

Self-assembly of escin molecules at the air-water interface

studied by molecular dynamics

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Abstract

Escin belongs to a large class of natural bio-surfactants, called saponins, which are present in more than 500 plant species. Saponins are applied in pharmaceutical, cosmetic, food and beverage industries, due to their variously expressed bioactivity and surface activity. In particular, Escin adsorption layers at the air-water interface exhibit unusually high surface elastic modulus (> 1100 mN/m) and high surface viscosity (ca. 130 N.s/m). The molecular origin of these unusual surface rheological properties is still unclear. We performed classical atomistic dynamics simulations of adsorbed neutral and ionized escin molecules to clarify their orientation and interactions on the water surface. The orientation and position of the escin molecules with respect to the interface, the intermolecular interactions, and the kinetics of molecular aggregation into surface clusters are characterized in detail. Significant differences in the behavior of the neutral and the charged escin molecules are observed. The neutral escin molecules rapidly assemble in a compact and stable surface cluster. This process is explained with the action of long-range attraction between the hydrophobic aglycones, combined with intermediate dipole-dipole attraction and short-range hydrogen bonds between the sugar residues in escin molecules. The same interactions are expected to control the viscoelastic properties of escin adsorption layers.

Keywords: saponin, escin, molecular dynamics, surface elasticity, adsorption.

Introduction

Saponins present a class of natural amphiphilic molecules, which contain a hydrophobic scaffold (called aglycone) and one to several hydrophilic oligosaccharide chains connected to the aglycone via glucoside bonds. Some saponins are used as essential components in vaccine adjuvants, while others exhibit anti-inflammatory, antimicrobial, anti-allergic, antiviral, cytotoxic, diuretic, hemolytic, and hypo-cholesterolemic activity [1-5]. The practical applications of saponins include also foam and emulsion stabilization in food, beverage and cosmetic products, cholesterol removal from foods (fats, milk, etc.), and micellar solubilization of vitamins and minerals in food additives [6-11].

To clarify the origin of various effects observed for saponin-containing systems and to optimize the applications of the latter, better understanding is needed for the relationship between the molecular structure, on the one hand, and the structure and properties of their micellar aggregates and adsorption layers, on the other hand [12,13]. Such an understanding can be significantly improved using the method of molecular dynamics (MD).

Escin is the main natural constituent of the saponin extract from the seeds of *Aesculus hippocastanum* (known also as *horse chestnut tree*) [14]. Efficient methods allow escin isolation and purification as a single molecular component [15] and it is sold as a market product. A number of bioactivity expressions of this saponin are detected, such as anti-inflammatory, blood vessel contracting and protecting properties [1]. Horse chestnut seed extract, containing around 20 wt % escin, is used in medicine for its venotonic effect, vascular protection, anti-inflammatory, and free radical scavenging properties.

This compound is also a surfactant with some unusual adsorption characteristics. Escin adsorption layers feature extremely high surface elastic modulus and very low gas permittivity, which is important for the properties of foams and emulsions stabilized by this natural surfactant [12,13]. These properties are sensitive to the pH of the water phase, because the sugar residue, attached to the aglycone, contains an ionizable carboxyl group, see Figure 1. For example, the surface elasticity decreases significantly upon pH increase, i.e., the adsorption layer formed from ionized escin molecules is less elastic than that composed of neutral ones [12]. The origin of this unusual surface behavior has not been clarified at the molecular level. Hence, escin is a bio- and surface-active substance, which deserves to be studied in more depth, given its multiple potential applications.

Several publications report MD simulations of saponin molecules [16-18]. However, this group of papers is focused mainly on the kinetics, self-assembly mechanism, and shape of micelles formed in bulk saponin solutions. In the study of Pedebos et al. [16,17] the stable conformations of two types of saponins in aqueous and in pyridine solutions are determined by unrestrained MD and shown to match well NOESY NMR measurements. One of these studies [17] reports for the first time the structure and conformation in aqueous solution of the saponin QS-21, extracted from *Quillaja saponaria* tree. This saponin is of particular interest, because it is used as essential component in the most efficient saponin-based vaccine adjuvants. A 3D atomistic model of QS-21, obtained by MD simulations, is proposed in the same study [17]. The MD modelling is combined with SAXS data to decipher the molecular conformation of QS-21 in the micelles and the forces leading to micellar growth upon concentration increase. In a separate study, Dai et al. [18] employ meso-MD and coarse-grained MD to discriminate the solubilization mechanisms of saikosaponin and Ginsenoside Ro (bioactive compound in ginseng). The molecules of Ginsenoside Ro form vesicles which, at low concentrations of saikosaponin, solubilize it in their surface layer, whereas mixed vesicles are obtained at higher saikosaponin concentrations.

Other works [19,20] report the results from MD simulations aimed at determining the populated conformations of saponins in solution and finding relationships between the molecular structure and certain bioactivity. Walkowicz et al. [19] study the saponin QS-21 and a series of its synthetic analogues and establish a crucial role of the conformation of the central glycosylic bond with the adjuvant activity. Lin and Wang [20] employ atomistic and coarse-grained MD to study the conformations of dioscin in water, decane, and water-decane, its interaction with cholesterol and its aggregation inside a lipid bilayer. Based on that, they propose a molecular mechanism for the hemolytic activity of this saponin, which consists of two steps: initial penetration of the saponin in the membrane and strong binding to cholesterol, followed by concentration-dependent curving of the membrane, subsequently leading to its destruction.

The MD method is suitable also for interpretation of the rheological characteristics of surfactant adsorption layers at interfaces, which is demonstrated on the example of the system 1,2-ethanebis(dimethyl dodecyl ammonium bromide- $C_{12}C_2C_{12}\cdot_2Br$) at the air/water interface [21]. It is determined experimentally that the dilatational modulus of $C_{12}C_2C_{12}\cdot_2Br$ is higher than that of DTAB

(Dodecyltrimethylammonium bromide), which is related to an interfacial force acting in a way as to compensate the external perturbation of the adsorption layer. The authors explain this effect by MD simulations, which show that the difference between the dilatational moduli of the two types of surfactants is due to the higher concentration of Br^- counter-ions in the vicinity of the nitrogen atoms of $\text{C}_{12}\text{C}_2\text{C}_{12}\cdot 2\text{Br}$ than around those of DTAB. These ions reduce the repulsion between the hydrophilic groups and enhance the adsorption of $\text{C}_{12}\text{C}_2\text{C}_{12}\cdot 2\text{Br}$ at the interface.

This literature overview shows that there are no data from MD simulations of saponin adsorption layers. Hence, a theoretical study in this direction would provide new insights into the behavior of these peculiar molecular systems. The current work presents a detailed classical atomistic MD description of the properties of escin molecules (single and in a cluster) adsorbed at the vacuum-water interface. The main goal is to elucidate the molecular characteristics and the intermolecular interactions for the neutral (ESC) and for the anionic (ESCA) forms of this saponin. A relation to the aforementioned rheological properties of the escin adsorption layers is sought, too.

Molecular Models and Computational Method

Escin (Figure 1) is selected for this study because highly purified, single-molecular-species extracts are available and can be studied experimentally, which allows construction of molecular models mimicking as close as possible the experimental conditions. In addition, escin is one of the saponins with highest surface elasticity [12, 26, 27].

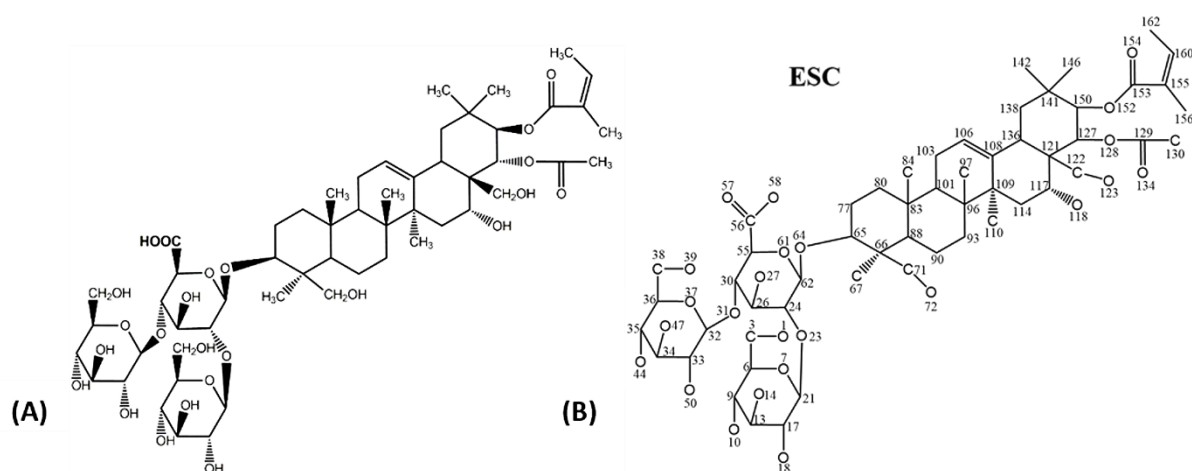


Figure 1. (A) Chemical formula of escin with cyclic hydrogens omitted for clarity. The carboxyl group containing a proton, which is removed in the anionic form, is denoted in bold. (B) Atom numbering of the non-hydrogen atoms in the neutral escin.

