Dynamics of hydration water in deuterated purple membranes explored by neutron scattering

K. Wood · M. Plazanet · F. Gabel · B. Kessler · D. Oesterhelt · G. Zaccai · M. Weik

Abstract The function and dynamics of proteins depend on their direct environment, and much evidence has pointed to a strong coupling between water and protein motions. Recently however, neutron scattering measurements on deuterated and natural-abundance purple membrane (PM), hydrated in H$_2$O and D$_2$O, respectively, revealed that membrane and water motions on the ns–ps time scale are not directly coupled below 260 K (Wood et al. in Proc Natl Acad Sci USA 104:18049–18054, 2007). In the initial study, samples with a high level of hydration were measured. Here, we have measured the dynamics of PM and water separately, at a low-hydration level corresponding to the first layer of hydration water only. As in the case of the higher hydration samples previously studied, the dynamics of PM and water display different temperature dependencies, with a transition in the hydration water at 200 K not triggering a transition in the membrane at the same temperature. Furthermore, neutron diffraction experiments were carried out to monitor the lamellar spacing of a flash-cooled deuterated PM stack hydrated in H$_2$O as a function of temperature. At 200 K, a sudden decrease in lamellar spacing indicated the onset of long-range translational water diffusion in the second hydration layer as has already been observed on flash-cooled natural-abundance PM stacks hydrated in D$_2$O (Weik et al. in J Mol Biol 275:632–634, 2005), excluding thus a notable isotope effect. Our results reinforce the notion that membrane-protein dynamics may be less strongly coupled to hydration water motions than the dynamics of soluble proteins.

Keywords Neutron spectroscopy · Hydration water dynamics · Purple membrane · Dynamical transition

Abbreviations

BR Bacteriorhodopsin
PM Purple membrane
EINS Elastic incoherent neutron scattering
MD Molecular dynamics
MSD Mean square displacements

Introduction

The purple membrane (PM) has provided biophysicists with an exceptional model of study. Natively forming a
two-dimensional crystal, PM contains a unique protein, viz. the light-driven proton pump bacteriorhodopsin (BR), and various lipid species. From the external membrane of Halobacterium salinarum, PM can be purified relatively easily in large quantities necessary for neutron scattering measurements. PM dynamics has been well-characterised by neutron scattering at various time and space scales for different hydration levels (Ferrand et al. 1993; Fitter et al. 1996, 1997; Lehnert et al. 1998) and different lipid contents (Fitter et al. 1998). Isotope labelling of the membrane has been successfully performed (Engelman and Zaccai 1980; Patzelt et al. 1997), and combined with incoherent neutron scattering experiments, revealed the dynamical heterogeneity of PM on the ps–ns time scale, with the core of BR found to be stiffer than its surroundings (Reat et al. 1998).

Hydrated PM undergoes a so-called dynamical transition at around 230–260 K, evidenced in incoherent neutron scattering experiments as a sudden increase in the atomic mean square displacement (MSD) as a function of temperature (Ferrand et al. 1993; Fitter et al. 1997; Lehnert et al. 1998). Neutron scattering experiments on soluble proteins (Doster et al. 1989; Fitter 1999; Roh et al. 2006; Wood et al. 2008), on another membrane protein (Pieper et al. 2007) and on RNA (Caliskan et al. 2006) revealed the same phenomenon at a temperature between 180 and 250 K.

The dynamical transition only occurs in sufficiently hydrated systems, and changing the solvent composition shifts the transition temperature (Caliskan et al. 2004; Cordone et al. 1999; Cornicchi et al. 2005, 2006; Paciaroni et al. 2002; Tsai et al. 2000). The term “slaving” has been used to express that the solvent imposes its dynamics on the macromolecule (Fenimore et al. 2002, 2004; Iben et al. 1989). In particular, molecular dynamics (MD) simulations have shown that the onset of translational diffusion of hydration-water triggers the protein dynamical transition in soluble proteins (Tarek and Tobias 2002; Tournier et al. 2003). In the PM, however, the onset of translational motion in hydration water occurs at 200 K (Berntsen et al. 2005; Weik 2003; Weik et al. 2005; Wood et al. 2007), i.e. well below the dynamical transition of membrane motions. Membrane-protein motions might be controlled by lipid motions (Kamihira and Watts 2006) rather than by water motions (Wood et al. 2007). Nevertheless, water is required to allow the conformational changes necessary for BR to function (Buchsteiner et al. 2007; Dencher et al. 2000; Weik et al. 1998).

