

Molecular basis of the osmolyte effect on protein stability: a lesson from the mechanical unfolding of lysozyme

Beata Adamczak*, Miłosz Wieczór*, Mateusz Kogut, Janusz Stangret and Jacek Czub

Osmolytes are a class of small organic molecules that shift the protein folding equilibrium. For this reason, they are accumulated by organisms under environmental stress and find applications in biotechnology where proteins need to be stabilized or dissolved. However, despite years of research, debate continues over the exact mechanisms underpinning the stabilizing and denaturing effect of osmolytes. Here, we simulated the mechanical denaturation of lysozyme in different solvent conditions to study the molecular mechanism by which two biologically relevant osmolytes, denaturing (urea) and stabilizing (betaine), affect the folding equilibrium. We found that urea interacts favorably with all types of residues via both hydrogen bonds and dispersion forces, and therefore accumulates in a diffuse solvation shell around the protein. This not only provides an enthalpic stabilization of the unfolded state, but also weakens the hydrophobic effect, as hydrophobic forces promote the association of urea with nonpolar residues, facilitating the unfolding. In contrast, we observed that betaine is excluded from the protein backbone and nonpolar side chains, but is accumulated near the basic residues, yielding a nonuniform distribution of betaine molecules at the protein surface. Spatially resolved solvent–protein interaction energies further suggested that betaine behaves in a ligand- rather than solvent-like manner and its exclusion from the protein surface arises mostly from the scarcity of favorable binding sites. Finally, we found that, in the presence of betaine, the reduced ability of water molecules to solvate the protein results in an additional enthalpic contribution to the betaine-induced stabilization.

Introduction

Protein folding and unfolding are fundamental processes for all living cells and as such have been extensively studied for several decades [1–4]. It is well known that multiple factors, including temperature [5], pH and pressure [6], can shift the equilibrium between the folded and unfolded states. A protein folding equilibrium can also be modulated by a class of organic co-solvents, often termed osmolytes, which are accumulated in cells in response to environmental stresses [7–9]. Osmolytes have been shown to affect the structural and thermal stability of proteins and other macromolecules [10,11], protein–protein and protein–DNA association [12] and the degree of reversibility of the folding/unfolding transition [9,11,13].

Urea, a byproduct of amino acid catabolism, is known to shift the folding equilibrium toward the unfolded state and to induce, at relatively high concentrations (>4 M), a complete protein denaturation [11,13–15]. Interestingly, even though urea disturbs protein structure and function, it is accumulated (up to ~600 mM) as a major osmolyte in certain cells and tissues, including the mammalian kidney medulla, to balance the high osmotic pressure of the extracellular medium [7]. This is possible because the damaging effect of elevated urea on proteins in these cells is counteracted by controlled accumulation of protecting co-solvents, such as methylamines, particularly glycine betaine and trimethylamine N-oxide (TMAO), as well as sugars and other polyols [7,9,16,17]. These co-solvents, referred to as compatible osmolytes, play a key role in the adaptation of cells to hypertonicity and other stresses, as they can stabilize proteins by shifting the folding equilibrium toward the native state, generally without affecting their cellular function [7,13,14,18].

Osmolytes are also commonly used in molecular biology and biotechnology, for example, to assist the proper folding of recombinant proteins by exploiting ‘chemical chaperone’ activity of osmoprotectants such as betaine or TMAO, or to facilitate refolding and activation of proteins expressed as insoluble inclusion bodies, as in the case of urea, guanidine hydrochloride and other denaturants [7,9,19].

Despite previous efforts, the molecular underpinnings of the co-solvent-induced protein stabilization and destabilization are not yet fully established [11,13,15,20–24]. Two basic/general mechanisms, ‘direct’ and ‘indirect’, or their specific combination [10,25,26] have usually been put forward to explain the effect of osmolytes on protein stability. According to the direct mechanism, osmolytes can perturb the equilibrium between the folded and unfolded states by preferential, usually favorable interactions, such as hydrogen

bonding, with the protein surface [15,20,27–31]. On the other hand, in the indirect scenario, co-solvent molecules alter the structural and dynamic properties of water, which can lead to weakening of the protein-stabilizing hydrophobic effect or to a decrease in the overall solvation capability of the mixture [22,24,32,33]. Indeed, many osmolytes are known to disrupt or enhance the natural water structure and hence are sometimes referred to as ‘structure breakers’ (urea) or ‘structure makers’ (betaine) [34,35].

Although the indirect mechanism has been previously invoked to account for the denaturing effect of urea [36,37], it is now more commonly believed that urea induces unfolding by interacting with a protein more strongly than with water, that is, by stabilizing the unfolded state in which the protein surface accessible for direct interactions with the solvent is larger [11,13,28,30,38–40]. The molecular mechanism behind this preferential interaction is, however, less clear. Recent data, obtained by measuring the transfer free energies of side chain and backbone units between water and urea solution [39,41] and by hydrogen-exchange experiments [38], suggest that the main enthalpic driving force for the urea-induced unfolding is provided by direct interactions between urea and the protein backbone. In contrast, preferential interactions of urea with model compounds determined by osmometry indicate that interactions with the protein side chains are at least equally important, contributing 60% to the total effect of urea [42]. This finding is also consistent with the molar volume and compressibility data for amino acids in urea solution [29] and with recent molecular dynamics simulations [13,28,30]. The observed propensity of urea to bind to the protein surface has been explained either by its ability to form hydrogen bonds — particularly with the backbone amide and carbonyl groups [38] — or by favorable dispersion interactions, mostly with the protein side chains [23,27,28,43].

Even less clear is the mechanism by which betaine and other methylamines shift the folding equilibrium toward the protein native state. According to the prevailing model, based largely on osmometry and transfer free energy measurements [39,40,42,44,45], the protecting co-solvents are preferentially excluded from the surface of the protein, especially from the vicinity of the backbone peptide groups [13,18,39,40,42,44,45]. This in turn gives rise to a force opposing the protein unfolding, as in the unfolded state it is more difficult for the (effectively repelled) co-solvent to avoid the unfavorable contact with the protein surface. Therefore, in the surface exclusion model, compatible osmolytes act as crowding agents by favoring more compact protein structures typical for the native state [46–48]. This also explains why the accumulation of protein-stabilizing osmolytes can sometimes be detrimental. For example, it has been shown that by promoting compaction, protecting osmolytes can induce aggregation of unstructured proteins [17,49]; they can also impair the functioning of certain enzymes, probably by restricting their conformational flexibility [50].

Importantly, while the standard Asakura-Oosawa model based on hard-core steric interaction predicted the excluded volume-induced stabilization to be entirely entropic in nature [51], recent theoretical considerations led to the development of a more sophisticated ‘soft-core’ model involving both entropic and enthalpic contributions to the stabilization free energy [52,53]. In this description, the overall nature of the stabilizing depletion force depends on the respective enthalpic and entropic components of the effective protein–co-solvent interaction potential. This idea sheds new light on the results of recent circular dichroism and computational studies, which have indicated that protein stabilization by protecting osmolytes is indeed at least partially enthalpy-driven and that this contribution might be mediated by water molecules perturbed by the co-solvent [46,54,55]. This perturbation can also be linked to enhanced water ordering, an effect reported in many previous studies and often put forward to explain the effective co-solvent–solute repulsion [20,34,35,56].

Here, we used molecular dynamics simulations to identify the mechanisms by which the presence of two different co-solvents, urea and betaine, affects the stability of the tertiary structure of lysozyme, a typical globular protein used in folding studies [57]. To provide a controllable model of protein denaturation, a mechanical force was used to increase the effective radius of a lysozyme molecule, enabling us to study the native state stability by relating the expended work to the extent of unfolding along a single conformational co-ordinate. The results of the simulations were found to be in general agreement with the available experimental data on the protein–osmolyte solutions. Encouraged by this agreement, we verified

and extended the existing models of osmolyte–protein interactions to present a consistent description of the atomistic-level driving forces that govern co-solvent exclusion and accumulation.

In particular, we observed that urea interacts favorably with almost all kinds of chemical moieties at the protein surface, and that this leads to its accumulation in an extended solvation shell around the protein molecule. Importantly, we noted that hydrogen bonding alone does not account adequately for protein–urea association, and that the dispersion interactions and water-mediated hydrophobic forces are crucial for the observed accumulation and, consequently, for driving denaturation in urea solutions. Our results also indicate that betaine is preferentially excluded from the vicinity of the protein backbone, giving rise to a sequence-independent penalty for the protein unfolding. Interaction energy decomposition suggests that this exclusion at least partially stems from the ligand-like behavior of betaine at the protein surface, providing us with a molecular-level interpretation of the major entropic contribution to the depletion force that promotes protein compaction, and a possible refinement of the canonical hard-sphere model of co-solvent exclusion. Finally, we note that, in the presence of betaine, the protein–water interactions are considerably weakened, and propose that the resulting decrease in protein solvation may account for the enthalpic contribution to the stabilizing effect of betaine.

