Interhelical interactions between D92 and C218 in the cytoplasmic domain regulate proton uptake upon N-decay in the proton transport of *Acetabularia rhodopsin II*

Jun Tamogami¹,*, Takashi Kikukawa²,³, Keisuke Ohkawa¹, Noboru Ohsawa⁴,⁵, Toshifumi Nara¹, Makoto Demura²,³, Seiji Miyachi¹,⁷, Tomomi Kimura-Someya⁴,⁵, Mikako Shirouzu⁴,⁵, Shigeyuki Yokoyama⁴,⁶, Kazumi Shimono¹,⁷,# and Naoki Kamo¹,²

¹College of Pharmaceutical Sciences, Matsuyama University, Matsuyama, Ehime 790-8578, Japan
²Faculty of Advanced Life Science, Hokkaido University, Sapporo 060-0810, Japan
³Global Station for Soft Matter, Global Institution for Collaborative Research and Education, Hokkaido University, Sapporo 001-0021, Japan
⁴RIKEN Systems and Structural Biology Center, Yokohama 230-0045, Japan
⁵RIKEN Center for Life Science Technologies, Yokohama 230-0045, Japan
⁶RIKEN Structural Biology Laboratory, Yokohama 230-0045, Japan
⁷Graduate School of Pharmaceutical Sciences, Toho University, Funabashi, Chiba 274-8510, Japan

*Correspondence: jtamoga@g.matsuyama-u.ac.jp

#Present address: Faculty of Pharmaceutical Sciences, Sojo University, Kumamoto 860-0082, Japan

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**Abbreviations**

AR, *Acetabularia* rhodopsin; BR, bacteriorhodopsin; ChR2, channelrhodopsin-2; PR, proteorhodopsin; ATR, all-trans retinal; PSB, protonated Schiff base; SB, deprotonated Schiff base; CP, cytoplasmic; EC, extracellular; H-bond, hydrogen bond; $\lambda_{\text{max}}$, absorbance maximum wavelength; $P_i$, $i$th photochemically-defined state determined by global fitting; DDM, n-dodecyl-β-D-maltoside; PC, L-α-phosphatidylcholine; ITO, indium-tin oxide; MES, MOPS, HEPES, CHES, CAPS, all abbreviations of Good’s buffers
Abstract

*Acetabularia* rhodopsin II (ARII or Ace2), an outward light-driven algal proton pump found in the giant unicellular marine alga *Acetabularia acetabulum*, has a unique property in the cytoplasmic (CP) side of its channel. The X-ray crystal structure of ARII in a dark state suggested the formation of an interhelical hydrogen bond between C218ARII and D92ARII, an internal proton donor to the Schiff base (Wada *et al.*, 2011). In this report, we investigated the photocycles of two mutants at position C218ARII: C218AARII which disrupts the interaction with D92ARII, and C218SARII which potentially forms a stronger hydrogen bond. Both mutants exhibited slower photocycles compared to the wild-type pump. Together with several kinetic changes of the photoproducts in the first half of the photocycle, these replacements led to specific retardation of the N-to-O transition in the second half of the photocycle. In addition, measurements of the flash-induced proton uptake and release using a pH-sensitive indium-tin oxide electrode revealed a concomitant delay in the proton uptake. These observations strongly suggest the importance of a native weak hydrogen bond between C218ARII and D92ARII for proper proton translocation in the CP channel during N-decay. A putative role for the D92ARII-C218ARII interhelical hydrogen bond in the function of ARII is discussed.
1. Introduction

Microbial rhodopsins are a family of photoreceptive membrane proteins broadly distributed in various microorganisms, such as archaea, eubacteria, and lower eukaryotes [1-4]. Notwithstanding their diverse functions, microbial rhodopsins possess common properties, which are often described as structural and photochemical similarities. These proteins are commonly composed of seven transmembrane α-helical bundles (helices A-G) and enclose an all-trans retinal (ATR) as a chromophore. ATR covalently attaches to a conserved lysine residue on helix G of the opsin, which forms a protonated Schiff base (PSB) linkage [5]. Thus, these proteins can be photoactivated by the absorption of visible light. Light irradiation induces the isomerization of the retinal from all-trans to 13-cis, initiating a stepwise cyclic photochemical reaction via several photointermediates called the photocycle. During the photocycle, microbial rhodopsins perform their respective functions, including ion pumping, channeling, and photosensing [3,6].

Among many microbial rhodopsins, bacteriorhodopsin (BR), which acts as a light-driven outward proton pump, is the most well-studied [7-10]. The proton transport of BR is essentially accomplished via three main groups associated with proton transfer events: a protonated/deprotonated Schiff base (PSB/SB), its proton acceptor D85BR, and donor D96BR, which are located at the center of the protein, the extracellular (EC), and the cytoplasmic (CP) transmembrane region of helix C, respectively. In addition, two groups on the EC and CP side modulate the unidirectional proton movement in BR. A proton-releasing complex, which is established by several amino-acid residues and water molecules in the EC channel, causes the ejection of a proton towards the external bulk medium during the first half of the photocycle and subsequently receives a proton
from D85\textsuperscript{BR} during the second half of the photocycle. Another crucial residue is T46\textsuperscript{BR} on helix B, the side chain of which forms a hydrogen bond with protonated D96\textsuperscript{BR} in the CP domain in the unphotolyzed state. This interhelical interaction contributes to the maintenance of the high $pK_a$ of D96\textsuperscript{BR} ($> \sim 11$ [11]) at the initial state together with the hydrophobic residues surrounding it [12]. Through the entry of water via the outward tilt of helix F into the CP region at the site of the photoproduct, D96\textsuperscript{BR} forms a new interaction with nearby water chains concurrently with the disruption of the H-bond with T46\textsuperscript{BR} [13]. This H-bonding rearrangement leads to the lowering of the $pK_a$ in D96\textsuperscript{BR} from greater than 11 to approximately 7 [14,15], with subsequent proton donation of D96\textsuperscript{BR} to SB. T46\textsuperscript{BR}, via this interhelical interaction, thereby regulates the function of D96\textsuperscript{BR} in the CP region and facilitates efficient proton translocation in BR, although this residue itself does not directly participate in the proton transfer events.

