ABSTRACT: Protein hydration shell dynamics play an important role in biochemical processes including protein folding, enzyme function, and molecular recognition. We present here a comparison of the reorientation dynamics of individual water molecules within the hydration shell of a series of globular proteins: acetylcholinesterase, subtilisin Carlsberg, lysozyme, and ubiquitin. Molecular dynamics simulations and analytical models are used to access site-resolved information on hydration shell dynamics and to elucidate the molecular origins of the dynamical perturbation of hydration shell water relative to bulk water. We show that all four proteins have very similar hydration shell dynamics, despite their wide range of sizes and functions, and differing secondary structures. We demonstrate that this arises from the similar local surface topology and surface chemical composition of the four proteins, and that such local factors alone are sufficient to rationalize the hydration shell dynamics. We propose that these conclusions can be generalized to a wide range of globular proteins. We also show that protein conformational fluctuations induce a dynamical heterogeneity within the hydration layer. We finally address the effect of confinement on hydration shell dynamics via a site-resolved analysis and connect our results to experiments via the calculation of two-dimensional infrared spectra.

INTRODUCTION

The hydration shell of a protein is known to have a critical influence on protein structure and function. In particular, the dynamic properties of the hydration shell play a role in biochemical processes including protein folding, enzyme function, and molecular recognition.1−3 A complete understanding of such processes therefore requires a detailed picture of protein hydration shell dynamics.

It has been shown both by experiments4−16 and simulations17−27 that the proximity of a biomolecule such as a protein perturbs water dynamics in its hydration shell. However, our understanding of this perturbation remains incomplete, and questions such as the magnitude and molecular origins of the perturbation are still actively discussed. Some studies, including, e.g., NMR4,6 and molecular dynamics25,26 results, indicate that, for the majority of the hydration shell, the water reorientation dynamics is moderately slowed down, by a factor of 2–3, compared to bulk water. This is in contrast to, e.g., time-resolved fluorescence spectroscopy,7,28 which suggests that a significant proportion of the water population is slowed down by at least an order of magnitude. In addition to the magnitude of the slowdown, its molecular origin is still not well established. The distribution of the retardation factor is known to be heterogeneous across the protein surface,8,10,25,29 and a complete understanding of this heterogeneity requires spatially resolved information on hydration shell dynamics, which to date has come from fluorescence spectroscopy,8,30 from NMR experiments,5,9,10,20,25−27,31 and from molecular dynamics studies.32,33,34 One such site-resolved computational study by one of us25 has shown that, for most water molecules within the hydration shell of the protein lysozyme, the dynamical perturbation is mainly due to an excluded volume effect dependent on local surface topology. A question then arises regarding the generality of conclusions drawn from a study of the hydration shell of any one protein.

Here, we expand the previous study of reorientational hydration shell dynamics recently presented for the enzyme lysozyme25 to three other proteins with very different sizes and functions, acetylcholinesterase, subtilisin Carlsberg, and ubiquitin, in order to examine the applicability of our previous results to any given protein. We use molecular dynamics simulations to access site-resolved information on hydration shell dynamics via a decomposition of the protein surface into sites of different chemical nature. We elucidate the molecular origins of the perturbation induced by each protein, using a theoretical framework previously established for water dynamics next to solutes, including proteins.25,33,34 We then go on to discuss the applicability of our conclusions to globular proteins in general, and to explore the effect of protein conformational fluctuations on hydration shell dynamics.

An understanding of the effect of confinement on protein hydration shell dynamics is also required, in order to provide a...
Table 1. Protein Molecular Weight, Secondary Structure in Terms of Helical and β-Sheet Composition, Surface Composition in Terms of the Total Time-Averaged OH-Bond Population of the Hydrophobe, H-Bond Donor or H-Bond Acceptor Sites, and Total Charge

| Protein             | Molecular Weight (kDa) | Secondary Structure | Relative OH-Bond Population | Total Charge |
|---------------------|------------------------|---------------------|----------------------------|--------------|
| ubiquitin           | 9                      | 23%                 | 34%                        | 68% 15% 17% 0 |
| lysozyme            | 14                     | 40%                 | 10%                        | 72% 15% 13% 0+8 |
| subtilisin Carlsberg| 27                     | 28%                 | 19%                        | 71% 13% 16% 0+1 |
| acetylcholinesterase| 59                     | 36%                 | 17%                        | 74% 14% 12% 0−9 |

The Journal of Physical Chemistry B

**METHODOLOGY**

**Simulation Details.** We performed molecular dynamics simulations of dilute aqueous solutions of four globular proteins, which cover a wide range of functions and molecular weights. This includes three enzymes: acetylcholinesterase (59 kDa), an esterase whose biological role is to break down the neurotransmitter acetylcholine, subtilisin Carlsberg (27 kDa), a serine protease (i.e., an enzyme that hydrolyzes peptidic bonds), and lysozyme (14 kDa), a glycoside hydrolase that breaks glycosidic bonds in bacterial cell walls.37 The fourth system is a regulatory protein, ubiquitin (9 kDa), which tags proteins for destruction and also directs protein transport.37,38

This choice was motivated by prior studies of hydration dynamics around these systems which employed different techniques and led to some conflicting conclusions.5,7,23,−25,27,31,35 and by the wide range of functions and sizes covered by these four proteins. Table 1 lists some of their key structural properties, and Figure 1 shows the great heterogeneity of their surface charge distributions.