To characterize the behavior of escin at the surface of water, MD simulations of four types of escin models are carried out (Figure 2): (1) a single nonionic surfactant; (2) a single anionic molecule, which is formed upon ionization of the carboxyl group present in one of the sugar residues, (3) 16 non-ionized molecules, and (4) 16 anionic molecules.

The initial geometry and the molecular mechanical parameterization of escin are described in the Supporting Information (Figures S1, S2; Tables S1, S2). In summary, the obtained atomic charges are added to the force field AMBER99 [22,23] which is used to simulate escin in combination with the model TIP3P [24] for the water molecules. The optimized geometry escin-1 (Figure S1) is taken as the starting structure for the atomistic MD simulations. In the first type of models, one neutral escin molecule is placed at the vacuum/water interface, its hydrophilic part being immersed within an ensemble of 12 326 explicit water molecules (Figure 2A). This model corresponds to a solution with bulk concentration of 4.5 mM, wherefrom the molecule had adsorbed onto the air-water interface. The area per molecule in such a system would be 52 nm². In this model, the adsorbed escin molecules would not interact with each other. The model contains also 2 Na⁺ and 2 Cl⁻ ions to mimic the experimental conditions with 10 mM NaCl added to the solution. The periodic box sizes are: $x = 7.18$ nm, $y = 7.18$ nm, $z = 17.18$ nm. The second type of model system encompasses one escin anion and is built on the basis of its neutral counterpart by just removing the carboxylate proton (marked in bold in Figure 1A) and adding one more Na⁺ as a neutralizing counterion.

The neutral and anionic models with 16 escin molecules (Figure 2B) also have identical starting configuration. They are built by translating the corresponding single-molecule model system into the XY plane, resulting in a regular initial spacing of the escin molecules at the surface. The sizes of the periodic boxes encompassing 16 hydrated escins are: $x = 16$ nm, $y = 6.4$ nm, $z = 17.18$ nm. In these larger models, the hydrophilic parts of the 16 surfactant molecules are immersed in a pool of 23 560 water molecules. The area per molecule in such an adsorption layer is 6.4 nm². This reproduces the conditions of a dilute adsorption layer, i.e., the model describes the initial stage of the adsorption experiments. Hence, the results presented in the current study should be regarded as the first step in unveiling the surface characteristics of escin adsorption layers. The simulations of the 16-molecule models are carried out also in the presence of 10 mM NaCl, which corresponds to 4 Na⁺ and 4 Cl⁻ ions

in the boxes. In the model with escin anions, additional 16 Na⁺ are present to neutralize the charge of the deprotonated carboxylic groups.

Periodic boundary conditions are applied in all simulations. 5 nm vacuum space is included along the Z axis (which is normal to the interface) both above and below the water layer to create the water-vacuum interface (Figure 2). The MD runs are at constant temperature of 293 K maintained by a Berendsen thermostat [25]. The periodic box volume and the number of molecules are also fixed. These conditions and concentrations are chosen to mimic the experimental studies carried out by Pasureva et al. [12] and by Golemanov et al. [26,27].

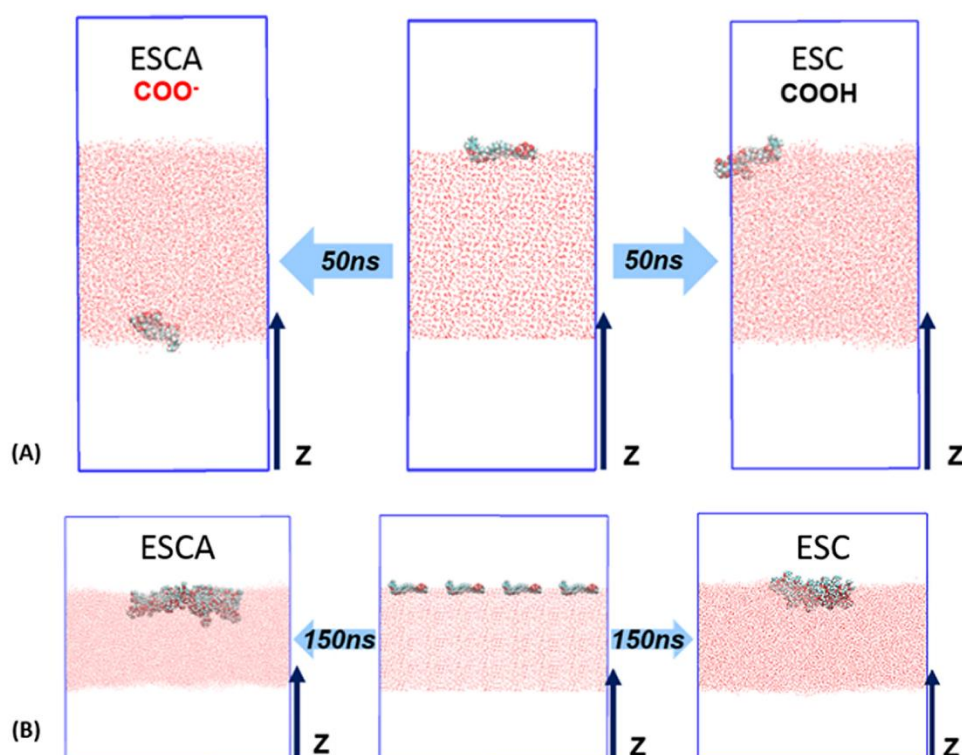


Figure 2. Simulated molecular models of: (A) a single-surfactant and (B) 16 molecules at the vacuum/water interface. The initial configurations (middle) are shown together with the final snapshots from the MD simulations of neutral (right) and anionic (left) forms of escin molecules.

The Lennard-Jones non-bonded potential is truncated at a cut-off distance of 1 nm with a switch function activated at 0.8 nm. Electrostatic interactions are evaluated in the monopole approximation with the method PME [28-30]. The direct part of the sum is truncated at 1.2 nm with a switch function

turned on at 1 nm. The time step for integration of the equations of motion is 2 fs and the leap-frog algorithm is used. The lengths of all H-containing bonds are constrained with LINCS [31] in the surfactants and with SETTLE [32] in the water molecules.

A standard protocol is employed for the MD simulations – energy minimization, followed by heating and equilibration of the systems. The evolution of the energy and temperature is followed to verify that the equilibrium has been reached. After that, production runs are generated with length of 50 ns for the single-surfactant models and of 150 ns for the 16-surfactant systems. In all trajectories the frames are recorded at intervals of 1 ps, i.e., the statistical analysis reported below encompasses 50 000 or 150 000 data points, respectively, unless otherwise specified.

To track the aggregation kinetics of escin in the 16-surfactant models, cluster analysis is performed with a limiting distance for similarity of 0.028 nm. Only the coordinates of the escin molecules are included in this analysis.

The simulations of the single-surfactant models are carried out with Gromacs 4.5.2 [33] and those of the 16-surfactant ones – with Gromacs 4.6.5 [34]. The package VMD 1.9.1 is used for visualization [35].