Here, we employed incoherent elastic neutron scattering and neutron diffraction to study hydration water in stacks of deuterated PM. The first part of this study extends previous neutron-diffraction measurements that showed intermembrane heavy water (D₂O) in the second hydration layers turns liquid-like at 200 K in flash-cooled natural-abundance PM (Weik et al. 2005). Here, the results have been reproduced for deuterated PM hydrated in light water (H₂O). In the second part, atomic MSD of hydration-water and PM dynamics were measured separately by using deuterated PM in H₂O and hydrogenated PM in D₂O, respectively. In complement to an earlier study on samples with high hydration (three to four water layers between successive PM patches at room temperature) (Wood et al. 2007), we here present data on the same samples, yet at a lower hydration level corresponding to only two water layers between successive membrane fragments. Again, dynamical transitions of hydration-water and of membrane motions are observed at 200 and at 260 K, respectively, indicating that hydration-water and PM motions on the ns–ps time scale are uncoupled below 260 K.

Materials and methods

Sample preparation

Two PM samples were prepared, a fully deuterated one (denoted D-PM) and a natural-abundance one (denoted H-PM). Both were purified from Halobacterium salinarum (H. salinarum) as described previously (Oesterhelt and Stoeckenius 1974). To produce fully deuterated PM, H. salinarum were grown on deuterated algal medium (Patzelt et al. 1997). Two hundred milligram samples of both D-PM and H-PM were used, as deduced from the BR concentration determined by the absorbance at 565 nm on the PM suspensions obtained after purification. D₂O in the deuterated PM sample (denoted D-PM-H₂O) and H₂O in the hydrogenated sample (denoted H-PM-D₂O) were exchanged against H₂O and D₂O, respectively, by three successive centrifugation steps. The two concentrated membrane suspensions were then placed on 4 × 3 cm² flat aluminium sample holders. The sample holders were placed in dessicators over silica gel for 12 h to achieve partial drying to approximately 0.5 g water/g membrane, as determined by weighing the holder regularly until it contained approximately 300 mg of sample. Vacuum was not applied to the dessicator at any stage of the sample preparation. The silica gel was exchanged for a saturated KNO₃ solution [in H₂O for the D-PM-H₂O sample and in D₂O for the H-PM-D₂O sample, respectively; yielding a relative humidity (r.h.) of 93%] for 7 days until the weight was stable. The D-PM-H₂O sample was then equilibrated over pure H₂O (100% r.h.) for 2 days, called a “high-hydration” (Hh) level, and is hereafter denoted D-PM-H₂O-Hh. The high-hydration level corresponds to about 0.3 g water/g membrane. The sample holder was sealed with indium and closed with an aluminium cover,
resulting in a sample chamber of 0.3 mm thickness. After temperature-dependent neutron diffraction experiments (see below), the D-PM-H$_2$O-Hh sample was opened and equilibrated for 3½ days over a saturated KCl solution (86% r.h.; in H$_2$O), leading to a “low-hydration” (Lh) level of 0.1 g water/g membrane. The sample was then sealed, closed again (0.3 mm chamber thickness) and denoted D-PM-H$_2$O-Lh. The H-PM-D$_2$O sample was equilibrated for 2 days over pure D$_2$O (100% r.h.) and then for 3½ days in a D$_2$O atmosphere at 86 % r.h. (resulting in 0.1 g water/g membrane), before it was sealed and closed (0.3 mm chamber thickness): it is denoted H-PM-D$_2$O-Lh. An estimate of the number of hydrogen and deuterium atoms in the different components of the H-PM-D$_2$O-Lh and D-PM-H$_2$O-Lh samples is given in Table 1.

Neutron diffraction as a function of temperature to determine lamellar spacing

As water evaporated from the membrane preparation, the PM patches oriented parallel to the surface of the sample holder, forming a stack of membranes with a characteristic repeat distance, called the lamellar spacing. The number of water layers intercalated between adjacent membrane fragments depends on the sample preparation protocol (r.h. and equilibration time) and can be determined by measuring the lamellar spacing using neutron diffraction. Lamellar spacings were determined on the diffractometer D16 at the Institut Laue-Langevin (ILL) by monitoring the first-order Bragg peak of the PM stack using a wavelength of 4.56 Å˚ (Weik et al. 2005).

