Proton transfer via a transient linear water-molecule chain in a membrane protein

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High-resolution protein ground-state structures of proton pumps and channels have revealed internal protein-bound water molecules. Their possible active involvement in protein function has recently come into focus. An illustration of the formation of a protonated protein-bound water cluster that is actively involved in proton transfer was described for the membrane protein bacteriorhodopsin (bR) [Garczarek F, Gerwert K (2006) Nature 439:109–112]. Here we show through a combination of time-resolved FTIR spectroscopy and molecular dynamics simulations that three protein-bound water molecules are rearranged by a protein conformational change that resulted in a transient Grotthuss-type proton-transfer chain extending through a hydrophobic protein region of bR. This transient linear water chain facilitates proton transfer at an intermediate conformation only, thereby directing proton transfer within the protein. The rearrangement of protein-bound water molecules that we describe, from inactive positions in the ground state to an active chain in an intermediate state, appears to be energetically favored relative to transient incorporation of water molecules from the bulk. Our discovery provides insight into proton-transfer mechanisms through hydrophobic core regions of ubiquitous membrane spanning proteins such as G-protein coupled receptors or cytochrome C oxidases.

FTIR spectroscopy | functional water molecules | proton-transport chain

The membrane protein bacteriorhodopsin (bR) (1) is an established system for the study of the functional role of protein-bound water molecules and is therefore a favorable model relative to other proton pumps and channels that bind and may also rely on active water molecules (2–4). High-resolution X-ray crystal structures resolving protein-bound water molecules are available for bR in addition to functional studies by time-resolved FTIR spectroscopy. Together they provide a detailed molecular reaction mechanism (1, 5–9).

bR performs a light-driven proton-pumping cycle. Proton release to the extracellular solvent and the involvement of protonated water molecules are well characterized (5, 8–10). However, description of a reprotonation pathway from the cytoplasm to the central proton-binding site, the Schiff base (SB), remains incomplete (for more details, see Fig. 1A). For the proton transfer from Asp96 to SB in the second cycle-half (12), only two water molecules (Wat501 and 502) appear in ground-state crystal structures. The region between these two residues is primarily hydrophilic (1). Two water molecules alone cannot span a Grotthus-type “proton wire” of hydrogen (H)-bonded water molecules (9, 14) between these positions. However, this hydrophobic barrier blocks proton transfer from the protonated SB (PSB) to the cytoplasmic proton uptake site in the release-cycle step, which would counteract the establishment of a proton gradient.

A bR conformational change during proton uptake is believed to allow water molecules to invade from bulk cytoplasmic water and build a transient proton-transfer chain (15–17). X-ray structures of various photocycle intermediates based on crystallized, mutated bRs at low temperature, show variability in numbers and positions of water molecules in this domain (15–17) (Fig. S1).

To further define how water molecules organize in this space in the WT protein, and to elucidate an active role for protein-bound water molecules in the reprotonation mechanism, we applied time-resolved FTIR spectroscopy to study the WT protein under conditions closer to the physiologically ones.

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Results and Discussion

Strongly H-Bonded Waters at the Uptake Site. IR signatures of water-molecule chains are expected to show up in the spectral regime of strong H bonds, between 3,000 and 1,600 cm⁻¹ (18). In this region we identified a broad spectral signature from 2,750 to 2,540 cm⁻¹, which appears transiently in the M intermediate (Fig. 1B, blue). The origin of this broad absorbance change is complex because absorbance changes of various strongly H-bonded water molecules within the protein overlap in this region (Fig. S2). The early disappearance of this band demonstrates H-bond breakage between Wat402 and the PSB in the pentameric arrangement at the release site (Fig. 1A, red) due to the all-trans to 13-cis retinal isomerization (5, 8). The absorbance of Wat402 is not restored until the last step of the photocycle, during O-BR relaxation, when the pentameric arrangement is recreated by Asp95 deprotonation (5, 9, 11) (Fig. 1A, green). Here we identified an additional absorbance change, which is overlaid in M (Fig. 1B, blue) and indicates the appearance of strongly H-bonded water molecules.

We hypothesized that a water-molecule chain at the uptake site causes the additional IR signature in the M intermediate. To assign this additional feature, we performed FTIR measurements on the V49A mutant (Fig. 1B, red). Val49 is located in the hydrophobic cavity at the proton uptake site just between Asp96 and the SB. The V49A mutation destabilizes the proton uptake of the SB and accumulates a greater proportion of the N-like intermediate compared to WT (17). In this mutant the additional infrared absorbance change in the M intermediate is no longer observed when compared to WT. The residual absorbance change in V49A appears to reflect only the H-bond change of Wat402 at the release site, involving cleavage by isomerization followed by the restoration of the ground state arrangement. The disappearance of this band in V49A compared to WT indicates that the additional absorbance change (Fig. 1B, blue) reflects the appearance of strong H-bonded water molecules in the M intermediate of WT between Asp96 and the SB.