*Acetabularia* rhodopsin (AR) is a light-driven proton pump found in a marine alga, *Acetabularia acetabulum*, whose characterization was first performed by Hegemann and colleagues [16]. Two AR homologue genes differing from the initially identified AR were subsequently cloned by Jung and colleagues: *Acetabularia* rhodopsin I and II (ARI and ARII, also abbreviated as Ace1 and Ace2, respectively) [17,18]. Previous electrophysiologial studies revealed that these two AR proteins function as an outward proton pump, like BR [17,18]. However, their detailed characterization has not progressed due to their relatively low expression in a general host, such as *E. coli*, in the absence of certain modifications [19]. Recently, a unique cell-free protein expression method developed by Shimono *et al.* [20] enabled the large-scale synthesis of the two ARs, allowing accelerated investigation of their photochemical and structural properties [18,21,22].
We previously determined the X-ray crystal structures of ARI and ARII [18,21]. In both, the residues corresponding to T46\textsuperscript{BR} were substituted by asparagine (N48\textsuperscript{ARI} and N45\textsuperscript{ARII}, respectively). The oxygen and nitrogen atoms on the side chains of these asparagine residues interact with two carboxyl oxygens of putative proton donor residues (D100\textsuperscript{ARI} or D92\textsuperscript{ARII}) through two H-bonds (see Fig. 1B and ref. [21]). Although the role of N48\textsuperscript{ARI} or N45\textsuperscript{ARII} has not been yet clarified, these interactions might provide stabilization of the protonation state of proton donors in the resting state, instead of the T46-D96 interaction in BR. In addition, a unique feature was found in the CP region of ARII. In the structure of the unphotolyzed ARII, at 3.2-Å resolution [18], C218\textsuperscript{ARII} on helix G was located in the neighborhood of D92\textsuperscript{ARII} on helix C (Fig. 1B), although the distance (approximately 3.9 Å) is somewhat far compared to the typical H-bond cutoff distance of 3.2 Å [23]. The residue corresponding to C218\textsuperscript{ARII} is usually a hydrophobic one in many microbial rhodopsins, thus, this is specific for ARII. If C218\textsuperscript{ARII} directly interacts with D92\textsuperscript{ARII} in the dark state, it is likely that a sulfhydryl group of C218\textsuperscript{ARII} forms a weak H-bond with a carbonyl oxygen (O\textsubscript{δ2}) on the side chain of protonated D92\textsuperscript{ARII}, which serves as a H-bond donor (Fig. 1C). From FTIR spectroscopic studies by Nack et al., a similar Asp-Cys interaction was identified in channelrhodopsin-2 (ChR2), a light-gated cation channel [24]. They proposed that this interhelical interaction on helices C and D acted as a molecular switch (called the DC gate) to control the lifetime of the conductive state in ChR2 [24]. However, the existence of a functional Asp-Cys interhelical interaction in the proton-pumping rhodopsins has not been yet reported. Since it is unclear whether the Asp-Cys interaction implied by the crystal structure of ARII in the dark state is formed even in the photolyzed state and functional, we herein investigated the effects of replacement of
C218ARII on the photoreactivity of ARII. The present study is the first report in which the effects of the Asp-Cys interaction on the photochemistry of a proton-pumping rhodopsin are described, and provides implications for its functional role.

2. Materials and methods

2.1. Sample preparation

Plasmids for the expression of D92NARII and two C218ARII mutants (C218AARII and C218SARII) were prepared by PCR using the QuikChange site-directed mutagenesis kit (Stratagene). In this mutagenesis procedure, a previously constructed expression plasmid for the cell-free protein synthesis of the wild-type ARII [22] was employed as a template. The sequences of the PCR product were confirmed using an automated DNA sequencer (Applied Biosystems). The expression and purification of ARII mutant proteins were performed as described previously [18].

2.2. Flash photolysis

Flash photolysis experiments were conducted using the same apparatus and procedure as previously described [25]. A laser pulse (Nd:YAG 532 nm, 7 ns, 5 mJ/pulse) was employed as a pump light. The temperature was maintained at 20°C with a thermostat (NCB-1200; Eyela). The solution for measurements contained 400 mM NaCl, 0.05% n-dodecyl-β-D-maltoside (DDM), and 6 mixed buffers (citrate, MES, HEPES, MOPS, CHES, CAPS, at 10 mM each). The medium pH was adjusted to be desired values by the addition of diluted HCl or NaOH solutions. The concentration of ARII mutant proteins in the sample solution was ~4-5 µM.

Global fitting analysis, assuming an irreversible sequential scheme including
quasi-equilibria [26], was performed for the obtained data by flash photolysis at pH 7.0. The details of this analysis were described in several previous publications [27-29]. Briefly, through simultaneous fitting using a multiexponential function for the dataset at wavelengths from 340 to 700 nm with a 10-nm interval, the optimal number of exponents in the fitting function was determined from the reductions in the standard deviations of the weighted residuals. From this analysis, the time constant ($t_i$) of the $P_i$-state, which represents the $i$th-appearing photochemically-defined state in the determined scheme, and the absorbance difference ($\Delta\varepsilon_i$) between $P_i$ and $P_0$ (the dark state) were calculated. Regression analysis of the data was performed using Microcal Origin software (Microcal Software) and Igor Pro software (WaveMetrics).

