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Adsorption of the Natural Protein Surfactant Rsn-2 onto Liquid Interfaces

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To stabilize foams, droplets, and films at liquid interfaces nature has evolved a range of protein biosurfactants. Compared to synthetic surfactants, these combine surface activity with biocompatibility and low solution aggregation. One recently studied example is Rsn-2, a component of the foam nest of the frog *Engystomops pustulosus*, which has been predicted to undergo a clamshell-like opening transition at the air-water interface. Using atomistic molecular dynamics simulations and surface tension measurements we study adsorption of Rsn-2 onto air-water and cyclohexane-water interfaces. The protein adsorbs readily at both interfaces, with adsorption mediated by the hydrophobic N-terminus. At the cyclohexane-water interface the clamshell opens, due to the favourable interaction between hydrophobic residues and cyclohexane molecules and penetration of cyclohexane molecules into the protein core. Simulations of deletion mutants showed removal of the N-terminus inhibits interfacial adsorption, consistent with the surface tension measurements. Deletion of the hydrophilic C-terminus also affects adsorption, suggesting that this plays a role in orienting the protein at the interface. The characterisation of the interfacial behaviour gives insight into the factors that control interfacial adsorption of proteins, which may inform new applications of this and similar proteins in areas including drug delivery and food technology and may also be used in the design of synthetic molecules showing similar changes in conformation at interfaces.

I. INTRODUCTION

A number of biological processes such as digestion, respiration, and biofilm formation occur at liquid interfaces. To assist in these processes a variety of proteins have evolved to function in these environments. Among these are protein biosurfactants, surface active proteins that act to reduce surface tension [1–3], and have the ability to stabilise foams and emulsions. Alongside their surfactant properties these have typically evolved to be biocompatible which has led them to exhibit structures that are quite different to the standard polar head/non-polar tail of most synthetic surfactants [4, 5]. Possibly the best-known example of biosurfactant proteins are the hydrophobins [3], small, amphiphilic proteins expressed by filamentous fungi (along with similar bacterial proteins [6]). These are characterised by a hydrophobic patch on one face, giving them an amphiphilic structure similar to surfactants or Janus particles [7]. Caseins are a class of micelle forming proteins [8], that are an example of intrinsically disordered proteins. These contain regions of hydrophobic and hydrophilic amino acids, having structures reminiscent of block copolymers. A final example are lung surfactant proteins [1], in particular surfactant proteins SP-B and SP-C. These are small α-helical proteins, with their surface activity arising due to the amphiphatic nature of their helices, that play a role in modulating the surface tension of pulmonary fluid.

An interesting example of biosurfactant proteins is Rsn-2 [2], a protein found in the foam nests of the tropical frog *Engystomops pustulosus* (previously *Physalemus pustulosus*). Rsn-2 was found to be the main surfactant component of this mixture with its removal inhibiting foam formation. Its sequence is unlike any other surfactant protein and it exhibits an amphiphilic character, with a hydrophobic region at the N-terminus (residue ids 1 to 9: LILDGDLLK) and a highly polar C-terminus (residue ids 87 to 96: RKDDDDDDGY). However, high-resolution NMR studies have shown that the amphiphilic character expected from the sequence and surface activity is not apparent in the solution structure. The solution structure of Rsn-2 (PDB ID: 2WGO) comprises two regular secondary structure features, consisting of an α-helix and a four-stranded β-sheet, joined by a flexible linker region and with flexible N- and C-terminal tails (Fig. 1). Unlike the hydrophobins, no hydrophobic patches were found on the protein surface, consistent with the lack of oligomerisation in solution [9]. Neutron reflectivity measurements found a layer thickness of ~8-10 Å [2], thinner than would be expected from its solution structure. Combined with the flexible linker region seen in the solution structure, MacKenzie et al [2] suggested that the protein opens up at the interface separating the helix and sheet regions, allowing the hydrophobic amino acids in the protein core to become exposed to air or oil. Further support for this comes from polarised IRRAS mea-

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measurements, which showed that the α-helix and β-sheet secondary structures remain intact and lie in the plane of the interface.

Moreover, both N- and C-terminal regions of the protein were shown to be more dynamic than the globular core and they may be able to participate in the interfacial adsorption. Indeed, previous experiments and simulations with a highly coarse-grained model [10] suggested that the N-terminal tail can interact with the interface before the protein core exposes its hydrophobic residues. While this is an appealing model for interfacial adsorption, the precise atomistic details for interfacial attachment and opening are still unclear, with the role of the N- and C-terminal regions being of particular interest.

