Hydrodynamics Across a Fluctuating Interface

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ABSTRACT

Understanding what happens inside the rippling and dancing surface of a liquid remains one of the great challenges of fluid dynamics. Using molecular dynamics (MD) we can pick apart the interface structure and understand surface tension. In this work we derive an exact mechanical formulation of hydrodynamics for a liquid-vapour interface using a control volume which moves with the surface. We show that surface dynamics, represented by the thermal fluctuations of the surface at the molecular scale, needs to be included in the hydrodynamic expressions for the exact balance of mass and momentum across the interfacial region.

We provide a mathematical framework to obtain the local definition of hydrodynamic fluxes at any point on the surface. These are represented not only by the flux of molecules and intermolecular interactions acting across surface, but also as a result of the local curvature and movement of the surface itself. By explicitly including the surface dynamics in the equations of motion, we demonstrate an exact balance between kinetic and configurational pressure normal to the surface. The hydrodynamic analysis makes no assumptions regarding the probability distribution function. Therefore, we provide a new technique valid for any system arbitrarily far from thermodynamic equilibrium giving an exact book keeping method to dissect the liquid-vapour interface. As a result, the presented equations can provide a theoretical basis for the study of problems such as bubble nucleation, droplet dynamics and liquid-vapour instabilities.

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I. INTRODUCTION

The liquid-vapour interface stands out as one of the great modelling challenges in engineering and the physical sciences. The first challenge concerns the observation that, while bulk fluids and gases are well described by continuum models, this approach breaks down at the interface, where large changes in the physical properties, localised to a very small region, invalidate any continuum assumptions. Molecular dynamics (MD) is an ideal model to study a problem of this type, modelling the liquid-vapour coexistence with no more than the solution of Newton’s law for a system of molecules.

The second challenge concerns the ability to translate the information at the molecular level to relevant continuum hydrodynamic properties. Thermal fluctuations, characteristic of the dynamics at the atomistic scale, are also present at the interface, blurring any property of interest. Capillary wave theory (CWT)\textsuperscript{12,29} provides a framework to describe these fluctuations, by representing the instantaneous shape of the fluid surface as an intrinsic surface in parametric form. Thermodynamic profiles across the interface represent the convolution of an assumed sharp intrinsic profile with the Gaussian distribution characterising the height of the intrinsic surface.

However, a mechanical route is also possible, based on instantaneous time evolving equations with no need for thermodynamics averaging, equivalent to the Newtonian mechanics which underpin molecular dynamics. Using the sharp intrinsic surface directly as a moving reference frame, mechanical equations can be obtained from the molecular data with no thermodynamic blurring.

The most important quantity describing the mechanical properties of the interfacial region is arguably the pressure tensor and its corresponding profile across the interface. However, the appropriate form of the molecular stress tensor has long been the source of debate, known to be nonunique, based on the interaction path\textsuperscript{30} and measuring reference frame\textsuperscript{11}, with some debate about the inclusion of kinetic terms\textsuperscript{10}. For a homogeneous system of particles with periodic boundaries, the virial pressure tensor is defined assuming a single value for the whole system\textsuperscript{25}. In inhomogeneous systems such as the liquid-vapour interface, a single homogeneous pressure is no longer sufficient and we require a local definition. The
seminal work of Irving and Kirkwood\textsuperscript{17} provides a localisation to give the pressure at a point in space. However, this is of limited practical use in molecular dynamics, as it includes the problematic Dirac delta functions, a direct consequence of the mathematical idealisation of a continuum. Several approaches have been applied to solve this, including mollifying the Delta function\textsuperscript{1,22,23}, integrating to get "Volume Average" (VA) stress\textsuperscript{9,21} or reformulating in terms of the pressure over a surface\textsuperscript{13,37,38}, known as the method of planes (MOP) pressure. Of these approaches, the pressure over the surface has the advantage of being conceptually the simplest form, namely the force acting over the surface divided by the surface area $\Pi = F/A$. It is also the only form to give exactly conservative equations, shown through the concept of a control volume\textsuperscript{34}.

The control volume equations are important in fluid dynamics, providing the basis for the conservative finite volume approach in computational fluid dynamics (CFD)\textsuperscript{10}. By expressing a weakened statement of the equations of motion, valid in an average sense over an arbitrary volume, they no longer demand a continuous field. This makes the control volume approach ideally suited to molecular dynamics, expressing everything as an average over a volume. This allows us to express the Irving and Kirkwood\textsuperscript{17} equations with integrated Dirac delta functions, in a form which demonstrates exact conservation in an MD system. For this work, we take advantage of an additional benefit of the control volume approach, the ability to define them with arbitrary shape. We choose the liquid-vapour interface $\xi$ to define the face of our control volumes, so the edge of our averaging grid follows the 2D surface. To define this surface, we apply the intrinsic interface approach first proposed by Chacón and Tarazona\textsuperscript{7}, fitting a set of Fourier components to the outermost molecules in a liquid cluster. A range of other approaches to define an interface have been proposed\textsuperscript{18,26,31,39}, and the surface pressure presented in this work can, in principle, be applied to any other surface, not just the Chacón and Tarazona\textsuperscript{7} functional form.

An assumption of CWT is that the density and pressures profiles are uncorrelated with the intrinsic surface from which the profiles are obtained. However, in order to balance the control volume equations of motion, it will be shown that it is necessary to explicitly include the pressure change due to the surface time evolution. The surface itself becomes part of the equations of motion, an addition which is seen to be essential to getting the correct form of
the equations of motion. Constant normal pressure over a liquid-vapour interface is predicted in the literature\textsuperscript{9} and is a requirement for momentum balance in an MD system\textsuperscript{14,37}, but has not observed in the literature plotting pressure near a surface\textsuperscript{5,32}. In this work, we show that the novel surface pressure equations, obtained by a formal derivation of the control volume equations gives the expected flat profile. This flat profile requires the pressure to include both the instantaneous normal to the intrinsic surface at every time and a term for the surface movement itself.

The structure of this manuscript is as follows, in the theory section II the mathematical form of the control volume is derived to give the expression for surface fluxes on a volume moving with the intrinsic interface. The next section outlines the details of implementation III, giving an overview of how to actually use these expressions in molecular simulations and the details of the setup. The result and discussion are included in section V before concluding remarks in section VI.

II. THEORY

We start with a high level description of the process in this section. The control volume formula for surface pressure are derived by starting from the definitions of Irving and Kirkwood\textsuperscript{17}. These link the continuum expressions for density, momentum and energy to their molecular equivalents at any point in space. The process is then a formal integral of these definitions over a volume in space, followed by the evaluation of their time evolution to get the mass and energy conservation as well as the momentum balance equations.

The volume integral follows the interface, with functional form obtained from the intrinsic surface method by fitting to the outer molecules at the interface. For the reader only interested in applying these equations, the key formula to get pressure are given in section III B as Eq. (45) and Eq. (46). These have the simple interpretation of a grid which deforms and moves following the liquid-vapour interface. We obtain all intersections of molecular interactions or particle trajectories with surfaces of the cells in that grid. The interactions and forces over a surface are in the form of a force over area, $F/A$, the surface pressure. When summed over all surfaces of any enclosed volume they exactly define the
change inside, a requirement of the validity of the conservation laws expressed in control volume form. A piecewise bilinear approximation of the moving interface is used to get surface crossings more efficiently, and mapping applied to simplify calculations. The contribution due to the movement of the interface itself is obtained by considering particles before and after movement.

A. The Irving Kirkwood Equations

The density at a given point in space can be obtained using Irving-Kirkwood’s procedure\textsuperscript{17}, defined here without the ensemble average\textsuperscript{10}, to give the instantaneous quantity obtained at any time in a molecular dynamics simulation,

\[
\rho(r,t) = \sum_{i=1}^{N} m_i \delta(r - r_i).
\] (1)

Here \(\rho\) is the continuum density at point \(r\) in three dimensional space and time \(t\). The sum on the right adds the mass \(m_i\) over all \(N\) molecules in the system, with the Dirac delta function \(\delta\) only non-zero when \(r\) is equal to \(r_i\), that is molecule \(i\) is located at point \(r\). We can define momentum,

\[
\rho(r,t)u(r,t) = \sum_{i=1}^{N} m_i \dot{r}_i \delta(r - r_i),
\] (2)

and energy,

\[
\rho(r,t)E(r,t) = \sum_{i=1}^{N} m_i e_i \delta(r - r_i),
\] (3)

in an analogous manner, with \(\rho u\) and \(E\) continuum momentum and energy respectively. Energy here is \(e_i = v_i^2 + 1/2 \sum_{j \neq i} \phi_{ij}\) based on half the energy of the intermolecular interaction.

B. Control Volume integral

The continuum approximation represent reality as a continuous field, obtained by taking the zero limit of an infinitesimal volume at each point. This same concept applied to a
discrete system results in, an infinitely thin and infinite large function at a point, the Dirac delta (formally a generalised function). The Dirac delta can be thought of as a useful placeholder representing the continuum assumption in a molecular system, but has limited use in practice, particularly in software implementation. A more tractable form of the Dirac delta function, for use in a discrete system, is obtained by the integration over an arbitrary volume to get the control volume form. This has the advantage that a conservative set of equations can be obtained in a molecular system. These are directly relatable to the equivalent control volume expressions in the continuum. More importantly for this work, the control volume shape can be chosen based on the geometry of interest. In this work, the volume is chosen to follow the intrinsic surface as it varies in time. The density at a point, Eq. (1), can be integrated over a volume as follows,

$$\int_V \rho(r,t) dV = \sum_{i=1}^{N} m_i \int_V \delta(r - r_i) dV, \quad (4)$$

similar for momentum Eq. (2),

$$\int_V \rho(r,t) u(r,t) dV = \sum_{i=1}^{N} m_i \dot{r}_i \int_V \delta(r - r_i) dV, \quad (5)$$

and energy Eq. (3),

$$\int_V \rho(r,t) E(r,t) dV = \sum_{i=1}^{N} m_i e_i \int_V \delta(r - r_i) dV. \quad (6)$$

To evaluate the integral of Eqs. (1-3), only the Dirac delta function must be integrated. The volume described by the triple integral is between four flat surfaces and two faces following the shape of the intrinsic surface $\xi$,

$$\int_V \delta(r - r_i) dV = \int_{x-}^{x+} \int_{y-}^{y+} \int_{\xi-}^{\xi+} \delta(x - x_i) \delta(y - y_i) \delta(z - z_i) dxdydz, \quad (7)$$

here the arbitrary volume is cuboidal in $x$ and $y$ directions denoted by plus and minus superscripts for top and bottom surfaces. The location of these surfaces are $x^+ \equiv x + \Delta x/2$ and $x^- \equiv x - \Delta x/2$ respectively with $\Delta x$ the $x$ width of the CV with centre at point $x$, with a similar definition using $\Delta y$ as width in $y$. The surface in the $z$ directions is described by
position $z^\pm \equiv z \pm \Delta z/2$ and $\xi(x, y, t)$, a continuous function of $x$, $y$ and time $t$, so surface position in $z$ denoted by $\xi^\pm(x, y, t) \equiv z(t)^\pm + \xi(x, y, t)$. In general, $\xi$ can be any function and we will make no assumption about its form until section [IV A] where a sum of trigonometric function will be used to describe the intrinsic surface between a liquid and vapour phase.

As the $\xi^\pm$ limits are a function of $x$ and $y$, the $z$ integral must be evaluated first,

$$\int_V \delta(r - r_i) \, dV = \int_{x^-}^{x^+} \int_{y^-}^{y^+} m_i \delta(x - x_i) \delta(y - y_i)$$

$$\times \left[ H(\xi^+(x, y, t) - z_i) - H(\xi^-(x, y, t) - z_i) \right] \, dy \, dx. \quad (8)$$

The Dirac delta is the Heaviside function upon integration, with the finite limits $\xi^-$ and $\xi^+$ inserted. The next two integrals over $x$ and $y$ use the sifting property of the Dirac delta, namely $\int \delta(x - a) f(x) = f(a)$ so the function $\xi(x, y, t)$ which describes the surface roughness becomes expressed in terms of molecular position,

$$\vartheta_i \equiv \int_V \delta(r - r_i) \, dV = \left[ H(x^+ - x_i) - H(x^- - x_i) \right]$$

$$\times \left[ H(y^+ - y_i) - H(y^- - y_i) \right]$$

$$\times \left[ H(\xi^+(x_i, y_i, t) - z_i) - H(\xi^-(x_i, y_i, t) - z_i) \right]$$

$$= \Lambda_{xi} \Lambda_{yi} \tilde{\Lambda}_{zi}, \quad (9)$$

where $\vartheta_i$, a function which selects molecules inside the Heavisides, called the control volume function. The $\Lambda$ notation is also introduced, a box car, or Bracewell function for each direction, so e.g. $\Lambda_{xi} = H(x^+ - x_i) - H(x^- - x_i)$ and the tilde on $\Lambda_{zi}$ indicates the function moves with the intrinsic surface $\tilde{\Lambda}_{zi} = H(\xi^+_i - z_i) - H(\xi^-_i - z_i)$ where the subscript $i$ on $\xi$ denoting the function is in terms of the molecular positions $\xi^+_i = \xi^+(x_i, y_i, t)$. As each $\Lambda$ is one if the particle is between the limits in that coordinate direction, the product $\Lambda_{xi} \Lambda_{yi} \tilde{\Lambda}_{zi}$ is therefore one if the particle is inside the volume; located between two intrinsic surface functions in the $z$ direction and bounded by two planes in the $x$ and $y$ directions. Any molecules outside of this volume will return a value of zero. This control volume described by $\vartheta_i$ has a constant width and height of $\Delta x$, $\Delta y$ respectively with the depth $\Delta z$ always constant at any $x, y$ location, as the same intrinsic interface function $\xi$ is used for top and bottom surfaces, so $\xi^+_i - \xi^-_i = \Delta z$.
C. Mass

The expression linking the control volume form of density in a continuum and molecular system of Eq. (4) can therefore be written concisely using Eq. (9) as,

\[ \int_V \rho(r, t) dV = \sum_{i=1}^{N} m_i \vartheta_i. \]  

(10)

That is, the mass of any molecule in the enclosed region between two intrinsic surfaces in \( z \) and four planes in \( x \) and \( y \) is contributing to the density in that control volume.