The flash photolysis data at the pH values other than 7.0 were collected at three characteristic wavelengths for ARII, 400, 520, and 610 nm, which primarily monitor the M-state, resting state, and red-shifted K- and O-states, respectively.

2.3. Measurement of absorption spectra at the dark state and determination of photointermediate spectra during the photocycle

The absorbance spectra in the dark state were measured using a spectrophotometer (V-560; Jasco). The obtained spectra contained the effects of the background scattering. Thus, this effect, which is represented as $A+B/\lambda^4$ ($\lambda$ in nanometers), was removed from each spectrum. To calculate the spectra of the $P_i$-states determined by the global fitting analysis, the scattering-removed spectra in the dark ($P_0$) were added to $\Delta\varepsilon_i$.

To estimate the absorbance spectra of the photoproducts, the obtained $P_i$-spectra were further analyzed using the following equation, represented by the sum of the skewed Gaussian functions [26,30]:

\[ \text{Absorbance} = \sum A_i \cdot \text{Gaussian}(x; \mu_i, \sigma_i) \]
Abs(\(\lambda\)) = \sum_{i=1}^{k} A_{i,\text{max}} \cdot \exp \left\{-\frac{\ln 2}{(\ln \rho_i)^2} \left[ \ln \left( \frac{10^7 \left( \frac{1}{\lambda_{i,\text{max}}} - 1 \right) \left( \frac{\rho_i^2}{\Delta \nu_i} - 1 \right)}{\rho_i} \right) + 1 \right]^2 \right\} \quad (k=2 \text{ or } 3),

where Abs (\(\lambda\)) is the absorbance, \(\lambda_{i,\text{max}}\) (in nanometers) is the absorbance maximum of the \(i\)th absorption band, \(A_{i,\text{max}}\) is the amplitude at \(\lambda_{i,\text{max}}\), \(\rho_i\) is the parameter of skewness in the \(i\)th absorption band, and \(\Delta \nu_i\) is the half-bandwidth of the \(i\)th absorption band (in reciprocal centimeters). The fitting of the data was performed using Igor Pro software (WaveMetrics).

2.4. Measurement of the photoinduced proton release and uptake

Experiments were performed using the same apparatus and procedure as described previously [22,31]. Distilled water suspensions of 100 \(\mu\)L volumes containing ARII mutant proteins (conc. \(\sim\)1-3 \(\mu\)M), which were reconstituted into 100-fold levels of L-\(\alpha\)-phosphatidylcholine (PC) from egg yolks (Avanti) at a molar ratio, were attached to the surface of an indium-tin oxide (ITO) transparent electrode (Techno Print Co.) under reduced pressure. A 300 W Xenon lamp with a cold-mirror was used as the light source for excitation of the proteins. When the light was used to irradiate the proteins for 2 ms through an IR filter (HA50; Toshiba) and a cut-off optical filter (Y44; Toshiba), the evoked photoinduced voltage changes were measured through an AC amplifier (MEG-1200; Nihon Kohden) with a low-cutoff electric filter, whose frequency was set to a high-pass 0.08 Hz. Medium containing 400 mM NaCl and 6 mixed buffers (citrate, MES, HEPES, MOPS, CHES, CAPS, at 1 mM each) was used for measurements. All
experiments were conducted at room temperature (~25°C).

3. Results

3.1. The importance of C218ARII as a preferable hydrogen bonding partner of D92ARII for modulation of overall photocycle kinetics

We first evaluated whether D92ARII (corresponding to D96BR) acts as a proton donor for SB. As anticipated, the neutralization of D92ARII (D92NARII) caused a remarkable slowing of the M decay, similar to D96NBR (Fig. S1A).

Next, we introduced two types of mutations into C218ARII to examine its role in the photocycle and accompanying proton transfer. One mutation was a C-to-A replacement to disrupt the interaction between D92ARII and C218ARII, if it exists. In another mutant, C218ARII was replaced by serine, which has a hydroxyl group as a side chain and is a potential amino acid residue that forms a H-bond with D92ARII. Figure 2 shows flash-induced absorption changes at three wavelengths (400, 530, and 610 nm) in the wild-type and the two mutants at neutral pH (~7). From global fitting analysis for the dataset at varying wavelengths, including these three traces, a sufficiently fit model for the wild-type was previously determined as a scheme containing five kinetically defined P-states [22]. However, the same analysis for C218AARII and C218SARII revealed that the best fit functions for their photocycles were composed of the sum of only four exponential terms.

As shown in Fig. 2, the completion of the photocycles in both C218ARII mutants was slower than that of the wild-type. The time constants of the rate-limiting reaction (P4-decay) in the photocycle of the two C218ARII mutants were 51.8 ms for C218AARII and 261.7 ms for C218SARII (Table 1), which are approximately 2.5- and 12.5-fold
slower than in the corresponding process of the wild-type (P₅-decay, τ₅ ~20.9 ms), respectively. Unexpectedly, the turnover rate of C218SARIII was approximately five-fold slower than that of C218AARIII. The difference of the kinetics between two mutants may be considered that the replacement of C218ARIII by serine did not disrupt the H-bond, but rather caused the formation of the distinct interaction from in the wild-type. Considering the higher polarity of the O-H group on the side chain of the serine compared with that of the S-H group of the cysteine, a more robust H-bond may be formed between D92ARIII and the introduced residue (S218ARIII) compared to the native one in the wild-type. Therefore, the stronger Asp-Ser interaction, at least at the dark state, may be disadvantageous for the more efficient proton-pumping photocycle of ARII, suggesting the preferability of cysteine as a weak H-bonding counterpart of D92ARIII rather than serine.