The Conformational Change at the Uptake Site. Formation of a deduced water chain depends on protein conformational changes during the photocycle, which would provide sufficient space for the chain. The largest conformational changes (19) occur in late M (20–22) or N (17, 23). A crystal structure showing sufficient space and water molecules between Asp96 and the SB to form a stable Grothuss connection was presented [Protein Data Bank (PDB) ID code 1P8U] (17), but this structure is of the invasive V49A mutant and may not reflect water distribution within the WT protein in the M intermediate due to steric alterations in V49A in the region of interest (17). In V49A an N’ intermediate is stabilized that shows conformational changes in helices C, F, and G. To obtain the corresponding N intermediate WT structure, we used homology modeling together with molecular dynamics (MD) simulations (Materials and Methods) and mutated Ala49 back to WT valine in silico. When comparing the conformational change in the simulated WT N intermediate to the ground state (9) in Fig. 2, we observed that the retinal was moved up toward Trp182. Additionally, the SB moved away from the release site and toward Asp96 at the uptake site (compare also Fig. S1).

The deprotonated SB must move from the proton release site in an early M state (M₁) to the proton uptake site in a late M state (M₃) to achieve vortical proton transfer through the protein. This is called the M₁-M₃ transition, but early M₄ and late M₅ absorptions overlap spectrally and are hard to distinguish kinetically (25). The L-M rate constant (IR absorbance changes in Fig. 1B) is dominated by the L-M₄ transition, whereas the L-M₅ and M₃-M₄ transitions are kinetically merged and not resolved (25). The low temperature stabilized X-ray structure intermediate of the D96N mutant (PDB ID code 1ICS) (15) shows an upward retinal movement and SB reorientation as expected for late M₅, as does the M structural of the E204Q mutant (PDB ID code 1F4Z) (16) (Fig. S1). We assume that our homology model of the WT N intermediate approximates the structure immediately after proton transport to the SB (“early N”), and that this simulated structure is similar to the M₄ structure. Asp96 in the homology model presented here is deprotonated as expected for WT N. This is in contrast to the N’ crystal structure of V49A, in which both SB and Asp96 are protonated (17), which does not represent a functionally relevant intermediate.

We hypothesized that the conformational change in the simulated WT N intermediate provides the space required for a water chain between Asp96 and the SB. The movement of the retinal C₁₃-methyl group shifts Trp182 toward the cytoplasmic protein side (Fig. 2, step 1) and pushes the extracellular end of helix F outward by about 1 Å. This is consistent with previous results (19, 21, 22). Trp182 movement in helix F cleaves the H bond to Wat501 (Fig. 2, step 2). The second hydrogen of Wat501 is bonded to the Ala215 backbone carbonyl group in helix G. There is a π-bulge in helix G at this residue so the Ala215 backbone carbonyl group is not H-bonded within the helix as usually expected (1) (Fig. 2). This arrangement is destabilized in M by retinal movement. Furthermore, retinal relaxation applies a different conformational restraint on the Lys216 backbone because of its side chain. As a result, hydrogen bonding within helix G changes from Lys216/Gly220 to Ala215/Gly220, and the carbonyl group of Lys216 moves out of the helix (Fig. 2, step 3 and Fig. S3). Other movements include a side chain rotation of Leu93 away from Ala215 and movement of the Phe219 side chain (Fig. 2).

These movements destabilize the previous hydrogen bonding positions for protein-internal water molecules and provide enough space for water molecules to arrange from Asp96 to the SB between helices C, F, and G (Fig. S3). In contrast to other investigations (19–22), the inward movement of Ala215 and outward movement of Lys216 do not move the end of helix G in our simulated structure. As shown previously (22, 26), the outward movement of helix F precedes the inward movement of helix G. Our simulated structure may reflect early F and G helix positioning. Alternatively, our simulation time may be too short to
show full helix G movement, which takes place on a microsecond time frame.

**Possible Water Arrangements at the Uptake Site by MD Simulations.**

We sought to determine the number of water molecules that are necessary to bridge Asp96 and SB. Intermediate crystal structures demonstrated the movement of water molecules into the hydrophobic barrier in M and N (15–17), but the positions and the number of water molecules differ greatly between these studies (8). In the N’ structure of V49A, alanine occupies less space than valine, such that additional water molecules may be present in the ground state as compared to WT. Using MD simulations we show the WT scenario in the late M/early N intermediate by keeping two to five water molecules between Asp96 and the SB (Fig. 3). In the simulations of the WT N structure, a linear chain of three water molecules provides a stable connection and a continuous proton pathway in late M/early N (Fig. 3). This three-water-molecule arrangement was not observed in the N’ structure of V49A, probably because of steric differences compared to WT.