Experimental

Seven primary systems were constructed to simulate the mechanical unfolding of lysozyme in pure water and in the presence of urea and betaine. In each system, a dodecahedron simulation box with a box vector length of 7.8 nm (corresponding to $\sim 335 \text{ nm}^3$) contained a single lysozyme molecule and solvent molecules corresponding to three betaine concentrations (1.0, 2.5 and 5.0 M), three urea concentrations (3.5, 5.0 and 7.0 M) and pure water (0 M). The specific numbers of water and osmolyte molecules are given in Supplementary Table S1. The force field parameters for urea, N,N⁰-dimethylurea and sarcosine (N-methylglycine) were obtained from the CHARMM General Force Field (CGenFF) [58], and the parameters for betaine were obtained by analogy using the CHARMM27 parameters for phosphatidylcholine (for the trimethylammonium and methylene groups) and for phosphatidylserine (for the carboxyl group) [59]. The choice of parameters for betaine was validated against the experimentally measured *m*-value, preferential hydration coefficient and osmotic pressure, and satisfactory agreement was obtained, as described in detail in Supplementary Figure S1 and Table S2 [45,55,60]. For water, the TIP3P [61] model was used. The initial structure of lysozyme was taken from the X-ray structure (PDB code: 1AKI), and the CHARMM27 force field [62] was used for the protein residues.

All MD simulations were performed with the GROMACS package [63]. Periodic boundary conditions were applied, and the particle mesh Ewald (PME) method [64,65] with a real-space cutoff of 1 nm was used to account for long-range electrostatics. The simulations were carried out in the NPT ensemble using a Berendsen-type temperature coupling [66] with a coupling coefficient of 0.1 ps and a reference temperature of 300 K, and Berendsen-type pressure coupling [66] maintaining the pressure at 1 bar with a coupling coefficient of 1 ps. During the simulations, bond lengths were constrained using the LINCS algorithm [67], and a typical time step of 2 fs was used. Initially, each system was equilibrated by carrying out 1 ms unconstrained simulations.

To unfold lysozyme in a controllable manner, an external potential was then applied to the radius of gyration (R_{gyr}), defined for the reference group consisting of each C_{α} atom in the protein backbone:

$$R_{\text{gyr}} = \sqrt{\frac{1}{N} \sum_{C_{\alpha}} (r_i - r_{\text{COM}})^2}, \quad (1)$$

where N is the number of C_{α} atoms, r_i is the position of *i*th C_{α} atom and r_{COM} is the position of the center of mass of the reference group.

During the enforced unfolding, R_{gyr} was changed from the initial value of 1.4 nm in the native state up to 3.0 nm, with a harmonic force constant of $7000 \text{ kJ mol}^{-1} \text{ nm}^{-2}$, and four pulling speeds were used: 1×10^{-4} , $1 \times$

10^{-5} , 5×10^{-5} and 5×10^{-6} nm ps $^{-1}$. To improve the signal-to-noise ratio in the unfolding curves, five simulation runs were carried out for each system at the lowest pulling speed.

Systems with 2.5 M sarcosine, 5 M N,N⁰-dimethylurea and 5 M scaled-charge betaine were prepared in an analogous manner, and only the lowest pulling speed of 5×10^{-6} nm ps $^{-1}$ was used. In the charge scaling simulations, all partial charges of the betaine molecule were scaled by a common factor of 1.2, 1.4 or 1.6. The total simulation time for data collection was 51 ms. In the hydrogen bond analysis, H-bonds were identified based on a donor–acceptor distance criterion of 0.35 nm and a 40° threshold for the hydrogen-donor–acceptor angle. Average H-bond energies were estimated using the formula by Espinosa et al. [68]:

$$E_{\text{HB}} = 2.5 \times 10^4 \exp(-36d(\text{H} \dots \text{O})) \quad (2)$$

where $d(\text{H} \dots \text{O})$ is the hydrogen-acceptor distance in nm and E_{HB} is the H-bond energy in kJ/mol.

Results and discussion

A common pathway of the enforced unfolding

To investigate the effect of typical denaturing and protecting osmolytes on the stability of the protein native state, we used nonequilibrium molecular dynamics to simulate mechanical unfolding of lysozyme in pure water and in the presence of increasing concentrations of urea and betaine. To this end, external steering forces were applied to all C $_{\alpha}$ atoms of the protein to increase its radius of gyration, R_{gyr} , from the initial value of 1.4 nm, characteristic of the native state, to a threshold value, in each case set at 3.0 nm (see Experimental for details).

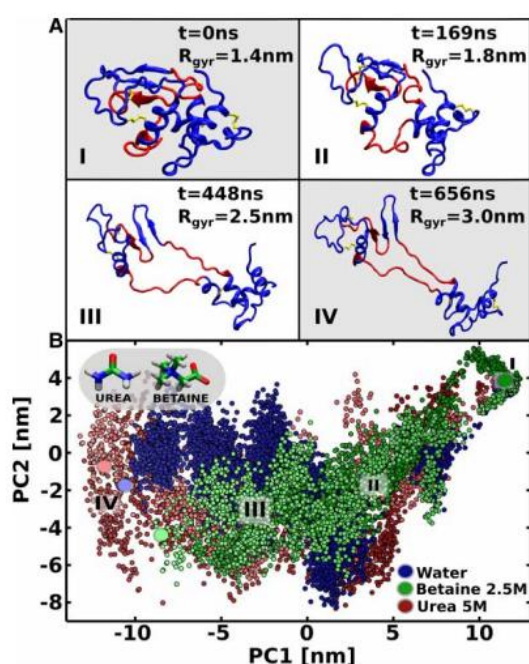


Figure 1. The mechanical unfolding of lysozyme.

(A) Snapshots of the simulated unfolding pathway obtained at the pulling speed of 5×10^{-6} nm ps $^{-1}$. The values of the radius of gyration in consecutive snapshots are: 1.4 nm — the native state; 1.8 nm — a rupture point; 2.0 nm — large changes in the lysozyme structure, i.e. the previously buried protein portion marked in red becomes exposed to the solvent; 3.0 nm — an arbitrary final state. (B) Unfolding trajectories in pure water (blue), in the presence of betaine (green) and in the presence of urea (red) projected onto the reduced subspace defined by the first two eigenvectors extracted from the unfolding in pure water. The initial and the final states are shown as large circles. All molecular images were created using VMD [69].

Inspection of the resulting trajectories shows that for all systems the unfolding process, illustrated in [Figure 1A](#), follows a common pathway: the protein splits into two globular subdomains, each held together by two disulfide bonds, and interconnected by two antiparallel, extended strands that upon unfolding become exposed to the solvent. To examine if this folded-to-unfolded conformational transition depends on the solvent composition, we projected the trajectories for all systems onto the plane formed by two first principal co-ordinates (PCs) obtained for the unfolding in pure water using principal component analysis. [Figure 1B](#) combines three such projections, depicting the unfolding pathways in pure water, 5.0 M urea and in 2.5 M betaine. As can be seen, in this projection the mechanically induced unfolded state of lysozyme in water (blue circle in [Figure 1B](#)) corresponds well to the endpoints of the respective unfolding pathways in urea and betaine solutions (red circle and green circle, respectively, in [Figure 1B](#)). As the first PCs by construction capture the large-scale unfolding modes, the proximity of the end-state projections suggests that, irrespective of the solvent composition, lysozyme evolves along a common enforced unfolding pathway adopting similar final structures. This conclusion finds further support in the dependence of the lysozyme solvent accessible surface area (SASA) and root-mean-square deviation on the radius of gyration, as shown in Supplementary Figures S2 and S3, respectively.

Work of unfolding provides a measure of the co-solvent-induced equilibrium shift

The force–extension curves (the steering force vs. R_{gyr}), shown in [Figure 2A](#), reveal that lysozyme unfolds in an apparent two-state fashion with one dominant activation barrier that is effectively lowered by the applied potential. Initially, up to the rupture point ($R_{\text{gyr}} \approx 1.7$ nm), which is relatively insensitive to the solvent composition, we observe a linear elastic response of the lysozyme molecule, with the structural deformation increasing proportionally to the applied force. The high energetic cost of the lysozyme unfolding during this initial stage is largely due to exposure of formerly buried hydrophobic side chains and other nonpolar segments to the solvent, as indicated by a large change in SASA (Supplementary Figure S2). After the rate-limiting barrier is overcome, the further increase in SASA is less pronounced and the steering force drops abruptly to much lower positive values, although the effect of high concentrations of urea and betaine, facilitating or opposing the unfolding, is still noticeable.

