
This version is available at http://strathprints.strath.ac.uk/47700/

Strathprints is designed to allow users to access the research output of the University of Strathclyde. Unless otherwise explicitly stated on the manuscript, Copyright © and Moral Rights for the papers on this site are retained by the individual authors and/or other copyright owners. Please check the manuscript for details of any other licences that may have been applied. You may not engage in further distribution of the material for any profitmaking activities or any commercial gain. You may freely distribute both the url (http://strathprints.strath.ac.uk/) and the content of this paper for research or private study, educational, or not-for-profit purposes without prior permission or charge.

Any correspondence concerning this service should be sent to the Strathprints administrator: strathprints@strath.ac.uk
Identifying interfacial molecules in nonplanar interfaces: the generalized ITIM algorithm

Marcello Sega∗
Tor Vergata University of Rome, via della Ricerca scientifica 1, I-00133 Rome, Italy

Sofia S. Kantorovich
Sapienza University of Rome, p.le A. Moro 4, I-00188 Rome, Italy and
Ural Federal University, Lenin Ave. 51, 620083 Ekaterinburg, Russia

Pál Jedlovszky
Laboratory of Interfaces and Nanosize Systems,
Institute of Chemistry, Eötvös Loránd University,
Pázmány stny 1/a, H-1117 Budapest, Hungary
MTA-BME Research Group of Technical Analytical Chemistry. Szt. Gellért tér 4, H-1111 Budapest, Hungary and
EKF Department of Chemistry, H-3300 Eger, Leányka u. 6, Hungary

Miguel Jorge
LSRE/LCM-Laboratory of Separation and Reaction Engineering,
Faculdade de Engenharia, Universidade do Porto,
Rua Dr. Roberto Frias, 4200-465 Porto, Portugal
Abstract

We present a generalized version of the ITIM algorithm for the identification of interfacial molecules, which is able to treat arbitrarily shaped interfaces. The algorithm exploits the similarities between the concept of probe sphere used in ITIM and the circumsphere criterion used in the $\alpha$-shapes approach, and can be regarded either as a reference-frame independent version of the former, or as an extended version of the latter that includes the atomic excluded volume. The new algorithm is applied to compute the intrinsic orientational order parameters of water around a DPC and a cholic acid micelle in aqueous environment, and to the identification of solvent-reachable sites in four model structures for soot. The additional algorithm introduced for the calculation of intrinsic density profiles in arbitrary geometries proved to be extremely useful also for planar interfaces, as it allows to solve the paradox of smeared intrinsic profiles far from the interface.
I. INTRODUCTION

Capillary waves represent a conceptual problem for the interpretation of the properties of liquid-liquid or liquid-vapor planar interfaces, because long-wave fluctuations are smearing the density profile across the interface and all other quantities associated to it. This is usually overcome by calculating the density profile using a local, instantaneous reference frame located at the interface, commonly referred to as the intrinsic density profile, \( \rho(z) = \langle A^{-1} \sum_i \delta(z - z_i + \xi(x_i, y_i)) \rangle \), where \((x_i, y_i, z_i)\) is the position of the \(i\)-th atom or molecule, and the local elevation of the surface is \(\xi(x_i, y_i)\), assuming the macroscopic surface normal being aligned with the Z axis of a simulation box with cross section area \(A\). During the last decade several numerical methods have been proposed to compute the intrinsic density profiles at interfaces\(^1\)–\(^6\). Despite several differences in these approaches, they are, in general, providing consistent distributions of interfacial atoms or molecules\(^6\) and density profiles\(^7\). Among these methods, \(\textit{itim}\)\(^4\) proved to be an excellent compromise between computational cost and accuracy\(^6\), but it is limited to macroscopically flat interfaces, therefore there is a need to generalize it to arbitrary interfacial shapes.

Before these works, albeit for other purposes, several surface-recognition algorithms have been devised, and will be briefly mentioned below. All of them are possible starting points for the sought generalization under the condition that, once applied to the special case of a planar interface, they lead to consistent results with existing algorithms for the determination of intrinsic profiles.

