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Photo-Induced Proton Transfers of Microbial Rhodopsins

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1. Introduction

Microbial rhodopsins are photoactive membrane proteins that are widely distributed over the microbial world. They commonly consist of seven transmembrane helices forming an internal pocket for a chromophore retinal, whose photo-induced isomerization triggers the respective photochemical reactions of these proteins. They are generally classified into photosensors or ion-pumps, but the members of both classes have individualities. In the case of photosensors, there exist a variety of signal-transduction modes, including interaction with other membrane proteins, interaction with cytoplasmic proteins, and light-gated ion channel activity. For ion-pumps, there exist outwardly directed H⁺ pumps and inwardly directed Cl⁻ pumps. In spite of these functional diversities, most microbial rhodopsins show photo-induced proton-transfer reactions among amino acid residues and the external medium. These reactions reflect the p*K*_a changes of some residues induced by the protein conformational changes during the respective photochemical reactions. To analyze these reactions, it is indispensable to detect the small pH changes of the external medium due to the proton release/uptake during the photoreaction cycles. Electrochemical cells using indium-tin oxide (ITO) or tin oxide (SnO₂) transparent electrodes are a powerful and convenient tool that enables such measurement in external media under a variety of pH conditions (Robertson & Lukashov, 1995; Wang et al., 1997; Koyama et al., 1998a; Tamogami et al., 2009; Wu et al., 2009). Here, we will describe the rapidly expanding family of the microbial rhodopsins and the application of the ITO method to their photoresponses.

2. Visual rhodopsins and archaeal rhodopsins

Rhodopsin is a protein in the retina of animals that works as a light sensor. Rhodopsin contains retinal (vitamin A aldehyde) as a chromophore and its absorption maximum is ~500 nm. Retinal binds with a specific lysine residue *via* a protonated Schiff base. Absorption of photons induces the isomerization of 11-*cis* to all-*trans* retinal and subsequent conformational changes including the deprotonation of the Schiff base (J.L. Spudich et al., 2000). Rhodopsin had been considered to be confined to animals until the findings from haloarchaea.

In the early 1970s, however, a retinal protein was discovered in the membrane of the highly halophilic archaeon *Halobacterium salinarum* (formally *halobium*). The natural habitats of *H. salinarum* are the Dead Sea, the Great Salt Lake and salt ponds. The newly discovered rhodopsin was named bacteriorhodopsin (BR), and it was shown to act as a light-driven proton pump (Oesterhelt & Stoeckenius, 1971). By using light energy, BR transports a proton from a cytoplasmic to an extracellular space, which produces the proton-electrochemical potential difference across the membranes, which in turn drives the synthesis of ATP *via* H⁺-ATPase. Light irradiation to BR induces the retinal isomerization from all-*trans* to 13-*cis*, while the isomerization of visual rhodopsins is from 11-*cis* to all-*trans*. In addition, the most significant difference from the visual rhodopsins is the existence of a so-called photocycle: light absorption leads to the excitation of the pigment, which decays to the original pigment *via* a variety of photo-intermediates. This linear cyclic photochemistry is called a photocycle. On the other hand, for most visual rhodopsins, the Schiff base linkage with the retinal is disrupted as a consequence of the photochemical reaction. The proton-transfer mechanism of BR has been intensively investigated so far. During the photocycle, the protonated Schiff base affords its proton to the extracellular space and receives another proton from the other side, *i.e.*, the cytoplasmic space. Thus, this cycle involves the alternation between the protonated and deprotonated states of the Schiff base.

Later, three additional retinal proteins were discovered in *H. salinarum*. These were halorhodopsin (HR) (Matsuno-Yagi & Mukohata, 1977; Schobert & Lanyi, 1982; Mukohata et al., 1999; Váró et al., 2000; Essen, 2002), sensory rhodopsin I (SRI) (Bogomolni & J.L. Spudich, 1982; Hazemoto et al., 1983; J.L. Spudich & Bogomolni, 1984) and sensory rhodopsin II (SRII, also called phoborhodopsin) (Takahashi et al., 1985; Tomioka et al., 1986; Wolff et al., 1986; E.N. Spudich et al., 1986; Marwan & Oesterhelt, 1987). BR and HR are light-driven ion pumps: HR is an inwardly directed Cl⁻-pump and BR is an outwardly directed H⁺-pump as described above. On the other hand, SRI and SRII act as receptors of phototaxis. In the cell membranes, these receptors form firm complexes with their cognate transducers. By utilization of these sensing systems, the cell moves toward light of the preferred wavelength ($\lambda > 520$ nm) where BR and HR can work, and escapes from the shorter wavelength light ($\lambda < 520$ nm) which may contain dangerous UV light. These retinal proteins also exhibit their own functions during the respective photocycles. Under the physiological states, HR and two SRs do not exhibit the proton-pumping activities. However, the photocycles of two SRs involve alterations between the protonated and deprotonated states of the Schiff bases, resulting in the proton releases and uptakes at the extracellular sides (Bogomolni et al., 1994; Sasaki & J.L. Spudich, 1999, 2000).

3. Microbial rhodopsins

About 30 years after the discovery of BR, archaeal rhodopsin homologues began to be identified in various microorganisms, including proteobacteria, cyanobacteria, fungi, dinoflagellates, and alga (J.L. Spudich & Jung, 2005). Thus, the microbial species containing the retinal protein genes inhabit a broad range of environments. At present, these rhodopsin homologues are called type 1 rhodopsins or microbial rhodopsins, and they define a large phylogenetic class spreading to all three domains of life, *i.e.*, archaea, bacteria and eukarya.

To distinguish them from rhodopsins in the microbial world, the rhodopsins in animals are called type 2 rhodopsins.