The D-PM-H$_2$O-Hh sample was characterised by a lamellar spacing of 62 Å˚ at room temperature, corresponding to an average inter-membrane water layer of 13 Å (the average thickness of a dry PM fragment is 49 Å). To determine its lamellar spacing after flash-cooling as a function of temperature upon heating (Weik et al. 2005), the sample holder was plunged into liquid nitrogen and then transferred into a cryostat pre-cooled at 100 K. The lamellar spacing was determined at 100 K and then on heating from 180 to 290 K. The time interval between successive data points was 28 min.

Subsequently, the D-PM-H$_2$O-Hh sample was opened, re-equilibrated to a lower hydration level (Lh) (see preceding section) and re-sealed. The lamellar spacing of the D-PM-H$_2$O-Lh and the H-PM-D$_2$O-Lh samples were then measured, and the samples were opened and re-hydrated/dehydrated as necessary until a lamellar spacing of 54 Å at room temperature was reached, corresponding to a 5 Å thick inter-membrane water layer. Identical lamellar spacings of the two samples ensured identical hydration levels. Between neutron scattering experiments, the samples were stored at $-20$ºC.

Elastic incoherent neutron scattering (EINS) experiments

Owing to their particular wavelengths and energies, thermal and cold neutrons are an important probe for macromolecular structure and thermal dynamics (Gabel et al. 2002). The incoherent cross section of hydrogen atoms is about 40 times larger than that of deuterium, and much larger than those of all other atoms present in biological macromolecules. Using deuterium labelling, it is therefore possible to focus on different components of a complex system. In the present work, studying fully deuterated PM in H$_2$O allowed us to predominantly measure hydration water dynamics. Natural-abundance PM hydrated in D$_2$O, on the other hand, provided information about membrane motions. Indeed, 60% of the total incoherent scattering cross section of D-PM-H$_2$O-Lh is due to the contribution of inter-membrane water. For H-PM-D$_2$O-Lh, the contribution of the D$_2$O is negligible, and protein (BR) and lipid make up 77 and 23%, respectively. In the estimations, we consider that 60% of the exchangeable hydrogen/deuterium atoms in bacteriorhodopsin exchange, and that all of the exchangeable hydrogen/deuterium atoms in the lipid head groups do so (Zaccay and Gilmore 1979).

So-called “elastic” scans (the elastic intensity only is recorded) were performed as a function of temperature on the IN16 backscattering spectrometer at the ILL, with an energy resolution of 0.9 μeV (full width at half maximum of the elastic peak) (Frick and Gonzalez 2001) and a wavelength of 6.275 Å. The $Q$-range accessible is 0.02–1.9 Å$^{-1}$, corresponding to movements on the angstrom (Å) scale. At the given instrumental resolution, hydrogen

| Table 1 | Number of hydrogen atoms in the different sample components, considering 1 BR molecule and associated lipid and water molecules for the H-PM-D$_2$O-Lh and D-PM-H$_2$O-Lh samples |
|-----------------|-----------------|------------------|
| H-PM-D$_2$O-Lh  | Number of deuterium atoms | Number of hydrogen atoms |
| BR              | 220             | 1,728            |
| Lipids          | 32              | 505              |
| Water           | 450             | –                |
| D-PM-H$_2$O-Lh  | 1,728           | 220              |
| BR              | 1,728           | 220              |
| Lipids          | 505             | 32               |
| Water           | –               | 474              |

The calculation has been performed assuming that all the exchangeable hydrogen atoms in the lipid head-groups, and 60% of those in the protein have exchanged (Zaccay and Gilmore 1979).
movements with characteristic times faster than 1 ns are monitored, which reflect the dynamics of the larger groups to which they are attached. A sudden drop in the elastic signal is indicative of an increase in movements faster than 1 ns on the angstrom (Å) length scale.

D-PM-H₂O-Lh and H-PM-D₂O-Lh samples were placed in a standard “orange” cryostat at room temperature at an angle of 135° with respect to the incident beam and then cooled to 50 K in about 2 h. Subsequently, the elastic energy was recorded on heating from 50 to 300 K at constant rates of 0.16 and 0.17 K/min for the D-PM-H₂O-Lh and the H-PM-D₂O-Lh samples, respectively. The acquisition was continuous on heating and the data were subsequently binned into points corresponding to a temperature step of about 5 K. The signal from the empty sample holder was subtracted, and the data were normalised to the intensity at 50 K.