Results and Discussion

Surface self-assembly of escin

It is revealing to trace first the trajectories of the molecules in the 16-surfactant models. The cluster analysis of the *neutral* surfactant model (Figure 3A) shows that up to 3 ns various short-lived clusters are formed, with their number decreasing upon progress of the simulation. After 3 ns, only two stable clusters are formed, each of them containing 8 escin molecules. At ca. 17 ns these clusters merge to yield a single 16-surfactant cluster, which remains stable till the end of the simulation at 150 ns. This is a very important outcome of our simulations because the spontaneous self-assembly of the adsorbed molecules indicates that strong, long-range attractive forces act between them. These forces lead to fast aggregation at the surface and prevent the formed cluster from breaking apart, which is in agreement with the experimentally registered fast formation of monolayers with high surface elasticity [12, 26].

Spontaneous self-assembly is observed also for the *anionic* escin (Figure 3B). However, its molecules aggregate much more slowly, compared to the neutral surfactant – a relatively stable 16-molecule cluster forms at 59 ns. Moreover, up to 150 ns this cluster often breaks into two smaller clusters of 12 and 4 surfactant molecules, which then re-assemble. This slower aggregation of the anionic form agrees qualitatively with the experimental finding that the energy of attraction between the adsorbed escin molecules is smaller in magnitude at pH = 8 when the escin molecules are ionized, as compared to pH = 4 when the escin molecules are neutral (unpublished results). Figure 2B shows the final frames from the 150 ns simulations. It is evident that the surfactants are aggregated together, leaving the larger part of the surface uncovered, irrespective of the ionization state of escin.

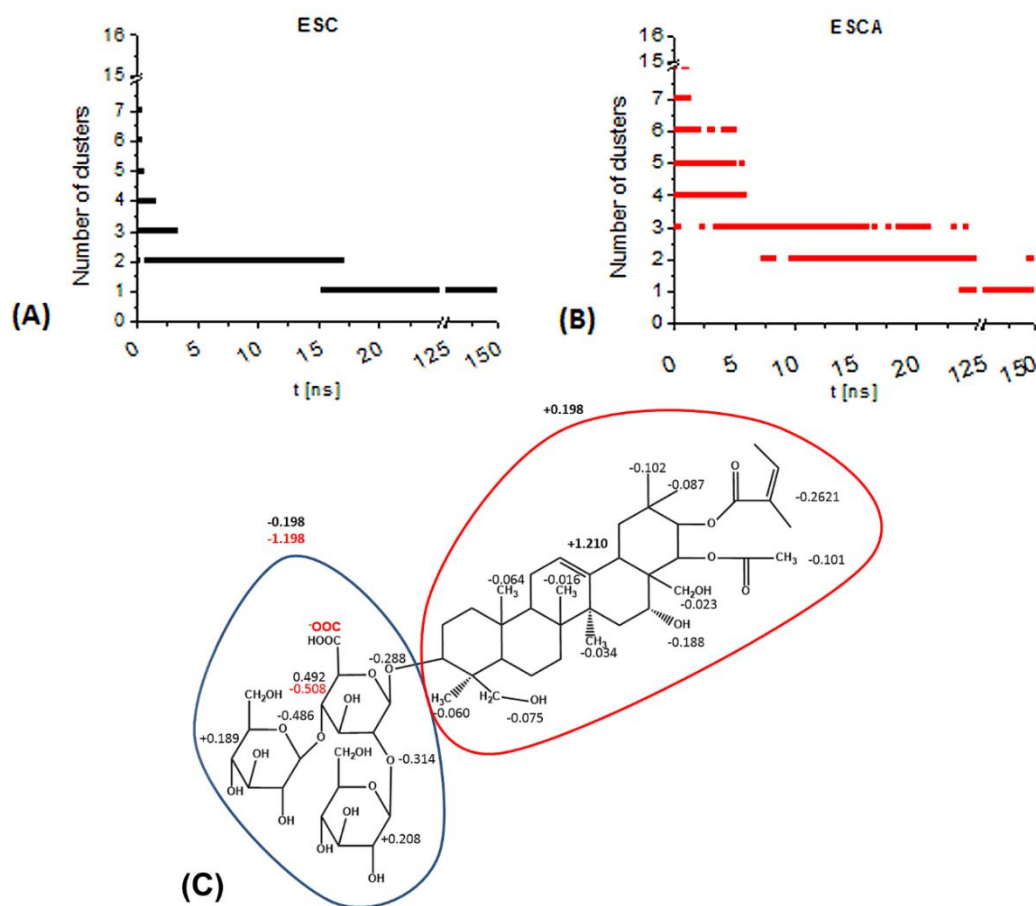


Figure 3. Evolution of the number of escin clusters in the 150 ns MD simulations of the 16-surfactant models of the (A) neutral and (B) ionized form; (C) RESP group charges of the different molecular fragments of neutral and anionic escin; the values, which do not coincide in the neutral and ionic form, are written in red.

The origin of the long-range attractive force driving cluster formation could be sought in the non-bonded interactions between the aglycone molecular fragments, which are not immersed in water (see the next section for the relative position of these fragments with respect to the air-water interface). To check this assumption, we calculated the group charges for the various parts of ESC and ESCA molecules – see Figure 3C. As expected, the calculated group charges are unequally distributed along the escin molecules. The most negative charges belong to the oxygen-containing functional groups in the hydrophilic part of the molecule (immersed in water) but the residues comprising the carbon-carbon double bond, the hydroxyl group, and the acetate residue in the aglycone are also negatively charged. In other words, the entire molecular periphery appears negatively charged, whereas the cyclic fragments in the aglycone center bear a significant positive charge – see Figure 3C.

Such charge distribution implies the appearance of electrostatic attraction between the oppositely charged molecular fragments in the neighboring surfactant molecules, when these molecules are properly oriented with respect to each other. This attraction is a kind of dipole-dipole intermolecular interaction (more precisely, multipole-multipole interaction) and could explain the fast formation of the molecular clusters at the interface. This picture implies also that the molecules in the forming cluster should assemble in a preferred mutual orientation, which maximizes this intermolecular attraction. In addition, upon closer approach of the aglycones, they interact via dispersion (London) type of attractive forces, which should be rather significant between such large molecular fragments. Rough estimates show that the total intermolecular energy between the neighboring aglycones could be well above $10 kT$ (where kT is thermal energy) if the aglycones are properly oriented with respect to each other.

For brevity, hereafter we denote the latter two types of forces (multipole-multipole and dispersion) as *van der Waals forces* between the aglycones. Note that the respective forces between the molecular fragments immersed in water are expected to be orders of magnitude smaller, due to the high dielectric constant of the water medium (which decreases the multipole-multipole interactions in accordance with Coulomb's law) and the reduced Hamaker constant for interaction across aqueous medium, as compared to the interaction across vacuum. Therefore, in the following discussion we consider negligible the van der Waals interactions between the escin molecular fragments immersed in the aqueous phase in the initial stages of cluster formation.

Expectedly, after deprotonation of the carboxyl group, the negative charge in the anionic ESCA molecules is localized mainly on the carboxylate anion with part of the charge transferred to its adjacent cycle (see the charges shown in red for ESCA in Figure 3C). This results in a large negative charge of the hydrophilic part of the anion, which is immersed in water, whereas the aglycone part bears virtually the same distribution of charges as that in the neutral ESC. This comparison between ESC and ESCA shows that a long-range van der Waals attraction between the aglycones is expected for both types of molecules – neutral and ionized. However, it would be superimposed over a long-range electrostatic repulsion between the charged hydrophilic fragments in the ESCA molecules. The latter electrostatic repulsion explains the two most important features in the evolution of the clusters of ionized ESCA molecules, namely, the slower aggregation kinetics and the lower stability of the largest cluster.

After the strong aggregation affinity of escin is revealed from the simulations, in the following sections we characterize the positions and orientation of the escin molecules with respect to the water surface and inside the clusters. The *shorter-range* interactions between the clustered escin molecules are also quantified. In the last section we combine all this information to explain the molecular processes, observed in the simulations, and discuss their possible relation to the experimental results about the rheological properties of escin adsorption layers.

Position of the amphiphiles relative to the interface

Analysis of the density profiles along the Z axis of the models provided the characteristic layer thicknesses in direction normal to the interface. The mass densities of the whole systems, escin, and water (Figures 4 and S3) are calculated for all models by block averaging the production runs in 10-ns sections. This resulted in mean standard error of the estimates of the Escin density of 0.01 kg.m⁻³ and 0.13 kg.m⁻³ for the small neutral and ionized models and of 0.06 kg.m⁻³ and 0.04 kg.m⁻³ for the large models, respectively.