Studies on BR, HR, SRI and SRII have shown that two amino acid residues corresponding to Asp85 and Asp96 in BR are key residues for the functional difference among these archaeal rhodopsins. As mentioned above, the Schiff bases of BR and two SRs become deprotonated during the early halves of the photocycles. Asp85^{BR} and the corresponding aspartates of SRs function as the proton acceptors from the Schiff bases. For HR, on the other hand, Asp85^{BR} is replaced by Thr, and thus HR can bind Cl⁻, a transportable ion, to the vicinity of the Thr and the protonated Schiff base. The other residue, Asp96^{BR}, functions as a proton donor for the reprotonation process of the Schiff base and contributes to acceleration of the turnover rate of the photocycle. This residue is not conserved in the two SRs and consequently their photocycles are much slower than that of BR. HR also undergoes a fast photocycle in the manner of BR but does not conserve this aspartate, because HR undergoes its photocycle without the deprotonation of the Schiff base.

On the basis of these findings, the physiological functions of newly found microbial rhodopsins have been deduced from the conservation states of the residues corresponding to Asp85^{BR} and Asp96^{BR}. In addition to these clues, the functions of some proteins have been confirmed experimentally by using the purified proteins and/or by observation of their photo-induced behaviors. From these analyses, the newly found microbial rhodopsins are now categorized as either ion-pumps (H⁺ or Cl⁻) or photoreceptors. In particular, many newly found pigments are categorized as H⁺ pumps. This fact suggests that the utilization of light energy *via* H⁺ pumps is widely adopted in the microbial world. For many H⁺ pumps, the pumping activities have been experimentally confirmed. Representative examples include proteorhodopsin (PR) from proteobacteria living throughout the world's oceans (Béjà et al., 2000); xanthorhodopsin (XR) from *Salinibacter ruber*, a highly halophilic eubacterium (Balashov et al., 2005); *Leptosphaeria* rhodopsin (LR) from *Leptosphaeria maculans*, a fungal pathogen to a plant (Waschuk et al., 2005); *Gloeobacter* rhodopsin (GR) from *Gloeobacter violaceus*, a cyanobacterium living in fresh water (Miranda et al., 2009); and *Acetabularia* rhodopsin (AR) from a gigantic unicellular marine algae, *Acetabularia acetabulum*, which reaches to 10 cm in height (Tsunoda et al., 2006). Recently, two clones of AR, named ARI and ARII, were isolated from the same organism (Lee et al., 2010). These were somewhat different from the original AR. For ARII, the H⁺-pumping activity was confirmed using *Xenopus* oocytes and the detailed photochemistry was examined with the help of a cell-free expression system (Wada et al., 2011; Kikukawa et al., 2011). On the other hand, photosensing rhodopsins have also been found in bacteria and eukarya. Although these represent only a minority of the newly found microbial rhodopsins, they exhibit a variety of signal-transduction mechanisms. Two SRs in the archaeal membrane relay the photosignals to the cognate transducer proteins embedded in the membrane. From the eubacterium *S. ruber*, SRI itself was found and has been extensively characterized (Kitajima-Ihara et al., 2008). Unlike these SRs, the photosensor called *Anabaena* sensory rhodopsin (ASR) from *Anabaena* sp. PCC7120, a cyanobacterium living in fresh water, is considered to relay the signal to the soluble protein (Jung et al., 2003). *Anabaena* does not have a flagellum and so does not show the phototaxis. Instead, *Anabaena* shows a photoresponse called chromatic adaptation. Thus, it is considered that ASR controls the biosynthesis of chromoproteins forming the light-harvesting complex. In addition to these, a new type of

photosensing rhodopsin called channelrhodopsin was found in *Chlamydomonas reinhardtii*, a green flagellate alga (Sineshchekov et al., 2002; Nagel et al., 2002; Suzuki et al., 2003). This is a light-gated ion channel and induces the phototile behavior of the cell. Thus, the world of microbial rhodopsins is now rapidly expanding.

4. The photo-induced proton transfer associated with the photocycle of microbial rhodopsins

Microbial rhodopsins have linear cyclic photochemical reactions called photocycles. For all microbial rhodopsins examined so far, the retinal Schiff bases are protonated in their dark states under physiological conditions. Except in the case of HR, a Cl⁻ ion pump, the Schiff bases become deprotonated during the photocycles independent of H⁺-pumping or photosensing rhodopsins. As described below, these primary and the subsequent proton-transfer reactions are closely related to the functions of both microbial rhodopsins.

4.1 H⁺-pumping rhodopsins

The illumination of the pigment protein leads to the excited state, which is relaxed thermally to the original pigment *via* various photochemical intermediates (Fig. 1A). The best-studied rhodopsin is BR (Haupt, 1999; Balashov, 2000; Heberle, 2000; Lanyi, 2004, 2006), and the H⁺-pumping mechanism of BR is described below. BR at the ground state and the intermediates K, L, M, N and O have been investigated with various biophysical methods. The photocycle comprises stepwise reactions of the thermal reisomerization of the photoisomerized 13-*cis* retinal to the initial all-*trans*, and the proton is transferred toward the higher pK_a residue accompanied with pK_a changes during the photocycle. Reflecting the differences in protein conformation and protonation states of some residues, the intermediates assume the respective absorption spectra. The photoisomerization from initial all-*trans* to 13-*cis* retinal is completed until the formation of K-intermediate. The subsequent proton transfer observed at around neutral pH occurs as the following sequence (see Fig. 1A and B). First, the deprotonation of the protonated Schiff base occurs in the formation of M-intermediate. The proton from the Schiff base is transferred to its counterion Asp85^{BR} and the subsequent proton release to the extracellular (EC) space occurs from the proton-releasing complex (PRC) consisting of Glu194^{BR}, Glu204^{BR}, Arg82^{BR} and water molecules. Next, the proton of protonated Asp96^{BR} locating at the cytoplasmic (CP) channel transfers to the Schiff base in the M-N transition, which lead to the reprotonation of the Schiff base. Then the deprotonated Asp96^{BR} uptakes a proton from the CP space in the N-O transition, which is accompanied by the reisomerization of retinal to the initial all-*trans* state. Finally, the proton of protonated Asp85^{BR} is transferred to PRC during the decay of O-intermediate, and then the protein returns to the original state. This series of proton-transfer reactions accomplishes the net proton transport from the CP to EC side.