Historically, the first class of algorithms addressing the problem of identifying surfaces was developed to determine molecular areas and volumes. The study of solvation properties of molecules and macromolecules (usually, proteins) might require the identification of molecular pockets, or the calculation of the solvent-accessible surface area for implicit solvation models\(^8\). Two intuitive concepts are commonly used to describe the surface properties of molecules, namely, that of solvent-accessible surface\(^9,10\) (SAS), and that of molecular surface\(^11,12\) (MS, also known as solvent excluded surface, or Connolly surface). The MS can be thought as the surface obtained by letting a hard sphere roll at close contact with the atoms of the molecule, to generate a smooth surface made of a connection of pieces of spheres and tori, which represents the part of the van der Waals surface exposed to the solvent. During the process of determining the surface, interfacial atoms can be identified
using a simple geometrical criterion. Many approximated\textsuperscript{13–24} or analytical\textsuperscript{11,12,25–30} methods have been developed to compute the MS or the SAS. In general, these methods are based on discretization or tessellation procedures, requiring therefore the determination of the geometrical structure of the molecule. Other methods which allow to identify molecular surfaces include the approaches of Willard and Chandler\textsuperscript{5} or the Circular Variance method of Mezei\textsuperscript{31}. Incidentally, the way the MS is computed in the early work of Greer and Bush\textsuperscript{15} resembles very closely the ITIM algorithm\textsuperscript{4}.

From the late 1970s, the problem of shape identification had started being addressed by a newly born discipline, computational geometry. In this different framework, several algorithms have been actively pursued to provide a workable definition of surface, and in particular the concept of $\alpha$-shapes\textsuperscript{32,33} showed direct implications for the determination of the molecular surfaces\textsuperscript{34,35}. The approach based on $\alpha$-shapes is particularly appealing due to its generality and ability to describe, besides the geometry, also the intermolecular topology of the system.

Noticeably, none of these methods – to the best of our knowledge – has ever been employed for the determination of intrinsic properties at liquid-liquid or liquid-gas interfaces. Prompted by the apparent similarities between the usage of the circumsphere in the alpha shapes and that of the probe sphere in the ITIM method, as we will describe in the next section, we investigated in more detail the connection between these two algorithms. As a result, we developed a generalized version of ITIM (GITIM) based on the $\alpha$-shapes algorithm. The new GITIM method consistently reproduces the results of ITIM in the planar case while retaining the ability to describe arbitrarily shaped surfaces. In the following we describe briefly the alpha shapes and the ITIM algorithms, explain in detail the generalization of the latter to arbitrarily shaped surfaces, and present several applications.

II. ALPHA SHAPES AND THE GENERALIZED ITIM ALGORITHM

The concept of $\alpha$-shapes was introduced several decades ago by Edelsbrunner\textsuperscript{32,33}. To date the method is applied in computer graphics application for digital shape sampling and processing, in pattern recognition algorithms and in structural molecular biology\textsuperscript{36}. The starting point in the determination of the surface of a set of points in the $\alpha$-shapes algorithm is the calculation of the Delaunay triangulation, one of the most fruitful concepts for compu-
tational geometry\textsuperscript{37,38}, which can be defined in several equivalent ways, for example, as the triangulation that maximizes the smallest angle of all triangles, or the triangulation of the centers of neighboring Voronoi cells. The idea behind the $\alpha$-shapes algorithm is to perform a Delaunay triangulation of a set of points, and then generate the so-called $\alpha$-complex from the union of all k-simplices (segments, triangles and tetrahedra, for the simplex dimension $k=1, 2$ and 3, respectively), characterized by a k-circumsphere radius (which is the length of the segment, the radius of the circumcircle and the radius of the circumsphere for $k=1, 2$ and 3, respectively) smaller than a given value, $\alpha$ (hence the name). The $\alpha$-shape is then defined as the border of the $\alpha$-complex, and is a polytope which can be, in general, concave, topologically disconnected, and composed of patches of triangles, strings of edges and even sets of isolated points. In a pictorial way, one can imagine the $\alpha$-shape procedure as growing probe spheres at every point in space until they touch the nearest four atoms. These spheres will have, in general, different radii. Those atoms that are touched by spheres with radii larger than the predefined value $\alpha$ are considered to be at the surface.

An example of the result of the $\alpha$-shapes algorithm in two dimensions is sketched in Fig. 1a. The ITIM algorithm is based instead on the idea of selecting those atoms of one phase that can be reached by a probe sphere with fixed radius streaming from the other phase along a straight line, perpendicular to the macroscopic surface. An atom is considered to be reached by the probe sphere if the two can come at a distance equal to the sum of the probe sphere and Lennard-Jones radii, and no other atom was touched before along the trajectory of the probe sphere. In practice, one selects a finite number of streamlines, and if the space between them is considerably smaller than the typical Lennard-Jones radius $R_p$, the result of the algorithm is practically independent of the location and density of the streamlines. The same is not true regarding the orientation of the streamlines; this is a direct consequence of the algorithm being designed for planar surfaces only. The basic idea behind the ITIM algorithm are sketched in Fig. 1b. A closer inspection reveals that the condition of being a surface atom for the ITIM algorithm resembles very much that of the $\alpha$-shapes case. Quadruplets of surface atoms identified by the ITIM algorithm have the characteristic of sharing a common touching sphere having the same radius as the probe sphere. In this way, one can see the analogy with the $\alpha$-shapes algorithm, the $R_p$ parameter being used instead of $\alpha$. The most important differences in the $\alpha$-shapes algorithm with respect to ITIM are the absence of a volume associated with the atoms, and its independence from any reference
frame. We devised, therefore, a variant of the $\alpha$-shapes algorithm that takes into account the excluded volume of the atoms.