The following quantities are evaluated from the density profiles: (1) *most probable* thickness of the escin cluster, as the FWHM of the Gaussian fit of ESC/ESCA peaks, and (2) *maximum thickness* of the escin cluster, as the difference of the Z-coordinates corresponding to the escin density at *three standard deviations* to the left and to the right from the Gaussian peak maximum (for sensitivity of the

estimates to the position of fixing this minimum density, refer to Tables S3 and S4 of the Supporting Information).

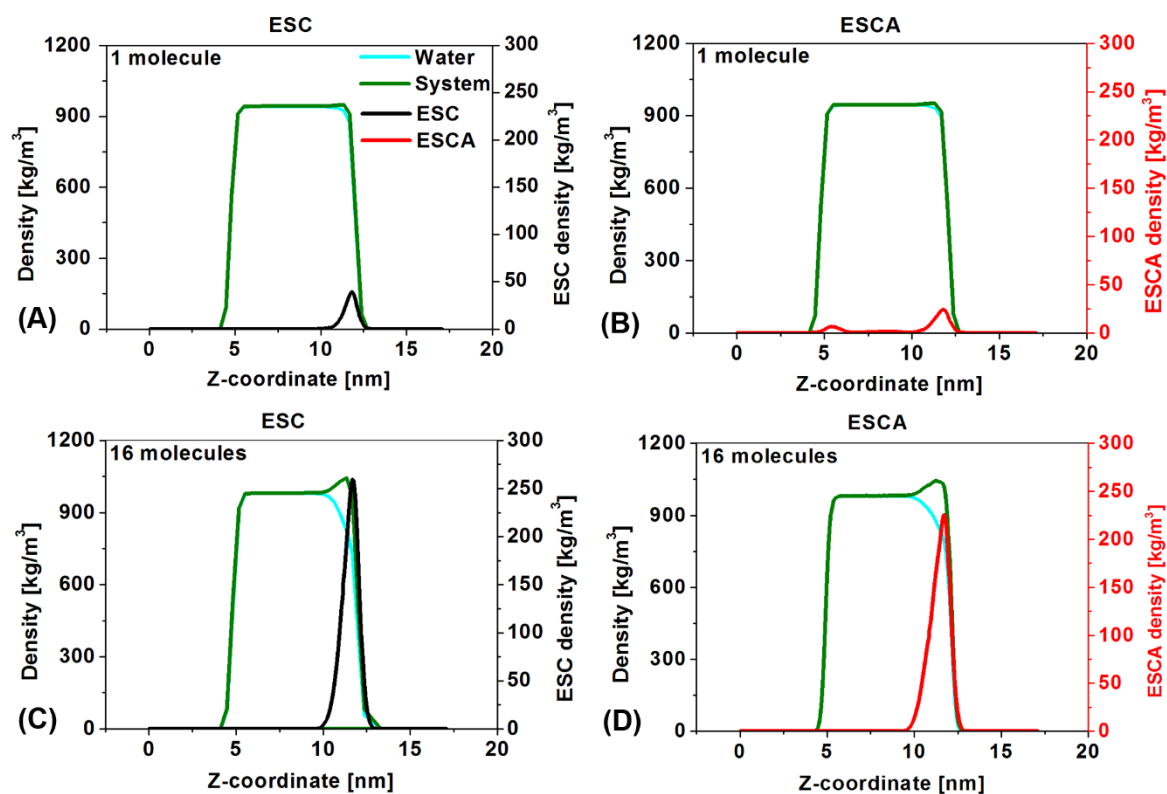


Figure 4. Density profiles in direction normal to the interface for the entire system, water, and escin in the models of: (A) single ESC, (B) single ESCA, (C) 16 ESC, and (D) 16 ESCA; the left ordinate is for the density of the system and water and the right one – for that of the surfactants.

The density profiles of ESC and ESCA differ significantly. The peak of neutral escin is located close to one of the water surfaces (where the molecules were initially placed), irrespective of the model size (Figures 4 and S3, left), which is a clear indication of the high surface activity and low hydrophilicity of the neutral amphiphile. The same peak for the ionic form is shifted towards the interior of the aqueous layer (Figures 4 and S3, right), which shows that the anionic surfactant is immersed deeper into water. Moreover, in the single-molecule model of ESCA, a second density peak for the surfactant is present, mirroring the main one, but at the opposite water surface (Figure 4, top right). This second peak is due to dissolution of the ESCA molecule in the water, followed by transfer to the second interface, as confirmed by visual inspection of the MD trajectory. This behavior outlines the anionic form of escin as being much more water-soluble, which is in agreement with the experimental results indicating that the anionic form of escin is more soluble than the neutral one [13].

The *most probable* thickness (projection along the normal to the interface) of the single ESC molecule is 0.88 nm and that of ESCA: 1.05 nm. The respective values for the 16-surfactant models are: 1.08 nm for ESC and 1.27 nm for ESCA. As expected, these thicknesses are larger for the clusters. This increase can be due to thermal fluctuations of the molecules in direction normal to the interface and/or to certain intermolecular interactions, which shift the individual molecules along the normal to the interface, so that the escin molecules adjust their positions with respect to the neighboring ones. The latter effect is expected to be more pronounced in a cluster of charged molecules, as it reduces the electrostatic repulsion between the neighboring molecules. ESCA has always larger thickness than ESC, irrespective of the model size, and this is in line with its higher hydrophilicity – the latter allows the molecules to submerge deeper into water and, hence, to occupy a broader interface region.

The *maximum* thickness (projection along the normal to the interface) of the surfactants in normal direction is as follows: 2.24 nm for single ESC, 2.68 nm for single ESCA, 2.75 nm for the ESC cluster, and 3.24 nm for the ESCA cluster. Again, the anions have larger thickness. This is an indirect reflection also of the packing of the molecules in the cluster, which is looser for the anions compared to the neutral surfactants. Also, the thickness increases when going from the single- to the 16-molecules models, which points to fluctuations/shifts in the positions of the molecules along the Z axis in the clusters.

To check the latter assumption, additional analysis is performed. It uses a reference atom in the surfactant molecule, which marks the location of these molecules with respect to the interface. The Z coordinate of the terminal carbon atom in the aglycone, connecting it to the sugar residue (denoted in green in Figure 5A) is selected for this purpose. The reason is that the aglycone is relatively rigid and the position of this specific atom can be used to represent the vertical displacement of the entire aglycone.

The evolution of this Z coordinate for each of the 16 molecules in the cluster is recorded for the last 50 ns of the trajectories (Figure S4) and a summarized distribution is provided in Figure 5B. One sees that the Z coordinate fluctuates on average in a range of 0.7÷0.8 nm, with some molecules featuring dispersion of the coordinate even above 1 nm. This result, combined with the significant width of the peaks in the histogram, confirms the assumed substantial mobility of the surfactants

normal to the interface, which is especially pronounced for the anions. To quantify precisely the position of the aglycone with respect to the air-water interface, the equimolecular dividing surface for water (EDS, denoted in blue on Figure 5B) is determined as the Z coordinate in the density profile of water (Figure 4) corresponding to water mass density of 500 kg/m^3 . The Z coordinate of the reference escin atom is almost exclusively below the EDS, by ca. 0.5 to 1 nm. This means that the aglycone is partially immersed in water, which is in line with the known significant solubility of escin in water [13].

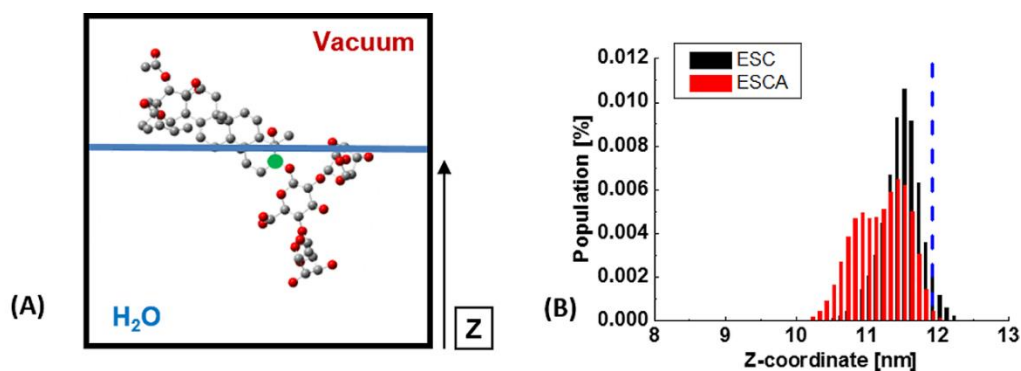


Figure 5. (A) Schematic representation of the selected reference aglycone carbon atom of escin and (B) distribution of its Z coordinate in the last 50 ns of the simulations of the 16-surfactant models; the blue dashed line denotes the equimolecular dividing surface.

