X-ray structure of human aquaporin 2 and its implications for nephrogenic diabetes insipidus and trafficking

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Human aquaporin 2 (AQP2) is a water channel found in the kidney collecting duct, where it plays a key role in concentrating urine. Water reabsorption is regulated by AQP2 trafficking between intracellular storage vesicles and the apical membrane. This process is tightly controlled by the pituitary hormone arginine vasopressin and defective trafficking results in nephrogenic diabetes insipidus (NDI). Here we present the X-ray structure of human AQP2 at 2.75 Å resolution. The C terminus of AQP2 displays multiple conformations with the C-terminal α-helix of one protomer interacting with the cytoplasmic surface of a symmetry-related AQP2 molecule, suggesting potential protein–protein interactions involved in cellular sorting of AQP2. Two Cd2+-ion binding sites are observed within the AQP2 tetramer, inducing a rearrangement of loop D, which facilitates this interaction. The locations of several NDI-causing mutations can be observed in the AQP2 structure, primarily situated within transmembrane domains and the majority of which cause misfolding and ER retention. These observations provide a framework for understanding why mutations in AQP2 cause NDI as well as structural insights into AQP2 interactions that may govern its trafficking.

Significance

Human aquaporin 2 (AQP2) is found in the kidney collecting duct, where it translocates water across the apical membrane and is crucial for urine concentration. AQP2 is regulated by trafficking between intracellular storage vesicles and the apical membrane, a process that is tightly controlled by the pituitary hormone arginine vasopressin. Defective AQP2 trafficking leads to nephrogenic diabetes insipidus (NDI), a water balance disorder characterized by large urine volumes, leading to dehydration. We have solved the X-ray structure of human AQP2 at 2.75 Å resolution. This structure deepens our molecular understanding of AQP2 trafficking, as well as serves as a structural scaffold for understanding why AQP2 mutations cause NDI.


The authors declare no conflict of interest.

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Data deposition: The atomic coordinates have been deposited in the Protein Data Bank, www.pdb.org (PDB ID code 4NEF).

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Water is the major ingredient of the human body, constituting 55–65% of our total body weight (1). Water homeostasis is maintained by the kidneys, which filter ~180 L of primary urine every day. Although most water is constitutively reabsorbed in the proximal tubules and descending limbs of Henle (2), the body’s water balance is fine-tuned by regulated water reabsorption, which takes place in the kidney collecting duct. Water reabsorption is mediated by aquaporins, membrane-bound water channels, of which seven of the 13 human isoforms have been located in the human kidney (3). Of these, human aquaporin 2 (AQP2) is present in the principal cells of the collecting duct, where it translocates water across the apical membrane and is crucial for urine concentration. AQP2 is regulated by trafficking between intracellular storage vesicles and the apical membrane, a process that is tightly controlled by the pituitary hormone arginine vasopressin. Defective AQP2 trafficking leads to nephrogenic diabetes insipidus (NDI), a water balance disorder characterized by large urine volumes, leading to dehydration. We have solved the X-ray structure of human AQP2 at 2.75 Å resolution. This structure deepens our molecular understanding of AQP2 trafficking, as well as serves as a structural scaffold for understanding why AQP2 mutations cause NDI.
tetramer and stabilize a specific conformation of the cytoplasmic loop D that is necessary for this interaction to occur. Radioligand binding studies show that oocytes expressing AQP2 bind Ca\(^{2+}\), supporting Ca\(^{2+}\) as the physiological ligand for these sites. Several interactions that lead to NDI when disrupted by mutation are also highlighted. Together, this structure provides insights into AQP2 mutations in NDI and opens new opportunities to study the structural mechanism behind membrane protein sorting, including ER recognition of misfolded proteins.

### Results

**Crystallization of AQP2.** Human AQP2 expressed in insect cells (17) has previously yielded 2D crystals diffracting to 4.5 Å resolution (18). Extensive efforts to produce better diffracting 3D crystals using full-length human AQP2 produced in insect cells failed to sufficiently improve the diffraction. AQP2 was therefore C-terminally truncated at Pro242 and expressed in *Pichia pastoris*. Pro242 corresponds to the last visible residue in the structure of human AQP5 (19), AQP2’s closest paralog with which it shares 63% sequence identity (19). With this truncated AQP2, significant improvement in diffraction quality was achieved. Complete data to 2.75 Å were collected on a single frozen crystal (Table 1), yielding a molecular replacement solution using a homology model based on human AQP5 (19). Iterative rounds of manual rebuilding and structural refinement resulted in a model with an R-factor and free R-factor of 20.2% and 22.5%, respectively (for typical electron density, see Fig. S2). Of particular significance, these crystals belonged to the space group P4\(_2\)\(_1\)\(_2\) with one tetramer in the asymmetric unit, whereby structural differences between individual protomers within the tetramer can be observed.