In the approach presented here the usual Delaunay triangulation is performed, but the $\alpha$-complex is computed substituting the concept of the circumsphere radius with that of the radius of the touching sphere, thus introducing the excluded volume in the calculation of the $\alpha$-complex. Note that this is different from other approaches that are trying to mimic the presence of excluded volume at a more fundamental level, like the weighted $\alpha$-shapes algorithm, which uses the so-called regular triangulation instead of the Delaunay one\textsuperscript{33}. In addition, in order to eliminate all those complexes, such as strings of segments or isolated points, which are rightful elements of the shape, but do not allow a satisfactory definition of a surface, the search for elements of the $\alpha$-complex stops in our algorithm at the level of tetrahedra, and triangles and segments are not checked. In this sense GITIM can provide substantially different results from the original $\alpha$-shapes algorithm.

The equivalent of the $\alpha$-complex is then realized by selecting the tetrahedra from the Delaunay triangulation whose touching sphere is smaller than a probe sphere of radius $R_p$, and the equivalent of the $\alpha$-shape is just its border, as in the original $\alpha$-shapes algorithm. The procedure to compute the touching sphere radius is described in the Appendix.

In the implementation presented here, in order to compute efficiently the Delaunay triangulation, we have made use of the quickhull algorithm, which takes advantage of the fact that a Delaunay triangulation in $d$ dimensions can be obtained from the ridges of the lower convex hull in $d+1$ dimensions of the same set of points lifted to a paraboloid in the ancillary dimension\textsuperscript{39}. The quickhull algorithm employed here\textsuperscript{40} has the particularly advantageous scaling $\mathcal{O}(N \log(\nu))$ of its computing time with the number $N$ and $\nu$ of input points and output vertices, respectively.

A separate issue is represented by the calculation of the intrinsic profiles (whether profiles of mass density or of any other quantity) as the distance of an atom in the phase of interest from the surface is not calculated as straightforwardly as in the respective non-intrinsic versions. For each atom in the phase, in fact, three atoms among the interfacial ones have to be identified in order to determine by triangulation\textsuperscript{7} the instantaneous, local position of the interface. This issue will be discussed in Sec. III for the planar, for the spherical or quasi-spherical and for the general case: here we simply note that we turned down an early implementation of the algorithm that searches for these surface atoms, based on the sorting
of the distances using $O(N \log N)$ algorithms like quicksort, in favor of a better performing approach, based on kd-trees$^{41,42}$, a generalization of the one-dimensional binary tree, which are still built in a $O(N \log N)$ time, but allow for range search in (typically) $O(\log N)$ time.

III. COMPARISON BETWEEN THE ITIM AND THE GITIM METHODS

We have compared the results of the ITIM and GITIM algorithms applied to the water/carbon tetrachloride interface composed of 6626 water and 966 CCl$_4$ molecules. The water and CCl$_4$ molecules have been described by the TIP4P model$^{43}$, and by the potential of McDonald and coworkers$^{44}$, respectively. The molecules have been kept rigid using the SHAKE algorithm$^{45}$. This simulation, as well as the others reported in this work have been performed using the Gromacs$^{46}$ simulation package employing an integration time step of 1 fs, periodic boundary conditions, a cutoff at 0.8 nm for Lennard-Jones interactions and the smooth Particle Mesh Ewald algorithm$^{47}$ for computing the electrostatic interaction, with a mesh spacing of 0.12 nm (also with a cut-off at 0.8 nm for the real-space part of the interaction). All simulations were performed in the canonical ensemble at a temperature of 300K using the Nosé–Hoover thermostat$^{48,49}$ with a relaxation time of 0.1 ps. A simulation snapshot of the H$_2$O/CCl$_4$ interface is presented in Fig. 2, where the surface atoms identified by the GITIM algorithm using a probe sphere radius of 0.25 nm are highlighted using a spherical halo.