The initial protein configurations were obtained from the crystallographic structures with PDB codes 4ARA (acetylcholinesterase), ISCN (subtilisin Carlsberg), 2LYM (lysozyme), and 1UBQ (ubiquitin). Each protein was solvated in a simulation box adapted to its size, containing between 4982 water molecules for ubiquitin, the smallest protein, and 28626 water molecules for acetylcholinesterase, the largest, corresponding to effective concentrations in the millimolar range, respectively, 1.8, 5.6, 5.3, and 10.3 mM for acetylcholinesterase, subtilisin, lysozyme, and ubiquitin. The proteins were described using the CHARMM22 force field with CMAP corrections.39

The water force field was SPC/E40 for lysozyme, subtilisin, and acetylcholinesterase systems and TIP4P/200541 for the ubiquitin system. These water force fields were chosen because they have been shown to correctly reproduce the dynamics of water at room temperature.36,42 However, of the two water models, only TIP4P/2005 provides a qualitatively correct description of water’s phase diagram.41 TIP4P/2005 was therefore used in one protein system in order to open the way to a future study of the temperature dependence of protein hydration shell dynamics. Since the comparison between results for the different proteins studied here is made via the ratio of hydration shell and bulk values, meaningful comparisons can be obtained from these different water models.

Simulations were performed using NAMD43 with periodic boundary conditions, at densities determined via equilibration in the NPT ensemble. Long-range electrostatics were treated using the particle mesh Ewald method. Switching functions were applied to nonbonded interactions from 10 Å, with a cutoff of 12 Å. Bonds between hydrogen and heavy atoms were constrained using the SHAKE and SETTLE algorithms. Simulations with pressure control used the Nosé−Hoover Langevin piston with a piston period of 100 fs and a damping time scale of 50 fs. Simulations with temperature control used the Langevin thermostat with a damping coefficient of 1 ps−1. All systems were equilibrated in the NPT ensemble at 300 K and 1 atm for at least 0.5 ns, followed by equilibration in the NVT ensemble at 300 K for at least 1 ns. Finally, production runs were between 4 and 20 ns long. Coordinates were output every 100 fs. Production runs for lysozyme, subtilisin, and acetylcholinesterase systems were in the NVT ensemble at 300 K with a 2 fs time step. The production run for the ubiquitin system was in the NVE ensemble with a 1 fs time step, and the resulting average temperature was 300 ± 2 K. Again, this
difference was due to the use of the ubiquitin system in a temperature-dependence study.

We also studied the effect of confinement on hydration shell dynamics using systems containing subtilisin Carlsberg in hexane solvent at three hydration levels. Simulation details were identical to those for subtilisin in aqueous solution. The hexane molecules were described using standard CHARMM parameters.39

**Analysis of Water Dynamics.** We analyze the molecular dynamics trajectories to provide a site-resolved analysis of water reorientational dynamics in the protein hydration shell, as outlined below.

We focus on the dynamics of individual water molecules and monitor the reorientation of a water molecule by following the dynamics of the water OH-bond vector $u$, via the second-order Legendre polynomial time-correlation function (tcf)33

$$C_2(t) = \langle \hat{P}_2[u(t) \cdot u(t)] \rangle$$

(1)

This is related to experimentally accessible values, namely, anisotropy decays from ultrafast infrared spectroscopy, and orientation relaxation times from magnetic relaxation techniques.33 After a sub-picosecond decay due to fast librational relaxation, the reorientational tcf is monoexponential for homogeneous systems such as bulk water at ambient temperature,33 while non-monoexponential is an indication of heterogeneity in the water dynamics. For tcf's calculated for a subset of water molecules with homogeneous water dynamics, the reorientation time $\tau_{\text{reor}}$ can be extracted via an exponential fit, performed here over the interval 2—10 ps in order to avoid contributions at short times from librational motions, and contributions at long times from water molecules which are no longer in the same environment as at the time origin.

The underlying mechanism of bulk water reorientation has been shown to be dominated by hydrogen-bond (H-bond) partner exchange via large-amplitude angular jumps from initial to final H-bond acceptors.33,34 It has been demonstrated that this is true not only in the bulk but also in the hydration shell of a range of solutes,33 including proteins.25 H-bond partner exchange is an activated process, passing through a transition state, and can usefully be seen as a chemical reaction. Jump kinetics can be followed via the cross time-correlation function33 between stable states44 I (initial) and F (final)

$$C_{\text{jump}}(t) = \langle n_1(t) n_F(t) \rangle$$

(2)

where $n_1 = 1$ if the OH bond is in stable state $X$ (i.e., forming a stable H-bond with the initial or final acceptor, respectively) and $n_F = 0$ otherwise. Absorbing boundary conditions are used in the product state in order to ensure that only the first jump from each initial H-bond acceptor is considered. The jump time $\tau_{\text{jump}}$ is the inverse of the rate constant for the H-bond exchange process, and can be found by fitting $1 - C_{\text{jump}}(t)$ with an exponential exp$(-t/\tau_{\text{jump}})$.36