We now use the control volume function to derive expressions for the fluxes over the surface of a volume, the so-called flux forms of the equations of motion. In the continuum, the control volume analysis involves taking the time derivative of the mass in a volume to obtain the flux over the surfaces of that volume:

\[ \frac{d}{dt} \int_V \rho(t) dV = - \oint_{S+} \rho \mathbf{u} \cdot dS^+ - \oint_{S-} \rho \mathbf{u} \cdot dS^- + \int_{S^+} \rho \mathbf{u} \cdot dS + \int_{S^-} \rho \mathbf{u} \cdot dS, \]  

(11)

where \( dS = n dS \) expresses an infinitesimal surface element \( dS \) with surface normal \( n \) and \( \oint \) indicates the integral is over all surfaces of the volume, which here represents six piecewise surface integrals. The molecular control volume has been shown previously derived for a uniform cuboid in space, a sphere, and is extended here for a general volume between intrinsic surfaces. Taking the time derivative of Eq. (10),

\[ \frac{d}{dt} \int_V \rho(t) dV = \sum_{i=1}^{N} m_i \dot{\vartheta}_i = \sum_{i=1}^{N} m_i \frac{d \vartheta_i}{dt} = \frac{d \Lambda_x}{dt} \Lambda_y \Lambda_z + \Lambda_x \frac{d \Lambda_y}{dt} \Lambda_z + \Lambda_x \Lambda_y \frac{d \Lambda_z}{dt}. \]  

(12)

The derivative of each of the three \( \Lambda \) functions will generate two terms for the top and bottom surfaces corresponding to the six surfaces of the volume. For example in \( x \) the derivative \( d \Lambda_x/dt = dH(x^+ - x_i)/dt - dH(x^- - x_i)/dt \) can be seen to give two terms for the top and bottom surface in the \( x \) direction. Consider just the top, or +, surface in \( x \),

\[ \frac{d}{dt} H (x^+ - x_i) = -\dot{x}_i \delta (x^+ - x_i), \]  

(13)

can be seen to be the particle’s \( x \) velocity \( \dot{x}_i \) localised by a delta function to the \( x^+ \) surface, the mass flux of a particle over the surface. The same process can be applied for \( x^- \) and \( y^\pm \).
surfaces. We obtain the top surface in $z$ from $d\tilde{A}_z/dt$ where both the molecular positions and the intrinsic surface $\xi_i^\pm$ depend on time,

$$\frac{d}{dt} \{ \xi_i^+ - z_i \} = \left[ \frac{d\xi_i^+}{dt} - \frac{dz_i}{dt} \right] \delta (\xi_i^+ - z_i)$$

$$= \left[ \dot{x}_i \frac{\partial \xi_i^+}{\partial x_i} + \dot{y}_i \frac{\partial \xi_i^+}{\partial y_i} + \frac{\partial \xi_i^+}{\partial t} - \dot{z}_i \right] \delta (\xi_i^+ - z_i). \quad (14)$$

The time derivative of $\xi_i^+(t) = z_i^+(t) + \xi(x_i(t), y_i(t), t)$ depends on particle positions $x_i$ and $y_i$ which are themselves a function of time, as well as the explicit surface time dependence. Each of the terms have a physical interpretation, the Dirac delta function is only non zero when molecules are crossing the surface, counting at the point of crossing 1) the $z$ velocity components of the molecule $\dot{z}_i$, 2) the surface curvature $\dot{x}_i \frac{\partial \xi^+}{\partial x_i}$ and $\dot{y}_i \frac{\partial \xi^+}{\partial y_i}$ times the $x$ and $y$ particle’s velocity at the location of a surface crossing and 3) the crossings due to surface time evolution itself $\frac{\partial \xi^+}{\partial t}$.

We evaluate Eq. (12) using the derivatives as shown in Eq. (13) and Eq. (14) to get the time evolution of density in a molecular control volume between intrinsic surfaces,

$$\frac{d}{dt} \sum_{i=1}^{N} m_i \dot{\vartheta}_i = - \sum_{i=1}^{N} m_i \left[ \dot{x}_i \left[ \delta (x_i^+ - x_i) - \delta (x_i^- - x_i) \right] \Lambda_y \tilde{A}_z 
+ \dot{y}_i \left[ \delta (y_i^+ - y_i) - \delta (y_i^- - y_i) \right] \Lambda_x \tilde{A}_z 
+ \left[ \dot{z}_i - \frac{d\xi_i^+}{dt} \delta (\xi_i^+ - z_i) \right] - \left( \dot{z}_i - \frac{d\xi_i^+}{dt} \right) \delta (\xi_i^- - z_i) \right] \Lambda_x \tilde{A}_z \right]. \quad (15)$$

To simplify this expression, we introduce notation for the Dirac delta terms analogous to the continuum surface element $dS$ used in the integral,

$$dS_{xi}^\pm \equiv \delta (x_i^\pm - x_i) S_{xi}; \quad dS_{yi}^\pm \equiv \delta (y_i^\pm - y_i) S_{yi}; \quad dS_{zi}^\pm \equiv \delta (\xi_i^\pm - z_i) S_{zi}. \quad (16)$$

with $S_{\alpha i}$ is the product of boxcar functions in the other two directions,

$$S_{xi} \equiv \Lambda_y \tilde{A}_z; \quad S_{yi} \equiv \Lambda \tilde{A}_z; \quad S_{zi} \equiv \Lambda \Lambda_y; \quad (17)$$

which can be seen to define an area in space between the surfaces, e.g. $S_{xi}$ is only one if a particle is between $y^+$ and $y^-$ in $y$ and $\xi_i^+$ and $\xi_i^-$. Using the definitions of Eq. (16) in the
time evolution Eq. (15),

\[ \frac{d}{dt} \sum_{i=1}^{N} m_i \dot{\vartheta}_i = \sum_{i=1}^{N} m_i \left( \frac{\partial \xi_i^+}{\partial t} \right) dS_{zi}^+ - \frac{\partial \xi_i^-}{\partial t} dS_{zi}^- \\
- \sum_{\beta \in \{x,y,z\}} \dot{\beta}_i \left[ dS_{\beta i}^+ - \frac{\partial \xi_i^+}{\partial \beta_i} dS_{zi}^+ + \frac{\partial \xi_i^-}{\partial \beta_i} dS_{zi}^- \right] \right) = - \oint_{S} \rho \mathbf{u} \cdot dS, \quad (18) \]

where \( \frac{\partial \xi_i^+/\partial z_i = 0 \) is used to write concisely and the last equality, linking continuum and molecuale expressions, follows from Eq. (11). The sum over all surfaces in Eq. (18), allows the continuum and molecular expressions to be compared surface by surface, for \( \beta = x \) and taking just the top + surface this is,

\[ \int_{S_x^+} \rho \mathbf{u} \cdot dS_x^+ = \sum_{i=1}^{N} m_i \dot{x}_i dS_{x_i}^+, \quad (19) \]

and for \( \beta = z \), against considering just the top surface,

\[ \int_{S_z^+} \rho \mathbf{u} \cdot dS_z^+ = \sum_{i=1}^{N} m_i \left[ -\dot{x}_i \frac{\partial \xi_i^+}{\partial x_i} - \dot{y}_i \frac{\partial \xi_i^+}{\partial y_i} + \dot{z}_i - \frac{\partial \xi_i^+}{\partial t} \right] dS_{zi}^+, \quad (20) \]

So, the \( z^+ \) surface flux of mass is made up of direct fluxes, surface curvature and surface evolution components which must all be evaluated. The shorthand notation for flux over each surface is introduced,

\[ \frac{d}{dt} \sum_{i=1}^{N} m_i \dot{\vartheta}_i = - \sum_{i=1}^{N} m_i \left[ \dot{\beta}_i^+ \cdot dS_i^+ - \dot{\beta}_i^- \cdot dS_i^- \right] = - \sum_{i=1}^{N} m_i \dot{\beta}_i \cdot dS_i, \quad (21) \]

where

\[ \dot{\beta}_i^+ = \begin{bmatrix} \dot{x}_i \\ \dot{y}_i \\ \dot{z}_i - \frac{\partial \xi_i^+}{\partial x_i} - \frac{\partial \xi_i^+}{\partial y_i} - \frac{\partial \xi_i^+}{\partial t} \end{bmatrix} \quad \text{and} \quad dS_i^+ = \begin{bmatrix} dS_{x_i}^+ \\ dS_{y_i}^+ \\ dS_{z_i}^+ \end{bmatrix} \quad (22) \]

and the \( \pm \) superscript is omitted in Eq. (21) when expressing the difference \( dS_i = dS_i^+ - dS_i^- \) or more generally could be a shorthand to denote flux over an arbitary surface.
D. Momentum

The same integration process used for density can be applied to get the momentum in a
control volume, written in integrated form as,

$$\int_V \rho(\mathbf{r},t) u(\mathbf{r},t) dV = \sum_{i=1}^{N} m_i \dot{r}_i \dot{\vartheta}_i. \quad (23)$$

The equivalent time evolution of momentum for a continuum control volume gives the surface
flux of momentum (convection) $\rho uu$ and pressure tensor denoted by $\Pi$,

$$\frac{d}{dt} \int_V \rho u dV = - \oint_s [\rho uu + \Pi] \cdot dS = - \oint_s [\rho uu + \Pi^k + \Pi^c] \cdot dS, \quad (24)$$

where the total pressure tensor can be split into kinetic pressure, $\Pi^k$, and configurational
pressure, $\Pi^c$, contributions. It is perhaps more natural to talk about the compressed state
of molecules in the configurational contributions as stress, but as this is simply the negative
of pressure, the term pressure is used here for both kinetic and configurational contributions.
We could also explicitly include the term for surface tension $\oint \gamma d\ell$ in these continuum
equations, but it is more convenient to include all terms in the pressure tensor and calculate the
surface tension from the stress tensors using the Kirkwood and Buff approach.

As with the mass equation, we evaluate the time evolution of Eq. (23) to obtain expressions
for the molecular surface flux and stresses, obtained from manipulation of the Dirac
delta functions,

$$\frac{d}{dt} \int_V \rho u dV = \frac{d}{dt} \sum_{i=1}^{N} m_i \dot{r}_i \dot{\vartheta}_i = \sum_{i=1}^{N} m_i \dot{r}_i \frac{d\vartheta_i}{dt} + \sum_{i=1}^{N} m_i \dot{r}_i \dot{\vartheta}_i. \quad (25)$$

The Kinetic term proceeds along the same lines as density, above, which gives,

$$\sum_{i=1}^{N} m_i \dot{r}_i \frac{d\vartheta_i}{dt} = - \sum_{i=1}^{N} m_i \dot{r}_i \dot{\vartheta}_i \cdot dS_i = - \oint_S [\rho uu + \Pi^k] \cdot dS. \quad (26)$$

We consider the Configurational term next,

$$\sum_{i=1}^{N} m_i \dot{r}_i \dot{\vartheta}_i = \sum_{i=1}^{N} F_i \vartheta_i = \frac{1}{2} \sum_{i,j}^{N} f_{ij} [\vartheta_i - \vartheta_j], \quad (27)$$
where the \( i,j \) notation is shorthand for a double sum over all \( i \) and all \( j \) indices. Equation (27) is the integral of the differences of two Dirac delta functionals, the infamous IK operator. Using the fundamental theorem of the calculus, this can be expressed in a much more convenient form,

\[
\vartheta_i - \vartheta_j = \int_V \left[ \delta (\mathbf{r} - \mathbf{r}_i) - \delta (\mathbf{r} - \mathbf{r}_j) \right] dV = \int_V \int_0^1 \frac{\partial}{\partial \lambda} \delta (\mathbf{r} - \mathbf{r}_\lambda) d\lambda dV = \int_0^1 \delta (\mathbf{r} - \mathbf{r}_\lambda) dV d\lambda \]

\[
= \int_0^1 \frac{\partial}{\partial \lambda} d\lambda = \int_0^1 \frac{\partial \mathbf{r}_\lambda}{\partial \lambda} \cdot \frac{\partial \mathbf{r}_\lambda}{\partial \lambda} d\lambda = \int_0^1 \mathbf{r}_{ij} \cdot \frac{\partial \mathbf{r}_\lambda}{\partial \lambda} d\lambda, \tag{28}
\]

where \( \mathbf{r}_\lambda = \mathbf{r}_i + \lambda \mathbf{r}_{ij} \), representing an integration along the line of interaction between \( \mathbf{r}_i \) and \( \mathbf{r}_j \). The intrinsic surface \( \xi(x, y, t) \) is independent of the dummy integral along the path between molecules, \( \lambda \), so we can change the order of integration allowing a volume integral of the Delta functions which follows the same process as Eq. (7) to Eq. (9) with molecular position replaced by point on line of interaction \( \mathbf{r}_\lambda \to \mathbf{r}_i \), giving,

\[
\vartheta_\lambda \equiv \left[ H (x^+ - x_\lambda) - H (x^- - x_\lambda) \right] \times \left[ H (y^+ - y_\lambda) - H (y^- - y_\lambda) \right] \times \left[ H (\xi^+ (x_\lambda, y_\lambda, t) - z_\lambda) - H (\xi^- (x_\lambda, y_\lambda, t) - z_\lambda) \right] = \Lambda_{x\lambda} \Lambda_{y\lambda} \tilde{\Lambda}_{z\lambda}. \tag{29}
\]

Here the control volume function selects molecular interactions through Heavisides to obtain the length of line in a control volume, by integrating along that line where any point is one when \( \mathbf{r}_\lambda \) is inside and zero otherwise. We denote the intrinsic interface function in terms of \( \mathbf{r}_\lambda \) as \( \xi_\lambda^{\pm} \equiv z^\pm + \xi(x_\lambda, y_\lambda, t) \). The functions \( \Lambda_{x\lambda}, \Lambda_{y\lambda} \) and \( \tilde{\Lambda}_{z\lambda} \) are analogous to the previous definitions with \( \mathbf{r}_i \) replaced by \( \mathbf{r}_\lambda \).