The neutral and the anionic forms differ in one more aspect. On one hand, the peak for the Z coordinates of ESCA is shifted towards the water phase, compared to that of ESC. This is in accordance with the analysis of the density profiles, which outlined ESCA as more hydrophilic. On the other hand, the distribution of ESC is unimodal, whereas that of ESCA is bimodal, see Figure 5B. This signifies that the aglycone of the ionic form has two preferred locations with respect to the interface, which may stem from the attempt of the anions to reduce the electrostatic repulsion between the neighboring molecules by displacing approximately half of them in normal direction.

The molecular *length* of the two forms of escin is estimated from the single-surfactant models as another characteristic structural parameter (Figures S5 and S6). The data show that the neutral molecule is slightly longer (2.22 nm) than the charged one (2.11 nm) and the length of the more stretched ESC molecule fluctuates in a narrower range. These lengths would coincide with the thickness of the escin layer if the surfactant molecules were perpendicularly aligned to the interface.

However, the lengths of both the neutral and the charged forms are much larger than this thickness, which may happen if the amphiphile molecules are tilted with respect to the interface (Figure 6). Therefore, evaluation of this tilt and some other characteristic angles is done next.

Interfacial orientation and intramolecular conformation of the surfactants

To assess the surface orientation of escin, the tilt of the aglycone (Figure 6A and S6), which is the most rigid part of the surfactant, is used. The tilt is evaluated as the angle closed between the Z axis and a vector interconnecting the two terminal carbon atoms of the aglycone cyclic part (see Figure 6A). This angle is calculated for each frame of the MD trajectories and the results are shown in Figure 6B-C.

The plot for single surfactant molecules in Figure 6B reveals that the angle is significantly smaller than 180° , i.e., the molecules are not perpendicular to the surface, which is in line with the results described in the previous section. The average value of the angle is similar for the two escin forms – 110° for ESC and 116° for ESCA, both profiles featuring fluctuations of ca. $\pm 15^\circ$. Rarely, both surfactants reach an angle of 180° , which corresponds to normal alignment relative to the surface. Much more frequently the aglycones are oriented parallel to the interface, i.e., at angles close to 90° .

The behavior of the *ionic* form during the last 10 ns is interesting, see Figure 6B. There, a seemingly drastic variation of the tilt is witnessed. This is actually due to the transfer of the anion to the opposite vacuum-water interface, as discussed above. Visual inspection of the trajectory shows that the ‘change’ of the tilt to ca. 65° is just a reorientation of ESCA at the second interface and in fact corresponds to the same alignment of the hydrophobic fragment with respect to this second interface. The difference in the angle values comes from reversal of the aglycone vector direction relative to the Z axis ($65^\circ = 180^\circ - 115^\circ$).

The evolution of the angle for all molecules in the 16-surfactant models is similar. Therefore, an illustrative curve for just one of these molecules is presented in Figure 6C. The tilt of the amphiphiles in the cluster practically coincides with that of the single surfactants. Hence, it can be concluded that this is the preferred orientation of the aglycone with respect to the surface, irrespective of the presence of neighboring surfactant molecules. Angles close to 90° are achieved also in the cluster, although seldom, with fluctuations of the angle similar to those in the single-surfactant systems. However,

angles of 180° are not populated in the cluster models. This is most probably due to the fact that, if the molecules align perpendicularly to the interface, their bulky sugar residues would disturb unfavorably the interaction with the neighboring surfactants because of steric hindrance. Hence, such orientation does not occur and this is the only effect on the tilt that stems from the cluster formation.

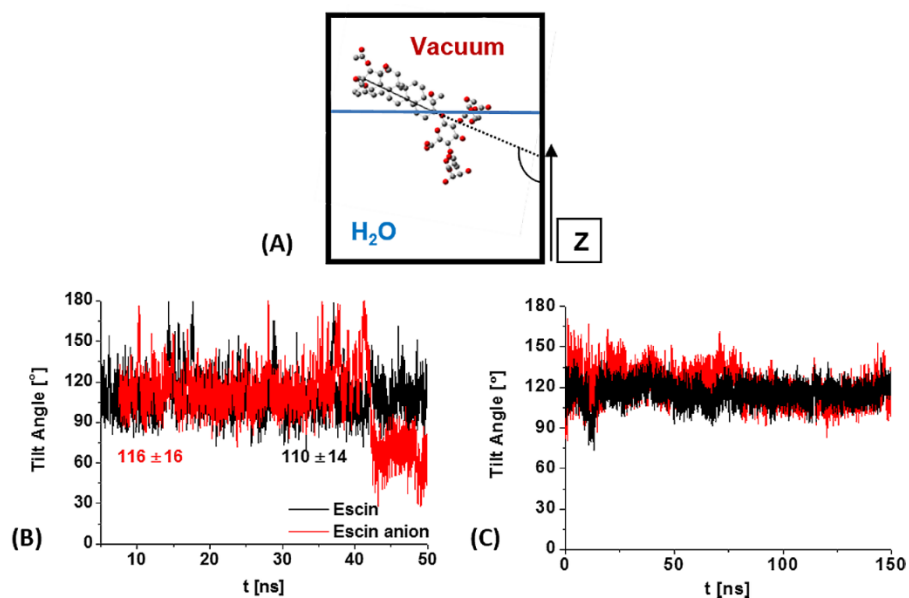


Figure 6. (A) Schematic presentation of the angle closed between the aglycone of escin and the Z axis; its evolution during the MD simulations of: (B) the single-surfactant and (C) the 16-surfactant models.

The main conclusion from the tilt angle analysis is that the preferred orientation of the aglycone relative to the surface is a robust characteristic, which is not affected by the presence of neighbors or by ionization of the escin at the studied low surface coverage.

The orientation of the sugar residues with respect to the aglycone is also analyzed to provide a comprehensive description of the amphiphile intramolecular conformation. The results from this analysis are presented in the Supporting Information, Figure S7. In summary, one branch of the sugar residue closes an angle of ca. 160° with the aglycone, independent of the charge state. The second sugar-aglycone angle is smaller and fluctuates less when it is close to 120° , while it is more flexible when it is being smaller than 120° . The hydrophilic part of the surfactant as a whole is rather mobile and undergoes spontaneous conformational changes at the studied nanosecond time scale.

Mutual orientation of the escin molecules within the clusters

The 16-surfactant systems allow us to analyze also the mutual orientation of the molecules in a cluster. The last 50 ns from the MD trajectories are used to quantify the mutual alignment of the amphiphiles because this orientation is already established in this last part of the simulations. The mutual organization of the neighboring surfactants therein is an indicator for the structure of escin surface domains/layers at low surface coverage. Therefore, the first step of the analysis is the identification of the closest-neighbor surfactants, which we denote as actual neighbors of each escin, in the neutral and anionic clusters (the procedure is described in the Supporting Information, Figure S8).

Figure 7 shows a histogram of the distribution of the number of actual neighbors of ESC and ESCA. Significant differences between the two systems are observed. The most frequent number of ESC actual neighbors is 5 or 6, while those of ESCA are only 2. The larger number of ESC neighbors indicates much better packing of the neutral molecules, stabilized by strong intermolecular attraction. In contrast, the small number of ESCA neighbors implies a looser packing, caused by the electrostatic repulsion between the ionized molecules.

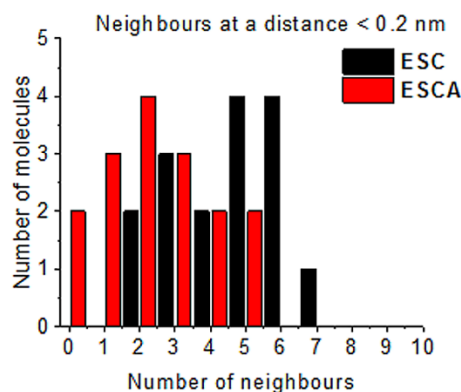


Figure 7. Number of neutral and anionic surfactants having a certain number of actual neighbors during the last 50 ns of the 16-surfactant simulations.