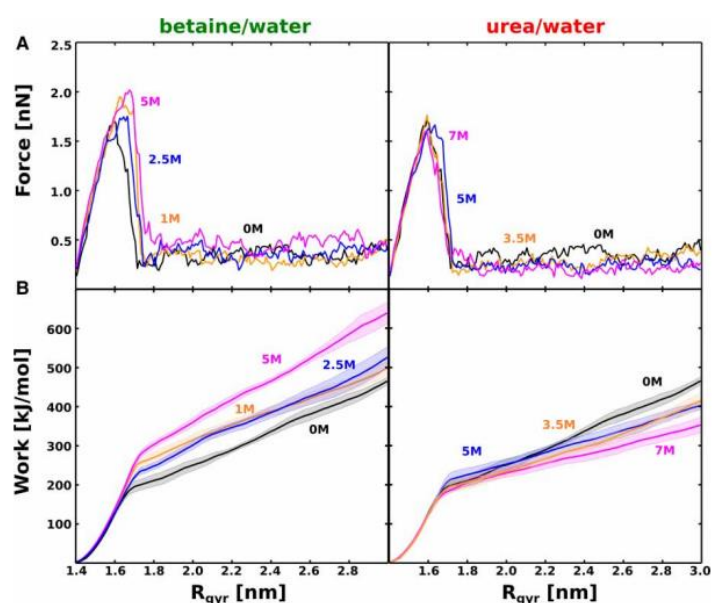


Figure 2. Pulling forces and unfolding work in steered MD simulations.

(A) The force–extension profiles for the lysozyme unfolding in the presence of increasing concentration of betaine and urea.

(B) The corresponding profiles of unfolding work as a function of R_{gyr} . All data correspond to the pulling speed of 5×10^{-6} nm ps $^{-1}$; for higher pulling speeds, see Supplementary Figures S4 and S5. All values were averaged over five independent simulations, with the shaded area representing the standard error of the estimate.

Figure 2B shows the amount of work required to unfold lysozyme in the presence of both co-solvents and in pure water, obtained at the lowest pulling speed of $5 \times 10^{-6} \text{ nm ps}^{-1}$ by integrating the steering force over R_{gyr} from the initial value up to 3 nm. Given the similar energy dissipation, the differences in the unfolding work with respect to pure water can be expected to reflect the actual perturbing effect of co-solvents on the protein folding equilibrium. Indeed, we found that, compared with pure water, more work is needed to unfold the lysozyme in the presence of betaine, by up to 40% with a 5.0 M solution, and the opposite is the case for the urea solutions, i.e. lysozyme becomes more susceptible to mechanical stress with the unfolding work decreasing by up to 20% with a 7.0 M urea solution. Therefore, these results indicate that the unfolding work computed using our model consistently captures the well-known fact that betaine stabilizes the native protein structure, while urea acts as a denaturing agent. It is, however, necessary to remark here that the value of the unfolding work itself — obtained with pulling speed orders of magnitude higher than in actual single-molecule experiments [70,71] — cannot be directly compared with the free energy of unfolding, as the expended work approaches free energy only in the quasi-static limit. Interestingly, in both osmolyte solutions, the expended work increases faster with increase in the pulling speed than in pure water such that the destabilizing effect of urea is effectively reversed in the limit of fast unfolding (Supplementary Figure S4). As both betaine and urea solutions have higher viscosity than pure water [72,73], this tendency suggests that whereas at higher speeds the steering force is dominated by viscous drag, at the lowest speed, the dissipation is small enough for the calculated work to reflect the osmolyte-induced effect on the folding equilibrium. It is worth noting that reduction in nonequilibrium effects is also seen from the increase in the number of H-bonds between the protein and urea with decrease in pulling speeds (see Supplementary Figure S5).

We additionally tested the above conclusions by performing simulations of forced unfolding of lysozyme in the presence of two other similar osmolytes, sarcosine (N-methylglycine) and N,N⁰-dimethylurea. The former stabilizes proteins more efficiently than betaine while having a slightly lower viscosity, and the latter is a more effective denaturant than urea while also being more viscous [72–74]. The unfolding work, shown in Supplementary Figure S6, is smaller in N,N⁰-dimethylurea than in urea solution, and larger in sarcosine than in betaine solution, consistent with the prevailing effect of osmolyte-induced (de-)stabilization and not with a dominating impact of solvent viscosity.

Hydrogen-bonding analysis reveals major differences between unfolding in urea and betaine solutions

It is expected that with increase in co-solvent concentrations, the amount of water available to solvate the hydrogen-bonding sites of the protein decreases in both the folded and unfolded state. Hydrophilic osmolyte molecules may compensate for this decrease by providing additional hydrogen bonds with the protein. However, co-solvents have different H-bonding properties compared with water, and such differences in solvation properties lead to positive or negative enthalpies of transfer of a protein from water to the osmolyte solution. Since the number of H-bonding sites on the protein surface generally increases during unfolding, these non-zero transfer enthalpies may contribute to stabilization or destabilization of the native state, respectively.

Accordingly, we determined the number of H-bonds formed between lysozyme and the solvent components as a function of the unfolding coordinate R_{gyr} (Figure 3A). Since the work of unfolding in each system relates to the differences in the number of H-bonds between the unfolded and folded states rather than the absolute numbers, in Figure 3B, we also compared the overall changes observed.

To make our results less sensitive to the conformational fluctuations of the numerous charged side chains on the protein surface, we focused the H-bonding analysis on the two antiparallel strands (Glu35–Asn65 and Ile88–Ala110) connecting the two lysozyme subdomains that split apart during the unfolding (Figure 1). To further facilitate the comparison, the intervening β -hairpin (Thr43–Gly54) responding differently in the betaine and urea solutions has been excluded. The chosen fragments, referred to as ‘exposed’ and shown

in red in Figure 1, are continuous polypeptide chains representative of the regions that are buried in the folded proteins and become exposed upon unfolding.

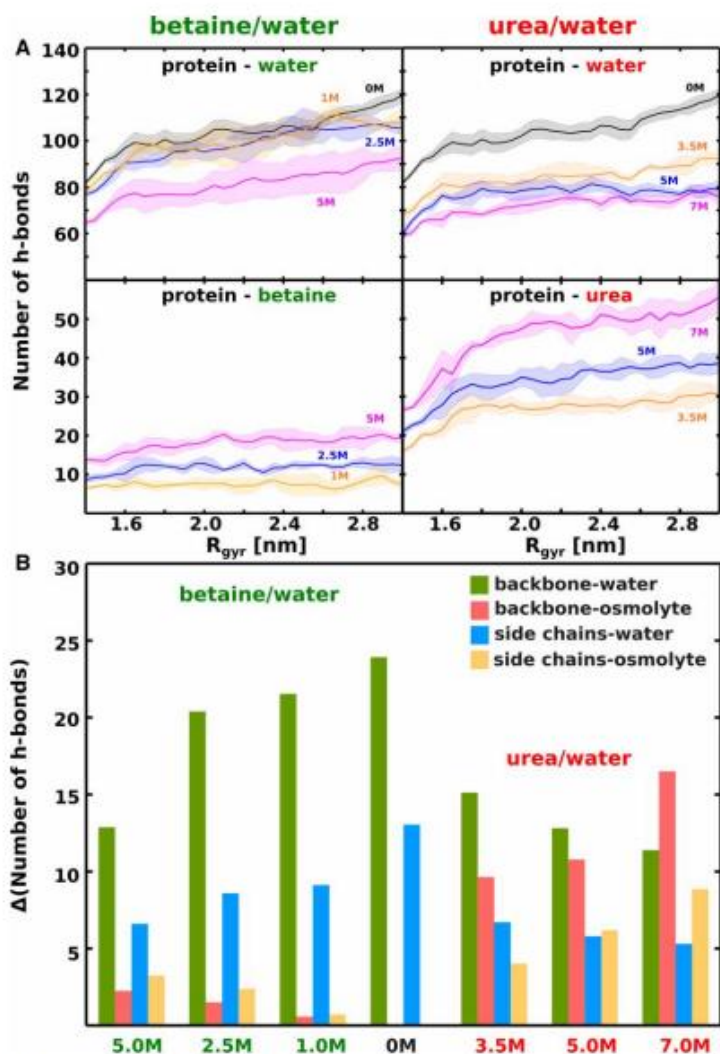


Figure 3. Unfolding-induced changes in hydrogen bonding.