As mentioned above, a light-driven proton pump has been found in many microorganisms belonging to bacteria and eukarya. These rhodopsins also undergo the photocycle including the intermediates similar to those of BR. However, there are several differences in the photocycles and proton transfers between BR and other H⁺-pumping rhodopsins. Examples are as follows. (1) PR from marine bacteria also goes through K, L, M, N and O (or PR') intermediates at around neutral pH (Dioumaev et al., 2002; Friedrich et al., 2002; Váró et al.,

2003). However, the proton movement from the proton donor residue (Glu108^{PR} corresponding to Asp96^{BR}) to the Schiff base and its subsequent proton uptake from the CP space occur simultaneously in the M-N transition (Dioumaev et al., 2002). For BR, these two proton movements occur separately in the M-N and N-O transitions. (2) For ARII from marine algae, the photocycle includes much larger reverse reactions between L, M, N and O than are seen in BR (Kikukawa et al., 2011). Although the reverse reactions are also present in BR, their rates in ARII appear to be much larger than in BR. A rapid reverse reaction might be disadvantageous for the unidirectional ion pumping. (3) For all microbial rhodopsins, the proton conduction channel is bisected by retinal. For BR, these two channels are severely isolated and, in the dark state, the CP half channel is kept under a highly hydrophobic condition. This asymmetric structure had been believed to be important for the ion pumping function. For several proton pumps such as XR and GR, however, this characteristic structure of BR is not conserved (Luecke et al., 2008; Miranda et al., 2009). Therefore, the newly found proton pumps seem to adopt a mechanism that is at least partly different from that in BR. Thus, it would be an interesting subject to clarify the essential mechanisms for the respective proton pumps.

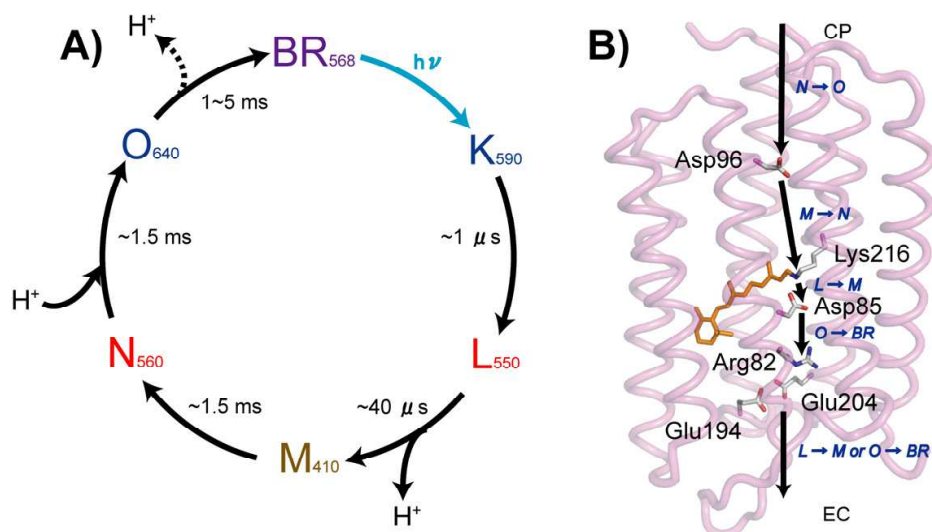


Fig. 1. The photocycle scheme (A) and the structure of BR (B) with the important residues involved in the proton-transfer reactions.

In (A), BR, K, L, M, N and O represent the unphotolyzed state and intermediates, respectively. Their λ_{\max} 's are given in the subscripts, and the lifetimes of the intermediates are also shown. The photoisomerization of the retinal by illumination triggers the stepwise photoreactions accompanied with the proton movements. In (B), these proton transfers at respective steps are indicated with arrows (see section 4.1 for the details). The timing of proton transfer depends on the pH of the medium. Above pH 5, the proton release occurs in L-decay while below at about pH 5, the proton release occurs in O-decay (arrow with a broken line) instead of L-decay. The structure in (B) was drawn from the PDB coordinate file 1C3W.

4.2 Photosensing rhodopsins

During the photocycles of almost all photosensing rhodopsins examined so far, the deprotonations of the Schiff bases occur much as for the H^+ -pumping rhodopsins. In many

cases, these deprotonations result in the proton release/uptake reactions with the external medium. Thus, the proton-transfer reactions also control the decay rates of some intermediates of photosensing rhodopsins.

The best characterized photosensing rhodopsins are SRI and SRII from archaea and bacteria. Some homologues of these SRs show the outwardly directed proton pumping activities when they exist alone in the membrane (Bogomolni et al., 1994; Sasaki & J.L. Spudich, 2000; Sudo et al., 2001; Schmies et al., 2001). Upon the complex formations with the cognate transducers, both the proton release and uptake occur at only the extracellular side (this is so-called proton circulation) instead of the vectorial proton transport. These facts mean that SRs possess proton-transfer machinery like BR and this machinery is probably sensitive to the protein conformational change relating with the signal transduction mechanism. Like the H⁺-pumping rhodopsins, the states having deprotonated Schiff bases are also called M-intermediates, and are the putative signaling states for SRs. The longer lifetimes of M-intermediates are considered to increase the signaling efficiencies. Thus, the reprotonations of the Schiff bases influence the signaling efficiencies.

Studies on two SRIIs from *Natronomonas pharaonis* (NpSRII) (Kamo et al., 2001; J.L. Spudich & Luecke, 2002; Pebay-Peyroula et al., 2002; Klare et al., 2004) and *H. salinarum* (HsSRII) (Sasaki & J.L. Spudich, 1998, 1999) have revealed the differences in the proton-transfer reactions associated with their M-intermediate decays. For NpSRII, the reprotonation of the Schiff base occurs by uptaking a proton directly from the bulk due to the lack of a proton donor to the Schiff base (corresponding to Asp96^{BR}). Therefore the M-decay in NpSRII is very slow as compared with BR and depends on the pH of the medium (Miyazaki et al., 1992). For HsSRII, on the other hand, there are two proton-transfer pathways in the decay of the M-intermediate. One is the pathway in which the proton comes directly from the bulk to the Schiff base, and the other is the pathway in which the proton comes from an unidentified X-H residue. Which proton pathway becomes the major component is dependent on the pH of the medium (Sasaki & J.L. Spudich, 1999; Tamogami et al., 2010). In addition, it has been reported that NpSRII possesses the H⁺-pumping activity (Sudo et al., 2001; Schmies et al., 2001) but HsSRII does not (Sasaki & J.L. Spudich, 1999, 2000). Thus, despite their identical physiological functions, NpSRII and HsSRII have several differences with respect to their photocycles and proton transfers. For these photosensing rhodopsins, therefore, it would be of interest to investigate the photo-induced proton-transfer mechanisms as well as their relations with the photosignaling transductions to the cognate transducers.