**Crystal Structure of AQP2.** AQP2 displays the conserved aquaporin fold, assembling as a tetramer with each protomer consisting of six transmembrane helices. Two half-membrane spanning helices, formed by loops B and E, constitute a seventh pseudotransmembrane segment (Fig. 1 A and B). Each protomer contains a water-conducting pore through which water is transported in a single file. Two conserved regions are of particular interest within this pore: the NPA region, containing two copies of the Asn-Pro-Ala (NPA) aquaporin signature motif, and the aromatic/arginine selectivity filter (ar/R), which determines transport specificity (20) (Fig. S3).

![Fig. 1.](image-url) Overall structure of human AQP2. (A) Overview of the AQP2 tetramer with half helices formed by loops B and E highlighted in yellow. (B) Overview of the AQP2 tetramer from the intracellular side. The color scheme for each of the protomers is used throughout the article (A, purple; B, magenta; C, pink; D, light pink).

When calculated using HOLE (21), all protomers show a similar pore profile, with the ar/R region constituting its narrowest point with an average diameter of 1.8 Å (Fig. S4).

**C Termini Show Significant Conformational Variability.** A striking difference between the X-ray structure of AQP2 and all other mammalian AQP structures is the highly variable position of the short C-terminal helix (Fig. 2A). This flexibility is likely to have hindered full-length AQP2 from yielding well-diffracting crystals. The C-terminal helix adopts a unique conformation in each of the four AQP2 protomers, and in one protomer (D), it is highly disordered. None of these conformations have previously been observed in mammalian AQP structures (19, 22–24), for all of which the C-terminal helix lies across the AQP cytoplasmic surface with only small variations between structures. In one protomer, the C-terminal helix interacts with a symmetry-related AQP2 molecule, forming one of several interactions that are important for crystal contacts. Specifically, four hydrophobic leucine residues (Leu230, 234, 247, and 240) that align on the same side of the C-terminal helix of protomer C are sheltered from exposure to the cytoplasm by this helix inserting into the cytoplasmic vestibule on the surface of protomer D of another AQP2 tetramer (Fig. 2B). This interaction is further stabilized by hydrogen bonds between side chains and main-chain carbonyls (Fig. S5).

An unusual flexibility in the AQP2 C terminus is likely to arise from two consecutive prolines (Pro225 and Pro226) that form a hinge region, whereas only one proline residue is present in the corresponding position in other mammalian AQP structures. In particular, the second proline may be functionally important in AQP2 trafficking, as the P226A mutant caused ER retention in Madin–Darby canine kidney cells, whereas the P225A mutant did not (25). However, whether this flexibility is important for phosphorylation and ubiquitination of the C terminus, events that are critical for AQP2 trafficking to and from the apical membrane, remains to be demonstrated.

**N Termini Adopt Two Distinct Conformations.** For the AQP2 N terminus, two distinct conformations can be seen (Fig. 2C), although this region is disordered before residue six in protomers B and C. In protomer A, the N terminus adopts a conformation allowing hydrogen bond interactions to form between Glu3, Ser82, and Arg85. This structural triad is also observed in human AQP5 (19) and in the spinach water channel plasma membrane intrinsic protein 2.1 (SOP1P2.1) (26), where for the latter it is proposed to play an important role in the gating mechanism. In contrast, protomer D of AQP2 adopts a conformation similar to that seen in AQP1, with transmembrane (TM) helix 1 extending an additional full turn into the cytoplasm. This conformation was also observed in the phospho-mimicking S115E mutant of AQP1, with transmembrane (TM) helix 1 extending an additional full turn into the cytoplasm.