3.2. Effects of replacement of C218ARIII on the rise and decay of the photoproducts revealed by spectral decomposition of the kinetic photocycle intermediates (Pᵢ-states)

A series of characteristic photoproducts for proton-pumping rhodopsins, such as K, L, M, N, and O, which are linked by reversible equilibrium reactions, appear during the photocycle of ARII [22]. Therefore, Pᵢ (i=1-5 (wild-type) or -4 (mutants)) spectra obtained through global fitting analysis contain a quasi-equilibrated mixture of several photointermediates. To estimate the contributions of each photoproduct in Pᵢ spectra at neutral pH, these spectra were separated into putative absorption bands of each species using skewed Gaussian functions (see the Materials and Methods section). The results of the spectral decomposition are shown in Fig. 3. The determined spectral parameters
for respective photointermediates, including their $\lambda_{\text{max}}$ values, are listed in Table S1.

As shown in panel (a) of Fig. 3, $P_1$ spectra in three ARII proteins contained a weaker absorption band at the wavelength region near 400 nm, as well as a red-shifted band. The former band presumably originated from the $\beta$-band [32], although three peaks of its fine structure were not clearly observed due to their small absorbance. The peak position of the latter band was different among the three proteins. The $\lambda_{\text{max}}$ of the red-shifted bands in the wild-type and C218S$^{\text{ARI}}$ was approximately 560 nm, whereas that of C218A$^{\text{ARI}}$ was located at a shorter wavelength (~540 nm). In addition, this band in C218A$^{\text{ARI}}$ was somewhat wide. Therefore, we speculated that $P_1$-states in the wild-type and C218S$^{\text{ARI}}$ represent only a red-shifted early photoproduct $K$, while that in C218A$^{\text{ARI}}$ is a mixture of two states, $K$ and most-likely $L$.

$P_2$ spectra possess two absorption bands (panel (b) in Fig. 3). Of these two bands, one is an absorbance band with a $\lambda_{\text{max}}$ at a shorter wavelength (~394 nm). Generally, this band, whose spectral property was nearly identical among the three proteins, was due to the $M$-state. The peak values of other absorption bands in C218A$^{\text{ARI}}$ and C218S$^{\text{ARI}}$ were blue-shifted compared to that of the wild-type. We tentatively assigned the band of the wild-type to $L$ in our previous publication [22]. However, in the present article, we conclude that this band should be reassigned to $K$ because its $\lambda_{\text{max}}$ (~560 nm) is rather close to the red-shifted band observed in $P_1$. Note that rigorous inspection reveals that the observed values slightly deviate from the calculated $K + M$ spectrum because of the existence of a small plateau absorption band located at approximately 450-550 nm (Fig. 3Ab), which may imply the presence of a small fraction of $L$. Therefore, we inferred that $L$ in the wild-type is difficult to accumulate, probably due to its fast decay or rapid reverse reaction to $K$. 

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In contrast, $P_2$-states in C218A\textsuperscript{ARII} and C218S\textsuperscript{ARII} have no K (Figs. 3Bb and 3Cb). Instead, $P_2$ contains a band with a peak at a shorter wavelength (~510-520 nm) than that of K. In general, this band may be assignable to L. Thus, the replacement of C218\textsuperscript{ARII} may lead to faster K-decay or not prominent L $\rightarrow$ K reverse reactions. The earliest kinetic photoproduct ($P_1$) in the D92N\textsuperscript{ARII} mutant also appeared to contain a large amount of L (Fig. S1B). Thus, the results of these mutational experiments suggest that interhelical H-bonding changes due to the replacement of either D92\textsuperscript{ARII} or C218\textsuperscript{ARII} affect the kinetics of K-L and/or L-M transitions.

In addition to the bands of M, $P_3$ and $P_4$ spectra had another red-shifted bands, and their bandwidth appeared to be somewhat broadened compared with the red-shifted K-bands in $P_1$ and $P_2$ (panels (c) and (d) in Fig. 3A). As reported previously [22], a long M-N-O equilibrium is created during the latter half of the photocycle in ARII. Therefore, these red-shifted bands may represent a mixture of N and O. Forward or reverse reactions may predominate, depending on the pH, in the equilibrium between M and N or N and O. Under acidic conditions (pH $\leq$ ~5.5), N was not accumulated because of the facilitated N $\rightarrow$ O forward reaction. Instead, two O-states (O\textsubscript{1} and O\textsubscript{2}) were produced [33]. The former O (O\textsubscript{1}) may not be observable at neutral pH for kinetic reasons. Thus, the latter O (O\textsubscript{2}) may correspond to the O-intermediate observed at neutral and basic pH values. We estimated a pure spectrum of O (O\textsubscript{2}) from the analysis using skewed Gaussian functions for the $P_4$-spectrum at pH 5.0 (Fig. S2A). The absolute spectrum of N was estimated from the same analysis for $P_4$ at pH 8.6, where N and O coexist but the fraction of N is predominant compared to O due to its slow decay (Fig. S2B). Using the calculated spectral parameters for N and O, except for their amplitudes ($A_{\text{max}}$), the red-shifted bands in $P_3$ and $P_4$-states were analyzed.
In P₃ of the wild-type, the fraction of M was the largest among the three photoproduct states, which may be attributed to the rapid reverse reaction from N to M. Since M and O are in a quasi-equilibrium state via N, the first and second phases in the biphasic decay of the absorbance change at 400 nm (ΔA₄₀₀), which may reflect the processes of the M(N)O-quasi-equilibrated state’s formation from M and its decay, respectively (Fig. 2A). Therefore, the more rapid M → N forward reaction led to the relatively larger amplitude of the first decay phase and the smaller amplitude of the second decay phase in the overall decay phase of ΔA₄₀₀. Conversely, the faster N → M reverse reaction led to the opposite result. Therefore, the larger amplitude of the second slower phase compared to that of the faster one in the decay of ΔA₄₀₀ of Fig. 2A signifies a faster N-to-M reverse reaction compared with the forward reaction, which is consistent with the observation of the large fraction of M in the MNO-quasi-equilibrated P₃-state (Fig. 3Ac).