We have used the ITIM and GITIM algorithms to identify the interfacial atoms of the water phase in the system, for different sizes of the probe sphere. In general, GITIM identifies systematically a larger number of interfacial atoms than ITIM for the same value of the probe sphere radius $R_p$, as it is clearly seen in Fig. 3. Remarkably, for values of the probe sphere radius smaller than about 0.2 nm (compare, for example, with the optimal ITIM parameter $R_p = 0.125$ nm suggested in Ref. 6), the interfacial atoms identified by GITIM show the onset of percolation. The reason for this behavior traces back to the fact that ITIM is unable to identify voids buried in the middle of the phase, as it is effectively probing only the cross section of the voids along the direction of the streamlines. This difference could explain the higher number of surface atoms identified by GITIM, as voids in a region with high local curvature (or, in other words, with a local surface normal which deviates significantly from the macroscopic one) will not be identified as such by ITIM. In GITIM, on the contrary, probe
spheres can be thought as inflating at every point in space instead of moving down the streamlines, and this is the reason why the algorithm is able to identify also small pockets inside the opposite phase.

It is possible to make a rough but enlightening analytical estimate of the probability for a probe sphere of null radius in the ITIM algorithm to penetrate for a distance $\zeta$ in a fluid of hard spheres with diameter $\sigma$ and number density $\rho$. Using the very crude approximation of randomly distributed spheres, the probability $p_0$ to pass the first molecular layer, at a depth $\zeta = \sigma$ is the effective cross section $p_0 = 1 - \frac{\pi}{4} \rho^{2/3} \sigma^2$, and that of reaching a generic depth $\zeta$ can be approximated as $p(\zeta) = p_0^{\zeta/\sigma}$, where $\kappa = \ln(1/p_0)/\sigma$ defines a penetration depth. Therefore, using a probe sphere with a null radius, ITIM will identify a (diffuse) surface at a depth $1/\kappa$, while GITIM will identify every atom as a surface one. For water at ambient conditions, the penetration is $\kappa^{-1} \simeq 0.186$ nm, a distance smaller than the size of a water molecule itself. This could explain why in Ref. 6, even using a probe sphere radius as small as 0.05 nm, almost only water molecules in the first layer were identified as interfacial ones by ITIM (see the almost perfectly Gaussian distribution of interfacial water molecules in Fig. 9 of Ref. 6).

Nevertheless, it is important for practical reasons to be able to match the outcome of both algorithms. It turns out that choosing $R_p$ so that the average number of interfacial atoms identified by both algorithms is roughly the same leads also, not surprisingly, to very similar distributions. The probe sphere radius required for GITIM to obtain a similar average number of surface atom as in ITIM can be obtained by an interpolation of the values reported in Fig. 3. An example showing explicitly the interfacial atoms identified by the two methods ($R_p = 0.2$ nm for ITIM and $R_p = 0.25$ nm for GITIM) is presented in Fig. 4: roughly 85% of surface atoms are identified simultaneously by both methods, demonstrating the good agreement between the two methods once the probe sphere radius has been re-gauged. The condition of identifying the same atoms as interfacial ones is much more strict that any condition on average quantities, like the spatial distribution of interfacial atoms or intrinsic density profiles. Hence, it is expected that a good agreement on such quantities can also be achieved.

The intrinsic density profiles of water and carbon tetrachloride are reported in Fig. 5, as computed by ITIM and GITIM, respectively, with the interfacial water molecules as reference. The procedure for identifying the local distance of an atom from the surface is in its essence
the same as described in Ref. 7. Starting from the projection \( P_0 = (x, y) \) of the position of the given atom to the macroscopic interface plane, the two interfacial atoms closest to \( P_0 \) are found (their position on the interface plane being \( P_1 \) and \( P_2 \), respectively). The third closest atom with projection \( P_3 \) has then to be found, with the condition that the triangle \( P_1 P_2 P_3 \) contains the point \( P_0 \). A linear interpolation of the elevation of \( P_0 \) from those of the other points is eventually performed, and employed to compute the distance \( z - \xi(x, y) \) which is used to compute the intrinsic density profile.

Efficient neighbor search for the \( P_1, P_2 \) and candidate \( P_3 \) atoms is implemented using kd-trees\(^42\) as discussed before. The two pairs of profiles are very similar, besides a small difference in the position and height of the main peak of the CCl\(_4\) profile (curves on the right in Fig. 5) and in the minimum of the water profile (curves on the left in Fig. 5) right next to the surface position, which are anyway compatible with the differences observed between various methods for the calculation of intrinsic density profiles\(^7\). The delta-like contribution of the water molecules at the surface is included in the plot in Fig. 5, and defines the origin of the reference system. Negative values of the signed distance from the interface correspond to the aqueous phase.