In this paper atomistic molecular dynamics (MD) simulations and biophysical analysis of deletion mutants are used to investigate the adsorption of Rsn-2 onto air-water and cyclohexane-water interfaces. Following early work on lipases [11] in recent years MD has been used to investigate the interfacial adsorption of a number of proteins, including several hydrophobins [7, 12–14], whey proteins, such as β-lactoglobulin [15, 16] and barley LTP [17], lysozyme [18], and myoglobin fragments [19], giving insight into their structures at interfaces and the relationship between protein structure and interfacial behaviour. Here we examine the mechanism of interfacial attachment of Rsn-2, and in the case of a water-cyclohexane interface, observe the hypothesised clamshell opening. Through consideration of mutant proteins with sections of the N- and C-termini deleted we investigate the role of these regions on interfacial attachment. Finally, we estimate the free energy contributions to the adsorption from the interactions between the protein and the two fluid phases and from the interfacial tension.

II. METHODS

A. Molecular dynamics simulations

All simulations were performed with the molecular dynamics software GROMACS 4 [20]. At the air-water interface the protein was modelled using the CHARMM27 force field [21, 22], whereas at the oil-water interface we used a version of the AMBER99sb-ildn force field [23] that includes a parametrisation for cyclohexane molecules [24]. To test the dependence of the results on the force field used, simulation of the wild type Rsn-2 at the air-water interface using the AMBER99sb-ildn force field was performed, giving results consistent with those from the CHARMM27 force field. In all cases water was represented according to the TIP3P model [25].

To prepare the system for the adsorption simulations, a protein configuration taken from the NMR structure is centred and solvated in a cubic box with side lengths set so that the protein is at least 1.2 nm away from the periodic image of the box (in oil-water simulations the size of the simulation box was further increased so that this condition is satisfied even upon unfolding). Counter ions were added to neutralise protein charge and the size of the box was increased along the z direction to create the two interfaces; in the oil-water system, the additional empty space is then filled with cyclohexane molecules. For each type of protein and interface considered we performed three production runs, with different random starting protein conformations from the NMR structure used for each (for the air-water interface we used frames 1, 11 and 21 for runs 1, 2 and 3 respectively, whereas for the oil-water interface we used frames 6, 14, and 16). The different NMR structures differ in the conformation of the N- and C-termini [2], with only small differences seen in the ordered regions. The system is equilibrated in the following steps: a steepest-descent energy minimisation procedure, a 50 ps run in the NVT ensemble and, only for the oil-water interface, a 100 ps run in the NAP$^z$ ensemble; in the last two steps the heavy atoms of the protein are restrained at their initial value using harmonic springs with a force constant equal to 1000 kJ mol$^{-1}$ Å$^{-2}$. Production runs were performed in the NVT ensemble for the air-water interface, and in the NAP$^z$ ensemble for the oil-water interface.

In all cases (including equilibration), the temperature was maintained at 300 K using the velocity-rescale thermostat [26], and in the oil-water system the pressure along the z direction was kept at 1 bar using the Parrinello-Rahman algorithm [27]. The equations of motion were integrated using a timestep of 2 fs. All bonds involving hydrogen atoms were constrained at their equilibrium position using the LINCS algorithm [28], whereas the geometry of water was constrained using the SET-TLE algorithm [29]. Electrostatic interactions were treated with the Particle mesh Ewald algorithm [30] with a Fourier grid spacing of 1.5 Å and a short-range cut-off of 10 Å. For simulations with the Amber force field short-
range Van der Waals interactions were switched to zero from 8 to 9 Å, while for CHARMM27 the Van der Waals interactions are cut-off at 12 Å.

Secondary structure assignments were made using the STRIDE algorithm [31]. The partition free energy ($\Delta F_{\text{partition}}$) in Table I was estimated as the sum of the air-water or water-cyclohexane transfer free energies [32] of each residue for which the side-chain centre of mass position is located inside the hydrophobic phase, using the condition $z > z_G$, where $z_G$ was computed from the Gibbs dividing surface for the air-water interface [33] and from the point at which the density of water equals the density of cyclohexane for the oil-water interface. The residues that form the hydrophobic core of the protein were not included in the sum, since we do not expect a significant gain in energy when these residues are transferred from the already hydrophobic environment of the core to the apolar fluid phase. These were identified as those that are not part of the N-terminal tail (id 1 to 16, which are known to be flexible from the NMR structure) and exposing a surface area lower than 30 Å in all conformations of the NMR structure. To estimate the surface area occupied by the protein at the interface, a grid of points is placed on the interface, defined by $z_G$, with the area being estimated from the number of grids points that were within a cut-off distance of $R_{vdw}^p + R_{vdw}^O$, from any protein atom, where $R_{vdw}^p$ is the van der Waals radius of the protein atom and $R_{vdw}^O = 1.4$ Å is the van der Waals radius of the water oxygen. All calculations presented in Table I used a grid spacing of 1 Å in the $x$ and $y$ directions (calculations with more grid points gave essentially identical results). The surface tension was taken from previous simulations [34] for the air-water interface, whereas for the water-cyclohexane interface it was computed directly from a molecular dynamics simulation in the absence of the protein using:

