Rhodopsins are the most universal biological light-energy transducers and abundant phototrophic mechanisms that evolved on Earth and have a remarkable diversity and potential for biotechnological applications. Recently, the first sodium-pumping rhodopsin KR2 from *Krokinobacter eikastus* was discovered and characterized. However, the existing structures of KR2 are contradictory, and the mechanism of Na⁺ pumping is not yet understood. Here, we present a structure of the cationic (non H⁺) light-driven pump at physiological pH in its pentameric form. We also present 13 atomic structures and functional data on the KR2 and its mutants, including potassium pumps, which show that oligomerization of the microbial rhodopsin is obligatory for its biological function. The studies reveal the structure of KR2 at nonphysiological low pH where it acts as a proton pump. The structure provides new insights into the mechanisms of microbial rhodopsins and opens the way to a rational design of novel cation pumps for optogenetics.

**INTRODUCTION**

The first microbial rhodopsin (bacteriorhodopsin) was discovered in the early 1970s in halophilic archaea (salt-loving microbes that live in saturated brines) (1). It was shown that bacteriorhodopsin is a light-driven proton pump used to generate a light-driven proton gradient as a source of energy for the cell (2). Soon after bacteriorhodopsin, two sensory rhodopsins (also known as slow rhodopsins) were found in the same microbe (*Halobacterium salinarum*) (3, 4). Then, again in the same organism, a fourth rhodopsin (5) was found that functions as an inward-directed chloride pump (halorhodopsin) that maintains the proper electrochemical balance (6). Because of their light activation via the chromophore retinal, all rhodopsins undergo a photocycle, which is connected to the function of these proteins. Therefore, rhodopsins played a key role in membrane protein research, extending our understanding of the molecular mechanisms of bioenergetics and transmembrane signaling and in the development of new biophysical, biochemical, and structural biology (x-ray crystallography and electron microscopy) methods, approaches, and potential applications (7).

In 2013, a new microbial rhodopsin was found in marine bacteria *Krokinobacter eikastus*, which can pump either lithium or sodium cations, but it pumps protons in the absence of these ions (8). This *K. eikastus* rhodopsin 2 (KR2) pumps sodium under physiological conditions (~500 mM NaCl and pH around 8 in the ocean) and belongs to a family of rhodopsins containing a unique NDQ motif, which is formed by the Asn**112**, Asp**116**, and Gln**123** triad. The unique residue in Na⁺-pumping rhodopsins, Asp**116**, was suggested to serve as a proton acceptor from the retinal Schiff base (RSB) (8). The photocycle of KR2 indicated that Na⁺ uptake occurs with the rise of the O intermediate. It was shown by Fourier transform infrared spectroscopy, however, that the ion already binds to the protein in the resting state at pH 8.0 and that the binding site is located distant from the RSB region. In 2015, crystal structures of KR2 were obtained simultaneously by Kato *et al.* (9) and our group (10). All structures were determined using crystals initially grown at a pH not higher than 5.6. The structures of KR2 solved by both groups at acidic pH 1.45 (pH 4.3) and 2.3 Å (pH 4.0) coincide, except for the presence of additional water molecules and the proposed earlier sodium ion bound to the protein surface in the model with higher resolution. The proteins in the crystals were in monomeric form. Optical properties of the protein are altered at acidic pH, and later in 2015, it was shown that KR2 pumps sodium ions and protons competitively and that the rate constant for Na⁺ uptake is much smaller than that of H⁺ (11). Under physiological conditions, when [Na⁺] is about 10⁷ times higher than [H⁺], the protein almost exclusively pumps sodium ions, so we will call this KR2 form as a Na⁺-pumping form. Determining the structure of Na⁺-pumping form is of high importance for understanding the mechanism of sodium translocation. Previously, this was approached in two different ways. In (9), the structure was obtained at pH 7.5 to 8.5, with crystals grown at pH 4.0 and soaked in buffer solution with pH 8.0 to 9.0 (9). The only difference between the pH 4 and pH 7.5 to 8.5 structures is in their Asp**116** side chain orientation. While at acidic pH, Asp**116** does not interact with the RSB, at high pH, its side chain is flipped to the cytoplasmic side, creating a strong hydrogen bond with protonated RSB. Thus, this structure was claimed to be the model of the ground state of the Na⁺-pumping form of KR2. In (10), the crystals were grown at pH 4.9 and 5.6 (10). KR2 forms pentamers in the crystals with a sodium ion bound at the oligomerization interface. Asp**116** is directly hydrogen bonded to RSB; however, it remains in the same orientation as at pH 4.3. In opposite, positions of the Asn**112**–Leu**74** pair in KR2 protomers vary within a pentameric assembly between two extreme conformations. The first one is similar to that in the monomeric form, with one water molecule...
in the Schiff base region. In the second, the Asn<sup>112</sup> side chain is flipped out of the RSB region, creating space for three water molecules between Asp<sup>116</sup> and Asp<sup>251</sup> residues. We named these conformations compact and expanded, respectively (12). Two different groups proposed two putative Na<sup>+</sup>-pumping mechanisms. In (9), it was suggested that after photon absorption, the chromophore isomerizes and the proton is moved from RSB to Asp<sup>116</sup>, followed by Asp<sup>116</sup> side chain flip toward Asn<sup>112</sup> and Ser<sup>70</sup> in the M state, creating space for Na<sup>+</sup> translocation near the deprotonated RSB to the Asp<sup>251</sup> and Arg<sup>109</sup> region in the O state. After that, the retinal reisomerizes and the proton is returned to RSB from Asp<sup>116</sup>. The importance of pentameric organization for Na<sup>+</sup>-pumping was stressed in (12), and the following mechanism was suggested. Photon absorption causes retinal isomerization, followed by H<sup>+</sup> transport to the Asp<sup>116</sup> side chain, neutralizing RSB in the M state. Na<sup>+</sup> passes the deprotonated RSB with the rise of the O state, and the ion uptake occurs, when Asn<sup>112</sup> is flipped out of the Schiff base region, so that the place for Na<sup>+</sup> binding site is created near Asp<sup>251</sup> and Asp<sup>116</sup>. After that, the proton is returned to RSB from Asp<sup>116</sup>, Na<sup>+</sup> is released to the extracellular side, and Asn<sup>112</sup> flips back toward the Asp<sup>116</sup> and Ser<sup>70</sup>.

