Diffusion at the liquid-vapor interface

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Abstract

Recently, the intrinsic sampling method has been developed in order to obtain, from molecular simulations, the intrinsic structure of the liquid-vapor interface that is presupposed in the classical capillary wave theory. Our purpose here is to study dynamical processes at the liquid-vapor interface, since this method allows tracking down and analyzing the movement of surface molecules, thus providing, with great accuracy, dynamical information on molecules that are “at” the interface. We present results for the coefficients for diffusion parallel and perpendicular to the liquid-vapor interface of the Lennard-Jones fluid, as well as other time and length parameters that characterize the diffusion process in this system. We also obtain statistics of permanence and residence time. The generality of our results is tested by varying the system size and the temperature; for the later case, an existing model for alkali metals is also considered. Our main conclusion is that, even if diffusion coefficients can still be computed, the turnover processes, by which molecules enter and leave the intrinsic surface, are as important as diffusion. For example, the typical time required for a molecule to traverse a molecular diameter is very similar to its residence time at the surface.
I. INTRODUCTION

Inhomogeneous systems present a number of features that make them intrinsically more complicated than bulk systems. The fact that the equilibrium state of the system depends on the position causes a number of physical quantities to be likewise dependent on the position (such as the molecular number density), or even ill-defined (such as the pressure tensor). This also applies to dynamical properties — the most important one of these, (self-)diffusion, is complicated by the fact that tracer molecules cannot be followed at will for any given length of time, since they will enter and abandon zones that have different dynamical properties.

It is therefore not possible in general to obtain a value for the diffusion coefficient $D$ by the well known Einstein relation for the mean standard deviation (MSD) of the displacements

$$\langle r^2 \rangle \rightarrow 6Dt.$$  \hspace{1cm} (1)

The direct application of such a formalism to an inhomogeneous fluid would result in an average result containing contributions from the bulk phases and the interfaces. Indeed, $D$ can be expected to have different values at different parts of the system. (The same problem would arise of course in the other main approach to $D$: by means of the Green-Kubo formula involving the velocity autocorrelation function.)

This is true in particular for the best-known inhomogeneous fluid system: the liquid-vapor interface. In this case, a liquid phase and its vapor are separated by an interfacial region which, on average, is flat. The spacial dependence is therefore limited in this case to one Cartesian coordinate, which we will take as $z$. For example, the mean density profile, $\rho(z)$, is obtained from simulation by defining slab in the $z$ direction (“binning”), and collecting occupation statistics for each of the slabs. This way, a profile is obtained that shows two plateaus at constant values corresponding to the liquid and vapor densities, and a typically monotonic interfacial variation between them. Theoretical approaches, from the pioneering van der Waals theory to the most recent density functional approximations, may be used to directly obtain $\rho(z)$, which depends only on the temperature, $T$, and on the molecular interactions. The density profiles may be much more structured in other, apparently more complex, systems like a dense fluid against a planar wall potential, but the apparent simplicity of the liquid-vapor interface hides a much deeper difficulty. The fluctuations of a free liquid surface have capillary wave (CW) modes with very low frequencies, and hence
low excitation energies for long wavelengths. In the absence of any external potential, the thermodynamic limit of a macroscopic free liquid surface becomes undetermined, and the interface would be fully delocalized by the long wavelength CWs. The Earth gravity field, which would be fully irrelevant for the thermodynamic properties of one-phase systems up to the scale of meters, becomes crucial to stabilize the liquid surface, damping the CW fluctuations for wavelengths larger than millimeters, and amplitudes larger than about one molecular diameter. Still, the mean density profile under Earth gravity conditions would be smoother than the one observed with typical computer simulations, which employ transverse box sizes in the range of $10 - 30$ molecular diameters (the interfacial width can be estimated to be about twice as large in the first case$^3$). Within that limited range for the transverse size the inclusion of the Earth gravity would be irrelevant, but it is already possible to observe the dependence of $\rho(z)$ with the transverse size of the simulation box$^{4,5}$.

The CW fluctuations are a severe nuisance for the study of the molecular diffusion at the liquid surface, since the molecular kinetics, relevant for any physicochemical process at the surface, is mixed with large collective fluctuations for the instantaneous position of the surface, which give a smooth and size dependent $\rho(z)$. The simplest estimations of surface diffusion properties consider the molecules within thin slabs placed in the region with inhomogeneous values of $\rho(z)$ in some cases separating diffusion in parallel and perpendicular components: Refs. 8, 10, 11 that deal with liquid-liquid interfaces, Refs. 12, 13, 14, 15, 16, 17, 18, 19, 20, 21 on liquid-vapor interfaces, and Refs. 22, 22, 23, 24, 25, 26, 27, 28, 29 on liquids (typically, water) adsorbed on, or confined in, different substrates. Another approach is to consider some operational definition of the outmost liquid molecules and track their dynamics for a limited time$^{30,31}$ Such procedures can only yield a coarse distinction between bulk and surface properties, given the obvious arbitrariness in the choice of parameters such as the surface slab width, the definition of outmost molecules, the complications associated with the diffusing molecules leaving and entering the different domains, and the blurring effect of the area-dependent CW fluctuations.