$$\gamma = 0.5 L_{\perp} (P_{\perp} - P_{\parallel}),$$  \hspace{1cm} (1)$$

where $L_{\perp}$ is the size of the simulation box along the direction perpendicular to the interface, $P_{\perp}$ and $P_{\parallel}$ represent respectively the components of the pressure perpendicular and parallel to the interface, and $\langle \rangle$ represents an average over time during an equilibrium simulation.

In Fig. 10, the number of contacts between the hydrophobic core and cyclohexane, and between α-helix and β-sheet are given by [35]:

$$N_{\text{contacts}} = \sum_{ij} \frac{1 - (r_{ij}/r_0)^6}{1 - (r_{ij}/r_0)^{12}},$$  \hspace{1cm} (2)$$

where $r_{ij}$ are respectively the distances between the $C_\gamma$ atoms of the residues in the core and the cyclohexane molecules, and the distance between the $C_\gamma$ atoms in the α-helix and those in the β-sheet we only include the large hydrophobic residues of the core, which possess a $C_\gamma$ atom. The sum runs over the appropriate pair of atoms, and $r_0 = 5$ Å. The switching function is 1 when a hydrophobic contact is formed (when the separation is small) and 0 where there is no contact (large separations).

For the clustering of protein orientations in Fig. 9, we employed the recent algorithm of search and find of density peaks [36]. In order to characterise the protein orientation, we considered the $z$-coordinates of residue ids 16 to 88 relative to the centre of mass of this region; N- and C-terminal tails have been excluded because of their flexibility. The distance between any pair of conformations was then defined as the root mean squared distance between the corresponding vectors of $z$ coordinates just defined. The three clusters in Fig. 9 have been obtained using a cutoff distance of 11 Å , but the identification of the clusters is robust with respect to changes in this value.

B. Recombinant Rsn-2 production

Recombinant Rsn-2 was produced in *E. coli* and purified as previously described[2]. Briefly, BL21[DE3] cells transformed with a pET28 based plasmid encoding Rsn-2 as a fusion protein with a thrombin cleavable N-terminal His$_6$-tag were grown in Luria-Bertani medium and induced with IPTG. The protein was purified from the soluble fraction of the lysed cells by Ni$^{2+}$-affinity chromatography and the tag removed by digestion with thrombin followed by a further Ni$^{2+}$-affinity chromatography step and size exclusion chromatography on a Superdex75 column (GEHealthcare). N- and C-terminal deletion mutants were produced in a similar fashion. After thrombin cleavage, the ‘wild type’ and truncated Rsn-2 proteins have an additional N-terminal tetrapeptide sequence, GSHM, before their first residue of the native sequence.

C. Surface activity measurements

Surface tension measurements were performed using a Kibron Microtrough S (Kibron, Espoo, Finland). The instrument was calibrated against air and pure water as per the manufacturer’s instructions and then zeroed against buffer. 500 µL samples were placed in a well of the multi-well plate immediately before measurement. The probe was immersed in the solution to ensure complete wetting, raised and then lowered until in contact with the surface before recording was initiated. The time taken from sample deposition to the start of recording was less than 10 s. Between samples the probe was flamed and between experiments, the multi-well plate was washed extensively with ethanol followed by ultrapure water and dried thoroughly. Protein samples were created by serial dilutions into 20 mM sodium phosphate, 50 mM sodium chloride, 1 mM sodium azide, pH 7.5 buffer from a stock solution in the same buffer. Of the protein concentrations tested, two were chosen to illustrate the differences between the Rsn-
2 variants. At 89 µM (equivalent to 1000 µg/mL for full length Rsn-2), surface activity is essentially fully developed within the dead time of the experiment allowing the ultimate effectiveness of the variants as surfactants to be assessed. In contrast, 445 nM is a near limiting concentration for the activity of wild type Rsn-2, with surface tension depression developing over several hundred seconds allowing the kinetics of its development to be compared for different variants.