5. Importance of measurements of the photo-induced proton transfer of microbial rhodopsins

As described above, the proton movements between the residues inside the protein as well as between the residue and the external space occur during the photocycles of most microbial rhodopsins. The protein conformational changes during the photocycles alter the pK_a of the residues and thereby cause the proton-transfer reactions. For H⁺-pumping rhodopsins, these proton movements directly couple with their functional mechanisms, and for the photosensing rhodopsins, these movements affect the signal transduction efficiencies by controlling the decay rates of some intermediates and reflect the conformational alterations by the complex formation with the cognate transducers. For respective proton-

transfer reaction, a proton moves from an amino acid residue having smaller pK_a to one having higher pK_a . Then, if the pK_a of the residue from which the proton is released to the external medium is larger than the pH in the medium, the proton cannot be released. For such a case, a subsequent proton movement (*e.g.*, a proton uptake from the medium) occurs prior to the release. This "traffic jam" of the proton movement actually occurs in the photocycle of BR (Balashov, 2000). The pK_a of its PRC at the proton-releasing state is about 6 (Zimányi et al., 1992; Balashov, 2000). Under an acidic pH sufficiently lower than pH 6, proton uptake is observed prior to the release (see Figs. 1A and 2B). Therefore, the extent of the pH-dependence of the proton release/uptake could be utilized to estimate the pK_a value of a residue that is important for the proton-transfer reaction. Similarly, the proton-transfer rate may afford information about some important amino acid residues. To analyze these reactions, it is necessary to detect a small pH change in the external medium due to the proton movements caused during a single photocycle.

6. Necessity of a device to measure rapid pH changes or proton-transfer rates at any pH

The photocycle of the ion-pumping rhodopsins completes in ~50-100 ms, and many photosensing rhodopsins have slower photocycle of ~sec. A pH glass electrode is too slow to respond to such pH changes. For such a rapid reaction induced by a flash, the pH change is usually obtained using pH-sensitive dyes whose absorbance depends on pH (Heberle, 2000). This is a convenient method and the rapid change is measurable. However, there is a weak point in that the medium pH should be restricted at pH near the pK_a values of the dyes. In other words, the measurements under various medium pH values cannot be performed with a single dye. In addition, the subtraction of the signal in the co-presence of a dye and a rhodopsin from that of the rhodopsin alone should be performed. If one wants to estimate the pK_a of an important amino acid residue, the pH profile of the magnitude and/or the rate of the proton transfer are indispensable. Hence, a device is needed for the detection of pH changes or proton-transfer rates at any pH. In the following sections, we will describe an electrochemical cell using an indium-tin oxide (ITO) or tin oxide (SnO_2) electrode and its application to the microbial rhodopsins. This electrochemical cell is a useful device for these measurements due to its high sensitivity and rapid time-resolution.

7. An ITO (or SnO_2) electrode works as a pH-sensitive electrode

Koyama and his coworkers first developed this method using BR (Miyasaka & Koyama, 1991; Miyasaka et al., 1992). They constructed a photo-electrochemical cell in which BRs were absorbed on a SnO_2 -coated transparent glass electrode, and detected the electric current evoked by constant illumination by using another SnO_2 electrode as a reference electrode. The origin of this electric current was assumed to be the charge displacement by BR (Koyama et al., 1994). On the other hand, Robertson and Lukashev suggested that the origin of this signal is the medium pH change caused by the photoinduced proton release and uptake in BR (Robertson & Lukashev, 1995). Actually, our group confirmed this suggestion by the following results of three experiments (Tamogami et al., 2009). 1) The equilibrium potential of an ITO electrode showed a linear relationship to pH (see Fig. 2A). 2) The amplitudes of photoelectrical signals decreased with increasing buffer concentration (see Fig. 2B). 3) HR from *N. pharaonis* (NpHR), an inwardly directed Cl^- pump, did not cause

a photoelectrical signal. HR is known to be converted into an H^+ pump in the presence of azide (Váró et al., 1996). In accordance with this, a photoelectrical signal was evoked from the HR-adsorbed electrode by the addition of azide (see Fig. 2C and Koyama et al., 1998b). Thus it was established that an ITO (or SnO_2) electrode works as a pH-sensitive electrode.

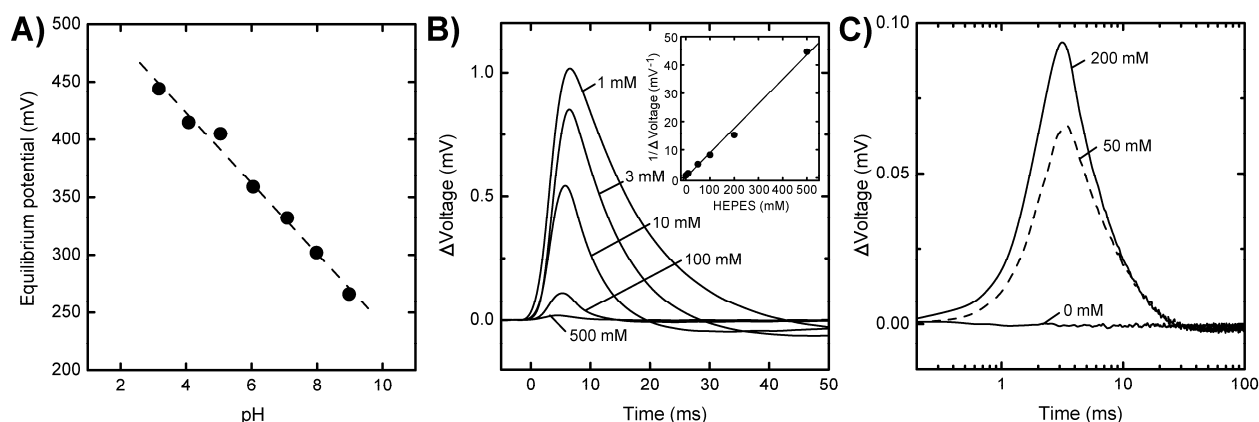


Fig. 2. An ITO (or SnO_2) electrode works as a pH-sensitive electrode.