Recently, it was shown by high-speed atomic force microscopy (HS-AFM) and circular dichroism spectroscopy that KR2 forms pentamers in a biological membrane under both acidic and neutral conditions. It was suggested that the monomeric form of KR2 is the artifact of crystallization, corresponding to neither the H<sup>+</sup> nor the Na<sup>+</sup>-pumping form of the protein (13).

Rational design of new cation pumps based on the KR2 structure has been actively performed, and it was shown that modification of the ion uptake cavity (particularly residues Asn<sup>61</sup> and Gly<sup>263</sup>) leads to a change of protein selectivity (9, 10, 14). Recently, it was shown that a single mutation of the α-bulge of helix G of Dokdonia sp. PRO95 Na<sup>+</sup>-rhodopsin [containing residues 253 to 257 in the case of KR2 (10) and also found in bacteriorhodopsin (15)] affects the selectivity of the protein (16). It was also proposed in (17) that Ser<sup>254</sup> of that α-bulge of KR2 matches the Asn<sup>110</sup> of the Na<sup>+</sup> binding pocket of the heterotrimeric guanine nucleotide–binding protein–coupled receptor δ-opioid receptor [Protein Data Bank (PDB) 4N6H]. However, no structural information on any sodium pump mutants is available at the moment.

Thus, although a lot of functional and structural information of KR2 and other Na<sup>+</sup>- and K<sup>+</sup>-pumping rhodopsins has been obtained, it is not yet clear what the structure of the ion-pumping form of KR2 is and what mechanism lies behind Na<sup>+</sup> translocation and the switch from the Na<sup>+</sup>-pumping form at physiological pH to the H<sup>+</sup>-pumping form at acidic pH. Resolving the existing discrepancy of structural data and deciphering the KR2 structure using crystals grown at pH 8.0 are crucial for revealing the sodium ion pumping mechanism and protein organization in its functional form. The need for a structure and a deep understanding of the mechanisms of cation pumping by microbial rhodopsins is also highly motivated by a recent demonstration that KR2 can be used as an optogenetic tool (9, 18). Thus, the structure will help to engineer new optogenetic tools.

**RESULTS**

**Structure of Na<sup>+</sup>-pumping form of KR2**

We present here the crystal structure of the KR2 ground state at 2.2-Å resolution obtained with the crystals grown at pH 8.0. The protein was crystallized using the in meso approach, similar to our previous work (10). The plate-like crystals of type I were red (fig. S5E) in color (maximum absorption at 528 nm), which corresponds to the absorption of light by the protein Na<sup>+</sup>-pumping form. These crystals contain one KR2 pentamer per asymmetric unit. Opposite to the previous results, all the five protomers in the asymmetric unit have the same conformation, which is similar to that of protomers A and D of the 4XTN model. Since the structure is pentameric, obtained at pH 8.0 and at a sufficient sodium concentration, we conclude that it represents the correct sodium-pumping conformation.