The fraction of N (λ_max ~526 nm) in P₃ was significantly smaller than that of O (λ_max ~562 nm) (Fig. 3Ac). This may be associated with a relatively faster N-decay at this pH in addition to the facilitated N-to-M reverse reaction described above. The red-shifted band in P₄ can be almost approximated as a spectral band of a single O (Fig. 3Ad), suggesting slight N-accumulation during the photocycle at neutral pH.

In contrast to the wild-type protein, P₃ and P₄ of the two C218ARII mutants included a relatively small fraction of M (panels (c) and (d) in Figs. 3B and C). The fractions of M and N in P₃ of C218AARII were almost comparable (Fig. 3Bc), whereas the fraction of N in P₃ of C218SARII was much larger than that of M (Fig. 3Cc). These results are approximately consistent with the ratios of the amplitude of faster and slower decay components of the ΔA₄₀₀ signals in Figs. 2B and C. Thus, the N → M reverse reaction in
these mutants was not as pronounced, unlike in the wild-type. Moreover, the fraction of N was much larger than that of O. With the transition from P₃ to P₄, the fraction of O in the wild-type slightly increased (Fig. 3Ac and d). In contrast, the fraction of N in the P₄-states of C218ÅRII and C218SÅRII remained predominant (Figs. 3Bd and 3Cd), implying a shift of the equilibrium between N and O towards N, i.e., a slower N-decay in mutants.

P₅ in the wild-type had an absorbance band with a $\lambda_{\text{max}}$ (~532 nm) close to that of the dark state (Fig. 3Ae), assigned to ARII’ previously [22]. In contrast, a P₅-state was not observed in C218ÅÅRII and C218SÅRII. The lack of detectable ARII’ in these mutants may be involved in the prolonged N-decay during their photocycles.

3.3. Differences in the photocycles of the two C218ÅRII mutants from the wild-type at various pH values

To further investigate the impact of the replacement at C218ÅRII on the photocycle, flash photolysis experiments for C218ÅÅRII and C218SÅRII were performed at varying pH values (Fig. S3). The representative results for two C218ÅÅRII mutants at four characteristic pH values are shown in Fig. 4. Different features in the photocycles of the mutants from that of the wild-type (shown as dotted, broken, and chain lines) included: 1) a faster K-decay (M-rise) in both mutants, which was observed particularly clearly at low pH values, 2) a slower O-decay under acidic conditions, 3) no observation of O at substantially lower pH than in the wild-type (7.5 (mutants) vs 8.8 (wild-type [22])) as an increase in pH, and 4) the formation of a M₅-like photoproduct (shown by the bold arrow in Fig. 4Bd) in the photocycle of C218SÅRII at alkaline pH values (8.5 and 9.0), as observed for proteorhodopsin (PR) from marine bacteria [34]. These points are
described in the Supplementary materials (see the captions of Figs. S4 and S5). In addition to the findings described above, the photocycle rates of two mutants are appreciably slower than those of the wild-type in all pH ranges. Although the reason for the slowing of O-decay at acidic pH values (2) is unknown, changes in the interaction of D92^ARII on helix C with C218^ARII on helix G by these mutations may influence the deprotonation of the PSB counterion D81^ARII, on the same helix C as D92^ARII, during the O-decay. Moreover, the appearance of unusual branch reactions with the formation of M_a in C218S^ARII (4) may suggest that the H-bonding partner of D92^ARII should be cysteine. Note that the similar photocycle branch in PR reduced the outward proton-pumping current [34].

3.4. pH dependence of multiple decay components of absorption changes at 400 nm

The decay of $\Delta A_{400}$ changes from mono- to bi- or triphasic with an increase in pH (Figs. 4 or S3). This tendency is similar to that of the wild-type [22]. As described previously [22], the appearance of this pH-dependent multiexponential decay is relevant for the formation of the equilibrium between M, N, and O with increasing pH. Because of the existence of this equilibrium between three photoproducts, it is difficult to separately estimate the rise/decay rates of respective states and examine the pH dependence of the kinetics of each. Thus, we attempted to investigate the pH dependence of these photoproducts from an analysis of the $\Delta A_{400}$ signals because the decay of $\Delta A_{400}$, which mainly reflects M-decay, is also influenced by the kinetics of the other photoproducts N and O. The upper panels (a) in Fig. 5 show close-ups of the $\Delta A_{400}$ for C218A^ARII and C218S^ARII at three characteristic pH values. The decaying phases of these signals were analyzed using the following equation with single, double, or triple
exponential terms:

\[ \Delta A_{400} = \sum_{i=1}^{n} A_i \exp\left(-t / \tau_i\right) \quad (n=1 \text{ or } 2 \text{ or } 3) \]

where \(A_i\) and \(\tau_i\) signify the amplitudes and time constants of the \(i\)th decay components. The decay time constants at respective pH values were estimated from the fitting noted above.