A) The relationship between the equilibrium potential of ITO and pH. B) The effect of buffer on the photo-induced signals in BR by the ITO electrode. The inset shows the plot of the reciprocal of the amplitudes of ITO signals against buffer concentrations. The experimental medium contains 400 mM NaCl and HEPES of each concentration (1, 3, 10, 100 and 500 mM) at pH 7.5. The excitation light (2 ms duration) was > 520 nm. C) The photo-induced ITO signals in NpHR in the presence and absence of azide. Experiments were performed in medium containing 133 mM Na_2SO_4 , azide at each concentration (0, 50 and 200 mM) and 1 mM HEPES at pH 7.0. The excitation light (2 ms duration) was > 440 nm. NpHR was expressed in the *E. coli* expression system, and then was reconstituted with Egg L- α -phosphatidylcholine. Other experimental setups are described in Fig. 3 and previous report (Tamogami et al., 2009). Panels A and B were adapted with permission from Tamogami et al., 2009, *Photochem. Photobiol.* Copyright 2009 The authors, Journal Compilation, The American Society of Photobiology.

8. Measurements of proton release/uptake by BR with a rapid time resolution

BR has been investigated in great detail. Thus, BR and its mutants are good references to show the relevance of our measurements using this electrochemical cell. Figure 3A shows a schemata of our electrochemical cell. This cell is essentially the same as that previously constructed by Koyama and coworkers with some modifications (Miyasaka et al., 1992). Illumination on the electrode-attached proteins induces the electrochemical potential change between the working and the counter ITO electrodes. Figure 3B indicates the flash-induced signals in BR. In this figure, the upward shift signifies the acidification of the medium near the working electrode. Thus, this shift signified the proton release from proteins to the bulk, while the downward shift signifies the proton uptake from the bulk to the protein. For BR, as described above, the sequence of the proton release and uptake can be altered depending on the external medium pH. For example, at pH 6.0 and 9.0, the proton release is followed by uptake, since the proton can be released from PRC at the early step of the photocycle (in the formation of M). The subsequent uptake occurs in the decay of N. On the other hand, at

pH 3.0, the proton uptake is followed by release. This may be interpreted as meaning that the proton cannot be released from PRC under this pH condition because the medium pH is lower than the pK_a of PRC in the proton-releasing state (M-intermediate). Thus, the proton uptake in the decay of N is observed first and the release occurs in the decay of O concomitantly with the deprotonation of Asp85^{BR} (see the arrow with a broken line in Fig. 1A). The same proton-transfer sequence was also observed in the PRC-lacking mutants, as shown in Fig. 4, where the data for a mutant of E194Q/E204Q^{BR} are summarized. For these PRC-lacking mutants, the proton uptake occurred first even at the neutral pH (see the trace labeled with "ITO signal" in Fig. 4). For wild-type BR at pH 4.5 (see Fig. 3B), the proton uptake occurred first and then overshoot of the signal was observed at around 35 ms. This may be interpreted as the mixture of the two populations of BR. The molecules exhibiting proton uptake first constitute the major population. Other molecules constituting a minor population exhibit proton release first. The final proton transfer corresponding to proton uptake (decay of the positive signal) caused by the minor population is slower than the proton release of the major population. Thus, the signal amplitude of this system can reflect the ratio of two kinds of molecules having different proton-transfer sequence.

In this measurement system, the time course of the photo-induced proton transfer can also be determined. Figure 4 shows the comparison of three types of signals measured for the E194Q/E204Q^{BR} mutant. These are (1) flash-induced absorbance changes of the mutant itself measured at three typical wavelengths; (2) a photo-induced pH change measured by the ITO system and; (3) the corresponding signal measured by a pH-sensitive dye, pyranine. The sign of the pyranine signal is opposite that of ITO. For pyranine, the upward shift signifies the proton uptake, while the downward shift signifies the proton release. As shown in this figure, the proton uptake and release agree well with the formation and decay of O, which are represented by the absorbance change at 660 nm. This is the typical proton-transfer sequence of the PRC-disabled mutant (Brown et al., 1995; Balashov et al., 1997; Dioumaev et al., 1998, Koyama et al., 1998a). The response of the dye is very fast and completely follows the pH change within this time range. The time course of the dye signal almost coincides with that of ITO signal. Thus, our current ITO measurement system can monitor the pH change in the time range of *ca.* 10 ms to several hundred milliseconds (Tamogami et al., 2009). This could be utilized to identify the intermediate accompanying the proton-transfer reaction.

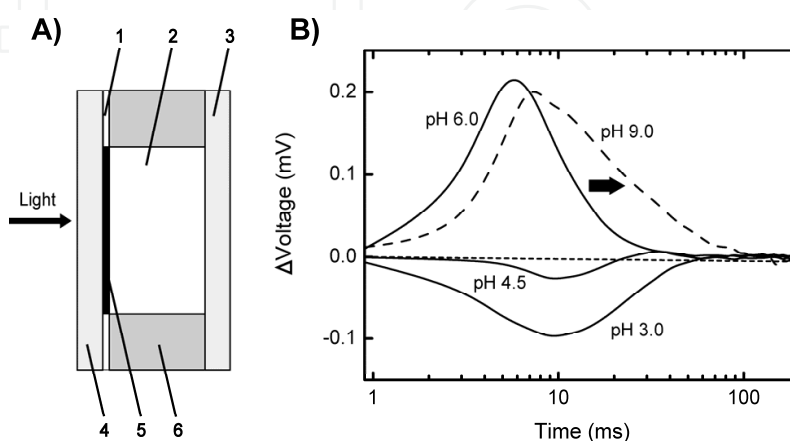


Fig. 3. The measurement of the photo-induced proton transfer in BR by the ITO transparent electrode.