The expression \( \mathbf{r}_{ij} \cdot \partial \vartheta_\lambda / \partial \mathbf{r}_\lambda \) in Eq. (28) is a sum over all three directions, considering the \( z \) component first,

\[
z_{ij} \frac{\partial \vartheta_\lambda}{\partial z_\lambda} = -z_{ij} \left[ dS_{z_\lambda}^+ - dS_{z_\lambda}^- \right], \tag{30}
\]

where the \( dS_{\alpha\lambda}^\pm \) term is defined following the convention used for Eq. (16) to be,

\[
dS_{x_\lambda}^\pm \equiv \delta (x^\pm - x_\lambda) S_{x_\lambda}; \quad dS_{y_\lambda}^\pm \equiv \delta (y^\pm - y_\lambda) S_{y_\lambda}; \quad dS_{z_\lambda}^\pm \equiv \delta (\xi^\pm - z_\lambda) S_{z_\lambda}, \tag{31}
\]
with \( S_{x\lambda} = \Lambda_{y\lambda} \tilde{A}_{z\lambda}, \) \( S_{y\lambda} = \Lambda_{x\lambda} \tilde{A}_{z\lambda} \) and \( S_{z\lambda} = \Lambda_{x\lambda} \Lambda_{y\lambda}. \) The derivatives of the \( x \) and \( y \) components are slightly more complicated due to the \( x_{\lambda} \) and \( y_{\lambda} \) dependency in \( \zeta \), so for \( x \) we have,

\[
x_{ij} \frac{\partial \vartheta_{\lambda}}{\partial x_{\lambda}} = -x_{ij} \left[ dS_{x\lambda}^+ - dS_{x\lambda}^- - \frac{\partial \xi_{\lambda}^+}{\partial x_{\lambda}} dS_{z\lambda}^+ + \frac{\partial \xi_{\lambda}^-}{\partial x_{\lambda}} dS_{z\lambda}^- \right].
\]

(32)

A similar result can also be obtained for \( y \). The derivative in \( x \) gives the same surface curvature terms \( \partial \xi / \partial x \) seen previously for molecular flux but in this case due to intermolecular interactions. Combining the six surfaces of the control volume, the configurational term of Eq. (27) can be written as,

\[
\frac{1}{2} \sum_{i,j}^N f_{ij} [\vartheta_i - \vartheta_j] = \frac{1}{2} \sum_{i,j}^N f_{ij} \int_0^1 \beta_{ij} \cdot dS_{\lambda} d\lambda = \oint_S \Pi^c \cdot dS,
\]

(33)

which is the total configurational pressure over all surfaces. Defining an analogous \( \beta_{ij} \) and \( dS_{\lambda} \) to the kinetic flux case,

\[
\beta_{ij}^\pm = \begin{bmatrix} x_{ij} \\ y_{ij} \\ z_{ij} - \frac{\partial \xi^+_{\lambda}}{\partial x_{\lambda}} - \frac{\partial \xi^-_{\lambda}}{\partial y_{\lambda}} \end{bmatrix} \quad \text{and} \quad dS_{i}^\pm = \begin{bmatrix} dS_{x\lambda}^\pm \\ dS_{y\lambda}^\pm \\ dS_{z\lambda}^\pm \end{bmatrix}
\]

(34)

We can therefore take any of the surfaces, for example the configuration pressure on the \( x \) surface is,

\[
\int_{S^+_x} \Pi^c \cdot dS_x^+ = \frac{1}{2} \sum_{i,j}^N f_{ij} \int_0^1 x_{ij} dS_{z\lambda}^+ d\lambda,
\]

(35)

and the \( z \) surface is,

\[
\int_{S^+_z} \Pi^c \cdot dS_z^+ = \frac{1}{2} \sum_{i,j}^N f_{ij} \int_0^1 \left[ -x_{ij} \frac{\partial \xi^+_{\lambda}}{\partial x_{\lambda}} - y_{ij} \frac{\partial \xi^+_{\lambda}}{\partial y_{\lambda}} + z_{ij} \right] dS_{z\lambda}^+ d\lambda.
\]

(36)

We are now in a position to write Eq. (25), the time derivative of the control volume in terms of the kinetic Eq. (26) and configurational Eq. (33) parts,

\[
\frac{d}{dt} \sum_{i=1}^N m_i \dot{r}_i \vartheta_i = - \sum_{i=1}^N m_i \dot{r}_i \beta_i \cdot dS_i - \frac{1}{2} \sum_{i,j}^N f_{ij} \int_0^1 \beta_{ij} \cdot dS_{\lambda} d\lambda
\]

\[
= - \oint_S [\rho uu + \Pi] \cdot dS
\]

(37)
which is the total stress over every surface of the control volume. Each of the faces of the control volume, top + or bottom − can be seen to define three of the components of stress tensor. Considering the top $x$ surface, with $dS^+_x$ and $S^+_{x\lambda}$ and assuming an average pressure on the surface area $\int_{S^+_x} \left[ \rho uu + \Pi \right] \cdot dS^+_x \approx \Delta S_x \left[ \rho uu_x + \Pi_x \right]$, with $\Pi_x = [\Pi_{xx}, \Pi_{xy}, \Pi_{xz}]^T$ so the pressure on the $x$ surface can be written as,

$$\rho uu_x + \Pi_x = \frac{1}{\Delta S_x} \sum_{i=1}^{N} m_i \hat{r}_i \hat{x}_i dS^+_x + \frac{1}{2 \Delta S_x} \sum_{i,j} f_{ij} \int_0^1 x_{ij} dS^+_{x\lambda} d\lambda,$$

(38)

which is identical to the flat surface obtained in previous work$^{34}$ and is consistent with a localised method of planes stress (see section III). The pressure on the $z$ surface is,

$$\rho uu_z + \Pi_z = \frac{1}{\Delta S_z} \sum_{i=1}^{N} m_i \hat{r}_i \hat{z}_i dS^+_z + \frac{1}{2 \Delta S_z} \sum_{i,j} f_{ij} \int_0^1 z_{ij} dS^+_{z\lambda} d\lambda,$$

(39)

where the equation describes the components of pressure on a control volume surface which is following the intrinsic interface, including terms due to curvature and surface evolution. Three connected faces form a tetrahedron which is consistent with Cauchy’s original definition of the stress tensor$^{24}$, giving,

$$[\Pi_x, \Pi_y, \Pi_z] = \left[ \begin{array}{ccc} \Pi_{xx} & \Pi_{yx} & \Pi_{zx} \\ \Pi_{xy} & \Pi_{yy} & \Pi_{zy} \\ \Pi_{xz} & \Pi_{yz} & \Pi_{zz} \end{array} \right]_{x \text{ surf} \ y \text{ surf} \ z \text{ surf}}$$

(40)

Section III will discuss how to implement this equation.

E. Energy

For completeness, the expressions for energy are stated here as they require no additional mathematics. The time derivative of Eq. (6) for energy results in a similar kinetic and
configurational term to the momentum equation,

\[
\frac{d}{dt} \sum_{i=1}^{N} e_i \dot{\varphi}_i = \sum_{i=1}^{N} \rho \frac{d\varphi_i}{dt} + \sum_{i=1}^{N} \frac{d}{dt} \varphi_i = \sum_{i=1}^{N} e_i \dot{\varphi}_i \cdot \frac{d}{dr} + \frac{1}{2} \sum_{i,j} \dot{\varphi}_i \cdot f_{ij} [\varphi_i - \dot{\varphi}_j]
\]

\[
= \sum_{i=1}^{N} e_i \dot{\beta}_i \cdot dS_i + \frac{1}{2} \sum_{i,j} \dot{\beta}_i \cdot f_{ij} \int_{0}^{1} \beta_{ij} \cdot dS \lambda d\lambda
\]

\[
= \oint_S [\rho \mathbf{u} \mathbf{E} + \mathbf{Π} \cdot \mathbf{u} + \mathbf{q}] \cdot dS.
\]

(41)

Where any surface of the control volume gives the energy equation in that coordinate direction. Isolating and obtaining contributions for heat flux \(q\) could proceed as outlined in previous work\(^{36}\), although careful consideration of the evolving surface and its contribution to heat flux may be required. This is left for future work.

III. IMPLEMENTATION

In this section, the mathematical equations are manipulated to obtain expressions which can be coded in an MD simulation. The similarity between the kinetic and configurational terms will be highlighted, as well as the similarity in operation required for both intrinsic and flat surfaces of the volume. It will be shown that the problem of obtaining the surface stress is reduced to obtaining the intersection of a line and a surface, a common problem in computer graphics and ray tracing. From equation (39), it is apparent the form of both Kinetic Curvature and Configurational Curvature terms are similar, with the difference between them the surface time evolution \(\partial \xi^+ / \partial t\), which we consider first.

A. Time Evolving Interface

This term, \(\partial \xi^+ / \partial t\), describes the change in mass, momentum or energy in a control volume as new molecules are absorbed or left behind when the intrinsic surface moves. This can be thought of as a 2D function sweeping through space and crossing the position of the particles. To evaluate this term, we first consider the time integration process applied in an MD simulation, i) particle positions at time \(t_1\) are used for the force calculation, ii)
the surface $\xi(t_1) \equiv \xi(x_i(t), y_i(t), t_1)$ is fixed at time $t_1$ and the evolution of the particles from $\mathbf{r}_i(t_1)$ to $\mathbf{r}_i(t_2)$ is used to get surface flux and iii) the particles are fixed at $t_2$ while the surface is evolved from $\xi(t_1)$ to $\xi(t_2)$. Considering the top surface in $z$, this proceeds as follows,

$$\int_{t_1}^{t_2} \frac{\partial \xi_i^+}{\partial t} dS^+_{zi} d\tau = \int_{t_1}^{t_2} \frac{\partial \xi_i^+}{\partial t} \delta \left( \xi_i^+(t) - z_i \right) S_{zi} d\tau = \int_{t_1}^{t_2} \sum_{k=1}^{N_{\text{roots}}} \frac{\partial \xi_i^+}{\partial t} \left| \frac{\partial \xi_i^+(t_k)}{\partial t} \right| S_{zi} d\tau$$

$$= \sum_{k=1}^{N_{\text{roots}}} \text{sgn} \left( \frac{\partial \xi_i^+(t_k)}{\partial t} \right) \left[ H(t_2 - t_k) - H(t_1 - t_k) \right] S_{zi}, \quad (42)$$

using the roots of the Dirac delta function given in the Appendix Eq. (A7). The derivative of the surface can be expressed, using the definition of the partial derivative in time with $\Delta t = t_2 - t_1$,

$$\frac{\partial \xi_i^+}{\partial t} = \lim_{\Delta t \to 0} \frac{\xi_i^+(x_i(t), y_i(t), t + \Delta t) - \xi_i^+(x_i(t), y_i(t), t)}{\Delta t} \approx \frac{\xi_i^+(t_2) - \xi_i^+(t_1)}{t_2 - t_1},$$

so, the expression $\text{sgn}(\partial \xi_i^+/\partial t) = \text{sgn} \left( \xi_i^+(t_2) - \xi_i^+(t_1) \right)$ as $t_2 > t_1$ and this term can be seen to simply determine the crossing direction. Obtaining the multiple potential roots, $t_k$, of a time evolving polynomial crossing position in space is a non-trivial exercise, but the expression $\text{sgn} \left( \xi_i^+(t_2) - \xi_i^+(t_1) \right) \left[ H(t_2 - t_k) - H(t_1 - t_k) \right]$ can be seen to be equivalent to a simple check if points $z_i(t_2)$ is crossed as the surface moves from $\xi_i^+(t_1)$ to $\xi_i^+(t_2)$, which can be achieved by the following functional,

$$\text{sgn} \left( \xi_i^+(t_2) - \xi_i^+(t_1) \right) \left[ H(t_2 - t_k) - H(t_1 - t_k) \right] S_{zi}$$

$$= \left[ H \left( \xi_i^+(t_2) - z_i(t_2) \right) - H \left( \xi_i^+(t_1) - z_i(t_2) \right) \right] S_{zi} \equiv \vartheta_t,$$  

(43)

Note that unlike the previous control volume functionals $\vartheta_i$ and $\vartheta_\lambda$, it is possible for $\vartheta_t$ to be negative.

B. Summary of Equations for Pressure

The new surface pressure equations derived in this work will be compared to the volume average (VA) form of pressure derived in previous work, obtained by integrating the Irving and Kirkwood expressions over a volume following an intrinsic surface,

$$\int_V \left[ \rho \mathbf{uu} + \mathbf{P} \right] dV = \sum_{i=1}^{N} m_i \mathbf{\dot{r}}_i \mathbf{\dot{r}}_i \vartheta_i + \frac{1}{2} \sum_{i,j} f_{ij} \mathbf{r}_{ij} \int_0^1 \vartheta_\lambda d\lambda.$$

(44)
A more general derivation of these volume average expression follows from the time evolution of Eq. (25), as shown in appendix C.