### Table 1. Crystallographic data and refinement statistics

<table>
<thead>
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<th>Data collection</th>
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<tr>
<td>Beamline/wavelength</td>
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<tr>
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<td>P4(_2)(_1)(_2)</td>
</tr>
<tr>
<td>Cell dimensions, a, b, c, Å</td>
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<tr>
<td>Resolution, Å</td>
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<tr>
<td>No. of reflections</td>
<td>33,066 (4,805)</td>
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<tr>
<td>Rmerge/Rmerge*</td>
<td>0.095 (1.059)/0.044 (0.630)</td>
</tr>
<tr>
<td>CC1/2</td>
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</tr>
<tr>
<td>I/ln</td>
<td>9.2 (1.3)</td>
</tr>
<tr>
<td>Completeness/redundancy</td>
<td>99.9 (99.8)/42.4 (3.7)</td>
</tr>
</tbody>
</table>

| Refinement |  |
| Resolution, Å | 50.0–2.75 (2.95–2.75) |
| Rwork/Rfree* | 0.202/0.225 (0.224/0.247) |
| Average B-factors, Å\(^2\) | 89.9 |
| r.m.s.d. from restraint target values |  |
| Bond lengths, Å/angles, ° | 0.009/9.13 |
| Ramachandran plot analysis, % | Favorable/allowed/disallowed 87.4/12.6/0.0 |
| PDB code | 4NEF |

*Number in parentheses is for highest resolution shell.

*Rfree was calculated for 5% of reflections randomly excluded from the refinement.

![Diagram](image-url) Overview of the AQP2 tetramer with half helices formed by loops B and E highlighted in yellow. (B) Overview of the AQP2 tetramer from the intracellular side. The color scheme for each of the protomers is used throughout the article (A, purple; B, magenta; C, pink; D, light pink).
SoPIP2;1, where the extension of TM helix 1 disrupts a Ca\(^{2+}\) binding site implied in gating (27). The reoccurrence of these two specific N-terminal conformations among regulated eukaryotic aquaporins suggests a structural principle governing their regulation. Indeed, the AQP2 N terminus is important for trafficking, as exchanging both the N and C termini of AQP1 for those of AQP2 was needed to transpose 1-desamino-8-D-arginine vasopressin–regulated shuttling, a characteristic of AQP2, onto AQP1 (25).

**AQP2 Binds Cd\(^{2+}\) at Two Sites.** Two prominent peaks, visible at 14 σ and 8.7 σ, respectively (σ is the SD of the map), were found in the AQP2 F\(^{obs}\)-F\(^{calc}\) residual electron density map. Corresponding difference density peaks were also seen in the F\(^{obs}\)-F\(^{obs}\) anomalous difference density map (9.4 σ and 4.5 σ, respectively), specifically indicating the presence of a heavy metal. Because CdCl\(_2\) was used as an additive during crystallization, these two peaks were assigned as Cd\(^{2+}\), with two Cd\(^{2+}\) ions built per tetramer (Cd1 and Cd2) at the cytoplasmic side of the membrane (Fig. 3A). Cd1 has full occupancy, binds at the membrane interface between protomers A and D, and is ligated by GluA155 of loop D and GlnD57 of TM helix 2, as well as two water molecules (Fig. 3B). Cd2 has partial (65%) occupancy, is located between loop B and the C-terminal tail in protomer C, and is ligated by HisC80, GluC232, and one water molecule (Fig. 3C).

Cd\(^{2+}\) binding clearly influences the structure of loop D, which adopts different conformations in all four protomers (Fig. S6A). The bonding of Cd1 displaces loop D of protomer A closer to the center of the AQP2 tetramer, where loop D of protomers A, B, and D all interact via hydrogen bonds between AsnA156, AsnD156, and GluB155 (Fig. S6B). In protomer D, this creates a binding pocket into which the C-terminal helix of a symmetry-related protomer C binds (Figs. 2B and 3A). Furthermore, the binding of Cd2 between loop B and the C terminus of protomer C helps position the C-terminal helix for this interaction (Fig. 3A). Thus, the binding of Cd\(^{2+}\) appears to be important for interactions between AQP2 tetramers within the crystal and explains why this additive was critical during crystal optimization.

**Oocytes Expressing AQP2 Bind Ca\(^{2+}\).** Although Cd\(^{2+}\) binding is clearly important for crystal formation, this does not necessarily imply functional significance. A regulatory role for divalent cations was suggested from the crystal structure of the gated spinach aquaporin SoPIP2;1 in which binding of Cd\(^{2+}\) was observed to control the conformation of the gating element loop D (26). For SoPIP2;1 it was speculated that the Cd\(^{2+}\) ion is replaced by Ca\(^{2+}\) in vivo, and independent studies showed Ca\(^{2+}\) almost completely abolishing the water transport ability of PIP aquaporins (28).