**FTIR Identifies a Third, Highly Mobile Water Molecule.** Computationally, three water molecules appear to arrange between Asp 96 and SB during the photocycle. However, in WT ground-state crystal structures, only two water molecules, Wat501 and Wat502, were identified in this region (Fig. S1, yellow). The Wat501 band is indicated at 3,670 cm⁻¹ in M, showing the appearance of a dangling, non-H-bonded water molecule (6) (Fig. 4A). This band has an additional shoulder at 3,658 cm⁻¹, which kinetically behaves differently in M and N (Fig. 4A). Therefore, it must represent two water molecules at the Wat501 position (27) (Fig. S2B). We have named these molecules Wat501a and Wat501b. In agreement, the

![four water molecules](image1.png)

**Experimental Evidence for the Water Chain Between Asp 96 and the SB.** Mutagenesis of each of the amino acids neighboring the proposed transient water chain in our late M/early N simulated structure (Fig. 5A) affects the additional IR signature between 2,750 and 2,540 cm⁻¹ in the M intermediate (Fig. 5C). These data support the presence of the water chain at the proposed position. In contrast to the release-site protonated water cluster, which has a spectral signature between 2,400 and 1,800 cm⁻¹ with decreasing intensity (5), we now observe a signature between 2,750 and 2,540 cm⁻¹ with increasing intensity representing the water chain (Fig. S4A). Quantum mechanics/molecular mechanics (QM/MM) calculations of IR absorbance bands on the release site (10) confirmed the spectral signatures from FTIR measurements (5). We think, that future QM/MM calculations on the linear three-water chain presented here, will confirm the previously undescribed spectral signature as well.

For a linear water chain in a hydrophobic environment, both strong H bonds and dangling H-bond absorbance bands should appear in the FTIR spectra (27–29). The observed dangling vibrational band of Wat501a and 501b in M and N (Fig. 4A) fit well into the calculated spectral range of dangling waters in a protein (30). Mutagenesis of neighboring residues along the chain influenced these dangling bands (6). This constitutes additional supporting evidence for the path of the linear water chain between Asp96 and SB.
in the L-M transition (Fig. 4). Water molecules to rearrange and form a functional chain between Asp96 and SB, thus preventing proton backflow. The cofactor-twist relaxation locations in the first half of the pump cycle during proton release, either on Asp96 by late M or on SB by early N.

Coincidently, with the formation of a water chain as early as M, we observe an H-bond change at Asp96 from 1,746 to 2,750 and 2,540 cm⁻¹ (red; see also Fig. 1B). We deduced that this reflected the participation of Asp96 in chain formation at M, but the conformation extending to SB through H bonding was not established before the late M/early N intermediate. As soon as SB bonds to the nearby water molecule, a proton is transferred from Asp96 to reestablish PSB. Late M and early N differ particularly in this proton shift along the water chain H bonds as expected for Grotthuss proton transfer (Fig. 5B), and the excess proton is located either on Asp96 by late M or on SB by early N.

In conclusion, we used time-resolved FTIR spectroscopy and MD simulations to resolve a transient, linear, proton-transport water chain in bR. Water molecules are stored at inactive storage locations in the first half of the pump cycle during proton release, thus preventing proton backflow. The cofactor-twist relaxation due to deposition of the SB in the M intermediate repacks amino acids between helices C, F, and G. This causes the inactive water molecules to rearrange and form a functional chain between Asp96 (proton donor) and the central proton-binding site in the proton uptake phase of the cycle. Together with the "proton diode" at the release site (9), this switch-like mechanism at the uptake site ensures efficient, directional, cross-membrane proton transport. The key event is the precise relocation of protein-bound internal water molecules, which appears to be energetically favorable to the invasion and extrusion of water molecules from the protein exterior because the necessary conformational and entropy changes are much smaller. We present a concept of protein-bound water molecules being actively involved in proton function as a model that may be applicable to proton-transfer mechanisms known to occur through other membrane proteins, including G-protein coupled receptors, cytochrome C oxidase (3), photosystem II (2), and hydrogenases (31).

Materials and Methods

B. Mutation, Expression, and Purification. The mutagenesis, expression, isolation, and purification of bR in Halobacterium salinarum are described elsewhere (32–34).

Sample Preparation and FTIR Measurements. A suspension of 600 μg of purified purple membrane sheets in 1 M KCl and 100 mM Tris at pH 7 was centrifuged at 200,000 × g for 2 h. The resulting pellet was squeezed between CaF₂ windows and transferred into an air- or vacuum-tight cuvette. Membrane sheets were performed with a modified Bruker IFS 66v/SVS (35) and a specially modified Bruker Vertex 80 spectrometer, recording data with 4 cm⁻¹ spectral resolution and a time resolution of down to 30 ns. The sample was kept at 20 °C during measurements. The photocycle was triggered by an excimer laser (Lambda Physics 305i) driven dye laser (Counamarin153).

