, nyr=11, dymn=0.01,
$end
$shear
sheart = 2.394,
$end
$hovertlist
xloc=1.7d0,
$end
$timestepadj
tmfac=1.5d0,
$end
$obstcl
nobs=0,
$end
$freesurf
nfrsr=1,
fa(1)=5.4, fa2(1)=-1,
fb1(1)=-95.4056834388, fb2(1)=-1,
fcl(1)=-1.04,
ifh(1)=1,
This input deck was used to compute the previously discussed "standard" case. For a detailed discussion of the input data, see reference 7.

B.2 Source Code of Main Subroutines for the Implementation of the XCSF Model

Two subroutines are listed. The first one serves the purpose of computing the fluid accelerations due to surface shear forces. The second one computes the surface shear forces using the XCSF method. Many other subroutines were modified or added to the original version of the code but they will be omitted in this paper for reasons of space.

```c
subroutine shrfrce

c======================================================================
cC======================================================================
cC Purpose -
cC compute fluid acceleration due to shear forces
C
C SRFFRCE is called by -
c
Cname name name name name name name name
C
C RIPPLE
C
C SRFFRCE calls the following subroutines and functions -
c
Cname source name source name source name source
C
C BC ripple
C======================================================================
c
#include "32bit.h"
#include "comdk2.h"
#include "iccgdkl.h"
data tiny /1.0d-25/, zero /0.0d0/
```
c

do 20 j=2,jml
   do 20 i=2,iml
      ij=(j-1)*imax+i
      ipj=i+j+1
      ipj=ij+imax

      rhiij=f(ij)*rhof
      rhoipj=f(ipj)*rhof
      rhoijp=f(ijp)*rhof
      rhoij=delx(i-1)*rhiij+delx(i)*rhoipj/dely(i)*delx(i+1)
      rhoipjdely=delx(j)*rhoij+delx(j+1)*rhoijp/dely(j)*delx(j+1)
      rhox=rhoc+tiny
      rho=1.0+rhox
      rho=1.0+tiny

      ftauxij=ftaux(ij)
      ftauipj=ftauy(ipj)
      ftauxipj=ftaux(ipj)
      ftauijdely=ftauy(ijp)

      fxrc=(delx(i)*ftauxipj+delx(i+1)*ftauxij)/(delx(i)+delx(i+1))
      if (gfnctn) fxrc=2.*rhox*fxrc/rhof

      fytc=(dely(j)*ftauyijp+dely(j+1)*ftauyij)/dely(j)+dely(j+1)
      if (gfnctn) fytc=2.*rhoy*fytc/rhof

      if (ar(ij).lt.1.0d-06) go to 10
      u(ij)=u(ij)+delt*fxrc/rho
      u(ij)=cvmgt(zero,u(ij),(rhox.lt.frctn*rhof).or.
         & (ar(ij).lt.1.0d-06))

      if (at(ij).lt.1.0d-06) go to 15
      v(ij)=v(ij)+delt*fytc/rho
      v(ij)=cvmgt(zero,v(ij),(rhoy.lt.frctn*rhof).or.
         & (at(ij).lt.1.0d-06))

      10 continue
   15 continue

20 continue

----
call bc
----
c
return
end
subroutine tension

Purpose -
Compute the volume forces due to surface tension

TENSION is called by -

name       name       name       name       name       name

---------       -------
RIPPLE      SRFEMIN

TENSION calls the following subroutines and functions -

name       source       name       source       name       source

---------       -------
BDYCELL       ripple      CRVATURE       ripple      SETARRY       ripple
ISMAX         cftmath     MOLLIFY       ripple

c********************************************************************
c
 implicit real*8 (a-h,o-z)
#include "32bit.h"
c********************************************************************

c###########
#include "comdk2.h"
#include "iccgdkl.h"
c###########

dimension ro(nxy)
dimension vnormx(nxy), vnormy(nxy)

data tiny /1.0d-25/, zero /0.0d0/

c.... zero-out the relevant arrays

---------
call setarry (ro,0.0d0,nxy)
call setarry (gradox,0.0d0,nxy)
call setarry (gradoy,0.0d0,nxy)
call setarry (tensx,0.0d0,nxy)
call setarry (tensy,0.0d0,nxy)
call setarry (fsv,0.0d0,nxy)
call setarry (ftilde,0.0d0,nxy)
call setarry (kappa,0.0d0,nxy)