To test whether AQP2 binds Ca\(^{2+}\), we performed a radioactive calcium-binding assay. In this assay, either control or AQP2-expressing *Xenopus* oocytes were generated and their membranes isolated and incubated in medium containing \(^{45}\)Ca\(^{2+}\). Following extensive washing, liquid scintillation counting revealed that significantly more \(^{45}\)Ca\(^{2+}\) was bound to membranes of AQP2-expressing oocytes than to those of water-injected controls (Fig. 3D). These data strongly suggest that AQP2 binds Ca\(^{2+}\) and, as for SoPIP2;1, imply that the observed Cd\(^{2+}\) binding sites in the AQP2 structure represent Ca\(^{2+}\) binding sites in vivo.

**Discussion**

**NDI-Causing Mutations in AQP2.** NDI is characterized by the inability to concentrate urine and can lead to severe dehydration unless water intake is increased. Fifty-one different mutations in the AQP2 gene have been identified in patients suffering from NDI (15, 16), most of which lead to autosomal recessive NDI whereby AQP2 is misfolded, retained in the ER, and rapidly degraded. In some cases, however, ER-retained mutants have been shown to retain some functionality (29–32), indicating that the degree of misfolding can be minor.

The locations of thirty-one NDI-causing mutations are identified in the AQP2 crystal structure, most of which are located within the transmembrane region (Fig. 4A). These are typically situated at helix–helix interfaces and presumably disturb packing interactions and correct folding and/or assembly of the AQP2 tetramer (Fig. S7A). A number of other mutations are found within the water-conducting pore, affecting key functional residues such as Asn68, Ala70, and Pro185 of the NPA region or Arg187 of the selectivity filter, or causing pore narrowing as in...
the case of V71M and V168M mutations (Fig. S7B). Precisely how these folding defects are recognized for retention in the ER and degradation cannot be deduced from the X-ray structure of only wild-type AQP2.

A small number of NDI-causing mutations are also found in loop regions. Within the extracellular loop C, mutation of Thr125 and Thr126 to methionine leads to autosomal recessive NDI (30, 31). These three threonines are situated close to an N-glycosylation site at Asn123. When compared against other mammalian AQP6s that lack this glycosylation site, the structure of loop C is distinctly different for AQP2 (Fig. 4B). Glycosylation at this site has been suggested to be important for protein sorting in the Golgi complex, as AQP2–N123Q is retained in this organelle when expressed in renal cells (33). In oocytes, however, expression of AQP2–T125M and T126M leads to ER retention (31, 32), whereas the glycosylation mutant N123Q does not (33), suggesting that it might not be the lack of glycosylation per se that causes ER retention but rather mutation-induced structural changes within loop C.

Within the cytoplasmic loop D, the D150E mutant has been identified in patients suffering from recessive NDI (34, 35). Our X-ray structure reveals that Asp150 mediates an interaction between loop D and the proximal end of the C-terminal tail, with Asp150 connecting to Pro225 via hydrogen bonds mediated by Arg152 (Fig. 4C). This interaction is conserved in other mammalian AQP6s that lack this glycosylation site (19, 22–24, 36). For AQP5, it was argued that disturbing this interaction might cause structural changes to the C terminus that flags the protein for trafficking (19). It is certainly plausible that the bulkier glutamate head group of the AQP2 D150E mutant would perturb this network of hydrogen bonds, resulting in structural changes, which could be recognized by the ER quality control as misfolded protein.

**Mutations Near Cd1 Cause ER Retention.** Our crystal structure of AQP2 revealed two unanticipated Cd2+ binding sites per tetramer (Fig. 3A). It is therefore striking that mutation of the Cd1 ligand Gln57 to proline, which disrupts this divalent cation binding site, has been identified in NDI patients and shown to cause ER retention in oocytes (37). It is plausible that the loss of a divalent cation at this site flags AQP2 for ER retention.

Mutation of Ser148, which is situated in a casein kinase II consensus site at the cytoplasmic end of TM helix 4, also results in ER retention (6). The structure reveals that Ser148 of promoter A interacts with the Cd2+ ligand Gln57 through a direct hydrogen bond (Fig. 4D). This structural arrangement is strongly reminiscent of the gated spinach aquaporin SoPIP2;1, for which an interaction between Ser115 and the Cd2+ ligand Gln31 plays an important functional role by regulating the opening or closure of the channel (Fig. S8) (26). In SoPIP2;1, phosphorylation of Ser115 disrupts this interaction and perturbs the Cd2+ binding site, thereby inducing channel opening through a conformational change of cytoplasmic loop D (26, 27). Should Ser148 of AQP2 be phosphorylated, this could be signaled through a change in protein structure triggered by disruption of this divalent cation binding site.