(A) Number of hydrogen bonds formed between the protein and the co-solvent or water molecules during the unfolding as a function of R_{gyr} . (B) The increase in the number of hydrogen bonds upon unfolding of lysozyme in pure water and in the presence of osmolytes. Four contributions are shown: backbone – water (green), backbone – osmolyte (red), side chains – water (blue), side chains – osmolyte (orange). All data correspond to the lowest pulling speed ($5 \times 10^{-6} \text{ nm ps}^{-1}$). All values were averaged over five independent simulations, with the shaded area representing the standard error of the estimate.

It can be seen from the left panels of Figure 3A that over the entire range of R_{gyr} , the total number of protein–water H-bonds in the betaine solution is only slightly reduced relative to that found in pure water (by up to 20% for the highest betaine concentration). At the same time, even though betaine can potentially form H-bonds with the protein via the carboxyl group, the number of such bonds remains relatively low during the unfolding. In fact, Figure 3B shows that the increase in the number of H-bonds formed by the protein with water is much larger than that with betaine and is also comparable with that seen in the absence of osmolytes, especially for the protein backbone for which 80–90% of the corresponding increase in pure water is observed. This reveals that the newly exposed H-bonding sites are solvated preferentially by water molecules and suggests that betaine is excluded from residues that form the protein core, in agreement with the experimental data [18,39,40,44,45].



The volume associated with unfavorable protein–co-solvent interactions — roughly proportional to the surface area — introduces a penalty for unfolding, as the solvent-exposed surface area is typically larger in the unfolded state; hence, exclusion from the protein surface has been suggested to account for the stabilizing effect of protecting co-solvents. The exclusion itself can basically occur through two main mechanisms. In the first scenario, betaine is excluded from protein surfaces by the effective repulsion, which may result either from the preference of a protein to interact with water or from the fact that incorporation of a bulky and highly polar betaine molecule into the protein surface requires satisfying certain steric and electrostatic constraints. Alternatively, since hydrogen bonds were found to be markedly shortened in the solvation shell of betaine [74,75], the exclusion effect can be ascribed to a strongly bound hydration layer that presumably forms around the osmolyte and reduces its interactions with the protein surface.

In stark contrast with betaine, the right panels of Figure 3 show that in urea solutions, it is the osmolyte that preferentially solvates the exposed H-bonding sites on the protein. Despite the fact that water molecules outnumber urea molecules by 4.5- to 12-fold, Figure 3B reveals that the number of new protein–urea H-bonds is comparable with that of protein–water (70–130% of the latter depending on the urea concentration). This suggests that urea interacts with the initially buried residues more favorably than water, and is significantly enriched at the protein surface, as observed experimentally [39,40,42,76]. The preferential protein–urea interactions overcompensate the entropy loss due to the urea/water concentration gradient and, therefore, are commonly believed to be responsible for the denaturing effect of urea [15,30,77,78]. The exact molecular mechanism of this preferential accumulation is, however, less clear [13,76].

In addressing this question, we first note that the increase in the total number of H-bonds between the protein and the solvent is only slightly greater (average by 3%) in the urea solution than in pure water (Figure 3). If we also take into account that the H-bonds formed by urea with the protein are on average longer and hence energetically less favorable than those formed by water (by ~ 10 kcal/mol, averaging over all concentrations employed), as shown by the H-bond energy analysis in Supplementary Figure S7 and Table S3, it can be concluded that, in urea solutions, direct H-bonds tend to stabilize unfolded protein conformations even less efficiently than in pure water. Therefore, contrary to some previous reports [26,38,79], H-bonding (polar) interactions alone are not sufficient to account for the reduced work of unfolding in urea solutions. Apparently, urea-induced destabilization occurs primarily due to nonpolar interactions including, as previously suggested, favorable urea–protein dispersion interactions [20,28,43] and/or weakening of the hydrophobic effect by urea accumulated at the protein surface [27,36,77,80].

To further verify this conclusion, we extracted representative structures from the unfolded ensemble generated in 5.0 M urea, replaced the co-solvent with water and ran 100 ns MD simulations with the backbone atoms positionally restrained; the reverse procedure, i.e. ‘resolution’ of the unfolded conformations generated in pure water with 5.0 M urea was also carried out (with all solvent molecules added anew in both cases). Changes in the H-bonding pattern upon solvent exchange, summarized in Supplementary Table S4, show that, in the presence of urea, the number of protein–solvent H-bonds is larger by only 2–3%, regardless of the protein conformation and a particular unfolding pathway. This result confirms that the overall ability of pure water to saturate the exposed H-bonding sites is very similar to that of urea solutions. Furthermore, the distribution of the total number of H-bonds between various types of residues is also quite similar for both solvents (with differences in the range of 0–7%).

Previous works have proposed that the effect of betaine and other methylamines on protein stability is partially enthalpic in origin and that it may stem from decreased protein solvation [46,52,53,55,81]. Accordingly, our data presented in Figure 3B show that the total number of new H-bonds, i.e. water–protein and betaine–protein combined, in the betaine solutions is reduced by $\sim 10\%$ compared with pure water, with no apparent dependence on betaine concentration. This decrease — leading to a less favorable solvation energy of the unfolded state — might be thought to account for the enthalpic cost of the unfolding in betaine solutions. To more directly test whether betaine solutions indeed have lower ability to solvate the exposed polar groups of the protein, we carried out for betaine an analogous ‘resolution’ procedure to that

described above for urea. We found (Supplementary Table S5) that, upon betaine addition, the number of H-bonds between the solvent and the backbone of the lysozyme unfolded in pure water decreases by ~10%, with almost all bonds (93.80%) formed by water molecules and the osmolyte contributing the remaining 2.20%. A similar decrease in the number of H-bonds is seen for the polar (6%) and hydrophobic (11%) side chains, while preferential H-bonding between betaine and positively charged basic residues partially compensates for this desolvation. These observations suggest that betaine present as a co-solvent measurably reduces the water activity, thereby disfavoring water–protein H-bonds. Since betaine itself is largely excluded from the vicinity of the protein and, therefore, cannot fully compensate for the loss of water–protein H-bonds, the resulting desolvation may possibly lead to enthalpic destabilization of the unfolded state with respect to the native one. It should be noted, however, that because H-bonds involving betaine are generally shorter (and thus stronger) than those involving water (Supplementary Figure S7 and Table S3), the overall destabilization effect can be less pronounced than would be expected from simple H-bond counting. Indeed, correcting for differences in average energy of H-bonds formed by water and betaine results in total H-bond energy changes that are only slightly (7–30 kcal/ mol) smaller in high betaine concentrations than in pure water.

Betaine exclusion as a local and urea accumulation as a more general phenomenon

Prompted by the nonuniform distribution of the solvent components around the protein, seen already in the H-bond analysis, we computed the local-bulk partition coefficient to provide a more detailed picture of the exclusion and accumulation phenomena at the protein surface. The coefficient reflects the deviation from a uniform spatial distribution of co-solvent molecules in the vicinity of the solute and was defined as follows:

$$K_p = \frac{N_o N_w^b}{N_w N_o^b} \quad (3)$$

with N_o and N_w being the number of osmolyte and water molecules within a defined distance from the solute (here taken to be 0.55 nm) [82], and N_o^b and N_w^b being the number of osmolyte and water molecules in bulk solvent. A value of 1 indicates equal concentrations of the osmolyte in the protein solvation shell and in bulk solvent, whereas values higher and lower than 1 correspond to enrichment and exclusion of the osmolyte from the vicinity of protein surface, respectively.

Figure 4 compares the changes in the coefficient during unfolding, computed separately for the backbone and side chains of the exposed part of the protein.

It can be seen that betaine and urea differ vastly in terms of their preference for the exposed backbone, consistent with the view that prevails in the literature [30,39,40,83]. Namely, in the solvation shell of the backbone, the fraction of urea molecules is ~1.5- (in the folded state) to 2-fold (in the unfolded state) higher compared with bulk solvent, while the concentration of betaine decreases by up to 50%.