Each KR2 protomer has seven transmembrane helices, named A to G, which are connected by three intracellular loops, three cytoplasmic loops, and an N-terminal α helix, capping the protein. Retinal is in the all-trans configuration and covalently bound to Lys<sup>255</sup> of the helix G via the Schiff base. The structure corresponds to the expanded conformation of KR2 (12). As in the previously determined structure of the expanded conformation of KR2 (10), there are three cavities present in the protomer (Fig. 1A).

One of the cavities is placed at the cytoplasmic side (Fig. 1B). This water-filled cavity protrudes from the protein surface looking from the hydrophilic part of the lipid bilayer to Glu<sup>123</sup>. Mutagenesis shows that this cavity is the ion uptake vestibule (9, 10, 14). There is, however, no notable difference between structures of the cavity and that of the protein in the monomeric form. Glu<sup>123</sup> is located at the level of hydrophobic/hydrophilic border of the protein surface and is separated from the Schiff base by the hydrophobic residues Val<sup>167</sup> and Leu<sup>160</sup> (Fig. 1B).

Second, there is a large cavity between the Schiff base and Arg<sup>109</sup>, which is the primary acceptor of the proton. The Ser<sup>254</sup> side chain is connected to the hydrogen bond network through Tyr<sup>218</sup> and Asp<sup>251</sup> side chains and therefore may play an important role in protein function (Fig. 1C).

Third, the ion-release cavity is placed in the extracellular part of the protein (Fig. 1D). Hydrogen bonds protrude from the Schiff base through Asp<sup>116</sup>, Asp<sup>251</sup>, Arg<sup>109</sup>, Gln<sup>244</sup>, Asn<sup>111</sup>, and water molecules to the ion-release cavity, separated from the bulk with the Arg<sup>243</sup>, Glu<sup>111</sup>, and Glu<sup>160</sup> triad.

**Pentamer oligomerization interface**

Each pair of the protomers interacts through transmembrane helices B, C, D, A’, and B’ with an extended hydrogen bond network (Fig. 2). The connection between protomers at the cytoplasmic side is completely mediated by water molecules (Wat506, Wat515, Wat488, and Wat434) (Fig. 2C). The hydrogen bond network at the extracellular side consists of two regions (Fig. 2, D and E). The view from outside of the pentamer shows tight connection of oxygen of His<sup>30</sup>′ with Asn<sup>112</sup> side chain and Tyr<sup>154</sup>′ oxygen through Wat401 and His<sup>30</sup>′ side chain with Tyr<sup>154</sup>′ from helix D and Gln<sup>26</sup>′ from the same helix A’, which is further bonded to Glu<sup>22</sup>′. Both side chain and backbone of Glu<sup>22</sup>′ interact with Asn<sup>105</sup>′, Asn<sup>′</sup>, and the backbone of the N terminus of the nearby protomer through water molecules Wat411, Wat412, Wat448, Wat459, and Wat460′.
and Wat491. Inside of the pentamer, Asn81′ side chain interacts with Gln80′, Ser29′, Thr83′, Thr33′, and Tyr36′ side chains through several water molecules. Backbone atoms of N terminus and BC′-loop are also hydrogen bonded via Wat514′, stabilizing the structure. In addition, Tyr108 is directly hydrogen bonded to Ser29′. An important part of interprotomeric connection is the sodium ion binding site, comprising Asp102 and Tyr25′ side chains, Phe86′ and Thr83′ main chain oxygens, and Wat420′ and placed at the extracellular side of the protein interface.

**Comparison with known KR2 structures**

The structure of the Na⁺-pumping form differs considerably from those obtained with the crystals grown at low pH. The structure is also very different from that obtained with crystals grown at pH 4.0 and soaked at pH 8.0 to 9.0 (Fig. 3) (9). It means that soaking of the crystals, which were grown at low pH, even at pH 8.0 to 9.0 cannot transfer the protein to Na⁺-pumping form in the case of KR2. A notable difference of monomeric and pentameric forms is the different orientation of the protein relative to membrane plane. In the monomeric form, the protein is tilted about 70° to the membrane plane. In opposite, the protein in the Na⁺-pumping pentameric form is oriented nearly perpendicular to the membrane plane. The considerable difference results in the position of ion uptake cavity relative to hydrophobic/hydrophilic membrane boundary. In Na⁺-pumping pentameric KR2, the cavity is placed in the hydrophilic part. In opposite, in monomeric KR2, the cavity is expected to be buried into the hydrophobic part (figs. S1 and S2).