An effect of putative phosphorylation of Ser148 of AQP2 was studied in oocytes by mutating this serine to alanine or aspartate to abolish or mimic phosphorylation, respectively (6). In these studies it was observed that the S148A mutant was properly targeted to the plasma membrane, whereas the S148D mutant was retained in the ER. Although no evidence for in vivo phosphorylation of Ser148 has been presented, it is intriguing that this serine-to-aspartate mutation, which almost certainly disrupts its interaction with the Cd2+ ligand Gln57, has such a clear effect on AQP2 trafficking, whereas replacement with alanine, which should not affect Cd2+ binding, does not. Moreover, an A147T mutation, the site adjacent to Ser148, has been identified in patients with autosomal recessive NDI, again due to ER retention of AQP2 (30). Although Ala147 does not directly interact with any Cd2+ ligand, mutation of this residue might perturb the Cd2+ binding site. Irrespective of its physiological significance, it is highly suggestive that missense mutation of three residues in close proximity to this divalent cation binding site causes ER retention. This indicates that this region is unusually sensitive to perturbations that result in the protein being viewed as misfolded by the ER quality control.

**Potential Role for Ca2+ Binding in AQP2 Trafficking.** Although Cd2+ is observed in the AQP2 structure because of its presence in the crystallization conditions, Ca2+ is a more likely physiological ligand. This argument is supported by our results from radioactive
binding assays demonstrating that oocytes expressing AQP2 bind significantly more Ca\(^{2+}\) than control oocytes (Fig. 3D). Binding of AVP to the V2R triggers, besides a rise in cAMP level, an increase in intracellular Ca\(^{2+}\) levels (38–42). Moreover, AVP-induced plasma membrane sorting of AQP2 is reduced by calmodulin and ryanodine receptor inhibitors, indicating that release of intracellular Ca\(^{2+}\) from ryanodine-sensitive stores is important for AQP2 targeting to the apical membrane (43, 44). Because the AQP2 Ca\(^{2+}\)-binding sites identified in the crystal structure are accessible from the cytoplasm, we hypothesize that transient changes in the intracellular Ca\(^{2+}\) concentration triggers changes in the structure of AQP2 and that these changes correlate with Ca\(^{2+}\)-dependent targeting of AQP2 to the apical membrane.

**LIP5 Interacts with the C-Terminal Helix of AQP2.** The presence of AQP2 in the apical membrane is affected not only by AVP-stimulated fusion of storage vesicles but also by the internalization rate upon cessation of the AVP signal. AQP2 internalization involves ubiquitination of Lys270 at the C terminus, which enhances endocytosis and targets AQP2 to multivesicular bodies (MVBs) for subsequent lysosomal degradation (19). A protein involved in MVB formation is the lysosomal trafficking regulator interacting protein 5 (LIP5). LIP5 is abundantly expressed in the collecting duct (45) and has been shown to interact with AQP2 and facilitate its lysosomal degradation (46).

Biochemical studies have localized the LIP5 binding site of AQP2 to residues 230–243, all of which are located within the AQP2 C-terminal helix (46). Moreover, LIP5 is known to bind proteins containing leucine-rich regions (47), and leucines 230, 234, 237, and 240 all align on the same side of the C-terminal helix in AQP2 (Fig. 2B). It is therefore intriguing that, for pro- tomer C, this leucine-rich region interacts with the cytoplasmic surface of a symmetry-related AQP tetramer. We conclude that the observed interaction arises from the propensity of the C-terminal helix to participate in protein–protein interactions and similar interactions are presumably important in LIP5-mediated trafficking of AQP2.

An involvement of this short C-terminal helix in protein–protein interactions has also been reported for the AQP0–Calmodulin interaction in which one Calmodulin molecule binds the C-terminal helix from two AQP0 protomers simultaneously (48, 49). This interaction, which results in reduced AQP0 water permeability, was shown to be dependent on four hydrophobic residues (Leu227, Val230, Leu234, and Leu237) aligning on the same side of the C-terminal helix, as is observed for Leu230, Leu234, Leu237, and Leu240 in AQP2 (49). Furthermore, these authors state that rotational movements of the C-terminal helices are necessary for the interaction to occur. Hence, exposed hydrophobic residues on the aquaporin C-terminal helix may well be a reoccurring motif involved in AQP–protein interactions.