(A) The structure of the photoelectrochemical cell constructed by using ITO electrodes. (1) A silicon sheet; (2) electrolyte plus buffer; (3) a counter ITO electrode; (4) a working ITO electrode; (5) a thin layer of the sample dried on the electrode surface; (6) a Lucite chamber. The emf change between the two ITO electrodes was picked up by an AC amplifier with a low-cut filter of 0.08 Hz which eliminated the baseline fluctuation. This limited the duration of observation time. (B) Flash light (2 ms)-induced signals in BR under varying pH values. Measurements were carried out in a solution containing 400 mM NaCl and 1 mM 6-mixed buffer (citrate/MES/MOPS/HEPES/CHES/CAPS) adjusted to the desired pH with HCl or NaOH. BR indicates the purple membrane prepared by a standard method (Becher & Cassim, 1975). The 6-mixed buffer was used because of its almost constant buffer capacities in a wide pH range. Panel B was adapted with permission from Tamogami et al., 2009, *Photochem. Photobiol.*. Copyright 2009 The authors, Journal Compilation, The American Society of Photobiology.

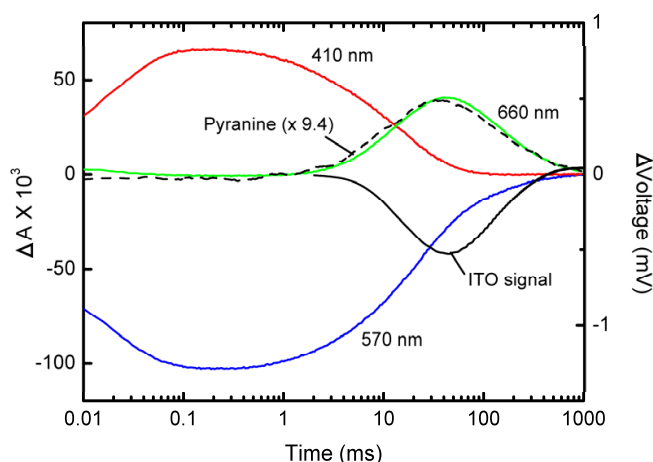


Fig. 4. Comparison between the flash-induced absorbance changes and proton-transfer signals in E194Q/E204Q^{BR}.

The red, blue and green lines represent absorbance changes of BR at 410, 570 and 660 nm, where the M-intermediate, unphotolyzed state and O-intermediate are mainly monitored, respectively. These traces were obtained by the flash photolysis spectroscopy performed by the procedure as described previously (Sato et al., 2003). The black broken and solid lines are the pyranine's signal (monitored at 450 nm) and ITO signal, respectively. Measurements of the absorbance changes of BR and pyranine were performed in the solution containing 400 mM NaCl plus 0.5 mM HEPES at pH 7.1 as described elsewhere (Tamogami et al, 2009). On the other hand, the ITO experiment was performed as shown in Fig. 3 in a solution containing 400 mM NaCl plus 1 mM 6-mixed buffer at pH 7.1. The protein sample was the purple membrane isolated from *H. salinarum* expressing this mutant.

9. Estimation of the pK_a values of important residues involved in the photo-induced proton transfer of BR

It is a pronounced advantage of the ITO electrode method that the measurements can be performed over a wide pH region. Since the buffer capacity is kept constant for the measuring pH range by mixing the buffering agents and the detected pH changes are quite small due to the very faint amount of the adsorbed protein, the measured voltage changes

are proportional to the numbers of protons moved by the photo-induced transfer. By analyzing these voltage changes, the pKa values concerned with the proton transfer can be estimated as described below. Figure 5A shows the peak magnitude of the voltage changes by BR (peak values in Fig. 3B) measured under different medium pH values. As shown in this figure, below pH ~ 5 , the proton uptake was followed by the proton release. On the other hand, above pH ~ 5 , the proton release was followed by the proton uptake. Corresponding to these two pH ranges, the pH profile consists of two bell-shaped functions having opposite signs. This suggests the contribution of the four pKa's of residues involved in the proton-transfer reactions. Then, this pH profile can be expressed by the following equation:

$$\Delta Voltage = -A \left(\frac{1}{1 + 10^{pKa_1 - pH}} \right) \left(\frac{1}{1 + 10^{pH - pKa_2}} \right) + B \left(\frac{1}{1 + 10^{pKa_3 - pH}} \right) \left(\frac{1}{1 + 10^{pH - pKa_4}} \right) \quad (1)$$

where A and B are constants used to adjust the magnitude of the response, and the four pKa values from pKa₁ to pKa₄ are assumed to increase in this order. A fitting analysis using Eq. 1 gave the following pKa values: pKa₁ = 2.6, pKa₂ = 4.1, pKa₃ = 6.1 and pKa₄ = 9.0. For BR, the pKa values involved in the photo-induced proton transfer had been estimated mainly by various spectroscopic measurements (Balashov, 2000). The pKa's determined by the ITO method agreed well with those previously reported. Therefore, the origins of the estimated pKa's were verified on the basis of previous reports. As a result, the pKa's from pKa₁ to pKa₄ were identified as the pKa's of Asp85^{BR} in the dark, PRC in the O-intermediate, PRC in the M-intermediate and PRC in the dark, respectively. In addition to this analysis using the peak voltages, the rates of the voltage changes were also informative. The proton uptake by Asp96^{BR} from the CP space, which coincides with N decay, becomes slow as the medium pH increases (see the traces at pH 6 and pH 9 in Fig. 3B). The rate constants of the proton uptake obeyed the Henderson-Hasselbalch equation with a single pKa, and the value of Asp96^{BR} during N decay was estimated at 7.8 (Tamogami et al., 2009). This value also agreed with the previously reported value (Balashov, 2000). These pKa values determined by the ITO method are considered those of key amino acid residues for the proton pumping function of BR. Therefore this method is useful for detecting the proton transfer directly and deducing the important pKa values.