The lower panels (b) in Fig. 5 show the plots of three decay time constants in C218AARII and C218SARII as a function of pH. The inset of Fig. 5Ab represents the corresponding plot of the wild-type. As seen similarly in these plots, \(\tau_1\) and \(\tau_2\) are almost independent of pH, although \(\tau_1\) appears to slightly decrease with acidification of the medium pH below approximately 6. Conversely, \(\tau_3\) apparently shows pH-dependence. From a comparison with the lifetime values of \(P_i\)-states at pH 7.0 (Table 1), \(\tau_1\), \(\tau_2\), and \(\tau_3\) in C218AARII and C218SARII are approximately consistent with the time constant values of \(P_2\), \(P_3\), and \(P_4\)-decay, respectively. As seen in Fig. 3B and C, the \(P_2\)-to-\(P_3\) conversion in C218AARII and C218SARII accompanies the transition from M (+L) to the quasi-equilibrated mixture of M, N, and O. Judging from a little bit of O in the \(P_3\)-states (Fig. 3Be and 3Cc), \(\tau_1\) in C218ARII mutants may primarily reflect the formation of the M, N-quasi-equilibrium state from M. During the M-to-N transition, the internal proton transfer from D92ARII to SB most likely occurs, which may account for the nearly pH-independent behavior of the \(\tau_1\)-plots. Along with the \(P_3\)-to-\(P_4\) transition with the time constant of \(\tau_2\), the fraction of N further increases at the expense of M. In the formed MNO-quasi-equilibrated \(P_4\)-state after this transition, N was most dominant. Therefore, the major determinant in the recovery of the original state from
the P$_4$-state ($\tau_3$) may be N-decay. Thus, the pH-dependence of $\tau_3$ implies that N-decay is accompanied by proton transfer from the external media. In the corresponding graphs of the wild-type, both of the third decaying phases of $\Delta A_{400}$ at near neutral pH (~7) (the inset of Fig. 5Ab) and the large fraction of N in the P$_3$ and P$_4$-states (Fig. 3Ac and d) were not observed because of the relatively faster N-decay at this pH value. However, an increase of the pH above 7.3 gave rise to the appearance of $\tau_3$, with the large accumulation of N in P$_4$ (lower panel (b) of Fig. S2B). It is noted that the pH-dependency of $\tau_3$ in the wild-type can be interpreted in the same way as in the case of C218A$^{\text{ARI}}$ and C218S$^{\text{ARI}}$, although N does not appear obviously in P$_4$ of the wild-type at pH $\leq$ ~7.

Comparison of the $\tau$-plots between the wild-type and C218$^{\text{ARI}}$ mutants revealed an important finding that in contrast to the similarity of the pH dependence of $\tau_1$ and $\tau_2$ among all proteins, there was the difference in the N-decay kinetics, which could be roughly estimated from $\tau_3$ values. The plots of $\tau_3$ in three ARII proteins showed a linear pH-dependence with almost similar slopes (approximately 0.55 for the wild-type, approximately 0.60 for C218A$^{\text{ARI}}$, and approximately 0.64 for C218S$^{\text{ARI}}$). The slopes of these plots were less unity. The reason for this shared slope reduction is unclear at present, but a possible explanation may be the difference in the pH between the protein surface and the bulk due to negative charges of proteins [35]. In contrast, the plotted absolute values of $\tau_3$ of C218$^{\text{ARI}}$ mutants were markedly different from those of the wild-type (shown as the broken line). The approximately 10-30-fold larger $\tau_3$ values of C218$^{\text{ARI}}$ mutants demonstrate a substantial delay of N-decay in these mutants. This is consistent with the conclusion drawn from the results of the spectral analysis of kinetic P$_3$ and P$_4$ states (panels (c) and (d) in Fig. 3).
3.5. Direct observation of delayed proton uptake with replacement of C218ARII

As described above, the observation of the pH dependence of \( \tau_i \) implies that the proton uptake from the media occurs upon N-decay and is delayed by the replacement of C218ARII. To further obtain evidence for this slowing of proton uptake, we measured the proton uptake/release in the C218ARII mutants during their respective photocycles and compared with that of the wild-type. Measurements at near neutral pH values led to the small magnitude of proton transfer signals with the initial proton uptake followed by release. This small signal is probably because of cancellation of opposite two proton transfer sequences which are composed of the first proton uptake followed by release and the reverse sequence [22]. With increasing pH above 7.5, the signal with the first proton release followed by uptake became predominant in the mixture of two signal sequences, resulting in that the sufficient large magnitude of signal was observed because of the lack of the cancellation [22]. Thus, we performed experiments at pH above neutral. Instead of the conventional method using pH-indicator dyes, we adopted an experimental technique using an ITO transparent electrode because this method more sensitively responses to the change of the external pH with the photoinduced proton uptake/release and can be applied to measurements over the wide pH range including alkali conditions. In addition, we previously confirmed that this experimental system can work as a time-resolved pH electrode at the millisecond timescale [31]. Note that experiments were conducted in samples reconstituted into lipids (egg-PC), while flash-photolysis measurements were done in detergent. Thus, when the data of two different experiments are compared, we should pay attention to the effects by different sample preparations, e.g., a difference in their time courses. Figure 6 shows
flash-induced ITO signals in C218AARI and C218SARI mutants at pH 8.5. In this figure, the proton release and uptake during the photocycle are represented by the upward and downward deflections, respectively. We found that the first proton release that probably occurs from E199ARI (corresponding to E204BR) is followed by uptake under this pH condition, in agreement with the order of proton transfer in the wild-type at the same pH value (gray solid lines) (also see ref. [22]). Comparison of time courses of the decreasing phases representing proton uptake in C218AARI and C218SARI with that of the wild-type revealed that the proton uptake in these mutants markedly slows down (also see time constants of the latter H+-uptake in each protein in Fig. S6). Thus, these observations are supportive of the delay of N-decay in parallel with proton uptake with the replacement of C218ARI.