IV. THE PROBLEM OF NORMALIZATION OF DENSITY PROFILES

Before applying GITIM to non-planar interfaces, one important issue has still to be solved, namely that of the proper calculation of intrinsic density profiles in non-planar geometries. In general, one uses one-dimensional density profiles (intrinsic or non-intrinsic) when the system is, or is assumed to be, invariant under displacements along the interface, so that the orthogonal degrees of freedom can be integrated out. When the interface has a non-planar shape, one needs to use a different coordinate system. In the case of a quasi-spherical object for example, one could use the spherical coordinate system to compute the non-intrinsic density profile, and normalize each bin by the integral of the Jacobian determinant, that is the volume of the shell at constant distance from the origin. In the intrinsic case, however, it is necessary to know at every time step the volume of the shells at constant distance from the interface.

The volume of shells at constant intrinsic distance can, in principle, be calculated at each frame by regular numerical integration, but this would require a large computing time and
storage overhead. Here, instead, we propose to employ an approach based on simple Monte Carlo integration: in parallel with the calculation of the histograms for the various phases, we compute also that of a random distribution of points, equal in number to the total atoms in the simulation. The volume of a shell can be estimated as box volume times the ratio of the number of points found at a given distance and the total number of random points drawn. We are following the heuristic idea that for each frame $j$ one does not need to know the volume of the shell $V_j(r)$ with a precision higher than that of the average number of atoms in it, $N_j(r)$. In addition, we assume that the surface area of the interface is large enough for the shell volume variations $\delta V_j(r)$ to be small with respect to its average value $\hat{V}(r) = \sum_j^N V_j(r)/N$. The average density

$$\rho(r) = \frac{1}{N} \sum_{j=1}^N \frac{N_j(r)}{V_j(r)}$$  \hspace{1cm} (1)

can be approximated as

$$\rho(r) \simeq \frac{1}{N} \frac{1}{V(r)} \sum_{j=1}^N \left[ N_j(r) - N_j(r) \frac{\delta V_j(r)}{\hat{V}(r)} \right].$$  \hspace{1cm} (2)

When the relative volume changes $|\delta V/V|$ are small, one can therefore simply normalize the histogram $\tilde{N}(r) = \sum_j^N N_j(r)/N$ by the average volume $\hat{V}(r)$ obtained by the Monte Carlo procedure, disregarding the terms of order $O(\delta V/\hat{V})$.

The correctness of our assumption is demonstrated incidentally by the application of this normalization once again to the planar case. The thin lines in Fig. 5 represent the intrinsic mass density profile of water and carbon tetrachloride, using the Monte Carlo normalization scheme instead of the usual normalization with box cross sectional area and slab width. Close to the interface, the Monte Carlo normalization gives results which are fully compatible with the usual method, showing that the accuracy of the volume estimate is adequate. On the other hand one can see that far from the interface the two profiles behave quite differently. The case with usual normalization decays slowly to zero: this effect is due to the presence of the second interface, whose profile is smeared again by capillary waves. The case with Monte Carlo normalization, on the contrary, shows that it is possible to recover the proper intrinsic density also at larger distances, and features such as the fourth peak at 2 nm, which are completely hidden in the normal picture, can be revealed. This shows that the use of the proper, curvilinear coordinate system is of fundamental importance also
for macroscopically planar interfaces. The calculation of the Monte Carlo normalization factors does not change the typical scaling of the algorithm, as it consists in calculating the histogram for an additional phase of randomly distributed points (which effectively behaves as an ideal gas). The better accuracy at larger distances, however, demonstrates that the use of the Monte Carlo normalization is much more efficient than the standard approach, as it requires much smaller systems to be able to extract the same information (e.g., to resolve the fourth peak in Fig. 5, an additional slab of about 2-3 nm would have been needed). In this sense, the Monte Carlo normalization procedure can be even beneficial in terms of performance.

V. EXAMPLES OF NON-PLANAR INTERFACES

A. DPC micelle

Dodecylphosphocholine (DPC) is a neutral, amphiphilic molecule with a single fatty tail that can form micelles in solution: these play a relevant role in biochemistry, especially for NMR spectroscopy investigations aiming at understanding the structure of proteins or peptides bound to an environment that is similar to the biological membrane\textsuperscript{50–53}. The molecular structure of DPC is shown in Fig.6. We have simulated for 500 ps a micelle of 65 DPC and 6305 water molecules using the force field and configurations from Tieleman and colleagues\textsuperscript{54}, and have calculated the intrinsic mass density profiles of both phases (DPC and water) using \textsc{gitim} and the Monte Carlo normalization procedure, with a probe sphere radius $R_p = 0.25$. The result of the interfacial atoms identification on the DPC micelle for a single frame is shown in Fig. 6, where water molecules have been removed for the sake of clarity, and interfacial atoms are highlighted as usual with a halo. The intrinsic mass density profile, calculated relative to the DPC surface, is reported in Fig. 7, with the DPC mass density profile shown on the left, and the water profile on the right.