The process of deriving equations valid for a molecular simulation, involving the integral over the surface $dS_i$ and $dS_\lambda$ functions, is given in appendix A. The equation for pressure on one of the flat control volume faces, here the $y^+$ surface, is shown in the appendix A to be,

$$
\int_{t_1}^{t_2} \left[ \rho \dot{u}_y + \frac{S_{\text{surf}}}{\Pi} \right] dt = \frac{1}{\Delta S_y} \sum_{i=1}^{N} m_i \dot{r}_i \cdot \frac{r_{i12} \cdot \mathbf{n}_y}{|r_{i12} \cdot \mathbf{n}_y|} \left[ H\left(\frac{y^+ - y_2}{y_{i12}}\right) - H\left(\frac{y_1 - y^+}{y_{i12}}\right) \right] \Lambda_x(t_k) \tilde{\Lambda}_z(t_k)
$$

$$
\frac{S_{\text{surf}}}{\Pi} \sum_{i,j}^{N} f_{ij} \frac{r_{ij} \cdot \mathbf{n}_y}{|r_{ij} \cdot \mathbf{n}_y|} \left[ H\left(\frac{y^+ - y_j}{y_{ij}}\right) - H\left(\frac{y_i - y^+}{y_{ij}}\right) \right] \Lambda_x(\lambda_k) \tilde{\Lambda}_z(\lambda_k).
$$

Equation (45) is written in this form to emphasize the expression is the molecular pressure tensor $m_i \dot{r}_i r_{i12}$ and $f_{ij} r_{ij}$ dotted with the surface normal $\mathbf{n}_y$. The $r_{i12}$ term is the evolution of a molecule $i$ from time $t_1$ to $t_2$. This equation is equivalent to the method of planes (MOP) form of stress, as demonstrated in the Appendix A, but localized to a control volume surface by $\Lambda_x \tilde{\Lambda}_z$. Recall that the Lambda functions are boxcar or Bracewell functions checking if a point is between two limits using two Heaviside functions $\Lambda_x(a) = H(x^+ - a) - H(x^- - a)$ and $\tilde{\Lambda}_z(a) = H(\xi^+ - a) - H(\xi^- - a)$. This control volume surfaces and associated normals are shown in Figure 1a) for an intermolecular interaction crossing the $y^+$ surface by green crosses. An example of the actual interactions from an MD simulation at a single timestep are also shown in Figure 1b) with blue and green crosses denoting the $x$ and $y$ surfaces, respectively. The expressions for the roots $t_k$ and $\lambda_k$ can obtained analytically for this flat surface case, with expression given in the Appendix A. The expression for stress on the
intrinsic surface \( z^+ \) is,

\[
\int_{t_1}^{t_2} \rho \mathbf{u}_z + \sum_{k}^{\text{surf}} \int_{z}^{\Pi} \Pi_z \, dt = \frac{1}{\Delta S_z} \sum_{i=1}^{N} m_i \mathbf{r}_{12} \cdot \hat{n}_z \sum_{k=1}^{N_{\text{roots}}} [H (1 - t_k) - H (-t_k)] \Lambda_x(t_k) \Lambda_y(t_k) \\
+ \frac{1}{\Delta S_z} \sum_{i=1}^{N} m_i \mathbf{v}_i \hat{n}_z \\
\sum_{i,j}^{\text{surf}} \sum_{k=1}^{N_{\text{roots}}} \left[ H (1 - \lambda_k) - H (-\lambda_k) \right] \Lambda_x(\lambda_k) \Lambda_y(\lambda_k),
\]

(46)

where, as in the flat surface, the expression for \( m_i \mathbf{v}_i \mathbf{r}_{12} \) and \( f_{ij} \mathbf{r}_{ij} \) are dotted with the surface normal. In this case this is the normal to the intrinsic interface \( \hat{n}_z(x,y) = \nabla \alpha(\xi - z) \). The normal \( \hat{n}_z \) is shown on the schematic of Figure 1a with the MD simulation showing the many interactions as red crosses on Figure 1b. The process of obtaining the intersection of the interaction line and the intrinsic surface is the main requirement to evaluate the expression of Eq. (46). The process is identical for time roots \( t_k \) and intermolecular \( \lambda_k \). Given the multiple possible crossings, a closed form expression is not possible, but any root finding process could be used. An approach based on approximating the surface as a set of bilinear patches is used here which allows an efficient root calculation from the ray-tracing literature\(^{28}\) to be applied. The computations are then fast enough to be part of a molecular simulation. Once the roots have been found, they can be put into the \( \Lambda \) functions to check if they are within the limits of a given control volume surface. In practice, integer division can be used to speed this process up, assigning the crossing to a cell, as discussed in the next section. Finally, the surface evolution term is obtain by checking if a particles positions \( \mathbf{r}_i(t_2) \) is in a given volume before and after the surface has evolved.
The result is a concise expression for the momentum flux over all surfaces, written as,

\[
\begin{align*}
\text{Accumulation} & \quad \sum_{i=1}^{N} [m_i \dot{r}_i(t_2) \partial_i(t_2) - \dot{r}_i(t_1) \partial_i(t_1)] = - \sum_{\alpha=\{x,y,z\}} \frac{1}{\Delta S_\alpha} \left[ \sum_{i=1}^{N} m_i \dot{r}_i \left( r_{i12} \left[ dS^+_{\alpha k} - dS^-_{\alpha k} \right] + \beta_1 \right) \right. \\
\text{Advection} & \quad + \left. \frac{\Delta t}{2} \sum_{i,j} f_{ij} r_{\alpha ij} \left[ dS^+_{\alpha \lambda k} - dS^-_{\alpha \lambda k} \right] \right], \\
\text{Forcing} & \quad \end{align*}
\]

(47)

where both sides are integrated over time with the force term shown integrated with the midpoint rule for simplicity \( \int dt \approx \Delta t \), consistent with the Verlet integration scheme used to propagate molecular positions and velocities. The function to get crossings on a surface is denoted for time \( s = t \) and space \( s = \lambda \) with,

\[
dS^{\pm}_{\alpha s k} = \frac{\tilde{n}_\alpha}{|\mathbf{r}_{12} \cdot \tilde{n}_\alpha|} \sum_{k=1}^{N_{\text{roots}}} \left[ H(1 - s_k) - H(-s_k) \right] \Lambda_\beta(s_k) \Lambda_\gamma(s_k),
\]

(48)

where the \( \beta \) and \( \gamma \) are the orthogonal directions to \( \alpha \), so if \( \alpha = x \), \( \beta = y \) and \( \gamma = z \), and the normal \( \tilde{n}_\alpha \) can be for a flat or intrinsic surface with roots \( s_k \) obtained for that surface. This same expression is valid for both kinetic and configurational terms on both curved and flat surfaces, with the simple interpretation of checking if a crossing is on the surface of a control volume.

The six surface pressures summed for any arbitrary control volume consist of terms due to intermolecular forces, labelled \textit{Forcing} in Eq. (47) as well as kinetic molecular crossings due to both molecular motion and surface movement labelled \textit{Advection}. These are exactly equal, to machine precision, to the change in momentum in the volume, called \textit{Accumulation}, demonstrated in section \( \Box \).

We now discuss the process of how to obtain these surface crossing terms efficiently as part of an MD simulation.
FIG. 1. a) A schematic showing surface normals $\mathbf{n}_y$ and $\mathbf{\tilde{n}}_z$ on an intermolecular interaction line $r_{ij}$ contributing to pressure on both the flat surfaces via Eq. (45) and the intrinsic surface with Eq. (46) and b) a random snapshot of an arbitrary control volume inside the liquid phase of an MD simulation, showing all the contributions used in the calculation of configurational pressure; with blue crosses on the $x$ surface, green crosses on the $y$ and red crosses for a bilinear patch of the intrinsic surface with the top $\xi^+$ surface coloured in light blue and the bottom $\xi^-$ coloured light red.

IV. METHOD

We use molecular dynamics (MD) simulation to model the liquid vapour interface, with a Lennard Jones potential,

$$\phi(r_{ij}) = \epsilon \left[ \left( \frac{\sigma}{r_{ij}} \right)^{12} - \left( \frac{\sigma}{r_{ij}} \right)^6 \right] - \phi(r_c); \quad r_{ij} < r_c,$$

(49)

with cutoff $r_c = 2.5$, which is shorter than required to give the correct surface tension but chosen to allow more efficient simulations. The force on particle $i$ is obtained from the sum of the gradient of this potential due to interaction with all $N$ other particles $F_i = \sum_{j \neq i}^{N} f_{ij} = \sum_{j \neq i}^{N} \nabla \phi(r_{ij})$ The simulations are run using the Flowmol MD code which has been validated extensively in previous work The time integration is achieved by velocity Verlet with a timestep $\Delta t = 0.005$.

$$r_i(t + \Delta t) = r_i(t) + \Delta t v_i(t + \Delta t/2)$$

$$v_i(t + \Delta t/2) = v_i(t - \Delta t/2) + \Delta t F_i(t)$$

(50)

20
The system is initialised as a liquid-vapour coexistence by creating an FCC lattice and removing molecules until the desired density is obtained. The system is periodic in all directions, with \( L_x = L_y = 12.7 \) while the surface normal direction \( L_z = 47.62 \). The middle 40\% of the domain is designated to be initialised as liquid with a density of \( \rho_l = 0.79 \) and the remaining domain is set to a gas density of \( \rho_g = 0.0002 \). This results in a system with \( N = 2635 \) molecules. The system is then run in the NVT ensemble, controlled by a Nosé Hoover thermostat at a temperature setpoint of \( T_s = 0.7 \) for 100,000 timesteps to equilibrate. The run is then restarted as an NVE ensemble and run for sufficient time to collect well resolved statistics.

In order to obtain the intrinsic interface, a cluster analysis is first used to identify the connected molecules defined to be a liquid cluster, shown in Fig 2 (a). In this simulation, the liquid region tends to be located in the centre of the domain and the cluster analysis identifies all connected molecules which are within the Stillinger cutoff length \( r_d = 1.5 \) from each other. A linked list is built of all molecules, before any which have fewer than three neighbours are discarded as not part of the cluster, giving \( N_\ell \) liquid particles. The cluster is then used to fit the intrinsic surface, as detailed in section IV A, with the fitting performed on the surface on the right-hand side of the cluster.

A. Intrinsic Surface

Up until this section, no assumption has been made about the functional form of surface \( \xi(x, y, t) \). To give a general form, the intrinsic surface method (ISM) approximates the liquid-vapour interface using a Fourier series representation,

\[
\xi(x, y, t) = \sum_{k < k_u} \hat{\xi}_k(t) \exp\left(2\pi i k \cdot r_\parallel\right),
\]

where \( \hat{\xi}_k(t) \) are the amplitudes associated with each wavevector, a function of time as they are refitted to the surface every time the molecules in the system evolve, with wave vector \( k = (\mu/Lx, \nu/Ly) \), parallel surface components \( r_\parallel = (x, y) \) and the number of wavelengths calculated from the system size \( k_u = \text{nint}(\sqrt{LxLy}/\lambda_u) \) based on \( \lambda_u \) the minimum wavelength of the surface, set to intermolecular spacing for a Lennard Jones fluid in this work, so \( \lambda_u \approx \sigma \).
as used in previous work\cite{23}. The surface fitting functions can be made simpler by expressing
in terms of just the real components,

\[ \xi(x, y, t) = \sum_{\mu=-k_u}^{k_u} \sum_{\nu=-k_u}^{k_u} a_{\mu\nu}(t) f_{\mu}(x) f_{\nu}(y), \quad (52) \]

where \( f_\mu(x) = \cos(k_x x) \), \( f_{-\mu}(x) = \sin(k_x x) \) and \( a_{\mu\nu} \) is the matrix of surface wavenumbers.

To fit the intrinsic surface, the square of the difference between surface molecules, called
pivots, and the intrinsic surface function \( \xi \) is defined to be,

\[ W = \frac{1}{2} \sum_{p=1}^{N_p} [z_p - \xi(x_p, y_p)]^2 + \psi \tilde{A}, \quad (53) \]

which is the function to be minimised as part of a least squares fit. An extra constraint \( \psi \)
is included to prevent overfitting by ensure the intrinsic area \( \tilde{A} \) does not become too large,
where \( \psi = 1 \times 10^{-8} \) in line with values in the literature\cite{24}. This is a linear algebra problem,
as discussed in Appendix \[B\]. The fitting of the interface is done in stages, with the first
fit to a subset of molecules of the cluster obtained by defining a grid, usually \( 3 \times 3 \) in \( x \)
and \( y \), although a finer grid can speed up fitting\cite{25}, and obtaining the outermost molecule
in each gridcell. The fitting to these \( N_p = 9 \) molecules is shown schematically in Fig 2b). The resulting zeroth interface guess, \( \xi^0 \), is then interrogated at the location of each
molecules with the difference sorted in order of \( \xi^0(x_i, y_i) - z_i \forall N_\ell \). The closest molecules are
then included and used to refit the surface, including the original 9 and incorporating an
additional one so \( N_p = 10 \). This can be done in batches to improve efficiency with a number
of the closest molecules added at the same time, giving identical results in most cases to the
process applied molecules by molecule. With the new surface, denoted by \( \xi^1 \), the process
is repeated and continues until a desired density of \( \rho_s = N_0/A \) is reached. The density of
\( \rho_s = 0.7 \) is shown to minimise molecular turnover in Chacón et al.\cite{8} so is used in this work.
When the number of pivots is \( N_p = \rho_s A \) is reached the process is stopped, with an interface
as shown in Fig 2c).