The differences are smaller in the environment around the side chains, with the urea content ~1.5 times higher than in the bulk, and the preferential interaction coefficient is roughly equal to 1.0 for betaine in all cases. This suggests that the exclusion observed for betaine arises only locally, that is, close to the backbone, while the accumulation of urea — albeit highest at the backbone — also extends into the side chain region. Notably, osmometric measurements reported recently for betaine and similar osmoprotectants (TMAO and sarcosine) also point to the conclusion that the stabilizing effect of osmolytes is due to their exclusion from the protein backbone rather than from the side chains [39,40,44,45,83]. Figure 4 also reveals that during the course of unfolding, and in particular close to the rupture point at 1.7 nm, urea becomes more abundant at the protein surface, as the preferential interaction coefficient is greater by ~0.5 for the unfolded than for the folded lysozyme. Betaine behaves differently, as the coefficient remains roughly constant during the folded–unfolded transition. These tendencies indicate that while urea has an even

stronger preference for the regions that become exposed to solvent upon unfolding — most notably, the backbone and hydrophobic side chains — betaine exhibits similar preference for the folded and unfolded ensemble. In the case of urea, the enrichment is then amplified by an increase in solvent-exposed surface area during protein unfolding.

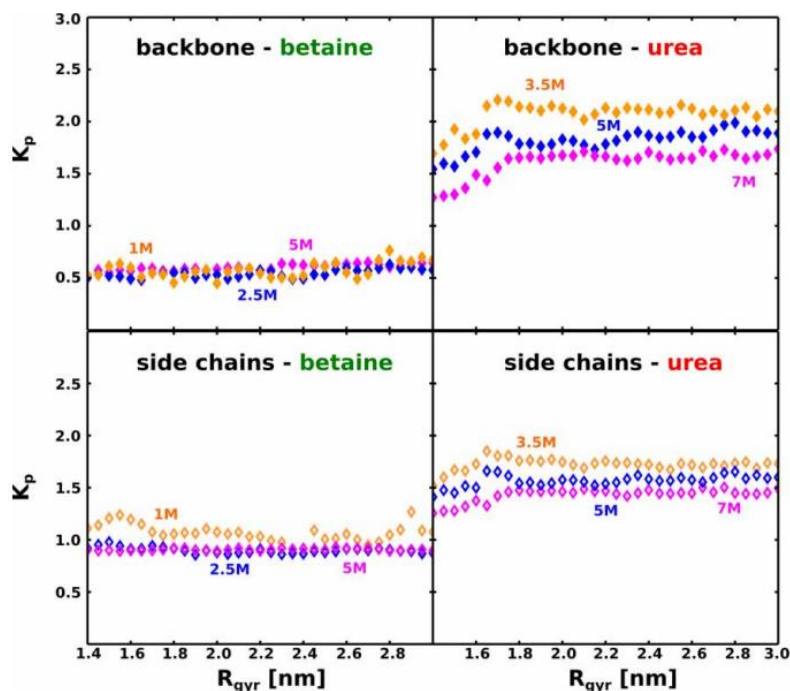


Figure 4. Analysis of exclusion and accumulation of osmolytes at the protein-solvent interface.

The local-bulk partition coefficient was calculated for the solvent molecules with respect to the protein (backbone and side chains) as a function of R_{gyr} , in different urea and betaine concentrations. All values were averaged over five independent simulations.

Since the popular approach to discuss osmolyte effects in terms of contributions from backbone and side chains [27,30,39–41] can be ambiguous due to the diverse chemical nature of the latter, we additionally dissected individual contributions from basic (b), acidic (a), polar (p), hydrophobic aromatic (h. ar) and hydrophobic aliphatic (h. al) residues to the overall K_p . This approach further facilitated a comparison of our results with available experimental and computational data [42,84–86]. Values of K_p determined for the folded and unfolded state are reported in Supplementary Table S6. Consistent with previous observations, urea accumulates at the protein surface, giving rise to K_p much larger than one. Strong urea enrichment is observed especially in the proximity of the backbone (K_p in the range of 1.44–1.90 for the folded state) and hydrophobic amino acids ($K_p = 1.44$ –1.90). A slightly lower accumulation of urea can be seen in the vicinity of polar ($K_p = 1.34$ –1.73) and basic ($K_p = 1.31$ –1.63) residues, with the solvation shell of acidic side chains being the least enriched in urea ($K_p = 1.11$ –1.46). Note that K_p values markedly larger than 1 typically decrease with concentration, as regions with high preference for urea saturate already at low concentrations, and the amount of co-solvent in their solvation shells does not change significantly with increase in bulk concentration.

On the other hand, interactions of betaine with the protein surface are mostly unfavorable ($K_p < 1$), with an exception of basic residues that were found to attract betaine molecules quite strongly ($K_p = 1.23$ –1.96). Among other individual residue types, betaine is strongly excluded from the backbone ($K_p = 0.62$ –0.83), hydrophobic ($K_p = 0.68$ –0.79) and acidic ($K_p = 0.75$ –0.84) residues, and roughly indifferent to the presence of polar side chains ($K_p = 0.86$ –0.94).

Notably, the K_p values for urea are in good qualitative agreement with a recent report by Holehouse et al. [86], where the action of denaturants was simulated in the context of small protein domains, even though the study used different force field parameters for urea (OPLS) from those in our case (CHARMM). Such

consistency indicates that the observed behavior of urea is relatively insensitive to the choice of force field. In turn, a comparison with the experimental data of Record and colleagues [42,84,85] reveals a substantial systematic offset in the computed values — most probably due to methodological differences, as the reference study used isolated small molecules to infer values of K_p for individual atom types, and the assumed additivity need not hold for a diverse environment of the protein surface. One significant outlier is the aromatic nonpolar side chains, which were found to strongly attract betaine molecules in the reference study (K_p of 1.62), while effectively repelling betaine in our simulations (K_p of ~ 0.75). We note that, even though the large discrepancy can be partially explained in terms of nonadditivity (in the unfolded state, the more isolated aromatic residues have larger K_p of up to 1.03), it is possible that existing nonpolarizable atomistic models do not describe the attractive nature of, for example, cation– π interactions sufficiently well. One would need to resort to *ab initio* explicit solvent models to rectify this issue.

To spatially resolve the observed interaction preferences, we calculated the radial preference function of osmolyte molecules around the unfolded state of lysozyme, for the whole protein, as well as separately for the backbone and side chains. The radial preference function was defined as the ratio of osmolyte and water molecules found at a distance r from the protein surface, divided by the corresponding ratio in the bulk solvent, according to a formula analogous to eqn (3):

$$f_o(r) = \frac{N_o(r) N_w^b}{N_w(r) N_o^b} \quad (4)$$

Thus, by construction, deviations from unity in the preference function can be interpreted in the same way as for the preferential interaction coefficient. Also, to investigate the exclusion and accumulation in the vicinity of different amino acid types, we calculated the preference function separately for acidic, basic, hydrophobic and polar residues in the intermediate osmolyte concentrations, that is, in 2.5 M betaine and 5.0 M urea solutions.

As seen in Figure 5, and consistent with the above discussion, betaine molecules are excluded from the exposed backbone. Within 0.5 nm of the backbone, the concentration of betaine is ~ 2 times lower compared with the bulk solvent, at least in 1.0 and 2.5 M solutions. The exclusion effect is least pronounced for the 5.0 M solution, indicating that, at higher concentrations, betaine has a stronger tendency to distribute more uniformly, possibly due to overlapping of the solvation shells of betaine molecules.

On the other hand, betaine molecules approach the exposed side chains easily, forming favorable contacts at two distinct intermolecular distances that correspond to two main modes of betaine–protein interaction, seen as peaks in Figure 5C. As the decomposed distributions suggest (Figure 5D), the highest peak at 0.3–0.4 nm almost exclusively corresponds to direct salt bridges between the carboxyl moiety of betaine and side chains of basic residues. Similarly, the second, much smaller peak at 0.5–0.6 nm can be largely attributed to the direct charge–charge interaction between the trimethylammonium group of betaine and acidic residues. Owing to the three methyl groups on the nitrogen atom of trimethylammonium, in this ionic pair, the positive charge is more delocalized and separated from the negative ion and hence, the interaction is significantly weaker. Interestingly, while the distribution for polar residues has two peaks that coincide with those for charged side chains, their height suggests that, in contrast with charged side chains, polar side chains (similar to hydrophobic) cannot attract and reorient betaine molecules efficiently. This, in turn, indicates a region-specific and strongly sequence-dependent orientational distribution of betaine at the surface of the unfolded protein, suggesting also that relatively few binding sites on the protein can accommodate a betaine molecule in an enthalpically favorable manner. As proposed above, this scarcity of osmolyte-binding sites could result in effective exclusion of betaine from the vicinity of the backbone, even though, occasionally, betaine–backbone association may be enthalpically favorable (see the Discussion of decomposed interaction energetics below).