The average separation of all possible surfactant pairs in the clusters is determined from analysis of the minimum interatomic distance (the procedure is described in the Supporting Information, Figure S8) and the outcome is summarized in Table S5.

As can be seen, the number of ESC surfactant pairs at a distance less than 0.2 nm is much larger than that of ESCA, 25 vs. 13, which is in agreement with the larger number of actual neighbors for ESC (Figure 7). On the other hand, the number of pairs, which changed their minimum distance around 0.2 nm (i.e. those in which the surfactant molecules rearranged during the last 50 ns of simulation), is similar for ESC and ESCA (11 and 10, respectively). This shows that both the neutral and the charged surfactant molecules undergo some restructuring inside the clusters. The more remote surfactant pairs, which have intermediate separation between 0.2 nm and 0.6 nm, are again much more for ESC, 17 vs. 6. These may be regarded as second neighbors indicative of liquid-type long range ordering, which is present only in the neutral cluster. In contrast, the remote pairs (separated by more than 0.6 nm) are much more for the ionized surfactants, 91 vs. 67, signifying the looser packing in the cluster of ESCA.

Hydration of the escin clusters by water molecules

The intra- and inter-surfactant hydrogen bonds (H-bonds) within the neutral and ionized clusters are studied next to clarify the role of these bonds for the observed packing. The analysis of the entire clusters, including also the hydrogen bonding between escin and water, is given in the Supporting Information (Figures S9 and S10). It discloses that the number of escin-escin hydrogen bonds (H-bonds) increases during the MD simulations, whereas that of escin-water H-bonds decreases. The latter processes are slower in the anionic system. To explain this, it may be assumed that a larger number of water molecules protrude among the ionized surfactants to screen the electrostatic repulsion between the charged hydrophilic groups, thus reducing also the overall strength of interaction between the ESCA molecules. This would result in looser molecular packing and lower surface elasticity, in conjunction with the experimental findings [12].

To verify the above hypothesis, an additional analysis is performed to determine the number of water molecules incorporated inside the surfactant clusters. For this purpose, the mass density profiles of water and escin in normal direction are generated (Figure 8) for a subset of structures (extracted at intervals of 5 ns from the last 50 ns of the MD trajectories), limiting the water profile to a distance less than 0.4 nm apart from the surfactants, which corresponds to including the water molecules from the first solvation shell only.

The plot in Figure 8 shows that the density profile of ESC (black curve) is narrower than that of ESCA (red curve), which confirms the tighter packing of the neutral molecules. Moreover, the peak of the anions is shifted towards water. Taking into account this shift and the position of the equimolecular dividing surface (EDS, the green line), the more expressed hydrophilicity of ESCA is confirmed, because a much smaller fraction of the anions can be found above the EDS, i.e., in the gas phase. The water profiles of the two systems are also different. The area encompassed by the dark blue water curve (177 units), corresponding to the water present in the ESC cluster, is significantly smaller than that below the light blue water curve from the ESCA system (239 units), which proves that there is much more water incorporated among the anionic surfactants, including also the space among the aglycones. This is corroborated by the larger number of water molecules (Table S6 of the SI) present inside the anionic clusters, which confirms the presence of more hydrating water molecules.

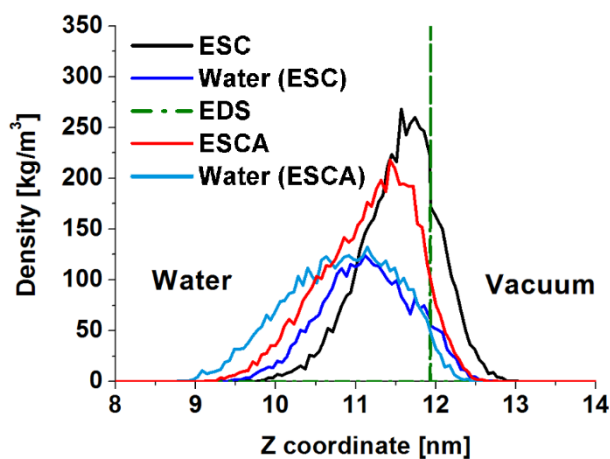


Figure 8. Mass density profiles of escin and water normal to the interface; the water profiles include only the water molecules, located at a distance < 0.4 nm from escin molecules (first solvation shell); the averaging encompasses a set of 10 frames spaced at 5 ns, taken from the final 50 ns simulations of the neutral and ionized 16-surfactant systems.

Interactions between the escin molecules

The analysis of the total number of escin-escin hydrogen bonds (Figure S9) does not yield an unequivocal average number of hydrogen bonds per surfactant because the molecules have varying number of neighbors. Furthermore, most of the escin molecules are at the boundary of the clusters and, hence, do not form the maximum possible hydrogen bonds with their neighbors. From experimental perspective, it can be expected that the elasticity of the escin layers would be related to the interactions

between well packed molecules, like those in the core of the cluster. These reasons prompted us to perform a more in-depth analysis of the local hydrogen bonding of the actual neighboring escin-escin pairs in the cluster core.

The results for two representative molecular pairs of this type (one pair from the neutral system and one pair from the ionized system) are shown in Figure 9. A representative geometry of the two pairs is displayed in Figure 9A showing that the two non-charged ESC molecules are very well packed, while the aglycones of the two charged ESCA molecules are not aligned well with respect to each other. The two molecules in each pair have minimum interatomic distance close to 0.2 nm during the entire simulation, combined with a significant number of intermolecular contacts (especially for the ESC-ESC pair), i.e., they are actual permanent neighbors. The ESC molecules in the neutral pair have ≈ 5 actual neighbors-per-molecule on average. For the charged paired ESCA molecules, the average number of neighbors is ≈ 1.5 per molecule.

One of the surfactants in the *neutral pair* forms more than 10 H-bonds on average with its five neighboring molecules, while the other surfactant – less than 5 H-bonds with its five neighbors (Figure 9B, left). Most of these H-bonds are strong (Figure 9C, left) – with length around 0.19 nm (cf. the Supporting Information). The H-bond length distribution of both molecules, however, contains an additional peak, centered at ca. 0.32 nm, especially intensive for one of the molecules in this pair. Such a characteristic length is too large to correspond to a typical H-bond. On the other hand, these forces exhibit strong angular dependence, which is a characteristic feature of the H-bonds. Therefore, these longer-range attractive forces have an intermediate nature between the classical H-bonds and the strong dipole-dipole interactions between the permanent dipoles of the interacting –OH groups in the molecules [36]. These H-bonds are formed when an appropriate steric match of the molecules is realized, which favors the proper orientation and the intermediate distance characterizing them [36]. They take place when the chemical structure of the particular molecules allows both appropriate molecular packing and suitable orientation of the groups involved in these bonds [36]. For convenience, hereafter we call these forces either “long-range H-bonds” or “specific dipole-dipole interactions”, bearing in mind their intermediate nature.

These intermediate forces seem to play a role for the stabilization of the two neutral ESC molecules in the pair. One of the molecules orients its proton-accepting and proton-donating groups in

a way, allowing the formation of a maximum number of ‘classical’ H-bonds with the other neighbors, while the second molecule in the pair maximizes these dipole-dipole interactions, at the expense of the classical H-bonds. To check this hypothesis, the H-bonds just between these two molecules are analyzed (Figure 9D,E). The data (the black columns) in Figure 9D show that ESC1 and ESC2 are connected by 3 H-bonds on average. Comparing this number to the one with all their neighbors, Figure 9B, it becomes evident that unequal number of H-bonds is present between each of these two molecules and its neighbors. From the 10 H-bonds ESC1 forms with its neighbors, 3 are with ESC2 and the remaining 7 – with the other four neighbors. Respectively, from the total of 4 H-bonds of ESC2, 3 are with ESC1 and only 1 bond with the other four neighbors. The bond length distribution of the three H-bonds formed between ESC1 and ESC2 (Figure 9E, black columns) is also non-standard. The peak at 0.19 nm has lower intensity, while the peak representing the longer-range dipole-dipole attraction is very intensive. Thus, we conclude that this additional dipole-dipole attraction between the sugar residues of two neutral neighboring escin molecules stabilizes the pair, beside the classical short-range H-bonds.