As usual, the delta-like contribution at $r = 0$ identifies the contribution from interfacial DPC atoms. In addition, we have calculated, for the first time, the intrinsic profiles of the orientational order parameters $S_1$ and $S_2$ of the water molecules around the DPC micelle. The two parameters are defined as $S_1 = \langle \cos(\theta_1) \rangle$ and $S_2 = \langle 3 \cos^2(\theta_2) - 1 \rangle / 2$, where $\theta_1$ and $\theta_2$ are the angles between the water molecule position vector (with respect to the micelle
center), and the water molecule symmetry axis and molecular plane normal, respectively. The orientation is taken so that $\theta_1 < \pi/2$ when the hydrogen atoms are farther from the micelle than the corresponding oxygen. The complete picture of the orientation of water molecules would be delivered by the calculation of the probability distribution $p(\theta_1, \theta_2)\textsuperscript{55,56}$, but here we limit our analysis to the two separate order parameters and their intrinsic profiles. Note that, since these quantities are computed per particle, there is no need to apply any volume normalization. The polarization of water molecules, which is proportional to $S_1$, appears to be different from zero only very close to the micellar surface. In particular, $S_1$ has a correlation with the main peak of the intrinsic density profile in the proximity of 0.4 nm. Water molecules located closer to the interface show a first change in the sign of the polarization and a subsequent one when crossing the interface. Farther than 0.25 nm inside the micelle, not enough water molecules are found to generate any meaningful statistics. Also the $S_2$ order parameter is practically zero beyond 0.6 nm, and again a correlation is seen with the main peak of the intrinsic density profile, and the maximum in the orientational preference is found just next to the interface, where $S_1 \simeq 0$, showing that water molecules are preferentially laying parallel to the interfacial surface.

B. Soot

One of the main byproducts of hydrocarbon flames, soot is thought to have a relevant impact on atmospheric chemistry and global surface warming\textsuperscript{58,59}. Electron, UV, and atomic force microscopy have revealed the size and structure of soot particles from different sources at different scales\textsuperscript{60–64}. In particular, soot emitted by aircraft is found to be made of several, quasi-spherical, concentric graphitic layers of size in the range from 5 to 50 nm\textsuperscript{60}. We have used four model structures ($S_{I1}, S_{I2}, S_{I4}$ and $S_{II}$ from Ref. 57) to demonstrate the ability of GITIM to identify surface atoms in complex geometries. In Fig. 8, the $S_{I1}$ model is represented in section as a triangulated surface (right), showing the four concentric layers, and in whole (left) showing the surface atoms as detected by GITIM using $R_p = 0.25$ nm. The histograms of the total number of atoms and of the surface ones, as a function of the distance from the center of the soot particles, are shown in Fig. 9 for the four different models, where it is seen how particles of the size of a water molecule have mostly access only to the inner and outer parts of the innermost and outermost shell, respectively, and cover them almost completely.
This finding is in a clear accordance with the results of the void analysis and adsorption isotherm calculations presented in Ref. 57.

C. Secondary cholic acid micelle

Bile acids, such as cholic acid are biological amphiphiles built up by a steroid skeleton and side groups attached to it. The organization of these side groups is such that hydrophilic and hydrophobic groups are located at the two opposite sides of the steroid ring. Thus, bile acids have a hydrophilic and a hydrophobic face (often referred to as the $\alpha$ and $\beta$ side, respectively) rather than a polar head and an apolar tail, as in the case of other surfactants like, for example, DPC. The unusual molecular shape leads to peculiar aggregation behavior of bile acids. At relatively low concentrations they form regular micelles with an aggregation number of 2-10, while above a second critical micellar concentration these primary micelles form larger secondary aggregates by establishing hydrogen bonds between the hydrophilic surface groups of the primary micelles$^{65,66}$. These secondary micelles are of rather irregular shape,$^{66,67}$ which makes them an excellent test system for our purposes.