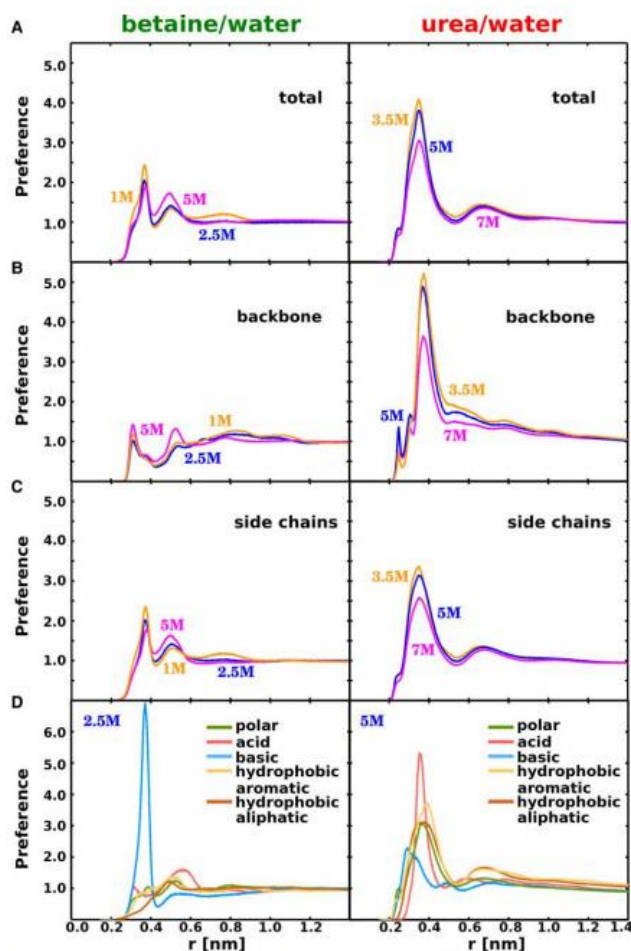


Figure 5. Radial preference functions of betaine and urea with respect to the protein.

Radial preference functions were calculated with respect to: (A) the whole protein molecule, (B) protein backbone and (C) protein side chains. (D) Radial preference functions decomposed into contributions from: polar (green), acid (red), basic (blue) and hydrophobic aromatic (orange) and hydrophobic aliphatic (brown) side chains in the intermediate osmolyte concentrations. In the chosen concentrations, betaine and urea occupy comparable volume fractions of the solvent.

Unlike betaine, urea strongly accumulates in the vicinity of the protein (Figure 5, right). Consistent with the already discussed hydrogen-bonding patterns, the urea is strongly attracted toward the backbone (local concentrations up to 3.5–5.0 times higher than in the bulk), and the enrichment is only slightly lower (up to 2.5–3.5 times) close to the side chains. Also, the peaks in the distribution functions are broader than in the case of betaine, indicating less constrained binding, i.e. protein-bound urea can assume more orientations than protein-bound betaine. Multiple factors contribute to the urea orientational and positional flexibility at the protein surface, as illustrated by the decomposed distributions in Figure 5D. For basic residues, the broader distribution reflects the possibility of forming a hydrogen bond to the charged head group and/or a hydrophobic contact with the aliphatic part of the side chain. On the other hand, the sharper and higher peak seen for acidic side chains is attributed to the fact that they interact with urea exclusively through hydrogen bonding, which, however, is entropically favored due to eight distinct ways of pairing two oxygen atoms of a carboxylic group with four hydrogen atoms of urea. The broad peak found for polar residues indicates that all kinds of donor–acceptor pairs and less specific contacts are formed in this case, which also contributes to the orientational freedom of urea at the protein–solvent interface. Finally, abundant contacts exist between urea and hydrophobic side chains, and these are nonspecific and spatially heterogeneous by nature, as confirmed again by the width of the distribution. In general, well-defined and wide peaks for uncharged residues (especially hydrophobic ones) indicate that urea indeed has a preference for the residues that form the core of the protein, as remarked above.

In the above discussion, we mentioned two potential mechanisms that relate the observed contact preference to the denaturing effect of urea: one based on weakening of the hydrophobic effect that drives protein folding [21,87,88], and another in which urea stabilizes the unfolded state by interacting favorably with the exposed protein core via van der Waals' forces [27–29,89]. Intriguingly, the distributions in Figure 5 show a decreasing but significant enrichment of urea that persists up to 1 nm from the protein surface, which suggests a picture of a diffuse urea-enriched region that surrounds the lysozyme molecule. One might expect this urea-rich shell to reduce water-mediated hydrophobic attraction between nonpolar protein surfaces, as the shell provides a local environment where water, urea and hydrophobic residues can favorably interact with each other. This supports the mechanism based on decreased hydrophobic forces; however, the other scenario, involving favorable protein–urea dispersion interaction, can still hold true, as they are not mutually exclusive and can in fact act synergistically in causing protein denaturation.

As also evident from Figure 5, at a given distance from the protein, the radial preference function can either increase or decrease with the osmolyte concentration. An increasing trend observed for certain regions of the protein surface means that, at low osmolyte concentrations, these regions are not readily solvated by the osmolyte, and the preference increases with increase in the bulk osmolyte concentration.

The regions that give rise to such an increasing trend have relatively low affinity for the osmolyte, and in the case of betaine, this corresponds almost to the entire protein surface — both the protein backbone and nonbasic side chains (as can be seen from comparing panels C and D on the left of Figure 5). In contrast, preference that decreases with the increasing osmolyte concentration implies a 'saturation' effect. In that case, increasing concentrations in the bulk barely affects the (already saturated) concentration in a local domain around the protein, meaning that the preference (being the ratio of the local and bulk concentrations) effectively decreases. As seen in Figure 5, for urea, these high-affinity regions correspond to the backbone and virtually all types of the side chains, as the preference function for the 7.0 M solution lies below the 3.5 M curve almost everywhere, except for the bulk solvent where the curves overlap by definition. For betaine, such high-affinity regions only exist at the basic side chains and at large distances (0.6–0.8 nm) from the protein, corresponding largely to the second solvation shell around the basic regions.

Analysis of interaction energies reveals the nature of osmolyte accumulation and exclusion

Recent theoretical considerations led to a conclusion that the overall nature of the stabilizing or destabilizing effect of osmolytes — either enthalpic or entropic — primarily depends on the contributions that dominate the effective osmolyte–solute interaction [53]. Hence, to investigate the nature of the driving forces behind the exclusion and accumulation of osmolytes, we first calculated the interaction enthalpies [$\Delta H_o(r)$ and $\Delta H_w(r)$] between a single osmolyte or water molecule and the environment, i.e. the protein and the rest of the solvent, as a function of the molecule's distance from the protein surface r (Supplementary Figure S8; see also captions in Figures 6 and 7 for details). From these profiles, we computed an effective interaction enthalpy profile by including the effect of solvent displacement, i.e. taking into account the fact that an approaching osmolyte molecule displaces a certain number of water and osmolyte molecules that would reside there otherwise. For betaine and urea, this effective enthalpy $\Delta H(r)$ was calculated as $\Delta H_o(r)(1 - \gamma_o(r)) - q\Delta H_w(r)(1 - \gamma_o(r))$, where $\gamma_o(r)$ is the volume fraction of the osmolyte at a given distance r from the protein and q is the ratio of average molar volumes of osmolyte and water (4.17 in the case of betaine and 2.53 for urea). Subsequently, to relate the enthalpy changes to the actual distribution of the molecules around the protein, we computed the corresponding free energy profiles for protein–solvent association as $\Delta G(r) = -kT \ln \rho(r)$, where $\rho(r)$ is the protein–solvent radial distribution function. The translational contribution to the free energy was subtracted by employing a normalization scheme based on the surface area of the protein, as measured by a probe of radius r , which led to the free energy profiles that are flat at large distances from the protein surface. Eventually, by subtracting $\Delta H(r)$ from $\Delta G(r)$, we computed the entropic contribution to the association free energy [denoted as $-\Delta S(r)$], which includes water-mediated

hydrophobic interactions and the abundance of solvent-binding sites at distance r that correspond to an average enthalpy of $\Delta H(r)$.