Classical capillary wave theory (CWT) gives a framework to interprete the effects of the CW fluctuations, and provides an accurate extrapolation of the $\rho(z)$ profile obtained in typical computer simulations to larger sampling sizes, including the effects of weak gravity fields. However, only over the last decade has CWT become a practical tool to extract the intrinsic molecular properties of a liquid surface, from the broad distributions produced by
the CW fluctuations. The theory assumes that an intrinsic surface (IS) may be defined, to describe the instantaneous boundary between the two coexisting phases, so that the molecular distribution referred to that surface would give an intrinsic density profile sharper that \( \rho(z) \) and, more importantly, independent of the transverse sampling size. Over the last decade, the increasing resolution in X-ray reflectivity data allowed the deconvolution of the Gaussian CW distribution out of the surface structure factor, and hence to obtain experimental results for the intrinsic profile in cold liquid metal surfaces, with a clear atomic layering structure.\(^{32}\) Similar results were obtained in computer simulations of simple fluid models,\(^{33,34}\) whenever the frustration of the freezing allowed to explore low temperatures, \( T/T_c < 0.2 \), relative to the critical one. As in experiments, the deconvolution is possible because even \( \rho(z) \) presents some layering at these low temperatures. These results indicate that the typical smooth shape of \( \rho(z) \) in a LV interface results from the convolution of the Gaussian CW fluctuations with a strongly layered intrinsic profile, such that it should be possible to identify, with reasonable confidence, the outmost molecular layer of the liquid phase. More recently, that concept led to the development of intrinsic sampling method in computer simulations,\(^{35,36,37}\) based on the operational definition of the CWT intrinsic surface as a geometrical locus for that first molecular layer, so that the intrinsic profile, and other molecular intrinsic properties of the liquid surface,\(^{38,39}\) may be extracted with accuracy and high resolution, without the CW blurring observed in \( \rho(z) \). The intrinsic profile calculated with such methods represent a direct, quantitative link between the generic framework of the CWT and the computer simulations of liquid surfaces. The result is a deeper structural understanding of the surface: the capillary wave fluctuations, which cause an area-dependent broadening of the mean profile \( \rho(z) \), are absent in the intrinsic one. These two profiles are, in general, remarkably different: between the plateaus corresponding to the liquid and vapor densities the intrinsic one is far from monotonic, showing a marked layering. In fact, it much more resembles the pair correlation function or the density profiles close to a hard wall.

The aim of our work here is to explore the application of the intrinsic sampling method to the analysis of the molecular diffusion at liquid surfaces. We may track down and analyze the movements of the surface molecules, i.e. those that define the IS. Thus we may obtain, with great accuracy, dynamical information on molecules that are “at” the interface, without the arbitrariness in the choice of a surface slab, and making it possible to separate the molecular diffusion on the surface from the fluctuations on the local position of the surface. The reader
is referred to the previous references for a complete description of the method, its variations, and their results. The only relevant aspect here is that for each instantaneous configuration of the system, this method selects a set of *surface molecules*, i.e. those identified as belonging to the *outmost liquid layer*, and called the IS *pivots* in the above references. Within the intrinsic disorder of a liquid surface we cannot expect that such molecular layer would have a sharp definition — it should indeed be regarded as a soft but robust concept: changes in the set of parameters used for the operational IS definition would produce small changes in the set of molecules which are identified as belonging to the surface, but (at least for $T/T_c < 0.8$) there is a clear optimal choice within rather narrow windows for all parameters that must be fine tuned, as commented at the end of the next section. In particular, the two-dimensional density of the first liquid layer, i.e. the number of surface molecules per unit area, is fairly well defined and provides useful information on the molecular structure of a liquid surface, which is blurred in the usual description in terms of $\rho(z)$.\textsuperscript{40}

We begin with a section on methodology, Section II, on both the simulation details and the way the IS is calculated. We then consider diffusion in Section III, divided in three subsections that go from the best known case, bulk diffusion, through diffusion parallel to the interface, to the more involved case of diffusion perpendicular to it. We further analyze our results in Section IV, where we consider systems at other temperatures and other transverse areas. We conclude with some remarks in V.

II. METHODOLOGY

We consider the standard Lennard-Jones fluid, in which molecules interact through a pairwise potential of the form

$$u(r) = \epsilon \left\{ \left( \frac{\sigma}{r} \right)^{12} - 2 \left( \frac{\sigma}{r} \right)^{6} - \left( \frac{\sigma}{r_c} \right)^{12} + 2 \left( \frac{\sigma}{r_c} \right)^{6} \right\},$$

with interactions truncated at a cutoff radius of $r_c = 3.02\sigma$.

In order to obtain dynamical information, molecular dynamics (MD) simulations are performed, using the software package DL\textunderscore POLY.\textsuperscript{41} The systems consist of a thick slab of liquid surrounded by vapor. We therefore obtain two LV interfaces. Since the two are independent (if the liquid is thick enough), the properties measured in both should be averaged in order to improve the accuracy, but for the sake of clarity we will just present
results obtained in one of the LV interfaces.

Our systems consist of 2592 molecules, generally (unless otherwise indicated) at a temperature of $k_B T = 0.678 \epsilon$, which is our estimate for the triple point temperature of the LJ fluid (at this cutoff, see Ref. 42 for a discussion of the effect of truncation on this temperature). The simulation cell is a square box of dimensions $L \times L \times L_z$, with (unless otherwise indicated) $L = 10.46 \sigma$ and $L_z = 90 \sigma$. Periodic boundary conditions are employed in all three directions. The time step is set at a reduced value of $dt = 4.56 \times 10^{-3} \sqrt{m/\epsilon}$. The systems starts from either a crystalline configuration or a system at other temperature and are equilibrated for $10^6$ time steps in the $NVT$ ensemble (with a Nose-Hoover thermostat with a time constant $10dt$).

After this period, configurations are obtained in the $NVE$ ensemble, in order to eliminate any possible spurious effects of the thermostat on the dynamics (in any case, we have checked these effects are typically negligible). For the results presented here, 50000 configurations are analyzed. We have found that only one configuration out of ten needs to be analyzed, since the dynamics is still slow for a time step of $10dt$. The analysis of the MSD is carried out in the standard way; more sophisticated treatments aimed at reducing computer storage are not needed in this case.