Compute cell densities and copy the raw color function, $f(ij)$, into the smoothed color, $ftilde(ij)$

do 50 j=2,jml
   do 50 i=2,iml
      ij=(j-1)*imax+i
      ro(ij)=f(ij)*rhof
      ftilde(ij)=ro(ij)
   50 continue

Reflect $ro(ij)$ and $ftilde(ij)$ in ghost cells

call bdycell(iml,jml,i,1,1,1,0.0d0,0.0d0,0.0d0,pbc,ro)
call bdycell(iml,jml,i,1,1,1,0.0d0,0.0d0,0.0d0,pbc,ftilde)

Optionally smooth the color function

if (smooth)
   & call mollify(2,iml,2,jml,1,imax,nsmooth,ro,ftilde,af)

Compute vertex-centered normal vectors

do 35 j=2,jml+1
   do 35 i=2,iml+1
      ij=(j-1)*imax+i
      imjm=ij-imax
      imj=ij-i
      im=i-1
      jm=j-1
      deltax=0.5*(delx(j)+delx(jm))
      deltax=0.5*(delx(i)+delx(im))
      rhot=(delx(i)*ro(imj)+delx(im)*ro(ij))/(delx(i)+delx(im))
      trhot=(delx(i)*ftilde(imj)+delx(im)*ftilde(ij))/(delx(i)+delx(im))
      &
      rhob=(delx(i)*ro(imjm)+delx(im)*ro(ijm))/(delx(i)+delx(im))
      trhob=(delx(i)*ftilde(imjm)+delx(im)*ftilde(ijm))/(delx(i)+delx(im))
      &
      rhol=(delx(j)*ro(imojm)+delx(jm)*ro(imoj))/(delx(j)+delx(jm))
      trhol=(delx(j)*ftilde(imojm)+delx(jm)*ftilde(imoj))/(delx(j)+delx(jm))
      &
      rhor=(delx(j)*ro(imojm)+delx(jm)*ro(imoj))/(delx(j)+delx(jm))
      trhor=(delx(j)*ftilde(imojm)+delx(jm)*ftilde(imoj))/(delx(j)+delx(jm))
      &
      gradrox(ij)=(rhor-rhol)/deltax
      gradroy(ij)=(rhob-trhob)/deltay

38
c.... store the non-unit surface normal vectors
cvnormx(ij)=gradrox(ij)
vnormy(ij)=gradroy(ij)
cgradftx(ij)=(trhor-trhol)/deltax
gradfty(ij)=(trho-trob)/deltay

35 continue

c.... Impose wall adhesion via an equilibrium contact angle in all obstacle cells
do 2350 j=2,jml
do 2350 i=2,iml
    ij=(j-1)*imax+i
    if (ijobs(ij).eq.0) go to 2350
    if (ijobs(ij).eq.1) thetaeq=cangleb
    if (ijobs(ij).eq.2) thetaeq=canglet
    if (ijobs(ij).eq.3) thetaeq=canglel
    if (ijobs(ij).eq.4) thetaeq=cangler
grdmgro=sqrt(gradrox(ij)**2 + gradroy(ij)**2 + tiny
grdmgft=sqrt(gradftx(ij)**2 + gradfty(ij)**2 + tiny
    tx=gradnwy(ij)
ty=gradnwx(ij)
csthtar=(tx*gradrox(ij)+ty*gradroy(ij))/grdmgro
csthtaf=(tx*gradftx(ij)+ty*gradfty(ij))/grdmgft
txr=cvmgt(-tx,tx,csthtar.lt.zero)
tyr=cvmgt(-ty,ty,csthtar.lt.zero)
txf=cvmgt(-tx,tx,csthtaf.lt.zero)
tyf=cvmgt(-ty,ty,csthtaf.lt.zero)
cgrdtwrx=grdmgro*txr
grdwry=grdmgro*tyr
grdnwrx=grdmgro*gradnwx(ij)
grdnwry=grdmgro*gradnwx(ij)
grdtwfx=grdmgft*txf
grdwfy=grdmgft*tyf
grdnwfx=grdmgft*gradnwx(ij)
grdnwfy=grdmgft*gradnwy(ij)
cgradrox(ij)=grdnwrx*cos(thetaeq)+grdtwrx*sin(thetaeq)
gradroy(ij)=grdnwry*cos(thetaeq)+grdtwry*sin(thetaeq)
gradftx(ij)=grdnwfx*cos(thetaeq)+grdtwfx*sin(thetaeq)
gradfty(ij)=grdnwfy*cos(thetaeq)+grdtwfy*sin(thetaeq)