We perform a site-resolved analysis of hydration shell reorientational dynamics and jump kinetics for each protein system studied here. The spatial resolution is performed as follows. The protein surface is divided into H-bond acceptor, H-bond donor, and hydrophobic sites. The hydration shell is defined as containing all water OH groups that are H-bonded to or within the hydrophobic cutoff of these protein surface sites, with each OH group in the hydration shell being assigned to a particular site at each time step. In cases of ambiguity in the assignment of an OH group, sites are given the priority acceptor > donor > hydrophobe, as this has been shown to be the order of greatest influence on water reorientational dynamics.25 Individual hydrophobic distance or H-bond distance and angle criteria are determined for each protein site from radial distribution functions between water oxygen or hydrogen atoms and amino acid atoms, calculated via molecular dynamics simulations of amino acids in aqueous solution. Typical criteria for the assignment of an OH group to a surface site (and therefore to the hydration shell) are $R_{\text{CO}} < 4.5 \text{ Å}$ for a hydrophobic site and $R_{\text{DA}} < 3.5 \text{ Å}, R_{\text{AH}} < 2.5 \text{ Å},$ and $\theta_{\text{HDA}} < 30^\circ$ for a H-bond donor or acceptor site, where C is a protein carbon atom, O is a water oxygen atom, A is a H-bond acceptor atom, D is a H-bond donor atom, and H is a hydrogen atom either in the protein or in water. For the calculation of jump tcf's using eq 2, tighter H-bond criteria are used to define stable H-bond states in the stable state picture.36 Typical values are $R_{\text{DA}} < 3.0 \text{ Å}, R_{\text{AH}} < 2.0 \text{ Å},$ and $\theta_{\text{HDA}} < 20^\circ$. We include only the first hydration shell in our analysis, as the perturbation induced by a biomolecule has been shown to fall off rapidly with distance from the surface.17,18,23

Individual reorientational and jump tcf's are then calculated for the subset of water molecules next to each site at the time origin, and individual reorientation and jump times extracted. Distributions of reorientation and jump times are constructed by weighting each time value by the OH-bond population next to that site. All other site-resolved values and probability distributions in this work are calculated or constructed in the same manner. For each system in aqueous solution, values characterizing bulk water dynamics are extracted from the subset of water molecules which are initially farther than 15 Å from the protein surface. Typical values of $\tau_{\text{bulk}}$ and $\tau_{\text{jump}}$ are 2.5 and 3.3 ps, respectively, at ambient temperature.36

# HYDRATION SHELL DYNAMICS OF FOUR DIVERSE GLOBULAR PROTEINS IN DILUTE AQUEOUS SOLUTION

The reorientational time-correlation function (eq 1) averaged over all water OH groups initially present in the hydration shell is shown in Figure 2 for all four proteins in aqueous solution. It is highly non-monoexponential in each case, revealing the heterogeneity of hydration shell dynamics, i.e., the presence of a broad distribution of relaxation times. One can distinguish two different types of heterogeneity, which we refer to as spatial and dynamical heterogeneity, and which we define as follows. Spatial heterogeneity arises from the chemical heterogeneity (the protein surface has, e.g., charged, polar, and nonpolar groups) and topological heterogeneity (the protein surface contains, e.g., troughs, pockets, and protrusions) of a static protein surface. As shown below, it is the main cause of heterogeneity in protein hydration shell dynamics. Dynamical heterogeneity arises from the dynamic nature of the protein as it samples its conformational space. In other words, a single protein surface site can induce a varying perturbation of water dynamics as the local conformation of the protein surface fluctuates. The following two subsections respectively address these two types of heterogeneity.

**Spatial Heterogeneity. Distributions and Mapping.** The spatial heterogeneity within a protein hydration shell arises from the great variety of exposed groups and local topologies at the protein surface, which leads to a broad distribution of reorientation slowdown factors $\rho_{\text{reor}}$ relative to the bulk situation, defined as

$$\rho_{\text{reor}} = \frac{\tau_{\text{reor,ss}}}{\tau_{\text{reor,bulk}}}$$

where $\tau_{\text{reor,ss}}$ is the reorientational time in the hydration shell and $\tau_{\text{reor,bulk}}$ is the time in the bulk.

$$\tau_{\text{reor,ss}} = \frac{1}{\rho_{\text{reor}}} \tau_{\text{reor,bulk}}$$

(3)

The hydration shell slowdown factor $\rho_{\text{reor}}$ can be calculated from the average of the reorientational tcf in the hydration shell $C(t)$ and that in the bulk $C_{\text{bulk}}(t)$

$$\rho_{\text{reor}}(t) = \frac{C(t)}{C_{\text{bulk}}(t)}$$

(4)

where $C_{\text{bulk}}(t)$ is the reorientational tcf averaged over all water OH groups initially present in the hydration shell.

$$\rho_{\text{reor}}(t) = \frac{\tau_{\text{reor,ss}}}{\tau_{\text{reor,bulk}}}$$

(5)

$$\tau_{\text{reor,ss}} = \frac{1}{\rho_{\text{reor}}} \tau_{\text{reor,bulk}}$$

(6)

$$\rho_{\text{reor}} = \frac{\tau_{\text{reor,ss}}}{\tau_{\text{reor,bulk}}}$$

(7)

$$\tau_{\text{reor,ss}} = \frac{1}{\rho_{\text{reor}}} \tau_{\text{reor,bulk}}$$

(8)

$$\rho_{\text{reor}}(t) = \frac{C(t)}{C_{\text{bulk}}(t)}$$

(9)

$$\tau_{\text{reor,ss}} = \frac{1}{\rho_{\text{reor}}} \tau_{\text{reor,bulk}}$$

(10)

$$\rho_{\text{reor}} = \frac{\tau_{\text{reor,ss}}}{\tau_{\text{reor,bulk}}}$$

(11)

$$\tau_{\text{reor,ss}} = \frac{1}{\rho_{\text{reor}}} \tau_{\text{reor,bulk}}$$

(12)

$$\rho_{\text{reor}} = \frac{\tau_{\text{reor,ss}}}{\tau_{\text{reor,bulk}}}$$

(13)