10. Application to other H⁺-pumping rhodopsins

This ITO method has been successfully applied to the newly found H⁺-pumping rhodopsins (Tamogami et al., 2009; Kikukawa et al, 2011). The panels B and C in Fig. 5 are the results for ARII from marine algae and PR from marine bacteria, respectively. Interestingly, the pH profiles of these three proton pumps are quite different. The prominent differences are as follows: (1) The pH profiles of the three rhodopsins commonly have bell-shaped negative peak areas, indicating that the proton uptake occurs prior to the release. However, the negative peak of PR is located at a quite higher pH (pH ~ 8) than the other two rhodopsins (pH 3 ~ 4). This reflects the difference of the most acidic pKa's, governing the pH where the proton uptake starts to occur. These pKa's correspond to those of aspartates (Asp85^{BR}, Asp81^{ARI} and Asp97^{PR}), the counterions of the respective protonated Schiff bases. This pKa for PR is about 7, which is much higher than the pKa's of about 2.6 of the other two rhodopsins. This high pKa for PR reported by using spectroscopic methods (Dioumaev et

al., 2002; Friedrich et al., 2002; Lakatos et al., 2003; Imasheva et al., 2004; Partha et al, 2005). Thus, we obtained the same result *via* direct measurements of the proton-transfer reactions. (2) As the pH increases, all three rhodopsins begin their proton releases prior to their proton uptakes.

The positive areas of the pH profiles correspond to this proton-transfer sequence. However, the starting pH's, which reflect the pK_a 's of the proton-releasing residues, are different. These pK_a values are about 6.1 for BR, 8 for ARII and 10 for PR, respectively. The higher pK_a 's of ARII and PR might reflect the absence of residues constituting the PRC of BR. ARII lacks a residue corresponding to Glu194^{BR}, one of two glutamates constituting the PRC. By using the mutant of ARII, we confirmed that another glutamate, Glu199^{ARII}, which corresponds to Glu204^{BR}, functions as the proton-releasing residue (unpublished data). On the other hand, PR lacks both glutamates. For PR, therefore, an unknown residue works as the proton-releasing residue. (3) The pH profile of ARII has a surprising feature: the magnitude of the proton release again increases with a further increase in pH above 10. This indicates that a certain residue, other than Glu199^{ARII}, starts to work as the proton-releasing residue at this pH range. These observations suggest that, despite the identical function, these proton pumps possess partially different mechanisms. Thus various interesting phenomena have been discovered by the experiments using the ITO method.

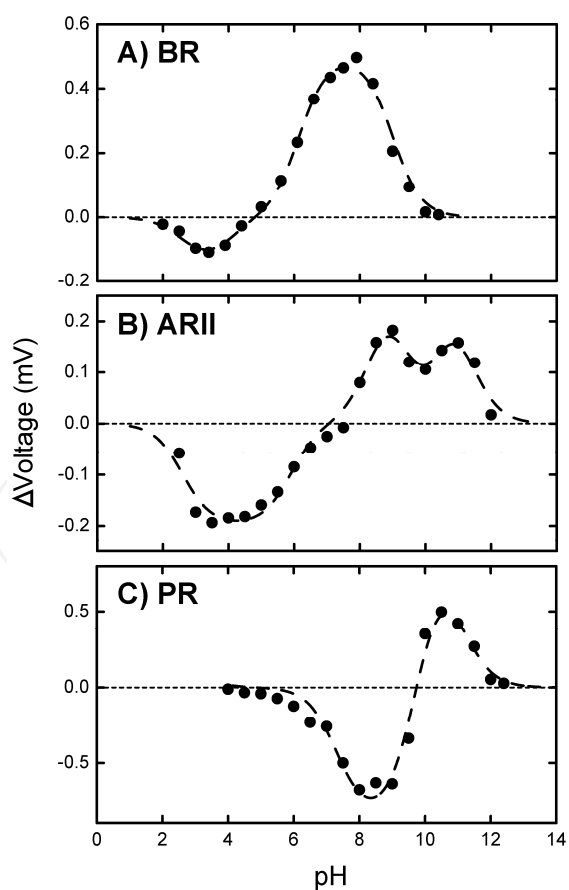


Fig. 5. Comparison of the pH profile of the photo-induced proton transfer among various microbial rhodopsins: (A) BR; (B) ARII; (C) PR.

The peak values of the photo-induced signals were plotted against the medium pH. The experimental conditions were identical to those described in Fig. 3. For BR, the purple membrane prepared by a standard method (Becher & Cassim, 1975) was used. ARII and PR were expressed in the cell-free system and *E. coli* expression system, respectively, and then they were reconstituted with Egg L- α -phosphatidylcholine. Panels A and C was adapted with permission from Tamogami et al., 2009, *Photochem. Photobiol.* Copyright 2009 The authors, Journal Compilation, The American Society of Photobiology. Panel B was adapted with permission from Kikukawa et al., 2011, *Biochemistry.* Copyright 2011 American Chemical Society.

11. Application to photosensing rhodopsins

This ITO method is also applicable to the proton transfers of photosensing rhodopsins. We have adopted this method for the archaeal sensory rhodopsins, NpSRII (Iwamoto et al., 1999), HsSRII (Tamogami et al., 2010) and a putative new class of photosensing rhodopsin called sensory rhodopsin III from *Haloarcula marismortui* (HmSRIII) (Nakao et al., 2011).

Consequently, we successfully determined the timings of the proton uptake/release during their respective photocycles. These results are attributed to the high sensitivity of this method as compared with an alternative method using a pH-sensitive dye. Most photosensing rhodopsins have slow photocycles (\sim sec). This slow turnover rate of the photocycle makes it difficult to adopt the pH-sensitive dye method for this measurement. The absorbance change due to the pH-sensitive dye is very small, and so a slight baseline fluctuation of the absorbance change results in a significant artifact. Especially for a long-term measurement corresponding to the slow photocycle, this baseline fluctuation becomes prominent.