III. RESULTS

A. Wild-type Rsn-2

We first consider the behaviour of the protein at an air-water interface. Consistent with experimental observations, simulations show that Rsn-2 adsors readily onto the interface, with this occurring within 50 ns for all three replicate runs (Fig. 2). From the centre-of-mass positions it can be seen that the protein can contact the interface without becoming permanently attached. For example for the first run the protein first contacts the interface within approximately 5 ns (Fig. 2B) but remains at the interface for less than 1 ns before diffusing back into bulk solution. The protein makes another transient contact with the interface before becoming attached for the remainder of the simulation at about 38 ns (Fig. 2D). In the adsorbed configuration most hydrophobic side-chains located in the N-terminal tail are exposed to the air phase. While the timescales differ for the other replicate runs similar behaviour is seen in all cases. Due to the small number of simulations performed we cannot state that transient contacts are a general feature of protein adsorption at interfaces. Similar short periods of transient adsorption prior to long-lasting adsorption has also been seen in simulations of β-lactoglobulin [15, 16].

At a water-cyclohexane interface the initial stages of the adsorption of wild type Rsn-2 are qualitatively similar to those observed at an air-water interface. Long lasting contacts are formed within 100 ns, and in all three runs the N-terminal tail is the part of the protein that contributes the most to the initial adsorption (Fig. 3). For instance in the first simulation run (Fig. 3B) at 12 ns the N-terminal tail interacts with the interface through residues L1 and I2, and from 24 ns (Fig. 3C) all hydrophobic residues in the tail (up to V16) have their side-chains exposed to the oil phase. The orientation of the protein at the interface can also change as a function of time and depends on the simulation run; for example from snapshots of the system observed from the direction perpendicular to the interface into the water phase, we can see that at 24 ns (Fig. 3D) both the helix and the β-sheet are perpendicular to the interface, whereas at 131 ns (Fig. 3E) they are parallel to the interface. This change in orientation corresponds to new contacts with the interface formed by the hydrophobic residues I17, L20 and F21 located at the beginning of the helix.

In this simulation run the protein also undergoes a large scale conformational change that is not observed in runs 2 and 3 or at the air-water interface. Starting from the formation of new contacts with residues L25, F29, V78 and P79, the cyclohexane molecules progressively invade the hydrophobic core of the protein and the distance between the β-sheet and the α-helix increases, as can be seen in the snapshot taken at 263 ns. This partial unfolding of the protein can be described as an unhinging of the helix with respect to the β-sheet, in which the core becomes directly exposed to the oil phase without making unfavourable contacts with water. The length of this simulation was extended up to 350 ns until the protein structure remained stable. This conformational change causes an increase in the area occupied by the protein at the interface, leading to a higher surface coverage for a given amount of protein. This is likely to be important for the biological role of Rsn-2 in foam stabilisation [9] and supports the mechanism suggested from previous experimental studies of Rsn-2 at an air-water interface [2].

The role of each part of the protein in the adsorption can be understood from the separation between the centre of mass of each residue and the interface as a function of time, as shown in Figs. 2G and 3G. For both air-water and oil-water interfaces it can be seen that the N-terminus is always close to the interface at the end of the simulation. The importance of the N-terminus can be explained by its high hydrophobicity (there are 6 leucines, 2 isoleucines, 1 proline and 1 valine within the first 17 residues of the protein) and flexibility. The latter is highlighted by the the high root mean square deviation with respect to the initial conformation (Fig. 5) and by the fact that the first 16 residues of the protein were determined to be disordered from NMR experiments [2]. Because of the flexibility of the N-terminus, the side chains of these residues are often exposed to the solvent, and they can easily adsorb at the interface. This behaviour is similar to the fly-casting mechanism observed in intrinsically disordered proteins [37]. Other regions of the protein can also form contacts with the interface. For instance, at the oil-water interface, in the third simulation run the protein adopts a perpendicular orientation with respect to the interface, different to the orientation observed in the other runs. This orientation appears to be stabilised by a hydrophobic contact with the interface formed by residue L47, which does not appear in runs 1 and 2. The hydrophobic residues at the beginning of the α-helix (L20 and F21) interact with the interface in runs 1 and 2 at the oil-water interface and in runs 1 and 3 at the air-water interface, and they seem to be responsible for the parallel orientation of the protein. Residues V78 and P79 are also close to the interface in most simulation runs, and their interaction anticipates the unhinging of the core observed in the first run at the oil-water interface.

The observation of different stable orientations at the interface might explain why the unhinging is a rare event. This suggests that this process has to proceed via the crossing of one or more free-energy barriers, correspond-
FIG. 2: Adsorption of wild type Rsn-2 at an air-water interface. (A–F) Snapshots of Rsn-2 adsorption observed in the first simulation run at times (left to right) 0, 5, 15, 38, 55 and 100 ns. (G) Distance of the protein centre of mass from the interface and residue-interface separations as a function of time. Cyan, red, and yellow lines denote first, second, and third runs respectively.