For each of these configurations, an IS analysis is carried out, as described in Ref. 37, requiring two operational parameters. One of them, $n_s$, has a very clear physical significance: the two-dimensional density of the IS, i.e. the number of surface molecules to be selected per unit area. Here we set $n_s \approx 0.8 \sigma^{-2}$, with a twofold motivation; on one hand, this value had already been identified as the most physical one from close inspection of the intrinsic profile for the LJ fluid. On the other, the same set of simulations described here can be used to independently obtain a value for this parameter, confirming this value as the best one from a kinetic analysis (see Ref. 40 for details on this procedure). The other parameter, $q_u$, fixes the maximum wave number to be used in the Fourier representation of the IS. This mathematical surface associated to each molecular configuration within the intrinsic sampling method will not appear explicitly this article, but still it is essential for the self-consistent procedure used to select the IS molecules. We have combined here a basis of 12 planar waves for each $x$ and $y$ direction, which with the transverse box size $L = 10.46 \sigma$ means a value of $q_u = 1.15 \times 2\pi/\sigma$, within the optimal range of choices as explained in the above references.
FIG. 1: Mean fraction of molecules which remain at the intrinsic surface for, at least, a time $t$, $N_s(t)/N_s(0)$, versus reduced time $t$, (units of $\sigma \sqrt{m/\epsilon}$). Dotted line: linear regression of the exponential decay at later times, shown to intercept $t = 0$ at about 0.8.

In each analyzed configuration we select a number of surface molecules, $N_s = n_sL^2$ (typically 88 is our optimal choice here) which self-consistently define the IS. The averages of the surface self-diffusion properties require us to follow the individual history of each of these molecules, i.e. to identify their permanence, exit, and possible reentrance in the IS list. All the reported surface diffusion properties are associated to the movement of molecules that stay continuously in the list of IS molecules, and this requirement sets a clear limitation on the available sampling times. In Figure 1, $N_s(t)$ represents the mean number of molecules which remain at the IS for, at least, a time $t$; the exponential decay $N_s(t) \sim \exp(-t/\tau)$, with a typical decay time, or residence time of $\tau \approx 4.3\sigma \sqrt{m/\epsilon}$ (at $k_B T = 0.678 \epsilon$), which we also list in Table I. This treatment mirrors the classical definition of the bulk residence time as the exponential decay time for the process by which neighboring molecules drift apart. It is also useful to keep in mind that for the case of Argon with the usual LJ parameters $\epsilon/k_B = 119.8K$ and $\sigma \approx 0.3405nm$ one unit of reduced time corresponds to 2.15 picoseconds. Thus, the decay time would be approximately 9.2ps for Argon.

This short value for the residence time implies that the sampling size would rapidly decrease (and the statistical noise increase) for large $t/\tau$. The extrapolation to $t = 0$ of this exponential decay at longer times indicates that about 80% of the molecules at the IS will leave it at the constant rate of $1/\tau$, associated to the long time exponential decay, while the remaining 20% of the molecules get out of the IS much faster, sometimes to undergo
FIG. 2: Mean density profile, $\rho(z)$ (solid line), density profile of surface molecules, $\rho_s(z)$ (dotted line), and density profile of old surface molecules, $n_0 P_0(z)$ (dashed line). The profiles are given in reduced units of $\sigma^{-3}$.

A rapid reentrance. These molecules may be regarded as those which are only marginally associated to the outmost liquid layer, e.g. those which may be interpreted either as a local intrusion of the IS towards the bulk liquid, or alternatively as a local extrusion of the next-outmost layer towards the surface. The disordered structure of the liquid makes the existence of such ambiguities unavoidable, and different recipes to identify the IS from the molecular positions could make different assignments to those molecules to be in, or out of, the list of surface molecules. That is what makes the concept of the outmost liquid layer a soft one, but at the same time a rather robust one, since any reasonable choice of the tunable parameters would agree in the selection of the large majority of the surface molecules. For the diffusion properties analyzed here there is a further advantage, since the relevant information to get the effective surface diffusion coefficients comes from the sampling of molecular displacements at the longest possible times, which automatically selects the properties of those molecules with long permanence at the surface, representing the outmost liquid layer with any sensible choice for the parameters in the IS definition. The only practical inconvenience of the rapid turnover of some surface molecules is that the time interval between analyzed configurations, $\Delta t$, has to be short enough to make sure that the estimation of $N_s(t)$ is not affected by further reduction — in practice that is achieved, at $k_B T = 0.678\epsilon$, with $\Delta t = 10dt$, when only about two and a half surface molecules are changed on average between consecutive configurations.
The mean profile of the whole system, $\rho(z)$, and that restricted to the surface molecules, $\rho_s(z)$, are presented in Figure 2. The later having a typical Gaussian shape which represents the fluctuations of the IS, and which becomes wider with increasing temperature and transverse sizes of the simulation box. Notice that despite the mean profile character of $\rho_s(z)$, all the selected surface molecules lie exactly on the instantaneous IS, we may therefore use them to follow exactly the self-diffusion of the molecules at the outmost liquid layer, without the coarsening effect of the CW fluctuations if we were selecting the molecules within a fixed surface slab. Moreover, we may keep track of the mean profiles for surface molecules which have continuous permanence in the IS list for more than a time $t$, thus defining a distribution $\rho_s(z,t)$, whose integral over $z$ is directly linked to the number of surface molecules older than $t$,

$$N_s(t) = L^2 \int dz \rho_s(z,t).$$

(3)

We may expect that any diffusion property sampled for relatively large times would correspond to molecules distributed as

$$\rho_s(z,t) = n_0 P_0(z) \exp(-t/\tau),$$

(4)

where the time dependences factorizes into the same exponential decay as $N_s(t)$, and we define a $z$-distribution of old surface molecules, $P_0(z)$, which is normalized to unity. The prefactor has been discussed above to be $n_0 \approx 0.8 n_s$, i.e. representing about 80% of the surface molecules. This is indeed the case, and in Figure 2 we compare the mean profile of the whole set of surface molecules, $\rho_s(z) \equiv \rho_s(z,0)$, and the distribution of old surface molecules normalized to $n_0$, i.e., $n_0 P_0(z)$. The distribution of old surface molecules is narrower, and asymmetric with respect to the whole one, and that may be interpreted in terms of the rate at which molecules are incorporated to, or deleted from, the list of surface molecules, as a function of their position $z$.