$$\tau_{\text{reor,ss}} = \frac{1}{\rho_{\text{reor}}} \tau_{\text{reor,bulk}}$$

(14)
This distribution has already been determined in the case of lysozyme,\textsuperscript{25} and we extend it here to our set of four diverse proteins. For each system, we focus on the first hydration shell, since it has been shown that the perturbation is very limited in the second shell for dilute conditions and only sites with a large charge density induce a perturbation extending beyond the first shell.\textsuperscript{45} Figure 3 shows that the four distributions are surprisingly similar. They all exhibit the same peak centered on moderate \( \sim 2 \) slowdown factors and a small amplitude tail at large slowdown factors. The value of the slowdown factor averaged over the fastest 90\% water molecules within the hydration layer obtained from our simulations ranges between 1.8 and 2.6 for the four proteins, in good agreement with recent magnetic relaxation dispersion (MRD) studies of three globular proteins, including ubiquitin, also studied here (2.0 at 290 K).\textsuperscript{5} These moderate slowdown factors are also consistent with a recent 2D-IR study of lysozyme hydration dynamics.\textsuperscript{10} The lengths of the molecular dynamics trajectories employed to compute the reorientational time-correlation function and the distribution of reorientation slowdown factors range between 4 and 20 ns, which is not sufficient to sample the full conformational space of these proteins. The impact of conformational fluctuations on hydration dynamics will be analyzed in detail below, but the comparison of results obtained respectively from two independent 20 ns simulations and from a shorter 4 ns run of lysozyme (Figure 4) shows that the 4 ns trajectory already provides a reliable determination of these quantities and that the results are very similar for two independent trajectories. Therefore, the differences observed in the results of two independent 20 ns trajectories and from a shorter 4 ns trajectory for (a) the second-order reorientational time correlation function \( C_2(t) \) (see Figure 2) and (b) the probability distribution of reorientation slowdown factors \( \rho_{\text{reor}} \) (see Figure 3).
in the results for the four proteins in Figures 2 and 3 do not originate from simulation variability.

Prior MD studies focusing on the mean residence time (MRT) of a water molecule within the protein hydration shell and an MRD investigation of water rotational dynamics have suggested that the distribution of water relaxation times within protein hydration shells can be described by a power law, \( p(t) \propto 1/t^\alpha \). The \( \alpha \) exponent of such a power-law fit was, for example, used to compare hydration shell dynamics across different proteins. An MRD study of dilute aqueous solutions of three globular proteins (ubiquitin, bovine pancreatic trypsin inhibitor, and \( \beta \)-lactoglobulin) yielded \( \alpha \) values ranging from 2.1 to 2.3, \( \sim 2.3 \) exponent was found for the MRT distribution computed for cytochrome \( c \). However, the MRT distribution from MD simulations of acetylcholinesterase yielded a much smaller power-law exponent of 0.84, which would indicate a much broader distribution of MRT. Since acetylcholinesterase is much larger than the proteins in the other studies listed above (see Table 1), this broader distribution could be caused by a size-dependent effect. However, our study of the reorientation time distributions shows that the acetylcholinesterase case is not different from the three smaller proteins (Figure 3), and a power-law fit of the tail of the reorientation time distribution (excluding internal water molecules) yields an exponent of \( \sim 2.3 \pm 0.1 \) in all four cases. The only difference is that the acetylcholinesterase distribution’s tail exhibits a slightly larger amplitude, as shown by the fraction of hydration shell water molecules whose slowdown factor is greater than 3, which is \( \sim 30\% \) in acetylcholinesterase and \( \sim 20\% \) for the three smaller proteins. We therefore propose that the low 0.84 value found previously for the acetylcholinesterase power-law exponent is due to the specific MRT definition used in that work, rather than to the protein size. It has been shown that different treatments of the transient escapes from the shell can critically affect the resulting MRT values. Our present reorientation results do not depend on such arbitrary choices and reveal no size-dependent effect over the range 9–59 kDa, apart from the trivial effect due to larger proteins containing more internal waters.

However, is it in fact meaningful to speak of a power law or any underlying analytical form for protein hydration shell dynamics? We first examine the power law \( 1/t^\alpha \) functional to describe the distribution of reorientation times. Figure 3b shows that a power law is an acceptable fit for intermediate slowdown factors (2 < \( \rho_{rer} \) < 10). However, the power law diverges at very low slowdown factors, and there is no clear power law behavior at long times in the reorientational tcf for all hydration shell water molecules. For the larger proteins subtilisin and acetylcholinesterase, the growing number of very slow internal water molecules leads to a plateau in the distribution at very large slowdown factor values, which cannot be properly described by a power law. On the basis of mode-coupling theory arguments, another functional that has been suggested to provide a good description of water relaxation dynamics within a protein hydration shell is the stretched exponential function \( \exp(-\|t/t^\beta\|) \) (see, e.g., refs 22 and 47). Figure 5 shows that a stretched-exponential functional form may appear to give a reasonable fit of the reorientational tcf (eq 1), at least at intermediate time delays. However, the corresponding probability distribution of reorientation times (i.e., the Laplace transform of the time decay) bears no resemblance to the distribution calculated explicitly from our simulations (Figure S). This confirms prior suggestions and clearly shows that a stretched exponential should only be regarded as a fit without any physical meaning. (The stretched exponential Kohlrausch function was shown to be reached only in the limit of very large wavevectors, i.e., for displacements much smaller than the intermolecular distance.) Therefore, while the stretched exponential and power-law fits remain a useful tool for analysis, our results unambiguously show that the global hydration shell dynamics is predominantly a sum of the dynamics of water molecules individually perturbed by local topological and chemical factors, with no simple underlying analytical form.