Here we analyze the surface of a secondary cholic acid micelle composed of 35 molecules, extracted from a previous simulation work$^{66}$ and simulated for the present purposes for 500 ps in aqueous environment. An instantaneous snapshot of the micelle is shown in Fig.10 (water molecules are omitted for clarity) together with a schematic structure of the cholic acid molecule. We calculated the density profile of water as well as of cholic acid relative to the intrinsic surface of the micelle by the gitim method. The resulting profiles are shown in Fig.11. The micelle has a characteristic elongated shape, which exposes a large part of its components to the solvent, so that roughly 80% of the micelle atoms are identified as surface ones. The small volume to surface ratio of the micelle is at the origin of the rather noisy intrinsic density profile for the micelle itself. The profile, in addition to the delta-like contribution at the surface, presents another very sharp peak located at a distance of about 0.18 nm inside the surface, due to the rather rigid structure of the bile molecule. The water intrinsic density profile, on the contrary, shows a marked peak at 0.25 nm, absent in the DPC micelle case, due to the presence of hydrogen bonds between water molecules and the hydroxyl groups of cholic acid.
VI. CONCLUSIONS

In this paper we presented a new algorithm that combines the advantageous features of both the ITIM method\textsuperscript{4} and the $\alpha$-shapes algorithm\textsuperscript{32,33} to be used in determining the intrinsic surface in molecular simulations. Thus, unlike the original variant, this new, generalized version of ITIM, dubbed GITIM, is able to treat interfaces of arbitrary shapes and, at the same time, to take into account the excluded volumes of the atoms in the system. It should be emphasized that the GITIM algorithm is not only able to find the external surface of the phase of interest, but it also detects the surface of possible internal voids inside the phase. The method, based on inflating probe spheres up to a certain radius in points inside the phase turned out to provide practically identical results with the original ITIM analysis for planar interfaces. Further, its applicability to non-planar interfaces was shown for three previously simulated systems, i.e., a quasi-spherical micelle of DPC\textsuperscript{54}, molecular models of soot\textsuperscript{57}, and a secondary micellar aggregate of irregular shape built up by cholic acid molecules\textsuperscript{66}.

Another important result of this paper concerns the correct way of calculating density profiles relative to intrinsic interfaces, irrespective of whether they are macroscopically planar or not. Thus, here we proposed a Monte Carlo-based integration algorithm to estimate the volume elements in which points of the profile are calculated, in order to normalize them correctly. The issue of normalization with the volume elements in macroscopically flat fluid interfaces originates from the fact that these interfaces are rough on the molecular scale, namely, at the length scale of the calculated profiles. We clearly demonstrated that using this new normalization the artificial smearing of the intrinsic density profiles far from the intrinsic interface can be avoided.

Two computer programs that implement, respectively, an optimized version of ITIM and the new GITIM algorithm, as well as the calculation of intrinsic density and order parameters profiles, are made available free of charge at \url{http://www.gitim.eu/}. The programs are compatible with the trajectory and topology file formats of the Gromacs molecular simulation package\textsuperscript{46}.
Acknowledgements

M.S. acknowledges FP7-IDEAS-ERC grant DROEMU: “Droplets & Emulsions: Dynamics & Rheology” for financial support, and György Hantal for providing the soot structures. Part of this work has been done by M.S. at ICP, Stuttgart University. P.J. is grateful for financial support from the Hungarian OTKA foundation under Project Nr. 104234. S.K. is supported by FP7-IDEAS-ERC Grant PATCHYCOLLOIDS and RFBR Grant mol_a 12-02-31374. Simulation snapshots were made with VMD$^{68}$.

VII. APPENDIX

Here, following Ref. 69 we derive the expressions for the radius \( R \) and position \( \mathbf{r} = (x, y, z) \) of the center of the sphere which is touching four other ones, having given radii and center positions \( R_i \) and \( \mathbf{r}_i = (x_i, y_i, z_i) \) \((i = 1, 2, 3 \text{ or } 4)\), respectively. The conditions of touching can be expressed with the following nonlinear system of four equations:

\[
|\mathbf{r} - \mathbf{r}_i|^2 = (R + R_i)^2.
\]  

By subtracting one of them from the other three (without loss of generality we subtract the one with \( i = 1 \)), the quadratic term, \( \mathbf{r}^2 \), will be eliminated and the system Eq.(3) would become linear with respect to \( \mathbf{r} \):

\[
\mathbf{M}\mathbf{r} = \mathbf{s} - R\mathbf{d},
\]  

where the matrix \( \mathbf{M} \) and the vectors \( \mathbf{d} \) and \( \mathbf{s} \) are defined as

\[
\mathbf{M} = \begin{pmatrix}
\mathbf{r}_1 - \mathbf{r}_2 \\
\mathbf{r}_1 - \mathbf{r}_3 \\
\mathbf{r}_1 - \mathbf{r}_4
\end{pmatrix},
\mathbf{d} = \begin{pmatrix}
R_1 - R_2 \\
R_1 - R_3 \\
R_1 - R_4
\end{pmatrix},
\]

and

\[
\mathbf{s} = \frac{1}{2} \begin{pmatrix}
r_1^2 - r_2^2 - R_1^2 + R_2^2 \\
r_1^2 - r_3^2 - R_1^2 + R_3^2 \\
r_1^2 - r_4^2 - R_1^2 + R_4^2
\end{pmatrix}.
\]