4. Discussion

In this study, we evaluated the influence of two types of substitutions at C218ARI, C-to-A or -S mutations, on the photocycle and accompanying proton transfer. These replacements gave rise to several minor alterations in the first half of the photocycle. In addition to the observation of faster K-decay at acidic pH (Fig. 4), the apparent detection of the fraction of L in P2-spectrum at neutral pH (Fig. 3) indicates that the kinetics change of K-to-L and/or L-to-M transition with the mutations. It was reported that the transition from K to L in BR is accompanied by important structural changes in the CP domain, such as reorientation of the N-H bond of PSB towards the CP side, upward movement of the C13 methyl group of retinal, and rotation of the side chain of L93BR, which are coupled with the vertical movement of a water molecule from the EC to the CP side [36]. Hence, the structural perturbation of helix C with the
mutation-induced interhelical interaction change between D92^ARI II and C218^ARI II might cause the change in the CP region dynamics for the early photoproducts K and/or L, resulting in effects on the kinetics of the formation and/or decay of these states. An example of an indirectly important role for an interhelical interaction in this function was previously reported in BR as well [37]. Perálvarez-Marín et al. reported that D115^BR on helix D forms an interhelical H-bond with T90^BR on helix C, and the disruption of the H-bond by the replacement of D115^BR or T90^BR to Ala renders the deprotonation of D96^BR during the photocycle impossible via the abolishment of the proper dynamics of helix C [37].

The most important effect of C218^ARI II replacement on the photoreaction was found in the second half of the photocycle. A major difference in the photocycles of the mutants was the restrained N-to-O transition in C218^ARI II mutants compared with the wild-type protein (Figs. 3 and 5). The replacement of C218^ARI II delayed the proton uptake upon N-to-O transition similarly for both mutants (Fig. 6) regardless of the polarity of the introduced residues (hydrophobic alanine or hydrophilic serine), suggesting that the retardation of N-decay by the replacement of C218^ARI II may not be merely attributed to a decrease in the proton conductivity of the CP channel. There are three conceivable factors that may induce the slowing of the N-to-O transition. The most feasible reason is the low proton affinity of D92^ARI II upon its reprotonation, hampering the quick uptake of a proton from the CP surface. In fact, in the T46V^BR mutant, which exhibited a slower N-decay similar to the present case, it was considered that the cause of the prolonged lifetime of N was the lowering of the pK_a of D96^BR in the photolyzed state with the abolishment of the interaction with T46^BR [15,38]. To investigate whether the similar decrease in the pK_a of D92^ARI II was caused with the
replacement of C218ARII, we evaluated the pK\textsubscript{a} values of D92ARII during N-decay (O-formation), which was defined as pK\textsubscript{a,N}, in the mutants. We estimated these pK\textsubscript{a,N} values from the pH dependence of the fraction of O. Here we assumed that the fraction of O is scarcely affected by the rate constants of the transitions prior to O-formation. In addition, it was assumed that the presence of a reverse reaction from O to N is negligible at acidic pH values (pH < ~7) because of relatively faster N-to-O transition. Under these assumptions, the fraction of O is theoretically determined by only rate constants of O-rise and -decay, and the fraction of pigment cycling [39]. The fraction of O can be roughly estimated from the maximum amplitudes of the absorbance changes at 610 nm (ΔA\textscript{610}) in the latter time region from ~0.2 ms. Thus, the ΔA\textscript{610}-normalized values in the wild-type ARII and two C218ARII mutants were plotted as a function of pH in Fig. 7. Since the O-decay rate is almost independent of pH at least in the pH region from ~4 to ~7 (see Fig. S7), the pH dependent behavior of these plots may be only explained by the pH dependence of the N-to-O transition kinetics, which is governed by pK\textsubscript{a,N}. As described below, this transition rate is regulated by other two factors (rates of switch of D92ARII and reisomerization of retinal) as well as pK\textsubscript{a,N}. These two factors are considered to be pH-independent and then the pH-dependence of the O-yield is determined only by pK\textsubscript{a,N}. From the plots in Fig. 7, the pK\textsubscript{a,N} values for the wild-type, C218ARII, and C218SARII were estimated to be 6.1, 5.3, and 4.7, respectively, whose magnitudes were in the order of the wild-type > C218ARII > C218SARII. We previously determined the pK\textsubscript{a} value of D92ARII in the wild-type upon H\textsuperscript{+}-uptake (N-decay) to be either 5.9 or 6.3 based on experimental results using the ITO method [22]. The close equality to the previous value indicates validity of the estimation by the present method. The order of the magnitudes of the pK\textsubscript{a,N} values among three proteins correlates well
with that of the turnover rates of their photocycles (see Fig. 2 or Table 1). During N-decay, the proton uptake occurs concurrently with the protonation of deprotonated D92ARII. Thus, the proton affinity of D92ARII at N (pK\textsubscript{a,N}) is one of the effective factors for the N-decay which is the rate determining step of the recovery to the initial state at neutral and above pH.

Although the changes of the D92ARII-C218ARII interaction by the replacements of C218ARII affected the pK\textsubscript{a} of D92ARII during N-decay, it is not clear whether C218ARII regulates the pK\textsubscript{a} of D92ARII at the photolyzed state in the same way as T46BR because the abolishment of the H-bond (C218AARII) caused the prolongation of the lifetime of N similar to T46VBR, but did not seem to induce much influence on the M-to-N transition (see Fig. 5Ab). In contrast, T46VBR led to more rapid M-decay as well as the delay of N-decay, resulting in the lack of observation of slow M-decay components by disappearance of the M-N quasi-equilibrium [15,38]. Therefore, we suppose that the perturbation of the H-bond of D92ARII with C218ARII might affect the D92ARII-N45ARII interaction, which led to the change of the pK\textsubscript{a} of D92ARII indirectly.