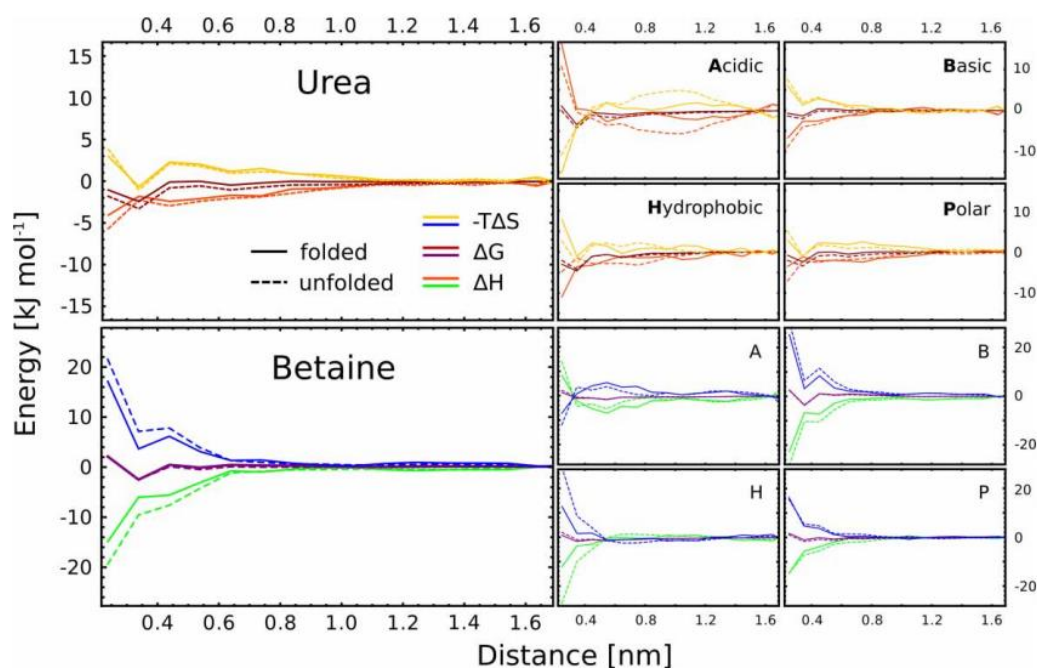


Figure 6. ΔG , ΔH and $-T\Delta S$ profiles for a single solvent molecule with respect to the protein surface.

The free energy (ΔG), enthalpy (ΔH) and entropy ($-T\Delta S$) changes were computed for the 2.5 M betaine and 5.0 M urea solutions, as a function of the distance from the protein surface.

Figure 6 shows the resulting profiles for osmolyte molecules, computed as a function of the distance from the protein surface in the systems with intermediate osmolyte concentrations (2.5 M betaine and 5.0 M urea). To distinguish between different types of amino acid environments at the protein surface, the profiles were also determined with respect to acidic, basic, polar and hydrophobic residues and, separately, with respect to the backbone (Figure 7). The original $\Delta H_w(r)$ and $\Delta H_o(r)$ profiles used to compute $\Delta H(r)$ in Figures 6 and 7 are shown in Supplementary Figures S8 and S9.

As can be seen from Figures 6 and 7, approaching the protein surface is enthalpically favorable for a urea molecule, consistently for the backbone and all residue types — except for acidic side chains, where the high cost of desolvation disfavors direct interactions with the osmolyte (see also Supplementary Figure S8). For the protein as a whole, the average effective enthalpic gain of roughly -4 – 6 kJ/mol, arising due to H-bonding and van der Waals' interactions, remains favorable in terms of free energy despite the cost of displacing water molecules from the protein surface (~ 2.5 kJ/mol corresponding to ~ 2.5 water molecules replaced by a single urea molecule) and the loss of translational entropy (< 3 kJ/mol in the range of concentrations studied). Therefore, the favorable direct interactions with the protein may be thought of as a major driving force behind the observed accumulation of urea at the protein surface.

To investigate these interactions in more detail and to examine their role in the unfolding process, we decomposed the solvation enthalpy change upon unfolding into individual contributions due to interactions between the solvent components and different types of residues. The results, shown in Supplementary Figures S10 and S11, clearly indicate that both electrostatic and van der Waals' interactions between the accumulated urea and the protein favor the unfolding process. As should be expected, this energy gain, seen in particular for the numerous hydrophobic and polar residues that become exposed to the solvent, is largely compensated by the loss of urea–water and urea–urea interactions (Supplementary Figure S12). The comparison with pure water (Supplementary Figure S10) shows, on the one hand, that the electrostatic contribution to the solvation enthalpy, related to the number and strength of protein–solvent hydrogen

bonds, is very similar regardless of the presence or concentration of urea. On the other hand, the van der Waals' interactions with the newly solvent-exposed side chains and backbone are much stronger for urea than for water and this difference results in the more negative solvation enthalpy change in the presence of urea. This observation supports the above argument that van der Waals' interactions rather than hydrogen bonds provide a critical enthalpic driving force for urea accumulation at the protein surface and hence for the urea-induced protein denaturation, which is consistent with previous reports [23,27,28,43].

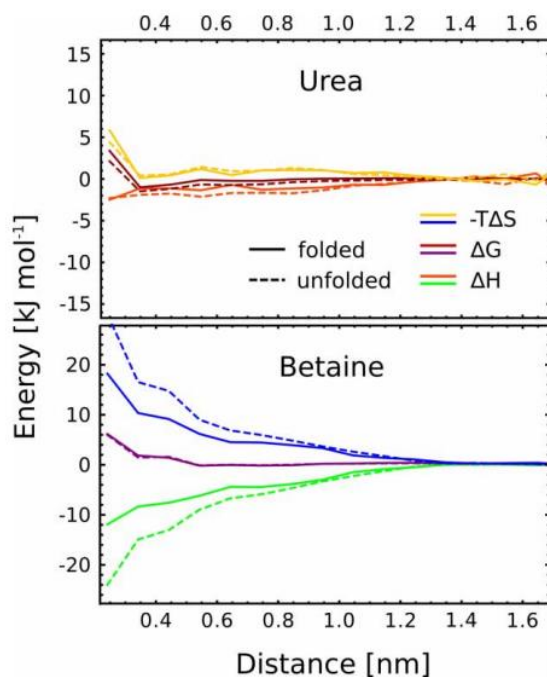


Figure 7. ΔG , ΔH and $-T\Delta S$ profiles for a single solvent molecule with respect to the protein backbone.

Free energy (ΔG), enthalpy (ΔH) and entropy ($-T\Delta S$) changes were for the 2.5 M betaine and 5.0 M urea solutions, as a function of the distance from the protein backbone.

Figure 6 further reveals that the enthalpic stabilization of the protein-bound urea is noticeably lower (by ~ 5 kJ/mol) in the vicinity of hydrophobic residues in the unfolded state and that, in this case, the broad minimum in the $\Delta H(r)$ curve is shifted by 0.4–0.6 nm toward the bulk solution. Even though the enthalpic driving force at hydrophobic patches favors the attraction of urea molecules toward the protein, the overall enthalpy change might not be sufficient to account for the preference of these regions for urea indicated by the $\Delta G(r)$ profile (see Figure 6). Indeed, a pronounced minimum in the $-T\Delta S(r)$ curve (yellow dashed line in Figure 6H) coinciding with the free energy minimum strongly suggests that urea accumulation around hydrophobic patches is in part also entropically driven. In fact, as already suggested above, a urea-rich region adjacent to the protein surface can be thought of as a buffer to reduce the (mostly entropic) cost of exposing hydrophobic residues buried in the core to the solvent. Also consistent with this explanation is the fact that the interaction enthalpy for a water molecule has a broad minimum at slightly larger distances (0.4–0.8 nm) from hydrophobic residues in the unfolded state, in the urea-rich region around the protein (Supplementary Figure S8). This finding supports the picture in which accumulated urea molecules, by interacting favorably with the protein surface and, at the same time, with water, create a local environment that effectively weakens hydrophobic forces holding nonpolar groups together in the native protein structure.

Interaction analysis also sheds light on the molecular underpinnings of the protein-stabilizing effect of betaine, and possibly other methylamines. First, Figure 7 demonstrates that even though betaine does not preferentially associate with the exposed protein backbone (see also Figure 5), unexpectedly, this association is accompanied by a favorable enthalpy change (ΔH in the range of -20 to -10 kJ/mol). This

enthalpy gain is apparently more than offset by the positive $-T\Delta S$ contribution, which makes the binding free energy unfavorable, supporting our claim that the exclusion of betaine results from the scarcity of binding sites capable of accommodating the ligand-like osmolyte molecule close to the backbone in an energetically optimal manner. Hence, our results point toward the first of two possible scenarios proposed above, where the exclusion mostly results from the effective repulsion due to an entropic factor that opposes protein-co-solvent association. However, while the canonical hard-sphere model assumes entirely steric nature of this repulsive force and hence a 'pure' excluded volume effect, in our description, the exclusion results mostly from the low availability of betaine-binding sites on the protein surface.