We have also investigated how these different reorientation times are distributed across the exposed surface of the protein. We have mapped individual reorientation times onto the protein surface for all four systems, as shown in Figure 6. Consistently with the distributions in Figure 3, we see that...
water molecules are moderately retarded throughout most of the hydration layer. More pronounced slowdown factors are observed especially in confined sites, e.g., the enzymatic active sites. We see a fairly uniform distribution of fast and slow dynamics across the hydration shell, with no large regions of similar water dynamics. This is in contrast to the “clustering” of water dynamics observed around one of the proteins, ubiquitin, in a recent NMR NOESY and ROESY study.31 However, we emphasize that these experimental results have been obtained under different conditions, since this technique requires encapsulation of the protein in a reverse micelle, while here the protein is studied in dilute aqueous solution.

Our present results can also be compared with those obtained by time-dependent Stokes shift (TDSS) spectroscopy and which suggest a large proportion of the hydration water population to be retarded by up to several orders of magnitude.7 In particular, for subtilisin Carlsberg, also studied here, TDSS experiments have measured a bimodal dynamics in the hydration shell, involving a sub-picosecond component with a 61% amplitude assigned to “bulk-like” water molecules, and a slower (∼38 ps) component with a 39% amplitude assigned to water molecules in strong interaction with the protein.7 The natural chromophore used in these experiments is the tryptophan residue Trp113, around which our present results do not reveal a pronounced slowdown of water dynamics. However, we note that our present study focuses on the dynamics of individual water molecules within the protein hydration layer, while TDSS is sensitive to collective motions affecting several water molecules and possibly of the protein itself, since they all influence the chromophore’s fluorescence energy.2 The large-amplitude slow component in TDSS decays may thus originate from coupled protein−water motions and water molecules displaced by slow conformational rearrangements of the protein.9,51,52

Extended Jump Picture. In order to explain the great similarity in hydration dynamics around proteins whose sizes, secondary structures, functions, and charge distributions are so diverse (see Table 1 and Figure 1), we now analyze the molecular factors governing the distributions of hydration shell dynamics.

In the case of lysozyme, it was recently shown that large angular jumps bring a dominant contribution to the overall reorientation dynamics of water molecules in the great majority of the hydration layer sites.25 We have therefore computed the distribution of jump slowdown factors ρjump, defined as

$$\rho_{\text{jump}} = \frac{\tau_{\text{hydrshell}}}{\tau_{\text{jump}}}$$

(4)

The jump time τjump is the inverse of the rate constant for the process of H-bond exchange by large-amplitude angular jumps (see the Methodology section). Figure 7 shows that these distributions are qualitatively similar to those of the reorientation slowdown, which suggests that the slowdown in the jumps brings a key contribution to the overall slowdown in hydration shell reorientation dynamics.

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The molecular origins of the jump slowdown can be identified and quantified using a picture considering the transition state for the process of H-bond exchange by large-amplitude angular jumps. For water next to hydrophobic sites on the protein surface, reorientation is slowed by the hindrance induced by the protein to the approach of a new H-bond acceptor. This can lead to a slowdown or an acceleration in reorientational dynamics, when

$$\rho_{\text{V}} = \frac{1}{1 - F}$$

(5)

where F is the fraction of jump transition state locations excluded by the presence of the protein, i.e., which overlap with the excluded volume of the protein atoms.53 For water hydroxyl groups initially H-bonded to acceptor sites on the protein surface, there is an additional perturbative effect arising from the free energy cost to stretch the initial H-bond with the protein to its transition state length, compared to the same free energy cost for a water−water H-bond. This can lead to a slowdown or an acceleration in reorientational dynamics, when...
the initial H-bond is respectively stronger or weaker than a water−water H-bond.54 This is referred to as the transition state H-bond (TSHB) effect.54 Finally, for water molecules accepting a H-bond from a donor site on the protein surface, reorientation is slowed via an excluded volume effect, as for water next to hydrophobic sites. Although such H-bonds can also vary in strength, they act on the water oxygen about which the angular jump occurs, and the influence of the resulting torque on the OH reorientational dynamics is negligible. Further details are given in a recent review on water dynamics33 and another on water dynamics in biomolecular hydration shells.2

In the case of lysozyme,25 it has been shown that the slowdown is due primarily to the excluded-volume (TSEV) effect arising from the local protein surface topology, with an additional free energetic effect for the slowest water molecules, related to H-bond acceptor strength (TSHB).25 These observations are used as a basis for understanding the molecular origins of the hydration shell reorientational dynamics of the three additional proteins studied here.

Application of Jump Analysis. Applying the extended jump picture to protein hydration shell dynamics can provide further insights into the nature of the molecular factors which cause the presence of the same two features in the distributions of reorientation times for the four proteins investigated here: a peak at moderate slowdown values and a tail at larger slowdown values.

We first focus on the peak in the distribution at $\rho_{\text{reor}} < 3$ (Figure 3), which contains, respectively, 83, 85, 80, and 70% of the hydration shell of ubiquitin, lysozyme, subtilisin, and acetylcholinesterase. Decomposing the $\rho_{\text{reor}}$ distribution into its contributions arising from water molecules perturbed by protein H-bond acceptors, H-bond donors, and hydrophobic groups shows that the peak corresponds principally to water molecules next to hydrophobic and H-bond donor sites. This is illustrated in Figure 8 for acetylcholinesterase, and similar results are found for the other proteins. Within the extended jump model, for these sites, water reorientation is moderately slowed down relative to bulk dynamics due to an excluded volume effect. The validity of the TSEV model53 is confirmed by Figure 9 which shows a strong correlation between $\rho_{\text{jump}}$ and the excluded volume slowdown factor $\rho_V$ for water next to hydrophobic and H-bond donor sites in all four proteins. This $\rho_V$ factor successfully rationalizes the slowdown for the vast majority of these sites, which in turn make up the majority (83−88%) of the hydration shell population. Deviations from the TSEV prediction occur only for deeply buried hydrophobic sites (values of $F$, the fraction of excluded transition state locations, close to 1) where the situation is no longer that of a water molecule at the interface between a solute and bulk water and where the TSEV model53 based on the approach of a new H-bond acceptor from the bulk no longer holds true.