Since we select a fixed number of surface molecules in each configuration, the loss of molecules from the IS layer, either towards the liquid or the vapor sides, is always compensated by the incorporation of new ones, and the time reversal symmetry of the MD guarantees that the inflow and outflow of surface molecules at a given value of $z$ are identical. We denote by $\nu_{io}(z)$ that input/output rate per molecule, which may be directly sampled along our simulations and are presented in Figure 3, together with $P_0(z)$, for ease of comparison. The results are fairly well compatible with rates being independent of the
FIG. 3: Distribution of the input/output rate per molecule, $\nu_{io}(z)$, normalized to the proper value set by Equation [6] (hence, reduced in units of inverse time, $\sqrt{\epsilon/m/\sigma}$). (solid line), together with $P_o(z)$ in reduced units of $\sigma^{-1}$, normalized to unity (dashed line).

previous permanence time of the molecule at the IS, so that

$$\frac{d}{dt} N_s(t) = -L^2 \int dz \rho_s(z, t) \nu_{io}(z)$$

which for large $t$ implies

$$\frac{1}{\tau} = -\frac{1}{N_s(t)} \frac{dN_s(t)}{dt} = \int dz P_o(z) \nu_{io}(z).$$

The shape of $\nu_{io}(z)$ is clearly composed of two contributions: a large peak corresponding to the exit/entrance of surface molecules to/from the bulk liquid, an a much smaller one corresponding to the exit/entrance of surface molecules to/from the bulk vapor. In the following section we show how to extract the surface diffusion coefficients from the information from the observed displacements of the surface molecules, and the information on their turnover distributions contained in $P_o(z)$ and $\nu_{io}(z)$.

III. DIFFUSION IN THE BULK, AND AT THE INTRINSIC SURFACE

We discuss diffusion for the particular choice of temperature $k_B T = 0.678 \epsilon$, in three different cases: bulk diffusion, diffusion parallel to the interface, and diffusion perpendicular to it.
FIG. 4: Square root of the mean standard deviation of displacements for different subset of molecules versus time, in reduced units: $\sigma$ for the square root of MSDs and $\sigma \sqrt{m/\epsilon}$ for time. Solid lines: $x$, $y$, and $z$ components in the bulk liquid (hardly distinguishable), long-dashed lines: $x$ and $y$ components for the intrinsic surface (hardly distinguishable), short-dashed line: $z$ component for the intrinsic surface. Dotted lines: prediction from the Maxwellian velocity distribution, Eq. (7), and fits to Einsteinian diffusion equations, Eq. (8), with two values of the self diffusion coefficient, one corresponding to the bulk liquid, the other to the intrinsic surface.

A. Diffusion in the bulk

In Figure 4 we plot the square root of the MSD for the $x$, $y$, and $z$ components of the displacement versus time for the bulk liquid phase. At short times all curves tend to the same line, with a slope of 2. This is the ballistic regime: times so short that collisions can be neglected. In this case the Maxwellian distribution for velocities directly provides a value for the MSD

$$\langle x^2 \rangle = \frac{k_B T}{m} t^2,$$

which is the dotted line in the graph.

At longer times a diffusive regime is reached, with the Einstein relation for each of the three Cartesian coordinates because of isotropy:

$$\langle x^2 \rangle \to 2Dt; \quad \langle y^2 \rangle \to 2Dt; \quad \langle z^2 \rangle \to 2Dt.$$

We indeed find no difference between the three components of the bulk. From linear interpolation of this line in the log-log graph one reads $D_b = 0.034 \sigma \sqrt{\epsilon/m}$, in good agreement
The time for the crossover between the ballistic and the diffusive regimes can be estimated from the crossing of the two linear regression lines. In this case this is \( t_c \approx 0.099 \sigma \sqrt{m/\epsilon} \). The corresponding mean displacement in each direction would be \( \Delta x_c \approx 0.081 \sigma \) if read from the intercept of the lines, or \( \approx 0.074 \sigma \) from the MSD curves at \( t_c \). We also list these numbers in Table II where we will also include results for other temperatures that will be discussed in the next Section. Since the density is \( \rho_{\text{liq}} = 0.83 \sigma^{-3} \), assuming a local coordination close to the FCC packing, the typical intermolecular distance would be \( d = (\sqrt{2}/\rho)^{1/3} \approx 1.2 \sigma \), and collisions would take place with typical displacements around \( \Delta x \approx d - \sigma_0 \), where \( \sigma_0 \) is the minimum of the LJ potential, \( 2^{1/6} \sigma \). This prediction yields \( \Delta x \approx 0.072 \sigma \), consistent with the values found.