From the wave-vector (Q) dependence of the elastic intensity (Q = 4πsin θ/λ, in which 2θ is the angle of the scattered neutron and λ is the neutron wavelength), the mean square displacement (MSD) of the investigated movements can be calculated according to the Gaussian approximation, which is valid for Q²<2:

\[ I(Q, \omega = 0) = I_0 \exp \left( -\frac{1}{6} Q^2 \langle u^2 \rangle \right) \]  

where \( I(Q, \omega = 0) \) is the normalised elastically scattered intensity and \( I_0 \) is the value of the scattering at Q = 0.

The earlier studies measured \( Q = 1.4 ~\text{Å}^{-1} \) arising from the spacing of deuterated lipids at 4.5 Å (Henderson 1975), which is not constant with temperature. The elastic intensity summed over all Q-values used for MSD determination, which was first used as a guide for interpreting MSD in neutron biological dynamics studies by Reat et al. (2000), is also presented as a function of temperature. Limited to the first term, the expansion of the exponential of Eq. 1 gives

\[ \sum_{Q} I(Q, \omega = 0) \approx I_0 \left( 1 - b\langle u^2 \rangle \right), \]  

where b is a constant depending only on the Q values over which the sum is performed. In a first-order approximation, the summed elastic intensity is therefore proportional to the MSD \( \langle u^2 \rangle \). The quantity was normalised to unity at low temperature by dividing the sum by the number of detectors over which it was performed.

Results and discussion

Lamellar spacing of flash-cooled D-PM-H₂O-Hh as a function of temperature

The samples described here are formed of stacked PM fragments, intercalated by a number of water layers at room temperature that can be adjusted according to different equilibration protocols (see “Materials and methods”). If the lamellar spacing at room temperature is larger than 54 Å, excess inter-membrane water leaves the membrane stack upon slow-cooling at temperatures around 260 K until a minimal spacing of 54 Å is reached and forms crystalline ice outside the stack (Lechner et al. 1998). Only two layers of water (corresponding to 54–49 Å = 5 Å) remain thus in the inter-membrane space below 260 K. Upon flash-cooling, however, 9 Å of water (three to four layers) can be trapped in natural-abundance PM hydrated in D₂O (Weik et al. 2005). On heating, two of the four water layers are observed to leave the stack at 200 K to form crystalline ice, and the lamellar spacing decreases to 54 Å. The decrease in lamellar spacing at 200 K has been attributed to an onset of translational mobility of inter-membrane water in the second hydration shells. Above 260 K, swelling of the membrane stacks is observed as crystalline ice melts and returns in the inter-membrane space. Samples with a lamellar spacing of 54 Å at room temperature do not exhibit major changes, and the spacing remains constant (to within 0.3 Å) as a function of temperature (Lechner et al. 1998). The different scenarios are schematised in Fig. 1.

The earlier studies measured D₂O in natural-abundance PM (Weik et al. 2005). Here, we examined the temperature-dependent lamellar spacing of the D-PM-H₂O-Hh sample to address possible isotope effects. Figure 2 shows the lamellar spacing of flash-cooled D-PM-H₂O-Hh upon heating, which exhibits a similar temperature dependence to the one of H-PM-D₂O-Hh measured earlier (Weik et al. 2005). The lamellar spacing of about 57.5 Å remains constant up to 200 K, then decreases to reach a minimal value of 54 Å at 250 K (only first hydration layers remain) and increases again above 260 K to reach the initial value of 62 Å at room temperature. The lamellar spacing of H-PM-D₂O-Hh (Weik et al. 2005) and D-PM-H₂O-Hh samples thus display an identical temperature-dependence within the experimental temperature error estimated to be about 5 K. Second hydration-layer water, H₂O or D₂O, thus turns liquid-like at 200 K.
A direct comparison of PM and hydration-water dynamics has been made recently by EINS using deuterated membranes hydrated in H$_2$O (the same D-PM-H$_2$O-Hh sample as the one examined in the previous section) and natural-abundance membranes in D$_2$O (Wood et al. 2007). A transition in the mean square displacements of hydration water at 200 K, attributed by MD simulations to the onset of water translational mobility, did not trigger a dynamical transition in PM that, in fact, took place at 260 K. The samples were characterised by a high-hydration level
corresponding to a lamellar spacing of 62 Å at room temperature (i.e., 13 Å of inter-membrane water) that is not constant as a function of (cryo-) temperature (see Fig. 1). Here, we present data on samples with a lower hydration level corresponding to a lamellar spacing of 54 Å (i.e., 5 Å of inter-membrane water), which varies by less than 0.3 Å between 300 and 50 K. Furthermore, upon heating and cooling, no crystalline ice is formed (Lechner et al. 1998).