Equation (4) has a unique solution if matrix \( \mathbf{M} \) is non-singular (the singularity of \( \mathbf{M} \) corresponds to the case when all 4 spheres are co-planar, which means that the unknown sphere
either does not exist, or is not unique):

\[ r = M^{-1}s - RM^{-1}d \equiv r_0 - Ru, \quad (7) \]

where \( M^{-1}s = r_0 \) and \( u = M^{-1}d \). Once Eq.(7) is substituted into the first of the constraints Eq.(3), it leads to the quadratic algebraic equation with respect to \( R \):

\[ (1 - |u|^2)R^2 + 2(R_1 - u \cdot v)R + (R_1^2 - |v|^2) = 0, \quad (8) \]

where \( v = r_1 - r_0 \). The solution of Eq. (8) can be found in the following form:

\[ R_{\pm} = \frac{-(R_1 - u \cdot v) \pm |R_1u + v|}{1 - |u|^2}. \quad (9) \]

If \( |u|^2 \) is not equal to unity (which corresponds to the case when the 4 spheres are tangential to one plane), then Eq.(9) provides two different solutions, and the positive one provides the radius \( R \) of the touching sphere as a function of the centre position \( r \). Eventually, the positions of their centres can be obtained by inserting \( R \) into Eq.(7). In the present manuscript in case of two possible solutions we choose the sphere with minimal radius.

\[ \text{Electronic address: sega@roma2.infn.it} \]


J. L. Bentley, Commun. ACM 18, 509 (1975).


Figures
FIG. 1: Left: example of the α-shapes algorithm on a set of points on the plane. The lines connecting the atoms represent the Delaunay triangulation (the triangles are labeled by numbers from 1 to 12). Solid lines mark triangles belonging to the α-complex, and dashed lines those which are not. The light-shaded atoms are those belonging to the α-shape, the border of the α-complex. Two atoms (in triangle 1) are outside the α-shape, and one (shared by triangles 9-12) is inside the α-shape. Right: schematic representation of the ITIM algorithm, applied to a single water molecule: the probe spheres (circles) are moved down the test lines (dashed lines) until they touch an atom.
FIG. 2: Simulation snapshot of a H$_2$O/CCl$_4$ system. The oxygen atoms at the interface between the H$_2$O phase (inner) and CCl$_4$ phase (outer) as recognized by the gitim algorithm are represented with an additional halo. Unconnected points belong to molecules which cross periodic boundary conditions.
FIG. 3: Average number of surface atoms identified by ITIM (squares) and GITIM (circles) as a function of the probe sphere radius.
FIG. 4: Water surface oxygen atoms in the H$_2$O/CCl$_4$ system in one simulation snapshot as recognized by GITIM exclusively (small spheres), ITIM exclusively (large spheres) or by both methods (sphere with halo).
FIG. 5: Intrinsic density profiles of water (curves on the left) and carbon tetrachloride (curves on the right) with respect to the water surface as computed with ITIM (solid curves) or with GITIM (dashed curves). The thin curves are computed using ITIM and the Monte Carlo normalization procedure described in Sec. IV.
FIG. 6: Right: schematic structure of a DPC molecule. Left: snapshot of a DPC micelle in water. Only the DPC constituents are shown for the sake of clarity. Atoms with a halo are those recognized by GITIM as surface ones.
FIG. 7: Upper panel: intrinsic density profiles of water (right) and of DPC (left). Lower panel: intrinsic profile of the orientational order parameters ($P_1$, solid line, $P_2$, dashed line). The vertical dashed lines marks the position of the interface.
FIG. 8: The $S_1^I$ soot model\textsuperscript{57} represented in section (right, triangulated surface) and in whole (left, wireframe) with the atoms identified by GITIM as surface ones highlighted using thicker, red elements. Besides surface atoms, also chemical bonds between surface atoms are highlighted, as well as five, six and seven membered rings (filled surfaces).
FIG. 9: Histograms of the atoms in the four soot models taken from Ref. 57. Each panel refers to a different structure (depicted with wireframe), and presents the distribution of all atoms (filled, darker area) and of surface atoms identified by GITIM (filled, lighter area), as a function of the distance from the center.
FIG. 10: Left: simulation snapshot of a secondary cholate micelle, with surface atoms highlighted. Right: the structure of the cholic acid molecule.
FIG. 11: Density profile of water (right) and cholic acid (left) in the secondary micelle.