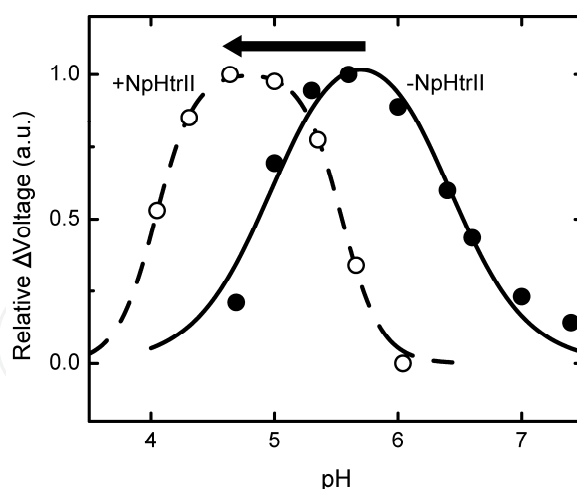


Fig. 6. The pH profiles of the photo-induced proton transfer in NpSRII in the presence and absence of NpHtrII₁₋₁₅₉.

The values of the data points at 10 ms after flash light (the duration of 4 ms) excitation were plotted against pH. The closed and open symbols are the plots in the absence and presence of NpHtrII₁₋₁₅₉, respectively. The NpSRIIs reconstituted with L- α -phosphatidylcholine were employed for the measurements. Added NpHtrIIs were truncated from the 1st to the 159th amino acid. The other experimental conditions were identical to those described in Fig. 3.

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For the three sensory rhodopsins examined so far, we confirmed that the rates of proton uptake reactions determined the decay of their respective M-intermediates, the putative signaling states. In addition to these, we also detected the alterations in the proton-transfer reactions of NpSR_{II} by the complex formation with its cognate transducer, called NpHtr_{II} (Iwamoto et al., 2004). Figure 6 shows the pH profile of the photo-induced proton transfer in NpSR_{II} in the presence and absence of NpHtr_{II}. As shown here, NpSR_{II} releases a proton prior to the uptake at this pH range. From the mutation analyses, the proton-releasing residue was identified as Asp193^{NpSR_{II}} (corresponding to Glu204^{BR}), which is located at the end of the EC channel. Then, the lower and higher pK_a's governing the bell-shaped pH profile were attributed to the pK_a's of Asp193^{NpSR_{II}} at the M-intermediate for release and at the dark state, respectively. As shown in this figure, the pH profile in the presence of NpHtr_{II} shifts to a lower pH compared to that in the absence of NpHtr_{II}, implying that the complex formation induces the conformational change of NpSR_{II} and leads to the pK_a changes of Asp193^{NpSR_{II}}. In the crystal structures, the significant conformational change around Asp193^{NpSR_{II}} was not observed by the complex formation (Gordeliy et al., 2002). Thus, the pK_a's of the proton-transfer reactions could be responsive to a small perturbation of the conformation.

The bell-shaped pH profile of NpSR_{II}, showing the first proton release, is shifted to acidic pH regions as compared with the positive peaks of the proton pumps shown in Fig. 5. This reflects the lower pK_a's of Asp193^{NpSR_{II}}, the proton-releasing residue. For NpSR_{II}, the binding of Cl⁻ to the vicinity of Asp193^{NpSR_{II}} was suggested by the ITO method (Iwamoto et al., 2004), ATR-FTIR measurement (Kitade et al., 2009) and the crystal structure (Royant et al., 2001), while this Cl⁻ binding is not known for the proton pumps examined so far. The physiological meaning of the much lower pK_a of Asp193^{NpSR_{II}} should be examined in a future study.

12. Conclusion and future perspectives

The electrochemical cell using ITO (or SnO₂) electrodes is a powerful and convenient device to detect the proton movements associated with photo-induced reactions of microbial rhodopsins and probably other photoactive pigments. Due to the high sensitivity and rapid response, this system enables us to follow the proton movements during a single photocycle under various buffer conditions. As described above, our current system cannot follow a reaction faster than 10 ms. The first proton movements of microbial rhodopsins appear to occur within 0.1-1 ms. Thus, the system response should be improved. On this point, we have already confirmed that the combination of nsec laser pulse with a homemade amplifier can accelerate the response to about 20 μs.

Early studies on the microbial rhodopsins concerned exclusively four rhodopsins in *H. salinarum*. BR in particular attracted much interest and was investigated in great detail. Thus, BR has been considered a kind of prototype of microbial rhodopsins. However, newly found microbial rhodopsins would seem to challenge the prototype status of BR, since they possess features not seen in BR, as described above. Microbial rhodopsins have been found in a wide variety of microorganisms living in various environments. Thus, it is reasonable to

consider that the newly found rhodopsins acquired their original mechanisms to adapt to their living environments. In this report, we showed the pH-dependent proton movements of BR, ARII, PR and NpSRII (Figs. 5 and 6). Even though only four rhodopsins were considered, their pH dependences were quite different. This might reflect the mechanical divergence of microbial rhodopsins. In the future, a detailed analysis of each rhodopsin will certainly be important. This should be achieved by using various amino acid mutants. Moreover, the proton movements of many more microbial rhodopsins should be examined using this electrochemical cell. From these studies, we could obtain deeper insights into the mechanistic principles of individual rhodopsins.

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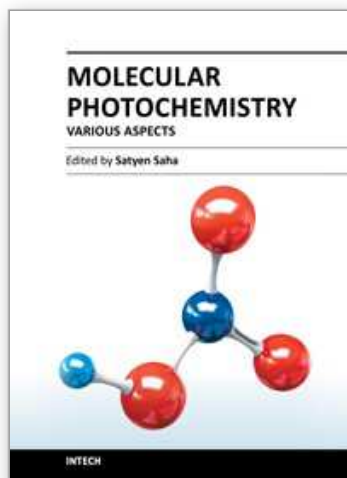
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There have been various comprehensive and stand-alone text books on the introduction to Molecular Photochemistry which provide crystal clear concepts on fundamental issues. This book entitled "Molecular Photochemistry - Various Aspects" presents various advanced topics that inherently utilizes those core concepts/techniques to various advanced fields of photochemistry and are generally not available. The purpose of publication of this book is actually an effort to bring many such important topics clubbed together. The goal of this book is to familiarize both research scholars and post graduate students with recent advancement in various fields related to Photochemistry. The book is broadly divided in five parts: the photochemistry I) in solution, II) of metal oxides, III) in biology, IV) the computational aspects and V) applications. Each part provides unique aspect of photochemistry. These exciting chapters clearly indicate that the future of photochemistry like in any other burgeoning field is more exciting than the past.

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