Once an intrinsic surface has been obtained, layers can be defined by using an offset \( \Delta z \)
to define binning regions. These can be used to bin quantities such as position, velocity or
fraction of intermolecular interaction to obtain density, momentum or pressure respectively
FIG. 2. A schematic outlining the process of defining the surface fluxes starting by a) getting the liquid cluster, b) fitting the initial surface to a number of outermost particles, with 3 show here of the $3 \times 3$ taken c) refining by including all closest molecules until surface density is a target value $\rho_s = n_0/A$, d) defining a set of layers with uniform offset $\Delta z$ from the surface e) Defining a grid by dividing the domain into a uniform grid of $\Delta x$ and $\Delta y$ cells and f) approximating the intrinsic interface as bilinear patches for each cell, with the calculation of crossings shown for flat surface as green crosses, the shaded region showing the cells between these flat surface crossings that are checked for intrinsic surface crossings with the two possible crossings shown by red crosses.

at fixed distance from the surface as shown in Fig 2d). This work extends this approach by using localisation to a grid of cells in the surface tangential directions, as shown in Fig 2e). This allows properties as a function of position on the interface to be evaluated and could be applied to explore surface transport, local evaporation and Maragnoi effects, among other things. In order to evaluate local quantities efficiently, the grid is replaced by a bilinear approximation shown in Fig 2f), which simplifies the calculation of the point of intersection shown by red crosses and the process of mapping to get binned contributions. Section IV C discussed this implementation in more detail.
B. Mapping of Points and Intersections

The control volume functions $\vartheta_i$ and $\vartheta_\lambda$ mathematically determine if a given point $\mathbf{r}_\alpha$ is located in any given cell as it moves with the intrinsic interface. In practice, a contiguous grid of cells is used, so instead of testing every point with every cell, the point locations are assigned to an appropriate cell using integer division. In pseudocode for this is just,

```plaintext
function get_cell (r)
    cell (1) = ceiling (r (1)/ cellside (1)) # Function $\Lambda_x$
    cell (2) = ceiling (r (2)/ cellside (2)) # Function $\Lambda_y$
    cell (3) = ceiling (r (3)/ cellside (3)) # Function $\Lambda_z$
    return cell
end function get_cell
```

where $r (\alpha)$ is $\mathbf{r}_\alpha$, the point in three dimensions and rounded division by cellside (\alpha) returns an integer which is the index of the cell where the point is located. Quantities can then be added to a cell using this index, including mass, velocity, energy when the point is a molecular location or stress and work when the point is on the intermolecular line of interaction. This can be seen to be a computational implementation of the boxcar functions $\Lambda_x$, $\Lambda_y$ and $\Lambda_z$ defined in Eq. (9), with no tilde in the $z$ direction as this is for a flat surface. The intrinsic surface form $\tilde{\Lambda}_z$ can then be implemented by a simple mapping of the point followed by rounding to obtain the cell index.

```plaintext
function get_intrinsic_cell (r)
    cell (1) = ceiling (r (1)/ cellside (1)) # Function $\Lambda_x$
    cell (2) = ceiling (r (2)/ cellside (2)) # Function $\Lambda_y$
    # Function $\tilde{\Lambda}_z$
    cell (3) = ceiling ((r(3) - xi(r(1),r (2)))/ cellside (3))
    return cell
end function get_intrinsic_cell
```

where $\xi$ is $\xi$ at each timestep, a function which returns the $z$ position of the intrinsic surface given inputs of position $x$ and $y$. These can be $x_i$, $y_i$, $x_\lambda$, $y_\lambda$, or $x_k$, $y_k$ shown in Figure 3.
where this mapping is applied to molecular positions, intermolecular interaction modelled in a piecewise manner or surface crossings, respectively. This has the convenient property that the intrinsic surfaces mapping can be switched on in a code by simply swapping a function pointer from the get_cell function to the get_intrinsic_cell function. For large domains in the surface tangential directions, the time taken by function $x_i$ to evaluate surface position can be prohibitive as it requires a sum over the two-dimensional Fourier surface, a calculation of order $O(4k_u^2)$ with $k_u$ scaling as $L$ (assuming $L_x = L_y = L$) so each surface evaluation is $O(L^2)$. The position of the surface at the location of every molecule is pre-calculated for efficiency to get density, momentum, temperature, potential energy and the kinetic part of the pressure tensor. Interactions between molecules must be obtained on the fly by discretising the line of interaction between two molecules and using the surface value at each segment of the line, as shown in Fig 3 by the dotted interaction lines. Some optimisations can be introduced, for example a quick estimate of the number of bins crossed between two points is used to minimise the number of segments in the line and the intermolecular stress and work calculation are performed at the same time. However, this still requires a larger number of evaluations of the intrinsic surface as each interaction crosses $O(20)$ bins, for an average of $O(100)$ interactions per particle (with $r_c = 2.5$) required for each of the $N$ particles. This is even worse for the control volume flux calculation, which requires the intersection of the intrinsic interface and the line of interaction, computational prohibitive as the intersection point is required to machine precision if we want to verify exact conservation of momentum. Even an efficient root finding algorithm with good initial estimate requires the surface to be evaluated many times to converge for every one of the $O(20)$ crossed surface by every one of the $O(100)$ interactions. These rough numbers clearly depend on bin resolution, cut off length as well as the density of liquid and vapour states, but even in low resolution cases appear to represent a limiting step computationally. A fast evaluation is therefore of paramount importance for intrinsic interface tracking calculation to be used as part of an MD simulation, achieved here using a bilinear approximation.
C. Bilinear Approximation for Surface Crossings

The surface is replaced by a piecewise continuous bilinear surface, sampling the intrinsic surface at an arbitrary accuracy. This uses a bilinear patch of the form,

\[ \xi^{BL}(x, y) = a_0 + a_x x + a_y y + a_{xy} xy \]  

(54)

where the coefficients are obtained by solving,

\[
\begin{bmatrix}
a_0 \\
a_x \\
a_y \\
a_{xy}
\end{bmatrix} = 
\begin{bmatrix}
1 & x_1 & y_1 & x_1 y_1 \\
1 & x_1 & y_2 & x_1 y_2 \\
1 & x_2 & y_1 & x_2 y_1 \\
1 & x_2 & y_2 & x_2 y_2
\end{bmatrix}^{-1}
\begin{bmatrix}
\xi(x_1, y_1) \\
\xi(x_1, y_2) \\
\xi(x_2, y_1) \\
\xi(x_2, y_2)
\end{bmatrix}
\]  

(55)

for each patch of the intrinsic interface. Once we have the expression for the bilinear surface, this can be used in the mapping for both particle position and lines between molecules. In order to get surface crossings, we use the fast bilinear-patch line intersection algorithm\(^{28}\) which equates the solution of a line of interaction \(r_s = r_1 + s r_{12}\) and the bilinear patch \(\xi^{BL}\) to obtain a quadratic equation. Solving this provides up to two crossings for a given surface, with some care required for zero or complex roots. As the surface is split into piecewise bilinear patches, this also has the advantage that each control volume has its own bilinear surface and the conservation check of Eq. (47) can be performed on each cell. The derivatives can also be obtained analytically \(\partial\xi^{BL}/\partial x = a_x + a_{xy} y\) and \(\partial\xi^{BL}/\partial y = a_y + a_{xy} x\), although these are discontinuous at the interface of contiguous bilinear patches. In practice this does not represent a problem for the surface pressures as the \(\Lambda_\alpha\) functions ensure all quantities are per cell. The advantage of supplementing the full intrinsic surface with the bilinear approximation, instead of directly fitting a bilinear surface, is the calculation of quantities from the full intrinsic surface \(\xi\) are available analytically, if needed, with the bilinear solution used as a starting point. Bicubic or higher order surface approximations could also be used but would not be able to take advantage of the fast interaction calculation\(^{28}\) for line and bilinear patch.

Determining the number of crossings of an intermolecular interaction and the 3D grid following the intrinsic surfaces (see Fig 2\(f\)) turns out to be a non-trivial exercise. This
FIG. 3. A schematic view of the three operations required to obtain density, momentum and pressures including i) binning molecular position, ii) binning lines between molecules approximated as a series of points and iii) obtaining the location of surface crossings. The two Heaviside functions are essentially a binning operation, checking if a point is between $\xi^-$ and $\xi^+$ with $\xi^\pm = z^\pm + \xi$, shown in red, which can be mapped to a simpler function on the right by subtracting the intrinsic surface value at the particle locations $\xi(x_i, y_i, t)$ or point on line $\xi(x_{\lambda}, y_{\lambda}, t)$. This mapping changes the calculation to a simple check if molecules are between flat surfaces $z^-$ and $z^+$ by integer division. Intersections on flat surfaces, denoted by a green cross, can be mapped in the same way to find the surface of the CV they apply, obtained by Eq. (45). Mapping is not possible for crossings of the intrinsic surface, denoted by a red cross before mapping, but a question mark after to emphasise a root finding must be employed.

is because it is not possible to predict a priori how many times the intermolecular lines will cross an arbitrary intrinsic surface split into interconnected $\xi^{BL}$ patches from just the knowledge of the starting and ending points. To solve this, for each interaction we first obtain all the crossings locations on all flat surfaces in $x$ and $y$, made possible as we have the closed form expression from Eq. (45). By ordering all intermediate flat surface ($x$ and $y$) crossings along the line of interaction, including the two particle positions at the end, we can take successive pairs and are guaranteed any intrinsic surface crossings between a pair must be on the same $\xi^{BL}$ patch. This is shown schematically in Fig 2(f) with green crosses denoting a pair of flat plane crossings, the shaded region shows the row of cells in between these two flat
crossings to be checked for intrinsic surface intersections and the red crosses denoting two found intrinsic crossings. As each grid volume must have at least two crossings or contain an end point (molecule position), by stepping in pairs we ensure all surface crossings have been obtained by this process. This has the added advantage that the coefficients of the bilinear surface, $\xi^{BL}$, can be loaded once for each row, the shaded region in Fig 2(f), and used to check all line-plane crossings with the efficient Ramsey et al.\textsuperscript{28} algorithm. The straight line to check is then between the current pair of flat surface crossings and patch $\xi^{BL}$ is shifted in multiples of $\Delta z$ to get the the successive bin surfaces between the $z$ coordinates of the surface crossings. The pseudocode for this process is as follows,

\begin{verbatim}
function surface_fluxes(r1, r2, quantity, fluxes)
    #Get all ordered crossing along line between r1 and r2
    crossings = get_flat_crossings(r1, r2)
    for i=1 to size(crossings)-1
        #Get cell indices for a pair of crossings
        cellA = get_intrinsic_cell(crossings(i))
        cellB = get_intrinsic_cell(crossings(i+1))
        #Check if same cell in both x and y
        #which means separated by rows of cells
        #as highlighted region in Fig 2(f)
        if (cellA(1) == cellB(1) & cellA(2) == cellB(2)
            #patch P is the z coordinate of the
            #4 corners of all cells in row
            P = get_patch(cellA)
            #Loop in z direction between cellA and cellB
            for j=cellA(3) to cellB(3)
                #Get all crossing for line r1 + s*r12 and patch P
                #with j*dz shift to surface of cell j
                rc = line_patch_intersect(r1, r12, P+dz*j)
                #Use crossing location to assign to cell
        
\end{verbatim}
cellS = get_intrinsic_cell(rc)
# Add flux quantity to array for mass, momentum or energy
fluxes(cellS(1),cellS(2),cellS(3)) += quantity
end for
end if
end for
end function update_surface_fluxes

V. RESULTS AND DISCUSSION

In this section, the results for pressure on the moving surface are presented using the new
surface flux equations derived in Sections II, summarised for implementation in III with setup
and methodology in IV. We start with a validation of the intrinsic density calculation and
parameterise the bilinear approximation for varying resolutions. Next, the novel surface flux
forms are shown to be exactly conservative, before being compared to the volume averaged
expressions. The full balance of the surface flux contributions is shown next, highlighting
the importance of surface evolution in the equations of motions and showing the normal
pressure should be constant over the interface. Finally, the calculation of the surface tension
using the surface pressure is discussed.

A study of bilinear resolutions is shown in Figure 4, with the density profile used to
assess the accuracy of the varying surface approximations. This is calculated as described in
section IV B by mapping the molecular positions based on the intrinsic interface and then
binning molecules to the appropriate volumes with width $\Delta z = 0.175$. The density for the
full intrinsic surface case is presented as a yellow line with $\lambda_r = \lambda_u = \sigma$ and compared to the
results of Chacón and Tarazona included as crosses in Fig 4 a). This surface with $\lambda_r = \sigma$
is then sampled to define a piecewise bilinear approximation of the surface. The cases with
5 and 10 bilinear bins per $\sigma$ are indistinguishable from the intrinsic surface and are omitted
in Fig 4 a). The case where each $\sigma$ unit has 2.5 bins, $\Delta r = 0.4$, is shown as a red line,
where despite some decrease in the peaks at zero, the density profile is largely identical. As
FIG. 4. The effect on intrinsic density of the bilinear approximation is shown for different resolutions defined as $\text{Resolution} = \sigma/\Delta r$ which is the number of times the square bilinear patches of side length $\Delta r$ fit in an intermolecular spacing $\sigma$ and for intrinsic surfaces the number of wavelengths $\lambda_r$ per $\sigma$ with $\text{Resolution} = \sigma/\lambda_r$. In $a)$ the full intrinsic solution is shown as a yellow line for the case where $\lambda_r = \lambda_u = \sigma$ compared to yellow crosses showing the solution from Chacón and Tarazona. Bin $\text{Resolution}$ of 10 and 5 are omitted as indistinguishable from the yellow line, red is a $\text{Resolution}$ of 2.5, green is 1.25 and dark blue is 0.63. The intrinsic interface with $\lambda_r = 4$ is shown for comparison in light blue. In $b)$ the $L^2$ Error $= \int_{-3}^{2} \rho_{\Delta r=\sigma}(z) - \rho_{\Delta r}(z)dz$ obtained from the sum over all $z$ values from $-3$ to $2$ for both bilinear (blue) with binsizes $\Delta r = \{1.6, 0.8, 0.4, 0.2, 0.1\}$ and intrinsic (red) cases $\lambda r = \{0.5, 2.0, 4.0\}$ are shown, where the $\lambda_u = 1.0$ is defined to be zero error. The fitted lines are of the form $A/\text{Resolution}$ with $A = 6.42$ for binsizes and $B/\text{Resolution}$ with $B = 2.46$ for the intrinsic wavelength.

increased sampling requires large memory requirements and slows calculation, the optimal case should provide good agreement for minimal resolution. At 1.25 bins per $\sigma$ shown in green on Figure 4 $a)$, the sampling clearly gives a smeared density peak and is deemed not sufficiently accurate to provide a good representation of the interface. The case of 0.625 bins
is also shown in light blue with the intrinsic interface for $\lambda_r = 4\sigma$ included for comparison in dark blue. This highlights that poor binsize Resolution has the same effect as a lower wavelength fitting of the intrinsic surface. The $L^2$ Error are shown in Figure 4(b) defined as the absolute sum of the density binning obtained for a given resolution, minus the $\lambda_r = \sigma$ case at every bin between $-3$ and $2$. The blue points are the varying bilinear cases, $\Delta r$, while the red points are different intrinsic minimum wavelength values, $\lambda_r$, including $\lambda_r = 2\sigma$ and $\lambda_r = 4\sigma$ giving a blurred density from underfitting as well as the case with $\lambda_r = 0.5\sigma$ which, perhaps counter intuitively, gives a less sharp density profile due to overfitting. The points for varying intrinsic and binsize resolution in Figure 4(b) are reasonably well fitted by the lines which are of the form $A/\text{Resolution}$ and $B/\text{Resolution}$ with $A = 6.42$ and $B = 2.46$, suggesting error will tend to zero in the limit of infinite resolution but slowly. Using bilinear approximations is therefore seen to main sufficient accuracy to obtain intrinsic quantities like density, with error over the whole plot negligibly small when $\Delta r < 0.2$ and reasonable with $\Delta r < 0.4$. The effect of changing bilinear resolution can be seen to be similar to using a larger minimum wavelength when fitting the intrinsic surface, with roughly a two fold difference in trends ($A/B \approx 2.6$) so we need twice the increase in $\Delta r$ resolution to match a change in $\lambda_r$. In this work, the presented plots use $\Delta r = 0.2\sigma$ or Resolution = 5 for the bilinear approximation to ensure measure quantities such as pressure are free from any artefacts.