We now turn to the distribution’s tail ($3 < \rho_{\text{reor}} < 20$), which is shown in Figure 8 to be mainly due to water molecules next to moderate to strong H-bond acceptor sites. Within the extended jump picture, these sites retard water reorientation via both the strength of the initial water−protein H-bond and an excluded volume effect.25 Although we do not explicitly quantify this effect here, it has already been shown to successfully rationalize water dynamics next to H-bond acceptor sites in both proteins and individual amino acids.25,54

Finally, the distributions for the larger proteins, acetylcholinesterase and subtilisin, have an even slower, low-amplitude
tail ($\rho_{\text{ex }}> 20, \text{see Figure 3b}$) arising from water molecules in internal or deeply buried sites. This is consistent with the fact that larger proteins are known to contain more internal water molecules. These extremely slow sites correspond to $\sim 2\%$ of the total hydration shell population in acetylcholinesterase and subtilisin and $<1\%$ in lysozyme and ubiquitin. The effect of these very slow internal water molecules is also seen in the reorientational tcf’s (Figure 2), where the amplitude of the tcf at long times scales with protein size.

Our analysis thus shows that the distribution of perturbation factors is dominated by an excluded volume effect, determined by local surface topology, i.e., the presence of pockets, protrusions, and clefts on the protein surface. The dominance of this effect is due in turn to a surface composition dominated by hydrophobic sites and H-bond donors. We now use these results to explain why the four proteins investigated here display very similar reorientational hydration shell dynamics, despite their very different biological functions, sizes, and secondary structures. While at first glance certain proteins appear to have quite specific shapes, including, e.g., the active-site cleft in lysozyme, the local protein surface topologies experienced by water molecules next to these four proteins are on average very similar, as shown by the distributions of excluded volume slowdown factors $\rho_V$ (Figure 10).

**Figure 10.** Probability distribution of excluded volume slowdown factors $\rho_V$ for the four protein systems in aqueous solution.

dynamical perturbation of the hydration shell, and an analysis analogous to that presented here showed that this arises from the reduced number of confined sites in the unfolded state.26

**Dynamical Heterogeneity due to Protein Conformational Fluctuations.** The results presented above demonstrate that a major factor causing the broad distribution of water dynamics within a protein hydration layer is its roughness, which leads to a great variety of local topologies. However, the shape of a protein is not constant in time because a biomolecule is a dynamical object, constantly sampling different conformations. Therefore, next to one given protein site, the perturbation induced on the surrounding water dynamics fluctuates when the local protein topology changes. This can lead to an additional, dynamical heterogeneity in hydration shell dynamics. Conformational changes in the protein surface can affect the water jump rate constant and hence its reorientational dynamics in different ways, for example, by changing the local excluded volume slowdown factor $\rho_V$ and for jumps between two protein H-bond acceptors by changing the positions of the two acceptors. Such conformational changes include, for example, hinge motions and pockets and clefts that fluctuate in size.

In order to assess the effect of conformational fluctuations on hydration shell dynamics, we use the lysozyme system, and calculate the normalized standard deviation $\sigma$ in the jump rate constant for each site over a 20 ns trajectory divided into five independent blocks, defined as

$$\sigma = \frac{\sqrt{\langle \tau_{\text{jump}}^2 \rangle - \langle \tau_{\text{jump}} \rangle^2}}{\langle \tau_{\text{jump}} \rangle} \quad (6)$$

where $\langle \cdots \rangle$ denotes an average over the blocks.

While a duration of 20 ns is certainly not sufficient to cover the full conformational space of the protein, it is already sufficient to sample many different conformations. This is demonstrated by performing a principal component analysis59 and projecting the trajectory on the first two principal components, which describe the greatest amount of variance in the protein heavy atom positions and which involve the hinge-bending motion of lysozyme.60 Figure 11 shows that different conformational basins are visited during the simulation. The 4 ns block size is somewhat arbitrary; however,
our goal is only to obtain a qualitative measure of the dynamical heterogeneity, and we showed in Figure 4 that this duration is sufficient to converge the distribution of water reorientation times.

The resulting values of the standard deviation $\sigma$ are mapped onto the protein surface in Figure 12. Larger values of $\sigma$ can be taken as a qualitative indication of increasing dynamical heterogeneity in the hydration shell dynamics at that surface position. Water at exposed or convex parts of the surface has relatively less heterogeneity in its dynamics, compared to the greater heterogeneity for water in partial confinement, such as in surface pockets, or in other locations subject to conformational fluctuations, such as in the pronounced active-site cleft in the upper right-hand part of the protein in Figure 12.

The correlation between $\sigma$ and the excluded volume fraction $F$ is quantified in Figure 13 for lysozyme surface sites. Exposed or convex parts of the protein surface correspond to low values of $F$ and display consistently low values of $\sigma$, which indicate a limited dynamical heterogeneity. Concerning sites in a concave surface environment or in partial confinement (high value of $F$), the dynamical heterogeneity covers a broad range of values, going from a very small dynamical heterogeneity for internal or deeply buried water molecules whose environment changes very little with time to very large values for other molecules including, e.g., those in the active-site cleft whose width fluctuates.