Closer look at the simulated molecules showed that the latter H-bonds involve several hydroxyl groups in the two terminal sugar residues of the neighboring escin molecules (see Figure S11). The two classical H-bonds involve the same terminal H-atom in the $-\text{CH}_2\text{OH}$ group of the ESC1 molecule (attached covalently to O-atom denoted by number 1 in Figure 1B). Around 30 % of the last 50 ns of the simulations, this H-atom is involved in a H-bond with mean distance of around 0.19 nm with the O-atom number 10 in the neighboring molecule ESC2. The energy of the H-bonds of such length is known to be in the range between 17 and 58 kJ/mol (7 to 22 kT per bond) [36]. Taking into account that this H-bond is active around 30 % of the time only, one could estimate its contribution into the molecular pair stabilization to be between 2 and 7 kT (around 4.5 kT on average). The same H-atom was involved around 10 % of the simulation time in another H-bond with O-atom number 14, belonging to another $-\text{OH}$ group in the sugar ring of ESC2. This gives an additional energy of attraction between the two molecules of 0.7 to 2.2 kT (around 1.5 kT on average). It seems that this H-atom flips between the two oxygen atoms in the neighboring ESC2 molecule by a minute conformational change of the $-\text{CH}_2\text{OH}$ group it belongs to. In total, these two H-bonds contribute ≈ 6 kT to the stabilization of this pair of escin molecules.

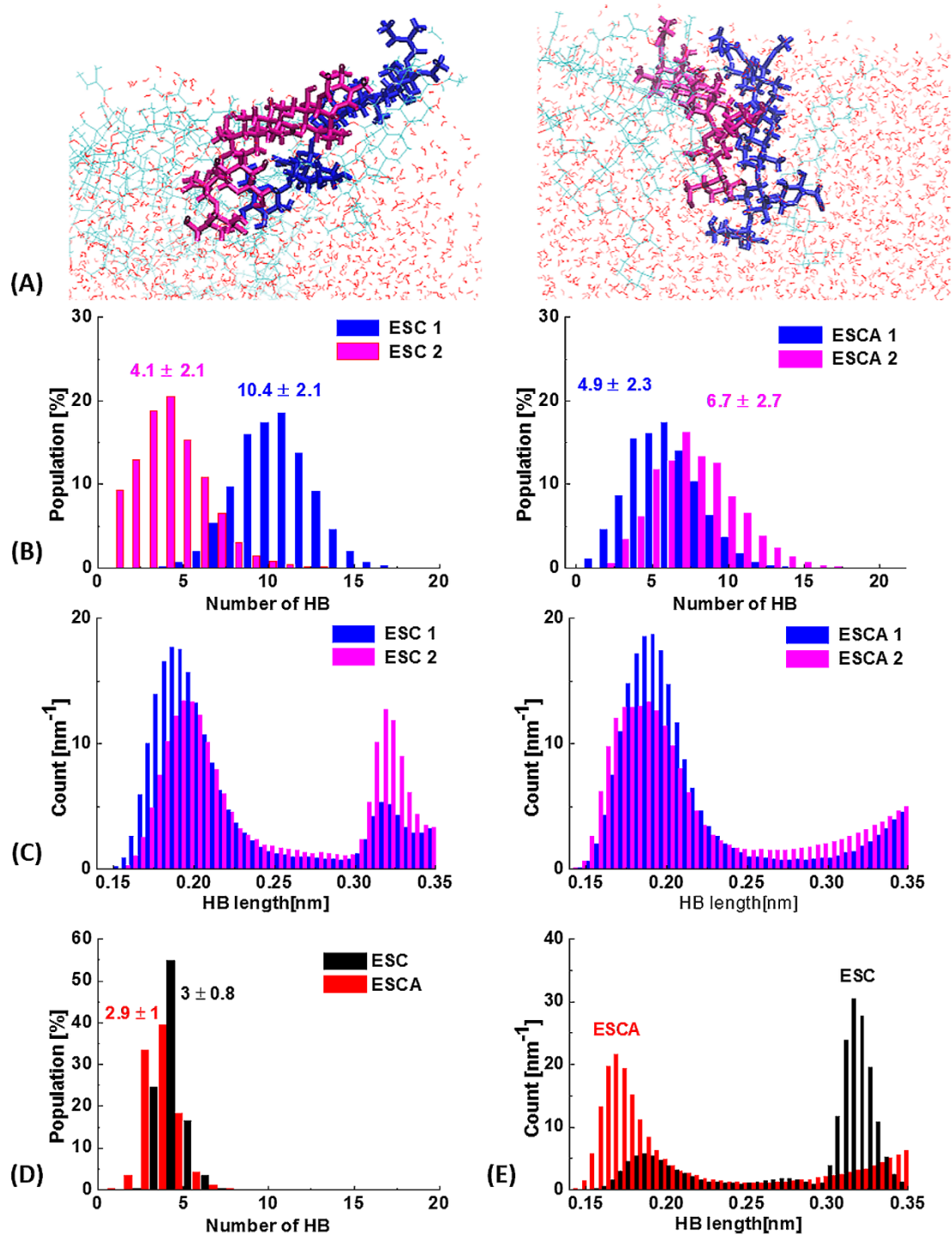


Figure 9. Analysis of the hydrogen bond distributions for well packed molecules from the ESC and ESCA clusters; (A) illustration of the molecular packing in the analyzed escin pairs; distribution of the (B) number and (C) length of the escin-escin H-bonds of the two molecules from the pair with all their actual neighbors; distribution of the (D) number and (E) length of the H-bonds within the escin pairs only.

The third H-bond is formed between the H-atom, attached to O-atom number 47 in ESC1 with the O-atom number 44 in the neighboring molecule ESC2 (see Figure 1B). The distance of this bond is longer, ≈ 0.32 nm, but it is preserved for the entire period of the last 50 ns, which shows that it is unexpectedly stable. Taking into account that this interaction is of dipole-dipole type and, hence, it should decrease as r^3 (where r is the dipole-dipole distance), we can estimate this bond to be an order of magnitude weaker than the H-bonds described above. Averaged over time, this third H-bond should give an additional contribution to the molecular pair stabilization of 1 to 2 kT . Note that this bond is formed between the other two terminal sugar residues in the neighboring escin molecules. Also, it may remain stable for a long time only if the interacting atoms are stabilized in the respective configuration by the overall packing of the ESC1 and ESC2 molecules – otherwise, this bond would be very labile and would live for less than 1 ns.

Similar analysis, performed for the closest neighboring pair of the ionized ESCA molecules (Figure 9D,E, red columns), led to different conclusions. The two anionic surfactants form around 6 ± 3 H-bonds with their actual neighbors (Figure 9B, right). On average, 3 of these bonds are between the two molecules in the close ESCA-ESCA pair, while the other 3 bonds are with molecules situated around this pair. The main H-bonds in the pair are realized between the H-atoms attached to O-atoms with numbers 44 and 47 (belonging to two neighboring hydroxyl groups in one of the terminal sugar residues) with the two oxygen atoms in the ionized carboxyl groups of the other ESCA molecule in the pair (viz. with the O-atoms with numbers 57 and 58). These H-bonds are particularly short, with length of around 0.17 nm only, which indicates that they are very strong – with energy of around 20 kT per bond [36]. Taking into account the fraction of the period, in which these bonds are active (50 %, 40 %, 20 %, and 15 % of the last 50 ns of the simulation, respectively), we can estimate that their contribution to the total interaction energy between the two paired ESCA molecules is very significant – around 25 kT . Long-range dipole-dipole interactions, at ≈ 0.32 nm, are also realized between the ESCA molecules in the pair, but much less than between the neutral surfactant molecules (Figure 9E), and their energy contribution is an order-of-magnitude lower. The H-bonds of the paired ESCA molecules with their neighbors are longer and, therefore, the cohesive energy of the ESCA pair with their neighbors is relatively low. Furthermore, between the charged carboxyl groups, separated at around 1.2 nm in the ESCA pair, we have an additional electrostatic repulsion. It is difficult to

estimate its energy, as the relative dielectric constant, ϵ , around the charged carboxyl groups is not known and could be much lower than that of bulk water, $\epsilon \approx 79$ [37]. Approximate estimates by the Coulomb's law, assuming $\epsilon \approx 10$, show that the energy of electrostatic repulsion should be above $5 kT$. In addition, as mentioned above, the aglycones of the two paired ESCA molecules are not aligned, they are not well packed and are immersed in water to a larger extent, as compared to the neutral ESC molecules. All these differences indicate lower van der Waals attractive energy between the aglycones of the charged ESCA molecules.