The present structure allows the reinterpretation of solid state nuclear magnetic resonance (NMR), time-resolved absorption spectroscopy, and other data that recently have become available (19–22). The structure shows that the Asp116 orientation is uniform within the pentameric assembly under physiological conditions, which is in agreement with the observed homogeneity of the retinal-binding pocket at pH 8.0 (19, 20). Table S4 shows that in pentameric KR2 at

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**Fig. 1. Architecture and cavities of KR2 protomers.** (A) Overall view. Helices F and G are not shown. (B) Detailed view of the cytoplasmic part of KR2 protomer. Helix G is shown with 90% transparency. (C) Detailed view of the Schiff base region. Helix B is shown with 90% transparency, and helix A is not shown. (D) Detailed view of the extracellular part of KR2 protomer. The hydrophobic membrane core boundaries were calculated using the PPM server (48) and are shown as solid horizontal lines. The cavities were calculated using HOLLOW (49) and are pink colored.

**Fig. 2. Overall architecture of KR2 pentamer and structure of the oligomerization interface.** (A and B) View from the cytoplasmic and extracellular sides, respectively. Only contacts between chains A (yellow) and E (green) are shown. (C) Cytoplasmic side of the oligomerization interface. (D and E) Extracellular side of the oligomerization interface.
Fig. 3. Comparison of Schiff base region of different KR2 structures. (A) Chain A of pentameric Na\(^+\)-pumping form (expanded conformation, pH 8.0) is shown in yellow. (B) Chain E of 4XTN model (compact conformation, pH 4.9) is shown in salmon. (C) The 4XTL model (one of two closely related to compact conformations pH 4.3) is shown in light blue. (D) The 3X3C model (closely related to compact conformation, soaked at pH 8.0 to 9.0) is shown in green. The red dashed ellipse shows double conformation of the Asp\(^{116}\) side chain. Red contoured arrows show the important displacement of the Asn\(^{112}\)-Leu\(^{74}\) pair (colored teal). Helix A’ of nearby protomer and fragments of lipid molecules are shown in orange for pentameric and monomeric models, respectively. The cavities are colored pink. The cartoon representation of helix A is hidden for clarity. A prosthetic group retinal is colored dark green.

Structural basis of the switch from Na\(^+\)- to H\(^+\)-pumping forms

It was shown previously that KR2 pumps sodium ions when their concentration is much higher than that of protons, that is, at least at pH 6.4 to 8.0 (11). This means that under physiological conditions, KR2 is in the Na\(^+\)-pumping form. However, no information on Na\(^+\) transport at acidic pH is available in literature. We studied the pumping activity of KR2 that was reconstituted into lipid vesicles at pH 8.0, 6.0, and 4.3 in the same manner, as described elsewhere (23, 24). The data indicate that KR2 is able to pump sodium ions at pH 6.0 and 8.0 and pumps protons at pH 4.3 (fig. S9). As the protein acts as the H\(^+\) pump at acidic pH even at physiological concentration of sodium, we define the low pH KR2 form as the H\(^+\)-pumping form. In addition, optical properties of the protein are altered with the decrease of pH (8), and structural rearrangements in the retinal-binding pocket were recently reported (19). Structural insights into the nature of transitions in KR2 with the pH change may help to better understand the mechanism of Na\(^+\) pumping. To discover the dependence of KR2 conformations on the pH, we have determined crystal structures of KR2 using crystals initially grown at pH 8.0 and soaked for 48 hours in buffer solutions with pH 4.3, 5.0, 6.0, and 8.0 and in the buffer solution described in (9) (table S1).

After soaking at pH 6.0, crystals remain red and diffract up to 2.7 Å, revealing the same structure as at pH 8.0 (fig. S7F). This result is in accordance with solid-state NMR structural studies of the retinal-binding pocket, where no substantial changes were observed between pH 6.0 and 8.0 (19).

After soaking at pH 4.3, the structure of the KR2 pentameric form could not be determined. The crystals changed their color from red to purple (absorption maximum at 550 nm) (fig. S7H). However, they lost their quality during the soaking procedure. To decrease the stress on crystals, we soaked them at pH 4.3 with a very gradual exchange of the buffer during 48 hours. However, even such gentle manipulations led to a loss of diffraction quality; the resolution dropped to 7 Å, and the mosaicity was extremely high. Thus, we decided to soak crystals at a higher pH.

The structure of KR2 that was obtained with the crystals soaked at pH 5.0 was solved to 2.6 Å. Crystals also changed their color from red to purple after soaking (fig. S7G). The location of most of the residues remains the same as in the model corresponding to pH 8.0. However, the region comprising residues 109 to 115 of helix C is slightly displaced. The distance between the Asp\(^{116}\) oxygen and the Schiff base nitrogen is increased to 3.2 Å, and at pH 8.0, it is only 2.8 Å. The weakening of Asp\(^{116}\) interactions with RSB around pH 5.0 was also observed with the solid–state NMR study of KR2 retinal-binding pocket and was assigned to the protonation of Asp\(^{116}\) (19). The change in protonation of Asp\(^{116}\) can also be indicated by the shift of absorption maximum of KR2 in crystals from 528 to 550 nm. At the same time, Leu\(^{74}\) and Asn\(^{112}\) side chains occupy two alternative conformations: The first is the same as that in the expanded conformation of KR2, and the second is similar to that of the compact conformation of KR2 (Fig. 4B and fig. S10, D and E). Consequently, the Schiff base cavity is reduced in volume in the KR2 fraction at pH 5.0 (Fig. 4B).