FIG. 3: Adsorption of wild type Rsn-2 at a cyclohexane-water interface. (A–F) Snapshots of Rsn-2 adsorption observed in the first simulation run corresponding, from left to right, to times 0, 12, 24 (side view and top view), 131 and 263 ns. (G) Distance of the protein centre of mass from the interface and residue-interface separations as a function of time. Cyan, red, and yellow lines denote first, second, and third runs respectively.

...ating to adopting the optimal interaction with the interface that will enable the unhinging motion of the core. The contribution of hydrophobic residues outside the tail might also decide whether adsorption is transient or longer lasting. For instance in the first simulation run at the air-water interface the initial attachment (at ~ 5 ns) involves purely the N-terminal region (up to residue 18) whereas the later adsorption (lasting for the remainder of the simulation) involves additional residues (K42-Y56 and V76-P79). As discussed above, for the
water-cyclohexane simulations the attachment of these residues to the interface, as well as aiding adsorption, may also precede the unhinging of the protein.

The secondary structure of the protein seen during the attachment of the interface (Fig. 4) remains largely unchanged, with both the long α-helix (residues 19 to 38) and the β-sheet (residues 45 to 88) remaining intact. The only partial loss of secondary structure is observed in the α-helix during the unhinging of the protein in the first run at the oil-water interface, and even in this case the conformation of this region is very close to the initial one. This maintenance of native structure is indicative of the fact that this protein has evolved to function at liquid interfaces so retains its structure here, in a similar manner to many other biosurfactant proteins such as the hydrophobins [3]. The conservation of secondary structure upon adsorption has also been observed from recent MD simulations of the hydrophobin HFBI [14], the whey protein β-lactoglobulin [15, 16], and a peptide derived from myoglobin [19].

The root mean square deviation (rmsd) of Rsn-2 (Fig. 5) shows the flexibilities of different parts of the protein, with the N-terminal tail being the most flexible. The structured core of the protein (residues 16 to 88) is typically very rigid, with the exception of the first run at the oil-water interface, where the large increase in rmsd illustrates the unhinging transition.

B. Effect of N and C-terminal deletions

To examine the contribution of the N and C-termini to the adsorption of Rsn-2 the surface activity of five mutants, ∆L1-L3, ∆L1-P15 (N-terminus), ∆Y96 and ∆D89-Y96 (C-terminus), and one double mutant, ∆L1-P15+∆D89-Y96, was measured by microtrough tensiometry. This was compared to that of wild type Rsn-2 at concentrations of 445 nM, to allow kinetic effects to be observed, and 8.9 μM, to assess the effect on overall surface activity (Fig. 6). Thus the specific contributions to surfactant activity of each of the motifs within the termini could be dissected. N-terminal deletions ∆L1-L3 and ∆L1-P15 delay the kinetics of surface tension depression at a concentration of 445 nM and reduced the overall activity by 3-5 mN/m at 8.9 μM. The ∆L1-P15 deletion slows the kinetics of Rsn-2 activity considerably more than the ∆L1-L3 mutant and reduces the decrease in surface tension by about 2 mN/m at the higher concentration.

Of the C-terminal deletions, the ∆Y96 mutation does not appear to have any significant effect upon surface ac-
tivity, and the ∆D89-Y96 reaches final surface tensions indistinguishable from WT Rsn-2 at both concentrations. However, at the lower concentration, ∆D89-Y96’s kinetics lie between those of the two N-terminal mutants. The ∆L1-P15+∆D89-Y96 double mutant displays the slowest kinetics of all the termini mutants, with activity only beginning to be evident after 600 seconds at the lower concentration. The minimum surface tension reached by this mutant (57.5 mN/m) is very similar to that of ∆L1-P15 alone, supporting the inference that the C-terminal mutants affect the kinetics of development of, but not the final surfactant activity achieved.

We therefore tested how deletions of the N-terminal first 3 (∆L1-L3) and first 15 (∆L1-P15) residues affect interfacial attachment (Fig. 7). Although the highly charged C-terminus does not directly interact with the interface, it may play a role in correctly orienting the protein, thus facilitating attachment. To test this hypothesis, we also studied mutants of Rsn-2 in which the last 1 (∆Y96) and last 7 (∆D89-Y96) residues were deleted. Figure 7 shows the centre of mass positions and residue separations for the ∆L1-L3, ∆L1-P15, ∆Y96 and ∆D89-Y96 mutants. All the mutants were studied at the air-water interface, whereas in the oil-water system we only considered the ∆L1-P15 mutant. For every mutant and type of interface we ran three independent simulations for 200 ns, a time that greatly exceeds the timescale of adsorption observed for the wild-type protein.