Having parameterised the effect of bilinear resolution, we move on to checking exact conservation for an arbitrary volume consisting of $10 \times 10$ bilinear segments of size $\Delta r = 0.4$. The surface is therefore of size $\sim 4 \times 4$ in $x$ and $y$, centred on the intrinsic surface molecules, so it is the flux either side of the interface at $\pm \Delta z/2 = \pm 0.0875$ that is recorded, along with contributions on the top and bottom due to interactions with the remaining molecules on the intrinsic interface. This is shown in Fig 5 where the terms in Eq. (47) are measured including the flux of molecules over the surface (Advection), both from molecules moving over the surface in red and the surface moving over molecules in yellow, plus the interaction force (Forcing) in green between the molecules all add up to the blue line for the change of momentum in the control volume (Accumulation). The Accumulation is shown in Fig 5 as a filled area under a curve to emphasise the fact that it is the integral of this area that
FIG. 5. Components of momentum labelled in Eq. (47) for a control volume of width $\Delta z = 0.175$ with $x$ and $y$ extents of 3.97, following the intrinsic surface with Advection including fluxes of molecules over the surface (red) and molecule crossings due to surface movement (yellow), Forcing the contributions due to forces between molecules crossing the volume surface (green) and Accumulation the resulting change in momentum inside the control volume which is exactly equal to Forcing + Advection (checked to machine precision).

determines the changing momentum inside the control volume. This average momentum is constantly changing due to forces acting on the volume as well as molecules entering and leaving. Exact conservation allows us to be sure that the implementation is correct, mollifies the issues associated with the non-uniqueness of the pressure tensor, providing an exact link between surface pressure on the volume and momentum change inside, as well as guaranteeing
all possible terms which could contribute to surface tension have been included. This exact conservation is valid for any arbitrary volume in space and is checked during all calculations in this work, providing a thorough validation of the mathematics and the implementation of the ray tracing on a complex moving surface. The contributions due to forcing can be seen to appear as a continuous line in Fig 5, as each molecule in the interface interacts with a neighbourhood of surrounding molecules within distance $r_c$. The sum of all interactions on all molecules in the control volume, as well as between molecules on either side of the $\sim 4 \times 4$ patch, varies continuously as relative proximity changes. The occasional large spikes due to molecules entering or leaving the volume represents an evaporation or condensation event. These occur both due to molecular motion and when the fitting process of the interface finds a closer molecule than the current set, which can manifest in either the interface moving past the molecule or the molecule moving over the interface which is no longer following. Molecules can also leave the volume in Fig 5 by diffusing along the intrinsic surface and leaving the 4 by 4 control volume. This detailed balance has potential applications in designing improvements to the intrinsic fitting process, for example to minimise evaporation events or track surface transport. The intrinsic fitting to a target density $\rho_s$ is equivalent to the mass control volume Eq. (21) having zero Accumulation and therefore ensuring zero net Advection for a control volume the size of the whole interface.

The new surface flux forms of pressure are compared to the volume average (VA) forms Eq. (44) in Figure 6. The volume average pressures are shown as points and surface pressures are shown by lines with kinetic pressure in red, tangential configurational pressure, $\Pi_T \equiv \frac{1}{2} \left[ \Pi^e_{xx} + \Pi^e_{yy} \right]$, shown in yellow and surface normal pressure $\Pi^c_N \equiv \Pi^c_{zz}$ in blue. The kinetic pressure term of Eq. (46) is split into a surface evolution component $\partial \xi / \partial t$ and the remaining kinetic term denoted with a prime, $\Pi^k_N$, so that,

$$\Pi^k_N = \Pi^k_N + \frac{\partial \xi}{\partial t}$$

and convection is assumed to be zero $\rho u u_z = 0$. The kinetic pressure calculated using the surface pressure definition is visually identical to the volume average one in Figure 6. Note the normal component of surface kinetic pressure is shifted by $\Delta z / 2$, as cell surfaces pressure is obtained on surfaces while volume average pressure is at the cell centres. The
FIG. 6. A comparison of pressure as a function of position over the surface tangential surface Eq. (45) and normal surface Eq. (46) (lines) compared to the volume average (points) pressure Eq. (44) from previous work\textsuperscript{5}. The kinetic pressure $\Pi_{k}^{S}$ and $\Pi_{k}^{T}$ are shown as red lines, where the prime on the normal kinetic component denotes it does not include the $\partial \xi / \partial t$ term and red points are $\Pi_{k}^{T}$. Density of particles shown as a grey line for reference, the tangential configurational pressure $\Pi_{c}^{T}$ is shown as yellow lines with $\Pi_{c}^{V}$ as yellow points and the normal component of configurational pressure $\Pi_{N}^{c}$ is shown as a blue line with blue points $\Pi_{N}^{V}$. The position of the interface is plotted as a black line with a white region to hide the peak due to the intrinsic interface.
density is shown in light grey for comparison, to highlight the kinetic pressure contributions are due to kinetic energy of molecules so directly correlated with their locations and identical in both surface and volume average measures. The similarity in tangential components of configurational pressure, shown by the yellow points and lines, is consistent with past work\textsuperscript{13}, which shows VA and MOP give identical results in the limit of small bins. As the tangential contributions are calculated on a flat surface, using the form of Eq. (45) they are expected to be identical to the VA pressure expressions. The normal components shown in blue is the only quantity that is seen to be different between surface and volume average pressure. This is a direct result of the form of Eq. (46) which calculates the pressure dotted with the surface normal $\mathbf{n}_z$ at the location of every interaction for the interface at every time. In contrast, the volume average tensor $f_{zij}r_{zij}$ is independent of the surface normal so remains aligned with the $z$ Cartesian axis. This also makes clear why the tangential components are the same, the surface pressure is shown to be dotted with the $\mathbf{n}_y$ in the flat surface case which aligns with the volume average $f_{yij}r_{yij}$ components so giving identical results. A similar observation is true for the other tangential component dotted with $\mathbf{n}_x$.

Using surface pressure normal to the instantaneous surface can be shown to be essential to the exact balance of momentum over the surface. To see this, in Figure 7 all contributions to the surface in Eq. (46) are plotted on the same graph. The configurational (blue, $\Pi^c_N$) and kinetic (red, $\Pi^k_N$) pressure terms are identical to the ones plotted on figure 6, but when the surface fluctuations (green, $\partial \xi / \partial t$) are added to give total kinetic pressure (yellow, $\Pi^k_N$), it can be seen that the resulting profile perfectly mirrors the configurational pressure. The result is the sum of the extended set of kinetic terms and configurational pressure gives a perfectly flat normal pressure over the surface, a required result for the interface to be stationary. Demonstration of a constant pressure profile near a wall in a molecular system was shown to be an important reason for using the VA or MOP form of pressure instead of the virial or IK1 expressions\textsuperscript{14,37}.

The shape of the configurational pressure profile $\Pi^c_N$ has a flat region from $z = 0.2$ to 0.8 before the first liquid peak at $z \approx 1$. The total kinetic part, $\Pi^k_N$, including $\partial \xi / \partial t$, in yellow exactly mirrors the configurational with this same flat region. It can be seen this flat region, that is, $\Pi^k_N + \partial \xi / \partial t = \text{Constant}$ from $z = 0.2$ to 0.8 results from an exchange
FIG. 7. All terms which contribute to the normal component of the total CV Pressure including configurational pressure $\Pi_{N}^{c}$ shown in green, kinetic pressure $\Pi_{N}^{k}$ shown in yellow and surface movement $\partial \xi / \partial t$ shown in blue, with the total kinetic contribution $\Pi_{N}^{k} = \Pi_{N}^{k'} + \partial \xi / \partial t$ shown in red. The total pressure $\Pi_{N} = \Pi_{N}^{k} + \Pi_{N}^{c}$, black line, is constant over the surface as required to ensure the surface is not moving. Note that this is not zero but a small finite value.

between contributions due to surface movement and contributions due to flux of molecules as we move in the $z$ direction. This is a consequence of the intrinsic surface fitting process, by choosing our reference frame to follow the interface molecules the measured kinetic pressure exposes the liquid structure in a similar way to the radial distribution function, in particular the liquid’s tendency to separate between molecular layers results in a drop to zero in the gap between the interface and first fluid layer. The interface movement term $\partial \xi / \partial t$ captures
FIG. 8. Calculation of the surface tension with normal minus tangential pressure $\Pi_N - \Pi_T$ as red lines for the full VA pressure, yellow lines for the surface configurational part only and blue lines the entire surface pressure including the surface evolution, where the minimum at $z = 0$ of the yellow and blue lines are $-1.2$ and $-4.9$ respectively. The corresponding cumulative integral of each curve to a given $z$ value, $\gamma(z) = \int_{-3}^{z} [\Pi_N(z') - \Pi_T(z')] dz'$, is shown in the same colour, with red circles VA pressure, yellow circles configurational surface and blue circles the full surface pressure.

the movement of the intrinsic interface, a reference frame which tracks the surface molecules, and that allows the plot to identify the liquid structure peaks observed. Put another way, the $\Pi^k$ term captures the structure inside the moving liquid cluster interface, the $\partial \xi/\partial t$ captures how that structure moves.

Finally, we consider the surface tension in Figure 8. In order to explore the various
contributions to surface tension from different parts of pressure, three different Kirkwood and Buff style formula are plotted,

\[
\begin{align*}
\gamma_{VA} &= \int_{-3}^{z} \left[ \Pi_{N} - \Pi_{T} \right] dz' \\
\gamma_{\text{Surf} c}^{z}(z) &= \int_{-3}^{z} \left[ \Pi_{N}^{c} - \Pi_{T}^{c} \right] dz' \\
\gamma_{\text{Surf}}^{z}(z) &= \int_{-3}^{z} \left[ \Pi_{N} - \Pi_{T} \right] dz' + \int_{-3}^{z} \frac{\partial \xi(z')}{\partial t} dz' 
\end{align*}
\] (57-59)

which includes the surface tension from the volume average pressure Eq. (57), the configuration part of the surface pressure Eq. (58) and from the full surface pressure Eq. (59).

In previous work, the contributions to surface tension from the volume average terms were discussed, and the red lines and circles in Figure 8 are presented as a basis for comparison. The configurational part is important as the normal component of configurational surface pressure was the only term which showed a difference when compared to the volume average pressure in Figure 6. As the kinetic normal and tangential components are identical for both volume average and surface pressure these cancel in the Kirkwood and Buff formula and the configurational terms contain the required contribution to surface tension. As a result, any difference between the surface tension calculated from the VA pressure and surface pressure would be expected to come from the \( \Pi_{N}^{c} \) term. The red and yellow lines in Figure 8 show the difference between these two pressure measurements is mostly located between the intrinsic interface and first layer in the liquid region from \( z = 0 \) to 1. The resulting integral to give surface tension from Eq. (58) (yellow circles) can be seen to still converge to the same value as the volume average Eq. (57) (red circles) in Figure 8. However, the surface pressure contributes more in the liquid region from \( z = -0.5 \) to \( z = 0 \), than the VA but a negative contribution from the intrinsic surface molecules themselves is observed at \( z = 0 \).

This is more pronounced for the full surface pressure, the blue line, to get the surface tension using Eq. (59), shown by blue circles in Figure 8. There is an almost linear contribution from \( z = -1 \) to \( z = 0 \) and an even larger negative contribution from the interface molecules at \( z = 0 \) when compared to the configurational part of Eq. (58). The interface control volume sits on the interfacial molecules themselves, so this peak represents the tangential forces between them, as well as a contribution due to the movement of the surface. The key
difference between the configurational Eq. (58) and the full surface pressure Eq. (59) is the inclusion of surface evolution in time \( \partial \xi / \partial t \). This can be seen to give a net zero contribution to surface tension in this equilibrium case, but redistributes giving a large negative contribution from the surface itself at \( z = 0 \) and equal positive contribution between \( z = \pm 1 \) in both the liquid and vapour region.

The surface pressure expressions have given further insight into the surface tension distribution while still integrating to the same overall value. However, this example of a flat equilibrium interface is the simplest possible case and all terms beside pressure would be expected to be contribute nothing to the surface tension. The real strength of the derived formulation is that it is valid away from equilibrium for any interface describable by a function of the form \( \xi(x, y, t) \). The control volume formulation provides exact conservation every single timestep for any arbitrary volume in space. This can provide pressure measurements and exact momentum balance in any dynamical system with complex time evolving interfaces, including bubble nucleation, film rupture or contact line dewetting.

VI. CONCLUSIONS

In this work, we derive a new formulation of surface pressure in a reference frame moving with the interface between a liquid and a vapour. This derivation starts from statistical mechanical definitions\(^{17}\) of density, momentum and energy equations but without ensemble average. The resulting equations are integrated over a volume fitted to the interface as it evolves in time, giving instantaneous surface fluxes on the moving surface. These surface flux equations are shown to include two extra contributions, one due to the instantaneous surface curvature at the point of surface crossings and one due to the movement of the surface in time. By including all contributions for curvature and surface movement, the equations can be shown to be exactly conservative in a molecular dynamics (MD) simulation. These extra terms are also shown to be essential to provide an exact force balance over the liquid-vapour interface. The derived equations are presented in a form which can be easily implemented in MD, requiring a ray tracing approach commonly used for computer graphics. Several insights into the pressure and surface tension are presented for the simplest case of a flat
interface between the liquid and vapour. Despite being tested in a simple system, the derived equations make no assumption other than mass conservation and Newton’s laws. As a result, they provide exact expressions for the equations of hydrodynamics which can take any local region on the surface and follow the instantaneous surface shape as it evolves. Therefore, the new equations are expected to have great potential applications to understand a range of hydrodynamics phenomena including Magagoni effects, growing bubbles, moving contact lines and deforming interfaces.