In summary, according to our functional and structural analysis at different pH, KR2 acts as a H\(^+\) pump at acidic pH and under the same conditions, a compact conformation of KR2 appears. Thus, we suggest that the compact conformation corresponds to the H\(^+\)-pumping form. Therefore, we conclude that the switch from the expanded to the compact conformation results in the corresponding switch from the Na\(^+\)- to H\(^+\)-pumping forms in KR2.
Effects of dehydration on KR2 crystals

A previously published structure of the pentameric form of KR2 at pH 4.9 (PDB: 4XTN) contains protomers in both compact and expanded conformations, when the structure at pH 5.6 (PDB: 4XTO) contains only protomers in the compact conformation. We assume that this inconsistency is mainly due to the crystals drying, as the crystallization drop was not covered with additional buffer solution while fishing the crystals in our previous work. To check this hypothesis and resolve the controversy, we fished wizened crystals initially grown at pH 8.0, which also turned from red color to purple with time, without addition of the buffer solution. After that, we added buffer solution to the same drop and let it equilibrate for 30 min. Crystal color turned back to the red within 1 min. Last, we fished the soaked crystals and solved the structure of both pentameric “dry” and “wet” forms at 3.0 and 2.6 Å, respectively. The structure leaves no doubt that in the dry form, all the protomers inside the KR2 pentamer are present in the compact conformation. We assume that this is coexistence in KR2 at pH 5.0. Arrows show the important displacement of the Asn112-Leu74 pair. Helix A* of nearby protomer is shown in orange. Helices A and B are not shown. The cavities are colored pink. Hydrogen bonds, stabilizing the compact conformation, are shown as black dashed lines. (C) 2Fo-Fc electron density maps of the double conformation of the Asn112-Leu74 pair in the KR2 model at pH 5.0. The maps are contoured at the level of 1.0σ.

Crystal structure of K⁺-pumping mutants

To better understand the mechanisms and to additionally verify the proposed key determinants of Na⁺ pumping, to and check whether they are also universal for pumping of other cations, we solved the structures of KR2 K⁺-pumping mutants.

Extensive mutational analysis of KR2 indicates that the S254A mutant has the capacity to pump not only protons and sodium but also potassium ions (fig. S4). Spectroscopic analysis also indicates the shift of maximum absorption wavelength from 525 to 545 nm in the case of this mutant. G263F was already demonstrated to pump K⁺ ions in our previous work (10). Here, we present structures of both monomeric and pentameric forms of the ground state of G263F and S254A mutants.

Monomeric blue-form structures of G263F and S254A were both determined at 2.0 and 2.1 Å, respectively, with the crystals grown at pH 4.3 and are nearly the same as that of monomeric wild-type KR2. The situation is the same with the pentameric form of the ground state G263F and S254A. Structures were determined using crystals grown at pH 8.0, and the resolution in this case was 2.4 Å for both G263F and S254A mutants. Structures are very similar to that of the Na⁺-pumping form of KR2. The only differences are in the mutation regions (figs. S5 and S6). In the case of the G263F mutant, ion uptake cavity is dramatically decreased and separated from the solvent by the bulky side chain of Phe263 in both the monomeric and pentameric forms.
Role of the pentamerization on function of KR2

As structure of KR2 depends dramatically on the oligomeric state, we decided to mutate the oligomerization interface to destroy the pentameric assembly and simultaneously check the functional activity of the protein variants. Recently, it was shown that the alterations at the oligomerization interface around His 30 affect the oligomeric assembly and charge protein selectivity and Schiff base–counterion interaction (8, 20, 28). Consequently, we generated H30L, H30K, and Y154F mutants of KR2, which were aimed at breaking the His 30–Tyr 154 hydrogen bond, located in the middle of the oligomerization interface and therefore stabilizing the pentamer.

Size exclusion chromatography (SEC) profiles of detergent-purified protein in Fig. 6A demonstrate dependence of the wild-type KR2 oligomerization on pH. It qualitatively corresponds to that in the crystals. At pH 8.0, SEC profiles of the mutants in comparison with the wild type indicate that for H30K and Y154F mutants, pentameric assembly is negligible when H30L was notably unstable and had a large peak corresponding to the protein aggregates (Fig. 6B). Absence of pentameric organization was also recently demonstrated for Y154A (in detergent micelles) but not for the H30A mutant (20). Pumping activity studies in Escherichia coli cell suspension were done as described elsewhere (23) and showed that Na⁺ translocation is notably decreased in the case of H30K and Y154F, when it is completely absent for H30L (Fig. 6C). Y154F and H30K were successfully crystallized at pH 8.0, and the structures were determined at 1.8 and 2.2 Å, respectively. The proteins form monomers in the asymmetric unit, and the space group is the same as that of the monomeric form of wild-type KR2. Both structures do not differ from the structure of the monomeric form of KR2, except for the mutation region (Fig. 6, D to F).