A decomposition of the probability distribution of $\sigma$ as a function of site type (Figure 14) shows that hydrophobic and H-bond donor sites dominate at lower dynamical heterogeneity, while H-bond acceptor sites dominate at higher dynamical heterogeneity. This arises from the fact that water molecules in concave surface environments, or in other words those most likely to experience dynamical heterogeneity, are often H-bonded to acceptor sites, since favorable energetics are required for a water molecule to enter a surface pocket or groove (as illustrated in the probability distribution of the excluded volume fraction $F$ decomposed as a function of site type in Figure 15).

In conclusion, in addition to the heterogeneity in hydration shell dynamics arising from the chemical and topological nature of a static protein surface, fluctuations in the surface conformation may lead to an additional, dynamical heterogeneity. The relative importance of these two types of heterogeneity in the hydration shell dynamics can be determined qualitatively via a comparison of normalized standard deviations in jump times. The magnitude of the spatial heterogeneity can be roughly quantified via $\sigma_G/\mu$, where $\mu$ and $\sigma_G$ are the average and standard deviation of a Gaussian fit of the main peak of the protein’s $\tau_{\text{jump}}$ distribution (we note that this underestimates the spatial heterogeneity by ignoring the $\tau_{\text{jump}}$ distribution’s tail). This measure gives a $\sigma_G/\mu$ value of 0.15–0.20 for the four proteins studied here. This can be compared to the magnitude of the dynamical heterogeneity as quantified by $\sigma$ (eq 6), which has a modal value of ~0.03 in the case of lysozyme (see Figure 14), 5–6 times smaller (this remains qualitative, since larger $\sigma$ values might be obtained when calculated on shorter independent intervals). We therefore stress that a simple, spatially resolved analysis as employed in the previous section is sufficient to capture and
rationalize the majority of the dynamic behavior of hydration shell water. However, considering dynamical heterogeneity may be important for understanding the behavior of small subsets of the hydration shell population, for example, in the hydration shell of proteins with marked conformational transitions such as hinge motions. Dynamical heterogeneity may also be important for protein hydration dynamics at low temperature.

HYDRATION SHELL DYNAMICS IN CONFINEMENT

The work presented here so far has considered proteins in dilute aqueous solution, as is the case in the majority of experimental and simulation studies of protein hydration shell dynamics. However, water dynamics in vivo occurs under conditions of macromolecular crowding, and certain experimental techniques employ high protein concentrations or conditions of confinement. An understanding of protein hydration shell dynamics is therefore incomplete without a consideration of the effects of confinement.

Description of Confined Systems. Many different types of confining situations exist, possibly with different impacts on protein hydration dynamics. Here, we focus on a protein and its hydration shell confined by an apolar organic solvent. We compare water dynamics in the hydration shell of subtilisin in three systems: the enzyme in aqueous solution, the enzyme with a monolayer of water (841 water molecules) in hexane solution, and the enzyme with approximately a half-monolayer of water (520 water molecules) in hexane solution. A monolayer is defined on the basis of the number of water molecules in the hydration shell of the enzyme in aqueous solution. Hexane is chosen because nonpolar organic solvents have been shown to conserve the enzyme hydration shell, in contrast to polar organic solvents, which “strip” water molecules from the enzyme surface.

The monolayer system is prepared so that the protein is initially surrounded by a uniform layer of water molecules. After equilibration, the protein surface is no longer completely hydrated. Instead, large patches of the surface contain no or only a scattering of tightly bound water molecules, and are in direct contact with the organic solvent, while other patches are completely hydrated, with several shells of water molecules. The same preparation method is used for the half-monolayer system, with the initial distribution of water molecules being as uniform as possible at reduced water content, and the same clustering of water molecules is seen after equilibration. As an example, the half-monolayer system after 5 ns of simulation time is shown in Figure 16. Since the distribution of hydrophobic and polar groups across the surface is approximately uniform and the hydrated and unhydrated surface patches are large, no particular correlation between hydrophobicity and hydration could be detected. Of note is the fact that the active site remains completely hydrated at both monolayer and half-monolayer hydration levels.

Effect of Confinement on Reorientational Water Dynamics. The distributions of reorientation times for the three hydration levels are shown in Figure 17. Decreasing the hydration level leads to a shift of the distributions toward larger slowdown factors and to a broadening of these distributions. This shows that confinement induces a retardation of water dynamics within the shell, and also that this slowdown is heterogeneous across the hydration shell; i.e., some sites are more slowed down than others.

In order to explore this heterogeneity on a site-resolved level, for any given surface site \( i \), we define the slowdown at hydration level \( h \) relative to the fully hydrated system in aqueous solution as \( \tau_{\text{rec}}(i)/\tau_{\text{bulk}}(i) \). Since the clustering of water on the protein surface is not identical in the two partially hydrated systems, the most meaningful comparison is between each of these systems and the fully hydrated system. These values are mapped onto
the protein surface in Figure 18. Water at the majority of protein sites is moderately slowed down upon confinement, by a factor of between 1 and 2 for $\tau_{\text{reor}}^{\text{monolayer/ bulk}}$ and between 1 and 3 for $\tau_{\text{reor}}^{\text{half monolayer/ bulk}}$. In general, the sites with the greatest slowdown are those next to parts of the protein surface which are completely dehydrated. This suggests that the water molecules whose dynamics is most retarded are those who experience an excluded volume for the approach of a new H-bond acceptor which is due not only to the protein surface but also to the apolar hexane solvent.