Figure 3 shows the elastic intensity summed over the $Q$-range examined (see “Materials and methods”) and the extracted MSD of D-PM-H$_2$O-Lh and HPM-H$_2$O-Lh. The elastic intensities (Fig. 3a) are very different for both samples, with inflections seen at approximately the same temperatures for which they were seen in the “high-hydration” samples (Wood et al. 2007): 120 and 260 K for the H-PM-D$_2$O-Lh sample, 200 and 260 K for the D-PM-H$_2$O-Lh sample.

Using the summed elastic intensities (where the error on each point is of the size of the points in Fig. 3a) as a guide, the atomic MSD of water and PM motions can be divided into regions with different characteristic temperatures. Below 120 K, MSD of both membrane (H-PM-D$_2$O-Lh) and water (D-PM-H$_2$O-Lh) are similar. Following an inflection at 120 K, the membrane’s MSD are clearly above the water’s MSD up to 260 K—the membrane has larger amplitude motions than its hydration water. The hydration water MSD evolves linearly with temperature to a higher temperature than PM, and undergo a broad transition at 200 K. At 260 K, the water MSD catches up with the membrane MSD and are almost identical to them up until 288 K. Above 288 K, it becomes impossible to calculate MSD for the water molecules, since their motions become too large to be resolved, and the Gaussian approximation is no longer valid.

Although the same transition temperatures are observed for the high-hydration samples studied here and the high-hydration samples measured earlier (Wood et al. 2007), small differences in the dynamics of water in the two hydration states can be seen. Water MSDs for the high-hydration and low-hydration samples are plotted for comparison in Fig. 4, where two temperature regions with significant differences in dynamics are evident. The first is between 150 and approximately 200 K, where the low-hydration sample displays larger MSD than the high-hydration sample. This indicates that the two inter-membrane water layers, present in both high- and low-hydration samples, are more mobile than ice in the same temperature region. The second temperature region where a large difference is observed is above 260 K, temperature above which in the case of high-hydration samples the ice present in the sample melts and flows back into the membrane stack and 13 Å of water is in the inter-lamellar space. Above 260 K, water in the high-hydration sample has much larger MSD than water in the low-hydration sample, implying that four layers of water confined in PM stacks show on average higher mobility on the ps–ns time scale than two water layers.

The main conclusion by Wood et al. (2007) is confirmed by the present experiments at a lower hydration level: the dynamical transition in the hydration water at 200 K does not trigger a transition in the membrane at the same temperature. Since BR contributes most to the scattering cross section of PM, it can be supposed that the absence of a transition at 200 K in the whole PM indicates that no transition occurs in the protein. The result suggests that, in contrast to soluble proteins, the dynamics of water and membrane proteins on the ns–ps time scale are decoupled below 260 K, possibly suggesting an important role of lipids in the dynamical interdependency of BR, lipids and water.

Acknowledgments We are grateful to Heloisa Nunes Bordallo, Bruno Demé and Bernhard Frick for their help during data collection at the ILL. We also thank Marek Koza and Douglas Tobias for fruitful discussions. Financial support by the CEA, the CNRS and the UJF is acknowledged, as well as a grant to MW from the Agence Nationale de la Recherche (project number JC05_45685). The study was supported by the EU DLAB program under contract numbers HPRI-CT-2001-50035 and RI13-CT-2003-505925.
Wood K, Caronna C, Fouquet P, Hauessler W, Natali F, Ollivier J, Orecchini A, Plazanet M, Zaccai G (2008) A benchmark for protein dynamics: ribonuclease A measured by neutron scattering in a large wavevector-energy transfer range. Chem Phys. doi: 10.1016/j.chemphys.2007.09.012

Wood K, Plazanet M, Gabel F, Kessler B, Oesterhelt D, Tobias DJ, Zaccai G, Weik M (2007) Coupling of protein and hydration-water dynamics in biological membranes. Proc Natl Acad Sci USA 104:18049–18054

Zaccai G, Gilmore DJ (1979) Areas of hydration in the purple membrane of Halobacterium halobium: a neutron diffraction study. J Mol Biol 132:181–191