These results evidence strong correlation between the oligomeric state and ion-pumping activity of KR2. The obstruction of the pentameric assembly, observed both in detergent and in crystals, leads to the decrease or complete absence of Na⁺ transport activity. Thus, we suggest that pentamerization is a key determinant of Na⁺-pumping activity of KR2.

DISCUSSION

Here, we presented a high-resolution structure of light-driven Na⁺-pump KR2 in its functional pentameric form. The structure of the KR2 Na⁺-pumping form suggests a putative ion translocation pathway. The first pore in the cytoplasmic part extending from the polar bulk to Gln 123 serves as a vestibule for sodium translocation. Photon absorption leads to retinal isomerization from the all-trans to 13-cis configuration. In M intermediate, the proton from RSB is moved to Asp 116, which is deprotonated and hydrogen bonded to RSB in the ground state. We suggest that the gate around Gln 123 opens with the M-to-O transition and Na⁺ is translocated to the Schiff base cavity where it is bound transiently in the O state. Then, with the decay of the O intermediate, the Arg 109 side chain is flipped to the extracellular side and opens the way for sodium translocation to the ion-release cavity. After that, the retinal reisomerizes and RSB reprotonates from Asp 116. Na⁺ is translocated to the ion-release cavity, and Arg 109 flips back, forming hydrogen bonds with the Asp 251 side chain as in the ground state. Last, the sodium ion is released to the bulk through the Arg 243, Glu 11, and Glu 160 cluster.

The structure of KR2 pentamer at pH 8.0 also shows that the five protomers in the pentamer are in the expanded conformation. The expanded conformations similar to that are found in the structures of all ion-pumping rhodopsins with the ion bound in the active center (12). We suggest that the expanded conformation is a key determinant of Na⁺ pumping and that, correspondingly, the large Schiff base cavity serves as a putative Na⁺ binding site in KR2 (29). The same enlarged cavity was found in the pentameric K⁺-pumping variants of KR2, stressing the universality of this determinant.

On the basis of our data, we propose a putative mechanism of the switch from Na⁺ pumping (at physiological pH) to H⁺ pumping (at acidic pH): The decrease in pH leads to a switch from the expanded to the compact conformations in pentameric KR2. This switch is possibly triggered by the protonation of Asp 116 at low pH, which leads to the rearrangements of residues 109 to 116 of helix C, including Asn 112. It results in the appearance of the compact conformation, in which the Asp 116 side chain is hydrogen bonded with the Asn 112 and Ser 109 but not with the RSB, as it was also shown by the NMR studies of the retinal-binding pocket of KR2 (19). We believe that the decrease of the Schiff base cavity in compact conformation obstructs Na⁺ passage through the protein but does not interrupt H⁺ transport as the proton is smaller than the sodium ion. Consequently, KR2 acts as a H⁺ pump at acidic pH. We estimate the “switch pH” to be around 5.0, which is in agreement with the conducted studies of KR2 (19, 22).

As our data indicate low functional activity of the protein at acidic pH, we can speculate that the compact conformation does
not correspond to the Na\textsuperscript{+}- and/or H\textsuperscript{+}-pumping form and may then be assigned as a low-pH inactive form of KR2. In this approximation, reduced pumping activity is provided by the existing fraction of the expanded conformation. The compact conformation may correspond to the structure of KR2 with protonated Asp\textsuperscript{116}. It was shown that D116N mutant depicts the wild-type KR2 with the protonated Asp and does not pump either H\textsuperscript{+} or Na\textsuperscript{+} (8). Additional accurate functional and structural studies are needed to better understand the KR2 behavior at low pH.

The similarity of KR2 monomeric form structures at all pH levels has three possible explanations: first, the tight packing of proteins in crystals, which blocks any conformational rearrangements of the protein; second, the absence of any structural changes in the protein ground state at different pH levels; and third, monomeric form is present only in the crystals grown under acidic conditions and does not at all represent the H\textsuperscript{+} or Na\textsuperscript{+} transporting conformation. The latter assumption was recently corroborated by HS-AFM studies of KR2 reconstituted into lipids, which showed that the pentamers, not the monomers, are formed in a lipid bilayer at both pH 8.0 and 4.3 (13). Our additional studies of KR2 mutants with an eliminated pentamer formation showed that, unlike in bacteriorhodopsin (30) and other rhodopsins, the oligomeric organization is vital for protein function. Therefore, we suggest that the pentamic architecture of KR2 is essential for the formation and stabilization of the expanded conformation and the positioning of the entrance of the cytoplasmic pore to the hydrophilic part of the membrane and therefore is functionally important for ion translocation by the protein.