### B. Diffusion parallel to the interface

Turning to the molecules at the interface, i.e. those selected as *surface molecules* by the intrinsic sampling method, their MDS, along the three Cartesian coordinates also plotted in Figure [here]. The ballistic regime at short times is the same for all three components, and the same as for the bulk; but at longer times the curves are very different from the bulk,
the $x$ and $y$, parallel, components remaining indistinguishable, the $z$ component differing. We focus on the parallel diffusion in this section, for which we could expect the first two relations of Eq. (8) to hold, but with a different value of the diffusion coefficient:

$$\langle x^2 \rangle \rightarrow 2D_\parallel t; \quad \langle y^2 \rangle \rightarrow 2D_\parallel t.$$  (9)

At long times, the MSD for the parallel components of our surface molecules show this expected linear growth with $t$ limit, and we find $D_\parallel = 0.13\sigma \sqrt{\epsilon/m}$, Therefore, the parallel diffusion coefficient is almost four times larger than the bulk one. This remarkable difference, with similar factors, had already been reported in works on water, ethanol, dimethyl sulfoxide, and liquid-liquid interfaces in LJ mixtures (although not all of these works discriminate different components of the diffusion coefficient) — we should remark that, on the other hand, very little change has been obtained for liquid-liquid interfaces in mixtures of water and other polar liquids. Two works consider the liquid-vapor interface of the LJ fluid: a value of $D_\parallel = 0.13\sigma \sqrt{\epsilon/m}$, at a similar temperature of $k_B T = 0.75\epsilon$, is reported in Ref. 12 (also included in Table I), in good agreement with our result. On the other hand, Ref. 20 find only a twofold increase over the bulk diffusion coefficient, even if their choice of temperature is again very close, $k_B T = 0.75\epsilon$.

The time for the crossover between the ballistic and the diffusive regimes is $t_c \approx 0.38\sigma \sqrt{m/\epsilon}$. The corresponding mean displacement in each is either $\Delta x_c \approx 0.32\sigma$ (from the intercept of the lines), or $\approx 0.23\sigma$ (from the MSD curves). This means that the same approximate increase with respect to the bulk by a factor of about three applies to the characteristic time between collisions, the characteristic length between collisions, and the diffusion coefficient (this is of course consistent with $\langle x^2 \rangle \propto Dt$).

As shown in Figure 4, the crossover from the ballistic to the diffusive behavior is quite different for the bulk and the surface molecules. The MSD in the bulk converges to the diffusive regime from above, i.e. after leaving the rapid ballistic regime, there is a time interval in which the relative growth of the MSD is slower than in the asymptotic diffusive regime. On the contrary, the parallel diffusion of the surface molecules shows a much smoother interpolation between the two limiting regimes, which may be interpreted as a signature of the stronger disorder in the correlation structure at the surface, causing a wider time distribution for molecular rearrangement leading to the diffusive regime. The MSD is always below the asymptotic diffusion, and with very little difference between the normal
(z), and transverse (x, y) directions up to \( t \sim 0.4\sigma \sqrt{m/\epsilon} \), and displacements \( \sim 0.2\sigma \), which are approximately half way between the ballistic prediction and the observed results for bulk for the same \( t \). These values are also listed in Table II under the heading of “split.”

The surface molecules reach the diffusive regime only for typical displacements larger than one molecular diameter \( \sigma \), which require a time similar to the residence time. If we compute the time needed to diffuse to a displacement of \( \sigma \) we find \( t_{\sigma} = 4.2\sigma \sqrt{m/\epsilon} \), a value very close to that of \( \tau \) (4.3 in reduced units). This means that only about one third of the molecules at the surface (\( \approx 0.8 \times e^{-1} \)) remain in it before diffusing a transverse distance similar to their diameter. We have to bear in mind the exponential decay for the number of molecules which remain at the surface after a time \( t \) — e.g. a molecule would likely move a distance 2\( \sigma \) on the surface in a time \( t \approx 4\tau \), while in the bulk it would need typical times 10 times larger to diffuse the same distance. However, less than two percent of the surface molecules (\( \approx 0.8 \times e^{-4} \)) would remain as such for \( t = 4\tau \) and longer times, so that the large value of \( D_{\parallel} \) has limited relevance for the actual surface kinetics. The turnover process of molecules from the surface to the bulk phases, and its reverse, would be at least as relevant as the diffusion on the surface. Therefore, it is most important to analyze that turnover process, beyond its simple description in terms of the typical time \( \tau \). In the next subsection we show how the intrinsic sampling method may also give an effective diffusion coefficient for the movement of the surface molecules in the z direction, in their wandering which will eventually take them out of the surface.

C. Diffusion perpendicular to the surface

The naive relation

\[
\langle z^2 \rangle \to 2D_{\perp} t
\]  \hspace{1cm} (10)

is bound to fail if we restrict the averaging to the molecules at the surface — these are limited in space to the region about which the IS fluctuates (both in position and in time). The MSD for the z coordinate will therefore tend to a constant value equal to the squared width of the surface molecules density profile. An intermediate diffusive regime between ballistic behavior and this final plateau may appear sometimes, but there is no reason to expect this in general. Indeed, this is not the case in the situation considered here, as is obvious from the dashed line in Figure 4.
We show how to go beyond this naive prediction in a series of steps. First, Einstein’s equation is more general than Eq. (10): this would be the second moment of a probability distribution:

\[ P(z, t) = \frac{1}{\sqrt{4\pi D \tau}} \exp \left[ -\frac{z^2}{4D \tau} \right]. \] (11)

This, of course, is the solution to the diffusion equation

\[ \frac{\partial P(z, t)}{\partial t} = D \frac{\partial^2 P(z, t)}{\partial z^2}, \] (12)

with an initial Dirac delta function distribution: \( P(z, t = 0) = \delta(z) \), and vanishing values of \( P \) and its flux for large \( \pm z \). In our MD simulations that \( P(z, t) \) would correspond to selecting molecules which at the time \( t = 0 \) were within a very thin slab in the \( z \) direction, and which in terms of their displacement \( z \), would diffuse to have a probability distribution \( P(z, t) \) after a time \( t \).