Accordingly, for the side chains, which are generally more accessible to the osmolyte molecules, no significant exclusion of betaine is observed in Figure 6 (flat free energy curve), even though in this case the corresponding enthalpy change is similar to (acidic residues) or smaller than (polar and hydrophobic residues) that for the backbone. It can be also seen from Figure 6 that the large minimum in the overall free energy profile of approximately -4 J/mol, corresponding to accumulation of betaine at ~ 0.35 nm, can be almost exclusively ascribed to the formation of salt bridges between the positively charged side chains and the carboxyl group of betaine.

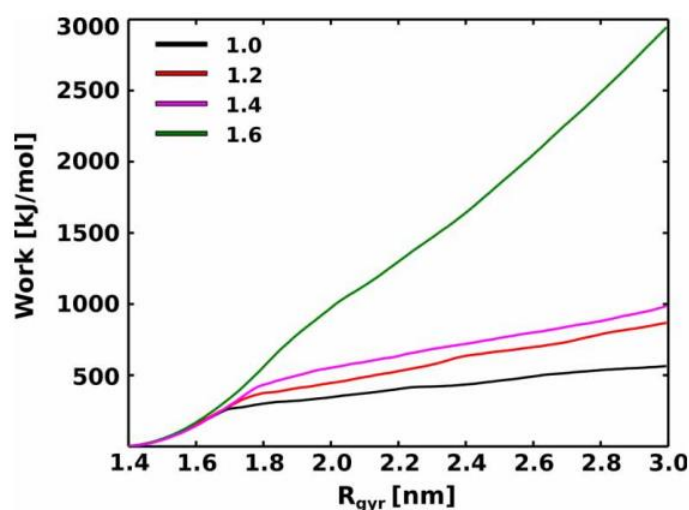


Figure 8. Changes in unfolding work upon the rescaling of partial charges in the betaine molecule.

The plot shows the amount of work required to unfold lysozyme in the presence of 5 M betaine with its partial charges scaled by a common factor ranging from 1.0 to 1.6, at the pulling speed of 5×10^{-6} nm ps⁻¹.

Aside from the dominant entropic contribution to the co-solvent-induced stabilizing effect, a growing body of evidence [46,54,81,90] points toward the existence of an enthalpic effect associated with the action of protecting osmolytes. This enthalpic contribution might arise due to changes in protein solvation, as already suggested by several researchers [81]. Indeed, in the H-bonding analysis, we found that betaine present as a co-solvent noticeably reduces the propensity of water to form hydrogen bonds with the protein, which is especially evident for the backbone, solvated almost exclusively by water. Supplementary Figure S9 provides additional insights into the observed reduced solvation effect, by showing that the enthalpic driving force for water-protein association is considerably smaller in the presence of betaine ($\Delta H_w \approx 0$) than in pure water ($\Delta H_w \approx -3$ to -2 kJ/mol). This difference indicates that the weaker hydrogen-bonding ability of water (or, more generally, the reduced activity of water) is caused by its strong interactions with betaine, consistent with the FTIR spectroscopy and molecular dynamics data showing a pronounced enhancement of the water structure upon the addition of betaine [74,91,92]. Interestingly, the analogous FTIR approach [21] and other experimental studies [23,28,87] have found that, unlike betaine, urea does not significantly affect the hydrogen-bonding network of water and is readily accommodated in liquid water structure. In agreement with these reports, the comparison of the original interaction enthalpy profiles ($H_w(r)$) for water

molecules in Supplementary Figure S9 shows that, in contrast with water-perturbing betaine, urea changes the energetics of interfacial water only very slightly ($\Delta\Delta H_w < 1$ kJ/mol).

Since the decreased solvation of the backbone and uncharged side chains in the presence of betaine necessarily disfavors the protein unfolding, the above findings support the notion that the betaine-induced stabilization is partially enthalpic in nature, as has been recently suggested [46,54,81,90]. Our data further indicate that the positive contribution to the enthalpy of unfolding is due to strong water–osmolyte interactions that reduce the ability of water molecules to solvate the newly exposed protein patches.

To additionally test the hypothesis about the enthalpic origin of protein stabilization by betaine, we scaled the partial charges of all betaine atoms by a common factor ranging from 1.0 to 1.6, and performed the mechanical unfolding simulations in 5 M betaine in the same manner as previously, at the lowest pulling speed of 5×10^{-6} nm ps⁻¹. As expected, the water-binding capacity increases with the polarity of the osmolyte molecule, as seen from the radial distribution functions in Supplementary Figure S13. Figure 8 shows that scaling up the charges led to a significant enhancement of the stabilizing effect of the co-solvent, as reflected in the increase in the unfolding work. This finding supports the notion that strong water binding might be partially responsible for the enthalpic contribution to the betaine-induced stabilization.

Conclusion

In this work, we studied the molecular origin of the perturbing effect of two osmolytes, urea and betaine, on the protein folding equilibrium. We simulated the mechanical unfolding of a lysozyme molecule in pure water and in aqueous solutions of both osmolytes, and found that our model correctly reproduced the stabilizing effect of betaine and the denaturing effect of urea, yielding larger and smaller values of unfolding work, respectively. Subsequently, to analyze the effect of osmolytes on the calculated work of unfolding, we investigated the molecular determinants of protein stability in the studied solutions. In particular, we traced the formation of new protein–osmolyte and protein–water hydrogen bonds during the course of unfolding. We also quantitatively assessed the magnitude and spatial patterns of osmolyte exclusion and accumulation at the protein surface, and complemented these results with a detailed analysis of the enthalpy of interaction and its changes upon denaturation for individual components of the systems studied.

We observed that, in our model, urea accumulates in the vicinity of the protein, which according to most previously reported experimental and computational results is directly associated with the denaturing effect [13,40,42]. Intriguingly, however, we found that the concentration of urea increases not only at the protein surface, but rather remains elevated at distances of up to 1.0–1.5 nm from the protein, indicating the presence of a diffuse urea-rich solvation sphere around the lysozyme molecule. Also, while accumulating urea plays a major role in solvating the protein core that becomes exposed upon denaturation, the overall strength of newly formed protein–osmolyte and protein–water hydrogen bonds in urea solutions is in fact similar to that observed in pure water. This indicates that, contrary to certain recent claims [38], hydrogen bonding cannot account solely for the denaturing effect of urea. Instead, our data point to two other driving forces for urea-induced denaturation. First, we noted that the unfolded state is indeed stabilized by favorable direct interactions between urea molecules and the residues forming the protein core. As revealed by an analysis of the enthalpic terms and the accumulation patterns, this stabilization arises because urea forms favorable contacts with the protein in multiple spatial arrangements, regardless of the type of residues involved. Besides acting as both H-bond acceptors and donors, urea molecules also interact favorably with nonpolar residues by dispersion forces, stabilizing the unfolded conformation in which those residues are exposed to the solvent. Second, the results indicate that the diffuse urea-rich shell that forms around the protein is capable of accommodating well both water molecules and nonpolar protein residues, effectively screening the protein from the water-mediated hydrophobic force that drives protein folding.

In the case of betaine, we found that the osmolyte molecules are excluded from the protein surface, which is commonly believed to lead to the stabilization of proteins by betaine and other compatible osmolytes. We also observed that the exclusion is most pronounced in the vicinity of the backbone, in agreement with recent findings [39,40]. At the same time, the zwitterionic betaine molecules were found to accumulate near charged, mostly basic, residues, giving rise to an inhomogeneous and variable distribution on the protein surface.

Our data also indicate that even though the protein–betaine association is overall favorable in terms of enthalpy, it occurs infrequently, betaine forms very few new contacts with the protein upon unfolding and the newly exposed protein surface, consisting largely of the backbone and hydrophobic side chains, is mostly solvated by water. Based on our findings, we postulate that betaine — a rather bulky molecule with a large dipole moment — exhibits a ligand-like behavior, i.e. it can only be accommodated in an enthalpically favorable manner at relatively few binding places within the protein–solvent interface. This scarcity of binding sites in turn results in an effective exclusion of betaine from the protein backbone and uncharged side chains that is entropic in nature. Such a picture provides a molecular-level reinterpretation of the canonical entropy-dominated model of protein stabilization by protecting osmolytes. Moreover, our data suggest that the presence of betaine in a solution decreases the enthalpic driving force for protein hydration, since water molecules interact favorably with betaine in bulk solvent, as proposed in a recent study [47]. The resulting weakening of protein solvation (when compared with pure water) can give rise to an enthalpic contribution to the stabilizing effect of betaine, already reported by several researchers [46,54,81,90].

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