An interesting question that remains is the oligomeric state of the sodium pumps in native membranes. Closely related bacterial proton pumps, proteorhodopsins, were shown to be pentameric or hexameric in crystals (31) and in native-like membranes (32), whereas the best-known archaeal proton pump, bacteriorhodopsin, is trimeric (33). The current understanding of the role of oligomeric organization of microbial rhodopsins is limited; many pumps are also functional as monomers (34).

For KR2, unfortunately, neither the amount of proteins per single K. eikastus cell nor their dissociation constant is known. Thus, the oligomeric state of KR2 in native membranes cannot be predicted, although we expect it to be pentameric because close correspondence between oligomer formation in crystallographic structures and in vivo was shown for other pump rhodopsins (13). Yet another interesting question is whether the KR2 protomers get excited simultaneously or only a fraction of protomers is excited under natural light conditions. For bacteriorhodopsin, it was shown that, depending on light intensity, one, two, or all three protomers may be excited at the same time within the trimer (35). However, there are no data for KR2 at the moment.

After solving the structure of KR2 in its functional form, we believe that this pump can be useful as an optogenetic tool. The construction of structure-based new variants is now possible. Here, the K\textsuperscript{+}-transporting variant might be of special interest because it could mimic, in a certain sense, the function of an outwardly directed K\textsuperscript{+} channel for the silencing of neurons by hyperpolarization of the cell in a minimally invasive manner.

**MATERIALS AND METHODS**

**Expression plasmid**

*K. eikastus* sodium-pumping rhodopsin gene (UniProt ID N0DKS8)–coding DNA was synthesized de novo. The nucleotide sequence was optimized for *E. coli* expression using the GeneOptimizer software (Life Technologies, USA). The gene was introduced into the pSCodon1.2 expression vector (Staby Codon T7, Eurogentec, Belgium) via Nde I and Xho I restriction sites. Consequently, the expressed construct harbored an additional C-terminal tag with a sequence LEHHHHHHH.

**Protein expression and purification**

*E. coli* cells of strain SE1 (Staby Codon T7, Eurogentec, Belgium) were transformed with the KR2 expression plasmid. Transformed cells were grown at 37°C in shaking baffled flasks in an autoinducing medium ZYP-5052 (36) containing ampicillin (100 mg/liter). When glucose level in the growing bacterial culture dropped below 10 mg/liter, 10 μM all-trans retinal (Sigma-Aldrich, Germany) was added, the incubation temperature was reduced to 20°C, and incubation continued overnight. Collected cells were disrupted in an M-110P Lab Homogenizer (Microfluidics, USA) at 172.3 MPa in a buffer containing 20 mM tris-HCl (pH 8.0), 5% glycerol, 0.5% Triton X-100.
(Sigma-Aldrich, USA), and deoxyribonuclease I (50 mg/liter) (Sigma-Aldrich, USA). The membrane fraction of cell lysate was isolated by ultracentrifugation at 90,000g for 1 hour at 4°C. The pellet was resuspended in a buffer containing 50 mM Na₂HPO₄/Na₂HPO₄ (pH 8.0), 0.1 M NaCl, and 1% DDM (Anatrace, Affymetrix, USA) and stirred overnight for solubilization. Insoluble fraction was removed by ultracentrifugation at 90,000g for 1 hour at 4°C. The supernatant was loaded on a Ni–nitrilotriacetic acid column (Qiagen, Germany), and KR2 was eluted in a buffer containing 50 mM Na₂HPO₄/Na₂HPO₄ (pH 7.5), 0.1 M NaCl, 0.5 M imidazole, and 0.1% DDM. The eluate was subjected to SEC on a 125-ml Superdex 200 prep grade column (GE Healthcare Life Sciences, USA) in a buffer containing 50 mM Na₂HPO₄/Na₂HPO₄ (pH 7.5), 0.1 M NaCl, and 0.05% DDM. Protein-containing fractions with the minimal A₂₈₀/A₃₈₂ absorbance ratio were pooled and concentrated to 60 mg/ml for crystallization.

**Liposome preparation**

Phospholipids (asolecin from soybean, Sigma-Aldrich) were dissolved in CHCl₃ (chloroform ultrapure, PanReac AppliChem) and liposome preparation malonate (pH 4.3) (Hampton Research, USA). The crystals were transferred from crystallization drop to the bath filled with 3.4 M sodium malonate at pH 4.3, 5.0, and 6.0. The crystals that were initially grown at pH 8.0 were transferred from the bath to avoid drying. The total time of soaking was not less than 48 hours.