The same experiment could be done selecting the molecules on the intrinsic surface at time \( t = 0 \), and following their \( z \) displacement with \( t \), but that would soon mix the surface and bulk diffusion as commented in the introduction. Alternatively, we may represent in \( P(z, t) \) only the \( z \) position of those molecules which have been continuously at the IS from \( t = 0 \). That probability distribution would not expand indefinitely, as required by the constant value of the MSD for the \( z \) component at long times in Figure 4. That effect can be ascribed to an external potential \( U(z) \) that constrains surface molecules to a particular location in space. Indeed, this potential should be identified with a potential of mean force (PMF): the mean-field description of the action of all other molecules on the one diffusing on the surface. Our diffusion equation would now be a Smoluchowski equation:

\[ \frac{\partial P(z, t)}{\partial t} = D \frac{\partial}{\partial z} \left[ \frac{\partial P(z, t)}{\partial z} + \frac{1}{k_B T} P(z, t) \frac{\partial U(z)}{\partial z} \right]. \] (13)

(application of Smoluchowski equations to diffusion in non homogeneous media can also be found in Refs. 12,28). The solution to this equation in the steady state would be an equilibrium distribution

\[ P_{eq}(z) \equiv P(z, t \to \infty) \propto \exp \left[ -\frac{U(z)}{k_B T} \right]. \] (14)

A final ingredient comes from the fact that molecules are continuously leaving the IS, a fact which may be incorporated through a loss term in the Smoluchowski equation:

\[ \frac{\partial P(z, t)}{\partial t} = D \frac{\partial}{\partial z} \left[ \frac{\partial P(z, t)}{\partial z} + \frac{1}{k_B T} P(z, t) \frac{\partial U(z)}{\partial z} \right] - \nu_0(z) P(z, t). \] (15)
where we have to plug the input/output frequency at each \( z \) position, \( \nu_{io}(z) \), discussed above and shown in Figure 3. The function \( P(z,t) \) solving this equation would not keep its normalization, since its \( z \) integral will decay with time, as the probability that an initially selected surface molecule would remain at the surface at time \( t \).

For very long times we expect that, independently of the way we initially select them, the remaining surface molecules will factorize as in Equation (4): \( P(z,t) \sim \exp(-t/\tau)P_o(z) \), in terms of the probability distribution for the \( z \) coordinates of old surface molecules, in Figure 2, and decaying with the typical time \( \tau \), so that (15) becomes

\[
D_\perp \frac{d}{dz} \left[ \frac{dP_o(z)}{dz} + \frac{P_o(z) U(z)}{k_B T} \right] - \left[ \nu_{io}(z) P_o(z) - \frac{1}{\tau} \right] P_o = 0, \tag{16}
\]

which, for a given \( D_\perp \) could be used to get the effective potential of mean force which is compatible with forms of \( P_o(z) \) and \( \nu_{io}(z) \) obtained from our MD simulations, and related to the turnover time \( \tau \) through (6). The potential of mean force could be easily obtained by integration of Equation (16):

\[
\frac{1}{k_B T} \frac{dU(z)}{dz} = -\frac{d}{dz} \log P_o + \frac{1}{D_\perp} \int_{-\infty}^{z} dz' \left[ \nu_{io}(z') - \frac{1}{\tau} \right] P_o(z'). \tag{17}
\]

Thus, the only unknown in Equation (15) is the parameter \( D_\perp \), which is required both to get \( U(z) \) and to set the scale in the time variation of \( P(z,t) \). If we integrate that equation from the equilibrium probability distribution for all the surface molecules, it would evolve in time towards the exponentially decaying distribution of old surface molecules, and we could get \( D_\perp \) through the comparison of the actual evolution of \( P(z,t) \) with the solution of the generalized Smoluchowski equation. A better estimation of \( D_\perp \) may be obtained if we start with a narrower probability distribution \( P(z,0) \), representing the surface molecules which are initially within a thin slab around a given \( z_0 \). The evolution of this probability distribution shows a first stage of broadening, dominated by the diffusion so that \( \langle (z-z_0)^2 \rangle \rightarrow 2D_\perp t \). In the later stage, the confining effects of \( U(z) \), and the losses regulated by \( \nu_{io} \) will saturate the MDS and lead to the mature distribution \( P(z,t) \sim \exp(-t/\tau)P_o(z) \), with a prefactor which would depend on the value of \( D_\perp \). The description of both the early and late stages with the same effective diffusion coefficient, which should also be fairly independent of the initial \( z_0 \) position of the chosen surface molecules, is a strong requirement to confirm the validity of the whole scheme, and hence the relevance of the best fitting value of \( D_\perp \) as a true normal surface diffusion coefficient.
In Figure 5 we show results for the time evolution of a peak initially at \( z_0 = -\sigma \) from the maximum of \( P_o(z) \) (its negative sign meaning toward the vapor side), with an initial width of 0.2\( \sigma \). The whole evolution of the \( P(z,t) \) can be traced, but we will just focus on its main features: its mean position, its MSD, and its decay (normalization). We try to fit these three curves with the numerical solution of Eq. (15), and we obtain a very good agreement fitting a perpendicular diffusion coefficient \( D_\perp = (0.10 \pm 0.01)\sigma \sqrt{\epsilon/m} \) that is comparable to, but smaller than the parallel one. Of course, the exponential decay of \( N(t) \) reproduces the previous value of \( \tau = 4.26\sigma \sqrt{m/\epsilon} \) (see dotted line in Fig. 5). We have checked that our results are largely independent of the initial position \( z_0 \), except for peaks very close to the liquid phase. Indeed, the procedure seems to fail somewhat on this side, presumably due to the more involved nature of the process close to the liquid, with “interstitial” molecules, fluctuating rapidly (ballistically) from being considered as belonging or not to the IS. Those are again the expected limitations of any attempt to locate the outmost liquid layer, and we should be ready to accept some degree of “softness” in that concept. Nevertheless, the whole procedure provides a fairly well defined value of the normal surface diffusion, with a value between those of the bulk and the tangential surface diffusion, and close to the later.