Discussion – relation between intermolecular forces and surface viscoelasticity

In this section we dwell briefly on how the revealed interactions between the escin molecules could explain the high viscoelasticity of the escin adsorption layers. The performed analysis showed two very important features of the studied escin clusters. First, they are formed very rapidly, which is an indication of a relatively strong and long-range attraction between the adsorbed molecules. Second, three essential types of complementary intermolecular interactions have been identified.

The first type of interaction takes place between the aglycone parts of the molecules. Due to the specific arrangement of the distributed positive and negative charges along the aglycone, a long-range electrostatic attraction acts between the escin molecules and orients them with respect to each other. The tilt of the aglycone molecules relative to the water surface can further facilitate the appropriate mutual orientation of the positively and negatively charged fragments of the aglycone, as shown in Figure 10 (see also the simulated molecules in Figure 9A). This attraction seems to be particularly pronounced for the non-charged ESC molecules, which are aligned very well with respect to each other.

For the ionic ESCA molecules, this long-range attraction is counteracted by the electrostatic repulsion between the charged carboxyl groups. The latter effect explains the slower kinetics of aggregation, the looser packing, and the larger fraction of water molecules inside the ESCA clusters (equivalent to the deeper immersion of ESCA molecules into the water phase). It explains also the observed two-peak distribution of the position of ESCA molecules normal to the interface, Figure 5B.

Inside the cluster of neutral escin molecules, preferred intermolecular distances corresponding to short-range (0.16 to 0.20 nm) and intermediate-range (0.30 to 0.35 nm) attractive H-bonds have been observed. Thus, we see that the clustering of the neutral molecules is governed by three complementary types of attractive forces, which cover a wide range of intermolecular distances. At close-packing, all these interactions combine in a synergistic way. The long-range van der Waals forces between the aglycones drive the molecule approach and orientation, while the intermediate and short-range H-bonds lock the molecular orientation and strengthen the attraction between the neighboring molecules. Note that all main fragments of the molecule, the aglycone and the two terminal sugar residues, are locked in the packed structure by the different types of interactions, explained above – see Figure 10 for illustration of this conclusion. Therefore, increasing the intermolecular distance in the adsorption layer of neutral escin molecules would require the overcoming of all these forces (from very small to large distances), which could explain the observed high *dilatational* elasticity of the escin adsorption layers [12].

Note that the relation between rheological properties and intermolecular interactions, incl. H-bonds, has been demonstrated qualitatively in other types of colloid systems. For example, in certain gels, formed by molecular self-assembly in solutions, the degree of viscoelastic recovery can be correlated qualitatively with the strength of H-bond interactions among the self-assembled molecules [38].

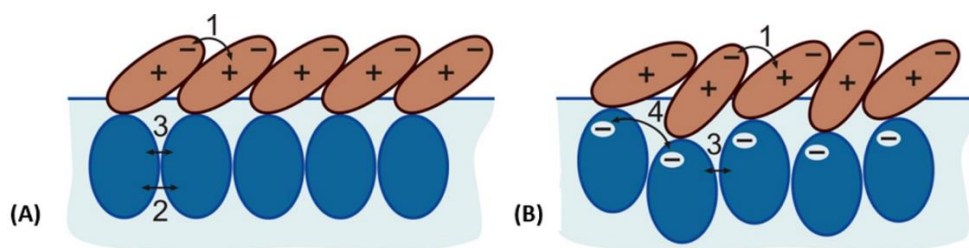


Figure 10. Schematic presentation of the arrangement of the escin molecules in a surface cluster for: (A) neutral molecules, and (B) ionized molecules. The main intermolecular forces governing the escin self-assembly are: (1) long-range attraction due to the inhomogeneous charge distribution in the aglycone and short-range dispersion (London) van der Waals forces between the aglycone fragments, (2) intermediate-range dipole-dipole/H-bond interaction, (3) short-range classical H-bonds, and (4) water-screened electrostatic repulsion between the charged carboxyl groups in the ionized escin molecules.

Furthermore, the strong dependence of the H-bonds and of the intermediate dipole-dipole interactions on the mutual orientation of the molecules, could explain also the relatively high *shear* elasticity of the escin adsorption layers [12, 27]. The shear elasticity is further enhanced by the relatively rigid structure of the aglycone, which implies that any rearrangement of the closely-packed molecules would be hindered by the limited flexibility of the aglycone scaffold.

The rearrangement of the molecules in sheared or dilated/compressed layers, at relatively large magnitudes of deformation, would require a series of bond-breaking and bond-forming events. These bond breaking-and-forming events lead to dissipation of energy, which would be observed as a viscous response of the deforming adsorption layers. Thus, the intermolecular forces between the escin molecules, described in the current study, could explain the observed, excessively strong viscoelastic response of the escin adsorption layers.

Finally, most of the reported phenomena are observed with the ionic form of escin as well. However, they are less pronounced due to the strong electrostatic repulsion between the charged carboxyl groups in the hydrophilic part of the molecule. This repulsion precludes the fine packing of the charged ESCA molecules in a homogeneous layer and, thus, it explains the much lower surface elasticity, measured with the respective adsorption layers [12].

Conclusions

Atomistic molecular dynamics simulations are carried out for escin (a representative of triterpenoid saponins) at the vacuum-water interface both for single and for clustered surfactant molecules in neutral or in anionic form. These simulations show that the escin molecules self-assemble very rapidly (for about 5 to 15 ns for the neutral form and within 60 ns for the anionic form), which indicates the presence of a strong, long-range attractive van der Waals forces between the aglycones of the adsorbed escin molecules. The aggregation kinetics of the ionic form is slower, due to electrostatic repulsion between the hydrophilic molecular fragments of the ionized escin molecules. Eventually, both the neutral and the anionic escin molecules form a single cluster at the water surface, which is much more compact for the neutral form. Unlike the neutral escin molecules, the charged

ones are loosely packed in the clusters and are displaced with respect to each other, in direction normal to the interface, to reduce the intermolecular electrostatic repulsion.

The preferred tilt angle of the escin aglycones, relative to the interface, is around 115°. This angle is not affected by the molecule ionization or by the presence of neighboring surfactant molecules in the cluster. The orientation of the sugar residues with respect to the aglycone fluctuates a lot and spontaneous conformational transitions occur, which outline significant flexibility of the sugar residues. Some molecules within the clusters are aligned with respect to each other, while regular ordering is not reached for the time of simulations (150 ns). The number of escin-escin hydrogen bonds varies significantly, depending on the intermolecular orientation. About 3 hydrogen bonds are formed on average between well packed escin neighbors, irrespective of their charge. Between the neutral escin molecules some of these bonds appear as specific attraction, intermediate in range and strength between the hydrogen bonding and dipole-dipole interaction.

Supporting Information

The following material is provided as Supporting Information: initial structures of escin (Figure S1); atom numbering of the non-hydrogen atoms in the neutral and ionized escin (Figure S2); rescaled density profiles of single molecules (Figures S3); Z-coordinates of the terminal atom of aglycone (Figure S4); schematic picture of measured molecule lengths (Figure S5); length of molecules, hydrophilic, or hydrophobic part (Figure S6); angles between aglycone and sugar chains (Figure S7); minimum distance and number of contacts for neighboring and non-neighboring molecules (Figure S8); number of inter- and intramolecular hydrogen bonds between escin molecules (Figure S9); number of intermolecular hydrogen bonds between escin and water molecules (Figure S10); representation of the hydrogen and oxygen atoms in the neighboring escin molecules between which H-bonds are formed (Figure S11); atomic connectivity information (Table S1); RESP atomic charges (Table S2); parameters from Gaussian fits of density profiles (Table S3); thickness of escin (Table S4); number of surfactant pairs found at different average minimum distance apart (Table S5); number of water molecules at a distance ≤ 0.4 nm from escin molecules (Table S6).

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