**Crystallization details and crystal preparation**

The crystals were grown using the in meso approach (38, 39), similar to our previous work (23, 40–42). The solubilized protein in the crystallization buffer was added to the monoooleyl-formed lipidic phase (Nu-Chek Prep, USA). The best crystals were obtained using the protein concentration of 25 mg/ml. The blue and red crystals were grown using the precipitate 1.0 M sodium malonate (pH 8.0), respectively (Hampton Research, USA). Crystallization probes were set up using the NT8 robotic system at the European Synchrotron Radiation Facility (ESRF), Grenoble, France (37). Absorption maxima were determined by fitting the absorption spectra with skewed Gaussian functions.

**Spectroscopic characterization**

Absorption spectra of KR2 in solution were collected using the UV-2401PC Spectrophotometer (Shimadzu, Japan). Absorption spectra of KR2 in crystals were collected at the ID-29S Cryobench laboratory at the European Synchrotron Radiation Facility (ESRF), Grenoble, France (37). Absorption spectra with skewed Gaussian functions.

**Structure determination and refinement**

Initial phases were successfully obtained in the C222₁ space group by MR using MOLREP (44) using the chain A from 4XTN structure as a search model. Initial phases for pentameric KR2 at different pH were successfully obtained in the C222₁ space group by MR using the crystal structure of KR2 at pH 8.0. Initial phases for monomeric KR2 at different pH were successfully obtained in the I222 space group by MR using the 4XTL structure as a search model. All models, which were used for MR, lacked side chains of residues Leu², Arg¹⁰⁹, Asn¹¹², Asp¹¹⁶, and Asp²⁵⁷ to avoid appearance of bias electron densities. The initial MR models were iteratively refined using REFMAC5 (45), PHENIX (46), and Coot (47).

**Measurements of the pump activity in E. coli**

*E. coli* cells of strain C41(DE3) (Lucigen, USA) were transformed with the KR2 expression plasmid. Transformed cells were grown at 37°C in shaking baffled flasks in an autoinducing medium ZYP-5052 (36) containing ampicillin (100 mg/liter) and induced at the optical density OD₆₀₀ of 0.6 to 0.7 with 1 mM isopropyl β-D-thiogalactopyranoside and 10 μM all-trans retinal. Three hours after the induction, the cells were collected by centrifugation at 3000g for 10 min and washed three times with unbuffered salt solution (100 mM NaCl or KCl and 10 mM MgCl₂) with 30-min intervals between the washes to allow exchange of the ions inside the cells with the bulk. After that, the cells were resuspended in 100 mM NaCl or KCl solution, respectively, and adjusted to an OD₆₀₀ of 8.5. The measurements were performed on 3 ml of stirred cell suspension kept at 1°C. The cells were illuminated for 5 min using the halogen lamp Intralux 5000-1 (Volpi, Switzerland), and the light-induced pH changes were monitored with a pH meter (Lab 850, SCHOTT Instruments, Germany). Measurements were repeated under the same conditions after addition of 30 μM CCCP.

**Measurements of the pump activity in liposomes**

The measurements were performed on 2 ml of stirred proteoliposome suspension at 0°C. Proteoliposomes were illuminated for 18 min using a halogen lamp (Intralux 5000-1, Volpi) and then were kept in the dark for another 18 min. Changes in pH were monitored using a pH meter (Lab 850, SCHOTT Instruments). Measurements were repeated for different starting pH and in the presence of 30 μM CCCP under the same conditions.
SUPPLEMENTARY MATERIALS

Supplementary material for this article is available at http://advances.sciencemag.org/cgi/content/full/5/4/eaav2671/DC1

Fig. S1. KR2 ion uptake pore.
Fig. S2. Comparison of orientations and cavities inside different KR2 structures.
Fig. S3. Comparison of extracellular regions of different KR2 structures.
Fig. S4. Activity tests of KR2 potassium-pumping mutants.
Fig. S5. Structural alignment of potassium-pumping KR2 mutants with the model of wild-type protein.
Fig. S6. Structures of potassium-pumping KR2 mutants.
Fig. S7. KR2 crystal soaking.
Fig. S8. KR2 dry and wet forms.
Fig. S9. Activity tests of KR2 at different pH.
Fig. S10. Electron density maps of KR2 structures.
Table S1. Summary information of crystal structures obtained.
Table S2. Data collection and refinement statistics of the wild-type KR2.
Table S3. Data collection and refinement statistics of KR2 mutants.
Table S4. Distance between Asp119 oxygen and RS8 nitrogen atoms in KR2 models.

REFERENCES AND NOTES


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