Remarkably, there is only one work, to our knowledge, that discusses normal diffusion in a liquid-vapor interface, Ref. 12. They report a value (also included in Table II) that is lower than ours, and closer to the bulk value. This is presumably due to their use of slabs, by which some of slower bulk diffusion is mixed with the normal diffusion. The remaining works all discuss liquid-liquid interfaces, providing very similar values for both diffusion coefficients at the interface,8,11 or lower values for the normal coefficient9,10 (even lower, in fact, than the bulk values). It is our feeling that this discrepancy stems mainly from the saturation of the curve corresponding to normal diffusion that has been discussed (short dashed curve in Figure 4): if a diffusion is computed for times shorter than the split time, a similar value for both coefficients will be found; but for longer times, an (effective) lower normal coefficient will be obtained.

IV. DISCUSSION

In order to support our main claims, we have performed simulations at other temperatures. We would like to consider temperatures higher and lower than the one considered.
FIG. 5: Diffusion of a narrow peak, comparing simulation results (solid lines) and predictions from Eq. (15) (dashed lines). Upper left: mean position (first moment of the distributions), in units of $\sigma$, lower left: MSD (second moment of the distributions), in units of $\sigma^2$, right: fraction of surviving molecules $N_s(t)/N_s(0)$ (normalization of the distributions). Dotted line in the later: rescaling of the dashed line to demonstrate we obtain the proper residence time. All graphs are versus reduced time $t$ (units of $\sigma\sqrt{m/\epsilon}$).

| model | $T^*$ | $\tau^*$ | bulk | IS | split | $\sigma$ |
|-------|-------|---------|------|----|-------|--------|
| SA    | 0.212 | 13.2    | 0.19 | 0.087 | 0.078 | 0.36   | 0.16 | 0.14 | 0.8 | 0.2 | 13.2 |
| LJ    | 0.678 | 4.3     | 0.099 | 0.081 | 0.074 | 0.38   | 0.32 | 0.23 | 0.4 | 0.2 | 4.2  |
| LJ    | 0.848 | 2.27    | 0.19  | 0.18  | 0.14  | 0.47   | 0.43 | 0.32 | 0.3 | 0.2 | 2.9  |

TABLE II: Table of relevant times and lengths for diffusion at different temperatures, in reduced units: $\sqrt{m/\epsilon}\sigma$ for time, $\sigma$ for lengths, $\epsilon/k_B$ for temperature. Listed: model type, temperature, residence time (these first two columns appear also in Table I), crossover time and displacements for the bulk, crossover time and displacements for the intrinsic surface, split time and displacement, time to cover a length of $\sigma$. For the crossovers, two displacements are listed, one from the intersection of the limiting lines, another from the curves themselves. Details in the main text.

Hence, we have collected results for the LJ liquid-vapor interface at a temperature 25% higher, $k_B T = 0.848\epsilon$. It is not possible in principle to lower the temperature below the triple point of LJ — hence, in order to observe the influence of a sizable drop in temperature, we have considered the soft alkali (SA) potential of Ref. 35. The parameters in this
model have a similar meaning to the ones in the LJ: $\sigma$ is the distance at which the potential vanishes and $\epsilon$ is related to the volume integral of the intermolecular potential function. Thus, the estimated critical temperature for this model is $k_B T_c \approx 1.43 \epsilon$, similar to the one of the LJ fluid ($k_B T_c \approx 1.33 \epsilon$). This is a potential engineered to result in a very low triple point temperature of $k_B T_t \approx 0.15 \epsilon$. We have considered a temperature of $k_B T = 0.212 \epsilon$. The lateral size is $L = 9.025 \sigma$, and the number of components employed in the IS Fourier analysis is 8. In addition, the optimum value of the surface density is, for this model $n_s = 0.7 \sigma^{-2}$. The curves showing the square root of the MSD as a function of time, for the two temperatures, are given in Figure 6.

The results for the residence time and for the diffusion coefficients are included in Table I. We see that with increasing temperature decay diffusion becomes faster and residence times shorter, as is to be expected. The ratio $D_\parallel/D_b$ is about 2.4 for the higher temperature (LJ fluid), since the increase in $D_\parallel$ is less drastic than that in $D_b$. For the lower temperature (SA fluid) this ratio is about 2: in this case the reduction in $D_\parallel$ turns out to be more drastic than that in $D_b$. This is probably due to the fluid phase being already very dense, while the molecules at the surface still have the option of hopping along it — surface diffusion is therefore reduced due to the prevalence of energetic attraction (exerted mainly from the bulk liquid) over entropy at lower temperatures.

We have also repeated the analysis of typical times and lengths for the curves in Figure 6 and collected this information in Table II. The most obvious correlation is between the time needed to diffuse a length of $\sigma$, $t_\sigma$ and the residence time. In all three cases, these two quantities are nearly equal. This corresponds to a natural assumption of a certain isotropy in the diffusion process: by the time a molecule at the IS has displaced a length of one molecular diameter, it is likely that the molecule has also left the IS (which is of course consistent with the IS being of molecular width). In other words, the product of $D_\parallel$ and $\tau$ should be nearly constant, which indeed is true, providing a typical length of $\sqrt{D_\parallel \tau} \approx 0.7 \sigma$ in all three cases.