Appendix A: Integrating the Equations

This appendix details the full mathematics of the process used to provide the equations for use in a molecular dynamics simulation. First to highlight the similarity between the configurational and kinetic part we integrate the kinetic expression over time in Eq. (39). This can be interpreted in two ways, either as part of time averaging or as part of the evolution of the system,

\[
\int_{t_1}^{t_2} \left[ \rho \mathbf{u} \cdot \mathbf{u} + \Pi_z^k \right] dt = \frac{1}{\Delta S_z} \sum_{i=1}^{N} m_i \dot{\mathbf{r}}_i \int_{t_1}^{t_2} \left[ \dot{x}_i \frac{\partial \xi^+_i}{\partial x_i} + \dot{y}_i \frac{\partial \xi^+_i}{\partial y_i} + \dot{z}_i \frac{\partial \xi^+_i}{\partial t} \right] dS^+_{z_i} dt
\]

\[
\Pi_z^c = \frac{1}{2} \frac{1}{\Delta S_z} \sum_{i,j} f_{ij} \int_{0}^{1} \left[ \dot{x}_{ij} \frac{\partial \xi^+_i}{\partial x_\lambda} + \dot{y}_{ij} \frac{\partial \xi^+_i}{\partial y_\lambda} + \dot{z}_{ij} \right] dS^+_{z_\lambda} d\lambda.
\] (A1)

The expression for pressure in Eq. (A1) can then be written as follows,

\[
\int_{t_1}^{t_2} \left[ \rho \mathbf{u} \cdot \mathbf{u} + \Pi_z^k \right] dt = \frac{1}{\Delta S_z} \sum_{i=1}^{N} m_i \dot{\mathbf{r}}_i \left( \int_{0}^{1} \left[ \frac{dx_\tau}{d\tau} \frac{\partial \xi^+_i}{\partial x_i} + \frac{dy_\tau}{d\tau} \frac{\partial \xi^+_i}{\partial y_i} + \frac{dz_\tau}{d\tau} \right] dS^+_{z_\lambda} d\tau \right)
\]

\[
\Pi_z^c = \frac{1}{2} \frac{1}{\Delta S_z} \sum_{i,j} f_{ij} \int_{0}^{1} \left[ \frac{\partial x_\lambda}{\partial \lambda} \frac{\partial \xi^+_i}{\partial x_\lambda} + \frac{\partial y_\lambda}{\partial \lambda} \frac{\partial \xi^+_i}{\partial y_\lambda} + \frac{\partial z_\lambda}{\partial \lambda} \right] dS^+_{z_\lambda} d\lambda,
\] (A2)

where we use the substitution \( \tau = [t-t_1]/[t_2-t_1] \) in the integral over time to get the same form as the interaction over path \( \lambda \), denoting \( \mathbf{r}_1 \equiv \mathbf{r}_i(t_1) \) and \( \mathbf{r}_2 \equiv \mathbf{r}_i(t_2) \) so \( \mathbf{r}_i(t) = \mathbf{r}_\tau = \mathbf{r}_1 + \tau \mathbf{r}_{12} \) with \( \mathbf{r}_{12} = \mathbf{r}_2 - \mathbf{r}_1 \) and \( \tau \) from 0 to 1. Now both expressions are in the form \( \mathbf{r}_\alpha = \mathbf{r}_1 + s \mathbf{r}_{12} \)
with \( \alpha = \{\tau, \lambda\} \) and \( s = \{\tau, \lambda\} \) allowing us to write the generalised control volume function,

\[
\vartheta_s = \left[ H \left( x^+ - x_{\alpha}(s) \right) - H \left( x^- - x_{\alpha}(s) \right) \right] \\
\times \left[ H \left( y^+ - y_{\alpha}(s) \right) - H \left( y^- - y_{\alpha}(s) \right) \right] \\
\times \left[ H \left( \xi^+_i - z_{\alpha}(s) \right) - H \left( \xi^-_i - z_{\alpha}(s) \right) \right] = \Lambda_x(s) \Lambda_y(s) \tilde{\Lambda}_z(s),
\]

(A3)

where each directional difference between two Heaviside functions is written using shorthand \( \Lambda_\beta \) with \( \beta = \{x, y, z\} \) and just the functional dependence on \( s \) shown. The tilde on the \( \Lambda_z \) term denotes the intrinsic interface component is included on this surface. Both integral expressions in Eq. (A2) are in the form of an integral of the derivative of Eq. (A3) with respect to \( s \). More generally, the total expression for all surfaces can be written concisely as,

\[
\int_0^1 \frac{\partial \vartheta_s}{\partial s} \, ds = \int_0^1 \frac{\partial}{\partial s} \left[ \Lambda_x(s) \Lambda_y(s) \tilde{\Lambda}_z(s) \right] \, ds = \int_0^1 \left[ \frac{\partial \Lambda_x}{\partial s} \Lambda_y \tilde{\Lambda}_z + \Lambda_x \frac{\partial \Lambda_y}{\partial s} \tilde{\Lambda}_z + \Lambda_x \Lambda_y \frac{\partial \tilde{\Lambda}_z}{\partial s} \right] \, ds,
\]

(A4)

for example, with \( s = \lambda \) the last term describes the difference between top and bottom configurational term in \( z \),

\[
\int_0^1 \Lambda_x \Lambda_y \frac{\partial \tilde{\Lambda}_z}{\partial s} \, ds = \int_0^1 \left[ \frac{\partial x_\lambda}{\partial \lambda} \frac{\partial \xi^+_\lambda}{\partial x_\lambda} + \frac{\partial y_\lambda}{\partial \lambda} \frac{\partial \xi^+_\lambda}{\partial y_\lambda} + \frac{\partial z_\lambda}{\partial \lambda} \right] dS^+_{z\lambda} - \left[ \frac{\partial x_\lambda}{\partial \lambda} \frac{\partial \xi^-_\lambda}{\partial x_\lambda} + \frac{\partial y_\lambda}{\partial \lambda} \frac{\partial \xi^-_\lambda}{\partial y_\lambda} + \frac{\partial z_\lambda}{\partial \lambda} \right] dS^-_{z\lambda} d\lambda.
\]

(A5)

For the \( x \) and \( y \) surfaces, the intersection of a plane and line can be obtained directly, considering the first term on the right-hand side of Eq. (A4),

\[
\int_0^1 \frac{\partial \Lambda_x}{\partial s} \Lambda_y \tilde{\Lambda}_z \, ds = \int_0^1 \frac{dx_\alpha}{ds} \left[ \delta \left( x^+ - x_{\alpha}(s) \right) - \delta \left( x^- - x_{\alpha}(s) \right) \right] \Lambda_y(s) \tilde{\Lambda}_z(s) \, ds.
\]

(A6)

We apply the property of the Delta function to express it as the sum of its roots,

\[
\delta \left( x^+ - x_{\alpha} \right) = \sum_{k=1}^{N_{\text{roots}}} \delta \left( s - s_k \right) \frac{dx_\alpha(s_k)}{ds}.
\]

(A7)

so positive surface in \( x \) can then be expressed,

\[
\int_0^1 \frac{\partial x_\alpha}{\partial s} \delta \left( x^+ - x_{\alpha}(s) \right) \Lambda_y(s) \tilde{\Lambda}_z(s) \, ds = \int_0^1 \frac{dx_\alpha}{ds} \sum_{k=1}^{N_{\text{roots}}} \delta \left( s - s_k \right) \Lambda_y(s_k) \tilde{\Lambda}_z(s_k) \, ds
\]

\[
= \frac{dx_\alpha(s_k)}{\partial s} \left[ \sum_{k=1}^{N_{\text{roots}}} \delta \left( 1 - s_k \right) - \delta \left( -s_k \right) \right] \Lambda_y(s_k) \tilde{\Lambda}_z(s_k).
\]

(A8)

\( i \) Direction of crossing  \( ii \) Crossing between limits  \( iii \) Crossing on \( yz \) CV Surface
The three annotated terms include  

\[ \frac{\partial x_\alpha(s_k)}{\partial s} = \frac{x_{12}}{\partial x_{12}} = \frac{\mathbf{n}_x \cdot \mathbf{r}_{12}}{|\mathbf{n}_x \cdot \mathbf{r}_{12}|}, \]  

(A9)

using \( dx_\alpha/ds = x_{12} \) so \( i \) is expressed in terms of the normal to the \( x \) surface \( \mathbf{n}_x = [1, 0, 0] \).

For part ii) of Eq. (A8), the Heaviside functions between integral limits is only non-zero if the point is on the line between \( r_1 \) and \( r_2 \). The expression for this root \( s_k \) on the flat surfaces is the intersection of a plane and a line, obtained by equating the surface \( x^+ \) and line \( x_1 + sx_{12} \) to solve for the value of \( s \) at crossing \( s_k = (x^+ - x_1)/x_{12} \equiv x_k^+ \) allowing the crossing between limits term of Eq. (A8) to be expressed as,

\[ [H(1 - s_k) - H(-s_k)] = H\left(\frac{x^+ - x_2}{x_{12}}\right) - H\left(\frac{x_1 - x^+}{x_{12}}\right) = \frac{1}{2} \text{sgn}\left(\frac{x_1 + s_k x_{12}}{x_{12}}\right) \left[ \text{sgn}(x^+ - x_2) - \text{sgn}(x_1 - x^+) \right], \]  

(A10)

which is the expression found in the method of planes form of stress and determines if \( r_1 \) and \( r_2 \) are on opposite sides of the plane. Finally, iii) the location of the crossing \( s_k \) on the flat \( x^+ \) surface gives a value of zero for \( \Lambda(s_k) \) when not within the limits of the control volume surface in the \( y \) and \( z \) directions, where the \( z \) CV surface moves as the intrinsic surfaces moves, shown in Fig 3. The location of crossing in each direction is,

\[ x_\alpha(s_k) = x^+, \; y_\alpha(s_k) = y_1 + s_k y_{12}; \; z_\alpha(s_k) = z_1 + s_k z_{12}, \]  

(A11)

and the \( \tilde{\Lambda}_z(s_k) \) function is,

\[ \tilde{\Lambda}_z(s_k) = H(z^+ + \xi(z^+, y_1 + y_{12}s_k) - z_1 - z_{12}s_k) \\
- H(z^- + \xi(z^+, y_1 + y_{12}s_k) - z_1 - z_{12}s_k). \]  

(A12)

The stress on the \( x^+ \) surface is therefore,

\[
\int_{t_1}^{t_2} \left[ \rho \mathbf{u}_x + \Pi_x^k \right] dt = \frac{1}{\Delta S_x} \sum_{i=1}^{N} m_i r_i \cdot \frac{\mathbf{n}_x}{r_{i12} \cdot \mathbf{n}_x} \left[ H\left(\frac{x^+ - x_{i2}}{x_{i12}}\right) - H\left(\frac{x_{i1} - x^+}{x_{i12}}\right) \right] \Lambda_y(t_k) \tilde{\Lambda}_z(t_k) \\

\Pi_x^c = \frac{1}{2} \frac{1}{\Delta S_z} \sum_{i,j}^N f_{ij} r_{ij} \cdot \frac{\mathbf{n}_x}{r_{ij} \cdot \mathbf{n}_x} \left[ H\left(\frac{x^+ - x_j}{x_{ij}}\right) - H\left(\frac{x_i - x^+}{x_{ij}}\right) \right] \Lambda_y(\lambda_k) \tilde{\Lambda}_z(\lambda_k),
\]
note the inclusion of the index $i$ so $\mathbf{r}_{12} \to \mathbf{r}_{i_{12}}$ to emphasise it is molecule $i$ which is evolving from time $t_1$ to $t_2$. The link to the common method of planes (MOP) expression in the literature can be seen using $x/|x| = sgn(x)$ and Eq. (A10) to give,

\[
\frac{\mathbf{r}_{12} \cdot \mathbf{n}_x}{|\mathbf{r}_{12} \cdot \mathbf{n}_x|} \left[ H \left( \frac{x^+ - x_2}{x_{12}} \right) - H \left( \frac{x_1 - x^+}{x_{12}} \right) \right] = \frac{1}{2} \text{sgn}(x_{12}) \text{sgn} \left( \frac{1}{x_{12}} \right) \left[ \text{sgn}(x^+ - x_2) - \text{sgn}(x_1 - x^+) \right] = \frac{1}{2} \left[ \text{sgn}(x^+ - x_2) - \text{sgn}(x_1 - x^+) \right]
\]

(A13)

so the expressions for stress on the $x$ surface are the well known MOP form,

\[
\int_{t_1}^{t_2} \left[ \rho \mathbf{u} \cdot \mathbf{n}_x + \mathbf{\Pi}_x^c \right] dt = \frac{1}{2} \frac{1}{\Delta S_x} \sum_{i=1}^{N} m_i \mathbf{r}_i \left[ \text{sgn}(x^+ - x_{i_2}) - \text{sgn}(x_{i_1} - x^+) \right] \Lambda_y(t_k) \tilde{\Lambda}_z(t_k)
\]

\[
\mathbf{\Pi}_x^c = \frac{1}{4} \frac{1}{\Delta S_z} \sum_{i,j} f_{ij} \left[ \text{sgn}(x^+ - x_j) - \text{sgn}(x_i - x^+) \right] \Lambda_y(\lambda_k) \tilde{\Lambda}_z(\lambda_k). \quad (A14)
\]

Obtaining the expression for crossing on the other flat surfaces $x^-$ and $y^\pm$ follow the process just outlined. It is complicated in the $z$ direction by the intrinsic surface, which we can evaluate as follows. The $z$ surface is the last term on the right of Eq. (A4),

\[
\int_0^1 \Lambda_x \Lambda_y \frac{\partial \tilde{\Lambda}_z}{\partial s} ds = \int_0^1 \Lambda_x(s) \Lambda_y(s) \frac{\partial}{\partial s} \left[ H \left( z^+ - \xi(s) \right) - H \left( z^- - \xi(s) \right) \right] ds
\]

\[
= \int_0^1 \frac{\partial}{\partial s} \left[ \delta \left( z^+ - \xi - z_\alpha \right) - \delta \left( z^- - \xi - z_\alpha \right) \right] \Lambda_x \Lambda_y ds. \quad (A15)
\]

Again, using the property of the Delta function to express the sum of the roots of intersection, the crossings of the intrinsic interface by the line of interaction,

\[
\delta \left( z^+ - \xi - z_\alpha \right) = \sum_{k=1}^{N_{\text{roots}}} \frac{\delta (s - s_k)}{|d/ds (\xi(s) - z_\alpha(s))|}
\]