We will first consider the adsorption at the air-water interface. Deletion of most of the N-terminal (∆L1-P15) is found to suppress adsorption completely over the length of the simulation. Even a smaller deletion (∆L1-L3) is found to decrease the affinity of the protein for the interface. While protein adsorption is found in runs 1 and 2, the attachment to the interface occurs largely through the remaining flexible tail. The centre of mass separation once the protein has adsorbed is thus typically larger than for the wild type protein. In run 3 the protein adsorbs at 150 ns, but the contact is only temporary and the protein diffuses back into bulk water after 30 ns.

When the highly charged C-terminus is mostly removed (∆D89-Y96), adsorption to the interface is not observed in any of the simulation runs. This result suggests that the more hydrophilic C-terminus aids adsorption, possibly by ensuring that the protein can achieve a favourable orientation to attach securely to the interface. Deleting only the final residue does not seem to have a large effect on the adsorption; the protein is able to adsorb to the interface within 200 ns in all three runs.

At the oil-water interface even the mutant expected to have the largest effect on the surface activity (∆L1-P15) adsorbs in all three simulation runs. This is due to the fact that there are still many exposed hydrophobic residues in the protein (e.g. V16, I17, L20, F21, V78, V79), and that these residues interact with the cyclohexane more favourably than with the air phase, because of their higher partitioning free energies [32] (in the next section we will give a more detailed description of the energetics of the adsorption). Furthermore, in the third simulation run we observe the same unhinging conformational change found in the wild type protein. In this simulation, the unfolding is preceded by the formation of hydrophobic contacts between the cyclohexane molecules and residues I17, L20, F21, L25, F29, V78 and V79, similar to the behaviour observed in the wild-type protein. The observation that the unhinging transition at the interface proceeds via the same pathway of hydrophobic interactions suggests that the precise orientation of the protein at the interface is fundamental for this transition.

Figure 8 shows that the adsorption does not affect the secondary structure of the mutants, and even the unfolding of the hydrophobic core causes only a minor decrease in the number of α-helical residues. Again this suggests that Rsn-2 has specifically evolved to maintain its secondary structure at the interface.

C. Clustering of protein orientations

In order to further characterise the adsorption of Rsn-2, we performed a clustering analysis (see methods section) of the configurations observed after adsorption for the interface types and Rsn-2 mutants where the protein...
is able to adsorb in all three simulation runs: wt and ΔY96 Rsn-2 at the air/water interface, and wt and ΔL1-P15 Rsn-2 at the oil/water interface. As suggested from visual inspection of the simulation snapshots, the clustering confirms that the protein can interact with the interface in a limited number of orientations (Fig. 9). When it is folded, the protein adsorbs either parallel or perpendicular to the interface. The former is facilitated by the hydrophobic residues V78 and P79, whereas the latter is stabilised by residue L47. The third cluster corresponds to the unhinged and partially unfolded conformation of Rsn-2, where all the hydrophobic residues on the β-sheet and on the α-helix are able to interact with the interface. It is interesting to note that the two possible parallel and perpendicular orientations of the folded protein can be found in all types of interfaces and Rsn-2 variants considered. In most simulation runs, the protein maintains the same orientation, signalling the presence of a free energy barrier to changing orientation.

D. What drives interfacial adsorption?

More insight into the driving force for interfacial adsorption can be found by considering separate contributions to the adsorption free energy, specifically from the partitioning of the hydrophobic residues into the apolar (air or cyclohexane) phase and the change in interfacial free energy caused by the adsorption of the protein at the interface. The change in free energy of the system due to interfacial adsorption can be estimated as [38, 39]:

$$
\Delta F_{\text{inter}} = \Delta F_{\text{partition}} + \Delta F_{\text{inter}} = \sum_{i, \text{apolar}} \epsilon_i - \gamma \Delta A.
$$

where $\epsilon_i$ is the free energy of transfer of residue $i$ from the water to the apolar phase, the sum running over the residues located in the apolar phase, $\gamma$ is the air-water or cyclohexane-water interfacial tension, and $\Delta A$ is the area of interface occupied by the protein. The first term, the partitioning energy $\Delta F_{\text{partition}}$, quantifies the gain (or loss) in energy when a residue is transferred from the initial water phase to the air or oil phase [38], and its optimisation tends to move hydrophobic residues into the apolar phase, and to keep hydrophilic residues into the water phase. The second term, the interfacial energy $\Delta F_{\text{inter}}$, represents the cost associated with the presence of an interface between two immiscible fluids [39]. As the protein occupies the interface, it lowers the energy of the system by reducing the interfacial area between the two fluids.