There seems to be some agreement in the split length, the approximate displacement beyond which the parallel and perpendicular diffusion curves split, with a value around $0.2 \sigma$ in all cases. Other correlations are seen to be model-dependent. For example, the trend for typical bulk crossover times and lengths differs in the SA model due to its shallower potential well. Indeed, the argument given above in terms of a local coordination close to the FCC
FIG. 6: Same as Figure 4 for a lower temperature of $k_B T = 0.212\epsilon$, for the SA model (left), and a higher one of $k_B T = 0.848\epsilon$ for the same model, the LJ fluid (right). Solid lines: $x$, $y$, and $z$ components in the bulk liquid (hardly distinguishable), long-dashed lines: $x$ and $y$ components for the intrinsic surface (hardly distinguishable, except at the longest times due to statistical noise), short-dashed line: $z$ component for the intrinsic surface. Dotted lines: prediction from the Maxwellian velocity distribution, Eq. (7), and fits to Einsteinian diffusion equations, Eq. (8).

packing is still valid for the higher temperature, yielding a value of $\Delta x \approx d - \sigma_0 = 0.12\sigma$ (to compare against $0.14\sigma$), but fails for the SA model, with a value of $d$ that is below the appropriate $\sigma_0 \approx 1.5\sigma$.

In addition to varying the temperature, it is natural to explore the dependence of diffusion with the interfacial area. On one hand, the CW framework leads us to think that there could be a change of $D_\perp$ with the area. On the other hand, if this quantity is a true intrinsic property of the surface, it should be area-independent. In order to explore this possibility, we have scaled the lateral length of our simulation box, $L$, by factors of $1/2$, $2/3$, $3/2$, and 2 (i.e., the area going from $1/4$ to 4 times the original). For the last two cases, we have increased the number of molecules to 8000 and 15625, respectively (otherwise, the liquid film would have been too thin); we have also enlarged the cell in the $z$ direction, with $L_z = 200$ for the smallest area in order to accommodate a thicker liquid slab. In order to keep the same resolution in the IS Fourier treatment, the 12 functions for the original area will now be 6, 8, 18, and 24, respectively.

As shown in Figure 7 functions $P_0(z)$ and $\nu_0(z)$ contract for smaller areas and stretch for larger ones, as is to be expected from the general broadening due to CWs. (In order
FIG. 7: Comparison of the different distribution functions for different lateral sizes. Left: input/output rate per molecule, $\nu_{io}(z)$; right: density profile for old surface molecules, $P_o(z)$. Solid lines: original lateral size $L = 10.46\sigma$ (same curves as in Fig. 3); dashed line: lateral size twice longer; dotted lines: lateral size twice shorter. All functions are normalized to unity.

to obtain these curves, it is important that the various properties of the IS should be computed by subtracting the $z$ value of the $q = 0$ (constant) mode — otherwise, bulk density fluctuations produce a spurious broadening of the distributions at smaller areas, contrary to the contraction that is found). Nevertheless, the repetition of the analysis by means of the Smoluchowski Equation (15), shows no measurable dependence of $D_\perp$ with the area within our 10% error bar. This supports the identification of $D_\perp$ as a true intrinsic property of the surface, and confirms the methods described. Indeed, simpler methods based on dividing the system in slabs or selected “outmost” molecules would have resulted in a clear dependence of $D_\perp$ with the area.

V. CONCLUSIONS AND FUTURE WORK

We have described an application of the intrinsic sampling method to the analysis of dynamical processes at the liquid-vapor interface. The main conclusion is that a liquid surface is a region of enhanced molecular mobility, with respect to that in the bulk liquid, but without strong anisotropy. The characterization of the normal diffusion constant for the molecules at the outmost liquid layer requires a much more elaborated method that for the transverse component, but at the end of the day we get similar values for $D_\perp$ and $D_\parallel$. 
both at high and low temperatures for the simple liquid models studied in this article. This fact is consistent with our results for the residence time, which governs turnover rate, i.e., the rate at which molecules enter and leave the intrinsic surface — a time that turns out to be comparable to the typical times for the threshold of diffusion. Thus, the typical time required for a molecule to travel a distance on the order of a molecular diameter is very similar to its residence time as a surface molecule. Therefore, even if diffusion coefficients can still be computed for molecules that stay at the surface for times long enough, the turnover processes are equally important when discussing the dynamical properties of the interface. We have discussed two main features of these processes: the overall residence time, and the input/output rate per molecule, $\nu_{io}(z)$, the spatial distribution function of the turnover rate.

The particular details of interfacial dynamics will be, of course, model-dependent. We next provide some relevant cases on which the method described here could be applied. References will be given to previous works on these systems, but we would like to emphasize that these works have employed the usual approach (by means of slabs), therefore the present approach (by means of the IS) could shed new light on the structure and dynamics of systems of considerable applied interest.

The most important applications of this technique would now be realistic models of complex fluids, such as the liquid-vapor interface of water.\textsuperscript{12,19} In this case, the more orderly nature of liquid phase, and the high surface tension, are likely to provide additional stability for the molecules at the interface, thereby leading to longer residence times and more surface molecules reaching the diffusion regime. We have already started an effort to study this system from the point of view presented in this article — in particular, the discrepancy in our value of the normal diffusion coefficient for the LJ fluid with results of Ref. \textsuperscript{12} suggests that the corresponding values for water will likewise differ. We also intend to employ this approach to clarify the controversy surrounding normal diffusion in liquid-liquid interfaces.\textsuperscript{8,9,10,11}

Similar systems of interest include aqueous solutions\textsuperscript{16,17,18,46} (the later is a useful review on ion solvation at liquid surfaces) and surfactants at interfaces.\textsuperscript{13,14,15,21} Surfactants will be usually located at the surface, and most will reach the diffusive regime. The relationship between structure and dynamics is specially interesting in systems with strong profiling, such as liquid metals,\textsuperscript{20} confined water\textsuperscript{22,23,25,27,28} and liquid-solid interfaces.\textsuperscript{29} Similarly, diffusion in amphiphilic bilayer structures\textsuperscript{24,25} micelles\textsuperscript{26} and microemulsions can be described in a
manner similar to the one presented here.

VI. ACKNOWLEDGMENTS

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