(A16)
allows Eq. (A15) to be written as,

\[
\int_0^1 \frac{\partial}{\partial s} (\xi - z_\alpha) \delta (z^+ - \xi - z_\alpha) \Lambda_x \Lambda_y ds
\]

\[
= \int_0^1 \frac{\partial}{\partial s} (\xi(s) - z_\alpha(s)) \sum_{k=1}^{N_{\text{roots}}} \frac{\delta (s - s_k)}{\partial s} (\xi(s) - z_\alpha(s_k)) \Lambda_x(s) \Lambda_y(s) ds
\]

\[
= \frac{\partial}{\partial s} (\xi(s_k) - z_\alpha(s_k)) \sum_{k=1}^{N_{\text{roots}}} \left[ H(1 - s_k) - H(-s_k) \right] \Lambda_x(s_k) \Lambda_y(s_k).
\]

i) Direction of crossing

ii) Crossing between limits

iii) Crossing on xy CV Surface

(A17)

The expression is broadly similar to the flat surface case of Eq. (A8), with a more complex expression for the direction of crossing in terms of the intrinsic surface i). This can be understood using a change of variable,

\[
\frac{\partial}{\partial s} (\xi - z_\alpha) = \frac{\partial r_\alpha}{\partial s} \cdot \frac{\partial}{\partial r_\alpha} (\xi - z_\alpha) = r_{12} \cdot \nabla_\alpha (\xi - z_\alpha),
\]

(A18)

which allows term i) to be written as,

\[
\frac{\partial}{\partial s} (\xi - z_\alpha) \left| \frac{\partial}{\partial s} (\xi - z_\alpha) \right| = \frac{r_{12} \cdot \nabla_\alpha (\xi - z_\alpha)}{\left| r_{12} \cdot \nabla_\alpha (\xi - z_\alpha) \right| \left| \nabla_\alpha (\xi - z_\alpha) \right|} = \frac{r_{12} \cdot \tilde{n}_z}{\left| r_{12} \cdot \tilde{n}_z \right|},
\]

(A19)

with \( \left| a \right| \) denoting vector magnitude, which must be positive so can be moved inside the absolute value on the denominator allowing the expression to be written in terms of the normal to the intrinsic surface,

\[
\tilde{n}_z \equiv \frac{\nabla_\alpha (\xi - z_\alpha)}{\left| \nabla_\alpha (\xi - z_\alpha) \right|},
\]

(A20)

this can be seen to be in the same form as the flat surface, obtaining the direction of the vector between \( r_1 \) and \( r_2 \) dotted with the surface normal. The crossing between limits of ii) and surface area term iii) are identical in form to the Eq. (A8) case, although \( s_k \) can no longer be obtained as a closed form solution as in the flat surface case of Eq. (A10).
So, the implementation of Eq. (A1) is therefore,

\[
\int_{t_1}^{t_2} \rho u_z dt + \Pi_z^k = \frac{1}{\Delta S_z} \sum_{i=1}^{N} m_i \mathbf{\dot{r}}_i \cdot \mathbf{\hat{n}}_z \sum_{k=1}^{N_{\text{roots}}} [H(1 - t_k) - H(-t_k)] \Lambda_x(t_k) \Lambda_y(t_k)
\]

\[
+ \frac{1}{\Delta S_z} \sum_{i=1}^{N} m_i \mathbf{\dot{r}}_i \partial_t
\]

\[
\Pi_z^c = \frac{1}{2 \Delta S_z} \sum_{i,j} f_{ij} \mathbf{r}_{ij} \cdot \mathbf{\hat{n}}_z \sum_{k=1}^{N_{\text{roots}}} \sum [H(1 - \lambda_k) - H(-\lambda_k)] \Lambda_x(\lambda_k) \Lambda_y(\lambda_k),
\]

(A21)

where again the use of \( \mathbf{r}_{i12} \) to emphasise this is per molecule. As the surface is no longer flat, the roots \( t_k \) and \( \lambda_k \) in this expression must be obtained using a form of algorithmic ray tracing, discussed in more detail in the body of the text.

Appendix B: Linear Algebra

The least square minimisation of the weighting function to fit the intrinsic surface has been discussed in the literature, most extensively in the supplementary material of Longford et al.\textsuperscript{[20]} However, a slightly different notation is given here to clarify a few steps, starting from,

\[
W(a_{\mu\nu}) = \frac{1}{2} \sum_{p=1}^{N_p} \left[ z_p \sum_{\mu=-k_u, \nu=-k_u}^{k_u} a_{\mu\nu}(t) f_\mu(x_p) f_\nu(y_p) \right]^2 + \psi \mathbf{\hat{A}}.
\]

(B1)

This can be seen to be a linear algebra problem by defining the vector \( \mathbf{f}(x, y) \) where each row is a wavevector,

\[
\mathbf{f}(x, y) = \begin{bmatrix}
    f_{-k_u}(x) f_{-k_u}(y) \\
    f_{-k_u}(x) f_{-k_u+1}(y) \\
    \ldots \\
    f_{k_u}(x) f_{-k_u}(y) \\
    f_{k_u}(x) f_{k_u}(y)
\end{bmatrix}
\]

(B2)
so the total matrix \( F \) can be defined by stacking matrices \( f^T \) for all pivot locations,

\[
F = \begin{bmatrix}
f^T(x_1, y_1) \\
f^T(x_2, y_2) \\
\vdots \\
f^T(x_{N_p}, y_{N_p})
\end{bmatrix}
\] (B3)

and defining \( z = [z_1, z_2, ..., z_{N_p}]^T \), \( a = a_{\nu\mu} \) and \( \tilde{A} = a^T a B \) with \( B = 4\pi^2 \text{diag}(\nu^2 + \mu^2) \) a matrix with values only on the diagonal. So, \( W \) in Eq. (53) becomes,

\[
W(a) = \frac{1}{2}||z - F a||^2 + \psi \tilde{A},
\] (B4)

and the minimum is obtained by setting the derivative with respect to \( a \) to zero,

\[
\nabla_a W = -F^T (z - F a) + \psi \nabla_a \tilde{A} = 0,
\] (B5)

and \( \nabla_a \tilde{A} = B a \) so the optimal value of \( a \) is obtained by solving this equation,

\[
a = (F^T F - \psi B)^{-1} F^T z.
\] (B6)

To maximise efficiency, LAPACK is used to solve this equation. In practice this means the matrix \( F \) of size \( M \times N_p \) with \( M = 4k^2_\omega \) wavenumbers and \( N_p \) pivot positions, is multiplied by its transpose (using e.g. Lapack DGEMM). The constraint is then applied by subtracting a matrix of size \( M \times M \) with just diagonal elements \( B = 4\pi^2 \psi [\mu^2 + \nu^2] \) that are non-zero. This can then be used in a linear algebra solver (e.g. Lapack DGESV) with right-hand side \( F^T z \) to get the values of \( a \).

Appendix C: A Derivation of the Volume Average Form

In this section, we consider how to obtain the volume average (VA) expressions from Eq. (25),

\[
\frac{d}{dt} \int_V \rho u dV = \sum_{i=1}^{N} m_i \ddot{r}_i \dot{\theta}_i = \sum_{i=1}^{N} m_i \dddot{r}_i \frac{d\vartheta_i}{dt} + \sum_{i=1}^{N} m_i \ddot{r}_i \dot{\vartheta}_i,
\] (C1)
where the continuum left-hand side is expressed using the divergence theorem,

\[
\frac{d}{dt} \int_V \rho u dV = -\oint_S (\rho u u + \mathbf{II}) \cdot d\mathbf{S} = -\int_V \frac{\partial}{\partial r} \cdot (\rho u u + \mathbf{II}) dV,
\]

while the right-hand side is expanded using Eq. (14), Eq. (27) and Eq. (28),

\[
\frac{d}{dt} \sum_{i=1}^N m_i \dot{r}_i \partial_i = \sum_{i=1}^N m_i \dot{r}_i \left( \dot{r}_i \cdot \frac{\partial \partial_i}{\partial r_i} + \frac{\partial \xi}{\partial t} \frac{\partial \partial_i}{\partial \xi} \right) + \sum_{i,j} f_{ij} \int_0^1 r_{ij} \cdot \frac{\partial \partial_{\lambda}}{\partial r_{\lambda}} d\lambda,
\]

by assuming \( \partial \partial_i \partial_r = -\partial \partial \partial_r \partial_r \) and \( \partial \partial_{\lambda} \partial_r = -\partial \partial_{\lambda} \partial_r \) and \( \partial \xi / \partial t = 0 \), we can express everything as a derivative in terms of \( r \),

\[
\int_V \frac{\partial}{\partial r} \cdot (\rho u u + \mathbf{II}) dV = \frac{\partial}{\partial r} \left[ \sum_{i=1}^N m_i \dot{r}_i \partial_i + \frac{1}{2} \sum_{i,j} f_{ij} r_{ij} \int_0^1 \partial_{\lambda} d\lambda \right],
\]

and so, comparing the expressions inside the derivative and assuming an average value for the volume yields the VA pressure given in previous work,

\[
\rho u u + \mathbf{II} = \frac{1}{\Delta V} \left[ \sum_{i=1}^N m_i \dot{r}_i \partial_i + \frac{1}{2} \sum_{i,j} f_{ij} r_{ij} \int_0^1 \partial_{\lambda} d\lambda \right].
\]

However, the assumptions \( \partial \partial_{\alpha} / \partial r_{\alpha} = -\partial \partial_{\alpha} / \partial r \) and \( \partial \xi / \partial t = 0 \) are not valid, as will be shown here for the configurational term, resulting in an extra term in the VA expression for a volume between curved interfaces. Pressure is defined in the Irving and Kirkwood\(^{17}\) process by collecting terms inside the gradient with respect to \( r \) and comparing forms to the continuum expression \( \partial / \partial r \cdot \mathbf{II} \). In the pointwise Irving and Kirkwood\(^{17}\) this uses the property of the Dirac delta \( \partial / \partial \lambda \delta (r - r_i) = -\partial / \partial \lambda \delta (r - r_i) \). For an integrated control volume between intrinsic surfaces, to see if the same process can be applied, we compare the derivatives, \( \partial \partial_{\alpha} / \partial r_{\lambda} \) and \( \partial \partial_{\alpha} / \partial r \). Starting with the derivative of \( \partial_{\lambda} \) with respect to \( r \),

\[
\frac{\partial \partial_{\lambda}}{\partial r} = i \frac{\partial \Lambda_x}{\partial x} \Lambda_y \Lambda_z + j \frac{\partial \Lambda_y}{\partial y} \Lambda_z + k \frac{\partial \Lambda_z}{\partial z},
\]

where e.g. \( \frac{\partial \Lambda_x}{\partial x} = \delta (x^+ - x_{\lambda}) - \delta (x^- - x_{\lambda}) \) and \( \frac{\partial \Lambda_{y}}{\partial z} = \delta (\xi^+_{\lambda} - z_{\lambda}) - \delta (\xi^-_{\lambda} - z_{\lambda}) \) as \( \xi^+_{\lambda} = z \pm \Delta z + \xi \) so \( d\xi^+_{\lambda} / dz = 1 \).

Next, consider the derivative of \( \partial_{\lambda} \) with respect to \( r_{\lambda} \)

\[
\frac{\partial \partial_{\lambda}}{\partial r_{\lambda}} = i \frac{\partial \Lambda_x}{\partial x} \Lambda_y \Lambda_z + j \frac{\partial \Lambda_y}{\partial y} \Lambda_z + k \frac{\partial \Lambda_z}{\partial z} \left[ i \frac{\partial \Lambda_z}{\partial x} + j \frac{\partial \Lambda_z}{\partial y} + k \frac{\partial \Lambda_z}{\partial z} \right].
\]
Noting that \( \partial \Lambda_x / \partial x = -[\delta(x^+ - x\lambda) - \delta(x^- - x\lambda)] = -\partial \Lambda_x / \partial x \) and similar for \( y \), while for \( z \) this is \( \partial \Lambda_x / \partial z = \delta(\xi_\lambda^+ - z\lambda) - \delta(\xi_\lambda^- - z\lambda) \), so \( \partial \Lambda_x / \partial z = -\partial \Lambda_x / \partial z \), so using these equivalences,

\[
\frac{\partial \vartheta}{\partial r} = -i \frac{\partial \Lambda_x}{\partial x} \Lambda_y \bar{\Lambda}_z - j \frac{\partial \Lambda_x}{\partial y} \Lambda_x \Lambda_y + \Lambda_x \Lambda_y \left[ i \frac{\partial \bar{\Lambda}_z}{\partial x\lambda} + j \frac{\partial \bar{\Lambda}_z}{\partial y\lambda} \right]
\]

where

\[
\frac{\partial \bar{\Lambda}_z}{\partial x\lambda} = \left[ \frac{\partial \xi_\lambda^+}{\partial x\lambda} \delta(\xi_\lambda^+ - z\lambda) - \frac{\partial \xi_\lambda^-}{\partial x\lambda} \delta(\xi_\lambda^- - z\lambda) \right],
\]

similar for \( y \) and using notation \( dS_\lambda^\pm = \delta(\xi_\lambda^\pm - z\lambda)\Lambda_x \Lambda_y \) so the full expression,

\[
\sum_{i,j}^N f_{ij} [\vartheta_i - \vartheta_j] = \sum_{i,j}^N f_{ij} r_{ij} \int_0^1 \frac{\partial \vartheta}{\partial r} d\lambda
\]

\[
= -\frac{\partial}{\partial r} \sum_{i,j}^N f_{ij} r_{ij} \int_0^1 \vartheta_i d\lambda + \sum_{i,j}^N f_{ij} r_{ij} \int_0^1 \left[ \frac{\partial \xi_\lambda^+}{\partial r\lambda} dS_\lambda^+ - \frac{\partial \xi_\lambda^-}{\partial r\lambda} dS_\lambda^- \right] d\lambda,
\]

(C10)

using \( \partial \xi_\lambda^\pm / \partial z\lambda = 0 \). The same argument follows for the kinetic part and the temporal evolution of the surface will also be included in the Extra Terms.

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