**Connection with Linear and 2D-IR Spectroscopy.** We then turn to linear and two-dimensional infrared spectroscopy in order both to explore further the H-bond dynamics of these systems and to demonstrate how our conclusions can be connected to experimentally obtainable values. 2D-IR spectroscopy is an ultrafast technique which is increasingly being used to probe water H-bond dynamics in a wide range of systems, including confining environments and the hydration shell of biomolecules such as DNA. In addition, 2D-IR spectroscopy has also been used to indirectly probe protein hydration shell dynamics via a vibrational probe covalently attached to the protein surface.

Here, we calculate the linear and 2D-IR spectra of the water stretch vibration. Because these spectra measure a signal collected from all water molecules in the system, for a protein in bulk aqueous solution, the hydration shell water signal would be swamped by the signal from bulk water. We therefore focus on two partially hydrated subtilisin systems, where all water molecules are close to the protein interface. Experimentally, isotopic mixtures such as HOD in H$_2$O are used to avoid the effects of intermolecular vibrational energy transfer. We thus calculate the spectra for the OD stretch of dilute HOD in H$_2$O. We employ the empirical map developed in ref 72 relating the vibrational frequency to the local electric field. The latter is obtained via an *a posteriori* treatment of classical molecular dynamics trajectories. While our calculations are based on a trajectory computed for a system containing pure H$_2$O, the effect on the calculated spectra has been shown to be negligible. Our choice to study the water OD stretch rather than the OH stretch is dictated by the necessity to isolate the mode under consideration as much as possible from other protein vibrational modes. While the water OH stretch frequency overlaps with the OH and NH protein bands, the OD stretch frequency range does not significantly overlap with the frequency range of major protein vibrational modes.

The resulting linear IR spectra for HOD in bulk H$_2$O and in the hydration shell of subtilisin at different hydration levels are shown in Figure 19. In hexane confinement, a blueshifted peak grows for decreasing hydration level. Such a peak had previously been observed experimentally at water–hexane interfaces and also in simulations of water next to model hydrophobic surfaces. It corresponds to dangling, non-H-bonded OD groups.

We now turn to the 2D-IR spectra, shown in Figure 20. 2D-IR spectroscopy provides detailed information on water H-bond dynamics, since the water stretch frequency is
a sensitive probe of the H-bonding interaction: it respectively shifts to the red and to the blue when engaged in a strong and a weak hydrogen-bond. The spectra show the correlation between excitation and detection frequencies of the water stretch after a given waiting time. The time evolution of the spectra is therefore a measure of the loss of frequency correlation, and hence gives access to time-resolved information on water dynamics. The spectra in Figure 20 clearly show that the frequency relaxation is slower in the confined hydration shell than in bulk water, and that this slowdown is even more pronounced when the hydration level decreases. This time evolution can be quantified using, for example, the center line slope (CLS), i.e., the slope of the positive peak’s crest along the horizontal excitation frequency axis, which provides an estimate of the frequency tcf. As shown in Figure 21, while in bulk water, the OD frequency decorrelates on a picosecond time scale, and in the confined hydration shell after 2 ps, a large frequency correlation is retained, leading to 2D-IR spectra which are still elongated along the diagonal (cf. Figure 20). This reflects the slower water H-bond dynamics in confinement described in the previous section. A slowdown in water spectral dynamics has also been observed in 2D-IR studies of water confined in other systems.

The 2D-IR spectra further provide a resolution of the linear IR bandwidth in terms of its homogeneous and inhomogeneous contributions, which respectively arise from the presence of rapid frequency fluctuations and of a static distribution of frequencies. The homogeneous width can be estimated as the full width at half-maximum (fwhm) of a Lorentzian fit of the 2D-IR spectra along the antidiagonal, while the inhomogeneous width can be determined from the fwhm of a Gaussian fit of the 2D-IR spectra along the diagonal. The time evolutions of the homogeneous and inhomogeneous widths are shown in Figure 22. They clearly show that upon confinement the inhomogeneous distribution of frequencies becomes broader due to the greater variety of local environments, while the homogeneous line width decreases due to the slower frequency dynamics. This explains a key difference between the 2D-IR spectra in Figure 20 for excitation frequencies on the blue edge. In bulk water, these blueshifted OD vibrations correspond to transient H-bond breaks quickly followed by the reformation of the H-bond, leading to a very fast frequency decorrelation. In the confined subtilisin hydration shells, these blueshifted frequencies arise from long-lived and weakly or non-H-bonded OD groups at the hexane interface, leading to a much slower frequency decorrelation.

These results demonstrate that the dynamics of the water H-bond network is slower at lower hydration level and that the distribution of relaxation times is broader. Our analysis shows
that these conclusions can be directly connected to experimentally accessible linear and 2D-IR spectra of such systems, and that these techniques could prove a valuable tool for the elucidation of their hydration shell dynamics.

### CONCLUDING REMARKS

Many different pictures have been suggested to describe and rationalize protein hydration shell dynamics. The spatial extent and magnitude of the perturbation induced by a biomolecule together with the origin of this perturbation have been extensively studied and discussed, with both local and longer-range effects being evoked. Here, we conclude that the reorientation dynamics of individual water molecules is only moderately perturbed in the hydration shell of subtilisin Carlsberg in hexane at different hydration levels.

In addition, we have shown that protein conformational fluctuations have a large impact on hydration shell dynamics, particularly at those parts of the protein surface which are concave or which cause a partial confinement of hydration shell water molecules. We have also evidenced a slowdown in hydration shell dynamics upon confinement which is heterogeneous across the protein surface, and demonstrated how the water dynamics in such a system can be explored via 2D-IR spectroscopy. Future work will extend the present approach to the study of translational dynamics of water molecules within protein hydration shells, which can be probed by NMR and neutron scattering techniques and which has been shown to display slowdown factors similar to those of reorientational dynamics.

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**Notes**

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