2350 continue

c.... Compute the mean curvature, kappa, using the smoothed color function, ftilde(ij)
c__________________
call curvature(2,iml,2,jml,1,imax,ftilde,gradftx,gradfty,
& delx,dely,ri,kappa,tiny)
c__________________
do 110 j=2,jml
    do 100 i=2,iml
        ij=(j-1)*imax+i
        ipj=i+j-1
        imj=i+j-1
        ipjp=ij+imax+1
        ijp=ij+imax
        ijm=ij-imax
        im=1-i
        ip=1-i
        jm=1+j
        jp=1+j
        c
        avnx=0.25*(gradrox(ipj)÷gradrox(ipjp)
        & +gradrox(ijp)+gradrox(ij))
        avny=0.25*(gradroy(ipj)÷gradroy(ipjp)
        & +gradroy(ijp)+gradroy(ij))
        vx=0.25*(vnormx(ipj)+vnormx(ipjp)
        & +vnormx(ijp)+vnormx(ij))
        vy=0.25*(vnormy(ipj)+vnormy(ipjp)
        & +vnormy(ijp)+vnormy(ij))
        c
        . . . compute the cell centered surface tension force
        c
tensin=sigma*kappa(ij)/rhof
        tensx(ij)=avnx*tensin
        tensx(ij)=cvmgt(zero,tensx(ij),ac(ij).lt.em6)
        c
tensy(ij)=avny*tensin
        tensy(ij)=cvmgt(zero,tensy(ij),ac(ij).lt.em6)
        fsv(ij)=sqrt(tensx(ij)**2 + tensy(ij)**2)
        c
        . . . compute the cell centered shear force
        c
        ftau(ij)=-(sheart)*vy/rhof
        ftau(ij)=cvmgt(zero,ftau(ij),ac(ij).lt.em6)
        ftau(ij)=(sheart)*vx/rhof
        ftau(ij)=cvmgt(zero,ftau(ij),ac(ij).lt.em6)
        ftau(ij)=0.0d0
        100 continue
        110 continue
        c
        ijmax=ismax(nxy,fsv,1)
        fmax=fsv(ijmax)
        ffloor=0.001*fmax
        do 200 j=2,jml
            do 200 i=2,iml
                kappa(ij)=cvmgt(kappa(ij),zero,fsv(ij).gt.ffloor)
            200 continue
        c
        9999 return
end
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**Computational Study of Surface Tension and Wall Adhesion Effects on an Oil Film Flow Underneath an Air Boundary Layer**

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**RIPPLE**, developed at Los Alamos National Laboratories, was used to compute the time development of an oil drop on a surface under a simulated air boundary layer. RIPPLE uses the volume of fluid method to track the surface and the continuum surface force approach to model surface tension and wall adhesion effects. The development of an oil drop, over a time period of approximately 4 seconds, was studied. Under the influence of shear imposed by an air boundary layer, the computed profile of the drop rapidly changes from its initial circular-arc shape to a wedge-like shape. Comparison of the time-varying oil-thickness distributions computed using RIPPLE and also computed using a greatly simplified numerical model of an oil drop flow (solution of Squires oil-drop equation which does not include surface tension and wall adhesion effects) was used to evaluate the effects of surface tension on FISF measurement results. The effects of surface tension were found to be small but not necessarily negligible in some cases.

**Surface tension, Skin friction, Computational fluid dynamics**

**Unclassified**