The values of $F_{\text{partition}}$, $\Delta A$, and $F_{\text{inter}}$ for Rsn-2 at both air-water and oil-water interfaces are presented in Table I. The interfacial free energies were calculated us-
ing simulation values of the interfacial tension; for the air-water interface \( \gamma = 52.3 \text{ mN/m} \) [34] and for the water-cyclohexane interface \( \gamma = 42.7 \text{ mN/m} \). At the air-water interface the change in interfacial free energy upon adsorption is significantly larger than the partition free energy. On the other hand, at the oil-water interface partitioning and interfacial contributions to the adsorption are comparable, especially for the wild type protein. This is explained by the fact that hydrophobic residues gain more energy from being transferred to the cyclohexane phase than to the air phase, for instance the partition free energy of leucine is 4.92 kcal/mol from cyclohexane to water and 2.28 kcal/mol from air to water [32]; some residues even switch from being hydrophobic to hydrophilic, e.g. the partition free energy of phenylalanine is 2.98 kcal/mol from cyclohexane to water and -0.76 kcal/mol from air to water [32]. This observation helps to explain firstly why N-terminal deletions are sufficient to affect the adsorption at the air-water interface but not at the oil-water interface, and secondly why we observe the unfolding and exposure of the protein core to the hydrophobic phase only at the oil-water interface.

Comparison of the surface areas shows that the protein occupies a significantly larger area at the water-cyclohexane interface than the air-water interface. This again is a consequence of the stronger partitioning of the hydrophobic residues out of the water. For the runs where unhinging is observed (wt run 1 and ∆L1-P15 run 3) the interfacial area is approximately two to three times larger than that measured for simulations where the protein remains in a closed conformation. The areas from the simulations for the unhinged proteins estimated using this method are smaller than those found from neutron reflectivity [2] (∼1412 Å²).

Finally, we analysed the unhinging of the hydrophobic core at the oil-water interface (run 1 for wild type Rsn-2 and run 3 for the ∆L1-P15 mutant) by looking at the number of contacts (see methods) formed between the cyclohexane molecules and the hydrophobic residues in the core of the protein (core contacts) and the contacts between the hydrophobic residues on the \( \beta \)-sheet and those on the \( \alpha \)-helix of the protein (\( \alpha - \beta \) contacts). Figure 10 shows that the adsorption of the protein corresponds to the formation of the first contacts with the cyclohexane molecules, while the \( \alpha - \beta \) contacts remain stable at their initial value. Then, around 180 ns for wild type Rsn-2 and around 65 ns for the mutant, we observe a further increase in the number of oil contacts and a simultaneous decrease in the number of contacts between the \( \alpha \)-helix and the \( \beta \)-sheet. These plots confirm what was already suggested from the observation of the simulation trajectory: the hydrophobic interface between the helix and the \( \beta \)-sheet is directly replaced with a new interface with the oil without having to first expose the residues to the

<table>
<thead>
<tr>
<th>Run</th>
<th>( \Delta F_{\text{partition}} ) (kcal mol(^{-1}))</th>
<th>( A ) (Å²)</th>
<th>( \gamma_{\text{sim}}A ) (kcal mol(^{-1}))</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Wild type</td>
<td></td>
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<td></td>
</tr>
<tr>
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<td>-0.80±0.09</td>
<td>142.4±1.3</td>
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<td>2</td>
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<td>5.31±0.06</td>
</tr>
<tr>
<td>3</td>
<td>-1.38±0.09</td>
<td>125.9±1.2</td>
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<td></td>
</tr>
<tr>
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<td>1.50±0.05</td>
</tr>
<tr>
<td>2</td>
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</tr>
<tr>
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<td>442.0±3.6</td>
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TABLE I: Average partition free energies, occupied surface areas, and interfacial free energies for Rsn-2 at air-water interface. Average values calculated over final 20 ns of simulations, uncertainties estimated from standard error.
unfavourable water solvent. This process is not expected to be significantly favourable nor unfavourable; therefore the unhinging appears mainly driven by the interfacial energy gained from the large increase in the area occupied by the protein at the interface (see Table I).