MD Simulations. The N model was created based on the N’ crystal structure of Schobert et al. (PDB ID code 1P8U) (17). Ala49 was mutated back to valine, thus recreating the WT sequence. Missing residues in the E-F loop were inserted based on the WT ground-state structure (PDB ID code 1QHJ) (36). These residues are also missing from other crystal structures, including 1C3W (1), 1M0M (37), and 1F4Z (16). The late M structure (PDB ID code 1C8S) (15) lacks more residues in this region. In silico mutation, followed by side chain minimization, was carried out with the MOBY program package (38). The applied protonations of amino acid chains matched the N intermediate. A bR trimer consisting of three ground-state molecules used as the starting structure previously (9, 39) was recreated from the crystallographic information using PyMOL (Schrödinger, LLC). The N’ intermediate may accumulate to a maximum of 40% in protein samples (17), so we replaced one of the ground-state structures with our N model by superposition of the transmembrane helix αC₃b atoms. The resulting trimer was then placed into a palmitoyloleylophosphatidylcholine membrane/saline environment and trajectories 20 ns in length were recorded, and water densities in the last 5 ns were analyzed, as described previously (39, 40).

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Fig. S1. Water molecule positions and retinal movements in different structures. Comparison of retinal movements and water position in the cytoplasmic half of bacteriorhodopsin (bR) as compared to WT [yellow, helices in blue, 1C3W (1)]. For comparison, the M intermediate crystal structure of mutant E204Q [1F4Z (2)] is shown in light orange and a late M of mutant D96N is shown in dark orange [1C8S (3)]. Movement of the C13 methyl group of the retinal is most pronounced in late M, as is the resulting movement of the indole ring of Trp182 and Lys216. Numbers and positions of water molecules differ, but in the M intermediate additional water molecules are moved from Wat501 into the vicinity of Asp96. None of these crystal structures show a chain of water molecules spanning Asp96 to Schiff base (SB).


Fig. S2. Decomposition of IR absorbance changes due to strong H-bonded water molecules. Time-resolved absorbance changes of bR WT from 3,000 to 2,540 cm⁻¹ (A, black line) show the (not time-resolved) rupture of the strong H bond of water 402 (B, red) to SB due to isomerization, and the appearance of strong H bonds in the M intermediate at the release site (1) (A and B, green). Dividing this spectral range into smaller domains reveals different absorbance changes. The domain from 3,000 to 2,800 cm⁻¹ (A, red line) shows the H-bond breakage of Wat402 (A, B, red) and the appearance of a strong H-bond at the release site (1) even more pronounced (A and B, green). The domain from 2,750 to 2,540 cm⁻¹ (A, blue line) shows an overlaid positive contribution already in the apparent L-to-M transition (marked in blue in A), which we attribute now to a transient water chain (B, blue), responsible for the reprotonation of SB by Asp96 in step 4.

Fig. S3. The π-bulge during molecular dynamics (MD) simulations. The π-bulge during MD simulations in the bR ground state and the simulated N structure. Hydrogen bond distances are indicated in yellow. In the simulations of WT bR (1), Lys216 and Gly220 form a stable helical hydrogen bond. Ala215 is oriented out of the helix and forms hydrogen bonds to Wat501a. In the simulated N structure, the π-bulge has switched: Ala215 forms a hydrogen bond to Gly220, whereas the carbonyl group of Lys216 is oriented out of the helix. However, the Ala215/Gly220 hydrogen bond in the simulated N structure is less stable than the Lys216/Gly220 hydrogen bond in the bR ground state, and Lys216 can transiently form a hydrogen bond with Gly220. This suggests that the switch of the π-bulge defect in N applies additional conformational stress to helix G and may transiently store energy from the retinal twist relaxation in the backbone. This may drive the back isomerization to all-trans retinal after reprotonation in N.


Fig. S4. Spectrally and kinetically resolved absorbance changes of strong H-bonded water molecules. M intermediate spectra are shown between 3,000 and 2,400 cm\(^{-1}\) (A) for WT (blue) and T89A (red), respectively. There is a clear additional band in the WT from 2,750 to 2,540 cm\(^{-1}\) as indicated in yellow (A). The time-dependent absorbance change between 2,750 and 2,540 cm\(^{-1}\) is shown in B. The additional band in the WT between 2,750 and 2,540 cm\(^{-1}\) disappears for T89A, and V49A (Fig. 1B). Its disappearance is better resolved by kinetics (B). An absorbance change in M is assigned to a linear water chain between Asp96 and SB.