In the case of AQP2, several proteins have been suggested to interact with the AQP2 C terminus (50–52). Whether or not these interactions involve hydrophobic interfaces within the C-terminal helix remains to be shown. Nevertheless, we suggest that the conformational variability of the AQP2 C terminus seen in the crystal structure allows for exposure of a variety of binding surfaces, whereby interaction with a wide range of proteins can occur.

**Conclusion**

A major challenge in membrane protein structural biology is to take the step from understanding the structure–function relationships of individual membrane proteins to understanding how these are regulated by posttranslational modification and protein–protein interactions. AQP2 is the most clinically studied member of the aquaporin family and is by far the best characterized in terms of biosynthesis, trafficking from storage vesicles to the apical membrane and vice versa, degradation, posttranslational modifications, protein–protein interactions, as well as mutations in patients suffering from NDI. The crystal structure of AQP2 provides structural insights into all of these phenomena. A deeper understanding of how AQP2 mutants are recognized and retained in the ER, or how other proteins such as LIP5 interact with the C-terminal helix, will require further structural studies of AQP2 including complex structures with bound interaction partners. Our structure of AQP2, recovered through rational modification of the AQP2 construct, both deepens our understanding of its function in the human kidney and lays a platform from which protein–protein interactions regulating a membrane protein channel activity by trafficking can be explored.

**Materials and Methods**

**Cloning, Overproduction, and Membrane Preparation.** C-terminally truncated human AQP2 was cloned and transformed into P. pastoris. Cells were grown in a 3 L fermentor typically resulting in 500 g wt cells/L culture after 24 h of MeOH induction. After harvesting using centrifugation, cells were broken using a French Press and membranes were pelleted. The resulting membranes were washed in an urea buffer and stored at -80 °C until further use. For further details, see SI Materials and Methods.

**Solubilization, Purification, and Crystallization.** Membranes containing AQP2 was solubilized in 2% (wt/vol) n-octyl-β-d-glucopyranoside (NG) by stirring for 1 h in an ice bath. AQP2 was purified and concentrated to a final concentration of 7.5 mg/mL (SI Materials and Methods). During the last purification step, the detergent was changed from NG to octylglucoside–neopentylglycol (OGNGP). Crystals were obtained using the hanging drop vapor diffusion technique by mixing protein and reservoir [20–30% (vol/vol) PEG400, 0.1 M Tris pH 8, 0.1 M NaCl, 0.15 M MgCl₂] in a 1:1 ratio, with 0.1 M CdCl₂ added to the drop only (SI Materials and Methods). Rod-shaped crystals appeared within a few days. Following cryoprotection (SI Materials and Methods), crystals were flash frozen in liquid nitrogen and stored for data collection.
Data Collection and Structural Determination. Complete data at 100 K from a frozen crystal were collected at European Synchrotron Radiation Facility (ESRF), Grenoble, France. Crystals belonged to space group P4₁ with one tetramer in the asymmetric unit (α = 119.11 Å, β = 119.11 Å, c = 90.48 Å, α = β = γ = 90°). Following processing and scaling of image data, the structure was solved by molecular replacement (SI Materials and Methods). Iterative rounds of refinement and manual model building (SI Materials and Methods) resulted in a model consisting of four protein chains (A–D) with the following residues—A2–240, B6–238, C6–241, and D2–233—with two C-telopeptides, one A100–1000, and 58 water molecules. The final R-factor and free R-factor for this model is 20.2% and 22.5%, respectively.

Radioactive Calcium Binding Assay on Total Membranes of Oocytes. Oocytes were isolated and injected with 10 nM AQP2 DNA in 50 nL water per oocyte or water for 3 d, after which membranes were prepared, incubated with ⁴⁰Ca²⁺, and washed (SI Materials and Methods). Following transfer to counting tubes, 4 mL of scintillation liquid was added and counted in Tri-Carb 299 TR liquid scintillation counter. High water permeability confirmed expression of AQP2. Each value consists of three independent groups of 10 oocytes. The experiments were performed three times. The difference between groups was tested by Student t test. A P value < 0.05 was considered statistically significant.

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