The effect of the N- and C-termini on interfacial adsorption was explained through simulations of Rsn-2 mutants with portions of these regions deleted. Removal of the entire N-terminal tail ($\Delta L1 - P15$) was found to prevent adsorption at the air-water interface across the entire length of the simulation. Smaller deletions also affected adsorption, with one of the $\Delta L1 - L3$ simulation runs failing to adsorb. C-terminal deletions have a weaker effect on the adsorption, with the $\Delta Y96$ mutant showing similar behaviour to the wild type protein. This is consistent with experimental dynamic surface tension measurements, where mutants with the $\Delta L1 - P15$ deletion are found to both have higher limiting surface tension and slower decreases in the surface tension, indicating that the speed and extent of interfacial adsorption is lower for these. The $\Delta L1 - L3$ mutant also exhibits a higher limiting surface tension compared to the wild type protein, although the effect is smaller than for $\Delta L1 - P15$ mutant, which again is consistent with the simulation studies. The fact that neither mutation at the C-terminus has any effect upon the final surfactant activity of Rsn-2 supports the idea that the C-terminus remains within the solvent and that the solvent exposed face is sufficiently hydrophilic without the poly-aspartic acid motif.

Compared to the air-water interface protein, deletions have a smaller effect at the cyclohexane-water interface. The $\Delta L1 - P15$ mutant still adsorbs at the cyclohexane-water interface, as the partition free energy for the remaining hydrophobic residues is more favourable at this interface.

The difference between adsorption at air-water and cyclohexane-water interfaces can be better understood through considering the individual partitioning and interfacial contributions to the free energy of the system: for the oil-water interface these are comparable to each other whereas the interfacial energy is more important for the air-water interface, due to the combination of higher surface tension for the air-water interface and lower air-water transfer free energies. While it is not directly involved in the adsorption, mutations involving C-terminal deletions also affect the adsorption at the air-water interface, potentially as this region helps orient the protein to enhance the attachment probability.

While in most of our simulations the protein remains adsorbed for a considerable period time we cannot exclude possible desorption events at later times. Indeed just as in the experimental system adsorption to the interface is a dynamic process to some occasional desorption would be anticipated. The change in interfacial free energy and partition free energies, which can be used
to approximate the adsorption free energy, show that for
the wild type protein these is much larger than $k_B T$ sug-
gestig that the protein is strongly attached to the inter-
face. This is particularly the case for simulations where
protein unhinging was seen, suggesting that this would
then make adsorption permanent.

The adsorption free energy may be calculated using
techniques such as metadynamics [40] or adaptive bias-
ing force [41]. These have been successfully used for short
peptides but it is very challenging to apply enhancing
techniques to such a larger proteins such as Rsn-2. To
date there is an example of replica exchange simulations of
a protein adsorbed at the interfaces [42], but the conver-
gence of these simulations has not been carefully evalu-
ated. Replica exchange metadynamics calculations have
been used to investigate the adsorption of short peptides
at interfaces [42] but the extension of these methods to
the larger proteins studied here is not trivial. The re-
results of our Rsn-2 simulations corroborate the previous
work using a simplified (Go) model for which exhaustive
sampling was possible [10], but also provide a more
detailed understanding of the initial stages of interfacial
adsorption.

By determining the residues involved in the adsorption
of Rsn-2 at liquid interfaces and examining the effects of
their removal experimentally we have provided a detailed
characterisation of its interfacial adsorption. This will
help inform new applications of proteins in areas includ-
ing food products [43] and drug delivery [44] and in the
design of synthetic molecules that exhibit similar confor-
mational changes at liquid interfaces [45].

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Appendix - Simulations at air-water interface using
Amber99sb-ildn force field

The protein centre-of-mass-interface and residue-
interface separations from simulations of Rsn-2 at an air-
water interface using the AMBER99sb-ildn force field are
shown in Fig. 11. This shows similar behaviour to sim-
ulations using the CHARMM27 force field (Fig. 2). The
protein makes transient contact with the interface after
about 10 ns. It then contacts the interface at approxi-
mately 26 ns, remaining attached for the remainder of the
simulation. As for the CHARMM27 simulations this at-
tachment again is mediated by the N-terminus. Further
residues (around residues 45 and 60) are in contact with
the interface, with these being further from the interface
during the initial, transient contact.

FIG. 11: Distance of the protein centre of mass from an
air-water interface as a function of time and
residue-interface separations from simulations using
Amber99sb-ildn force field.

As for the Charmm-27 simulations at an air-water in-
terface, as well as simulations at a water-cyclohexane
interface the secondary structure remains largely un-
changed during adsorption at an air-water interface
(Fig. 12).

FIG. 12: Protein secondary structure as function of
time at an air-water interface from simulations using
Amber99sb-ildn force field.

This demonstrates that the behaviour of the protein is
similar using both CHARMM27 and AMBER99sb-ildn
force fields, suggesting that the observed behaviour is
not dependent on the force field used. The qualitative
results are similar for the simulations with different force
fields, with the occupied surface area (61 ± 0.6 Å²) and
partition free energy (-1.22±0.05 kcal/mol) being close
to the CHARMM27 values (Table 1).