The second possible determinant for the N-decay kinetics may be the accessibility change of D92ARII between SB and the CP bulk [40]. In the case of BR, D96BR connects to SB but not the intracellular medium upon the M-to-N transition, blocking the misdirected migration of a proton of D96BR towards the CP bulk. At the next stage (upon the transition between two N-states (N\textsubscript{1} and N\textsubscript{2}) [41,42]), this connection changes into the CP surface to take up a proton from the cytosol. Owing to this switching mechanism, the back-flow of a proton from PSB to D96BR is prevented, although the “switch-off” of the connection between PSB and D96BR is essentially achieved by the inversion of the pK\textsubscript{a} values in these two residues. Thus, the accessibility-switching
machinery of D96BR may assist the efficient reprotonation of D92ARII from the CP bulk at the N-state and facilitate the subsequent transition from N (13-cis) to O (all-trans). A second factor may be the change in the interaction between D92ARII and C218ARII, which may be involved in the accessibility switch of D92ARII at N-state.

The third possible factor is the rate of thermal reisomerization of retinal after reprotonation of D92ARII. This rate could be also affected by the replacement of C218ARII.

Considering three possible determinants for N-decay described above, the transition from M to O in ARII may be summarized as follows:

(1) Proton transfer from protonated D92ARII to SB (M → N1).

(2) Accessibility switch of deprotonated D92ARII facing SB into the CP bulk and formation of connectivity with the external media, which are caused by the influx of water with the conformational change in the CP channel (N1 → N1').

(3) Formation of the below dissociation equilibrium: D92− + H2O ⇌ D92H + OH− (N1' ↔ N2). The presence of water molecules may originate from the (further) opening of helix F. The position of this equilibrium depends on pKa,N and pH around D92', which is assumed to be that of the CP medium.

(4) Protonated D92ARII (D92H) facilitates the isomerization of retinal from 13-cis to all-trans, implying that N2 transforms to O via D92H (N2 → O). This transition occurs with retaining the equilibrium between N1' and N2. Thus, during the transition of N1' to N2, OH− is produced and goes out to the CP medium, resulting in the alkalization in the medium (H+-uptake).
For representing the N → O transition (processes (2)-(4) in the above scheme), here we introduced the following approximation for the kinetics of the N-to-O transition mainly observed as the third decay component of ∆A_{400} (τ₃). Since the overall kinetics of N-decay is probably determined by three factors described above: i) the switching rate of D92^{ARII} (denoted as kₕ), ii) the fraction of protonated D92^{ARII}, and iii) the rate of the cis-to-trans isomerization of retinal (denoted as kᵢ), this transition rate can be described by

\[
k₃ = kₕ \times \left( \frac{[\text{D92H}]}{[\text{D92H}]+[\text{D92}^-]} \right) \times kᵢ
\]

Since kₕ and kᵢ are most probably independent of the external pH, the product of these denotes k_N. Then, this equation is rewrote to

\[
k₃ = kₕ \frac{1}{1+10^{nₐ(pH-pK_{ₐ,N})}}
\]

where pK_{ₐ,N} represents the pKₐ of D92^{ARII} upon the N-to-O transition as defined above. In this equation, kₕ ( = kₕ ⋅ kᵢ) is independent of the protonation state of D92^{ARII} and corresponds to the maximal N-decay rate as is similar to the equation proposed by Balashov et al. [39,40]. nₐ stands for the Hill coefficient, which is relevant to the slope of the τ₃-plot. This equation can be also expressed using time constant τ as follows:

\[
τ₃ = τₕ \left[ 1+10^{nₐ(pH-pK_{ₐ,N})} \right]
\]
Under the condition of $pK_{a,N} \ll pH$, which is satisfied in the $\tau_3$-plots of three ARII proteins (see lower panels (b) in Fig. 5), the above presented function on $\tau_3$ is approximated as

$$\tau_3 \approx \tau_N \times 10^{n_a(pH-pK_{a,N})} = \frac{\tau_N}{10^{n_a pK_{a,N}}} \times 10^{n_a \cdot pH}$$

Since the lower panels of Fig. 5 are plotted as logarithm of $\tau_3$ versus pH, the values of $\log \frac{\tau_N}{10^{n_a pK_{a,N}}}$ should be estimated from the y-intercepts. In addition, the slopes are almost equal among three proteins. Therefore, the shifts of y-axis originate from the difference of the y-intercept values. We calculated the $\tau_N$ values in three proteins using respective y-intercepts, $n_a$ (slope), and $pK_{a,N}$ values determined from Fig. 7 (assuming the pH-independency of $pK_{a,N}$). The differences of the $\tau_N$ values between the wild-type and mutants, $\frac{\tau_{N,C218A}}{\tau_{N,\text{wild-type}}}$ and $\frac{\tau_{N,C218S}}{\tau_{N,\text{wild-type}}}$ were estimated to be ca. 2.0 and 2.8, respectively. According to the above definition ($k_N = k_s \cdot k_i$), $\tau_N (-1/k_N)$ signifies a constant including the time constants of the switch of D92 ARII (denoted as $\tau_s$) and reisomerization of retinal (denoted as $\tau_i$). Therefore, these differences may reflect the delay of either the switch in D92 ARII or reisomerization of retinal (or both) by the replacements of C218 ARII.

As described above, we showed that there are main two factors for the slowness of the photocycle rates (especially the slow decay of N-intermediate), which are caused by the changes of $pK_{a,N}$ and $\tau_N$, respectively. If the $H^+$-uptake upon the $N_1$-to-$N_2$ transition is due only to the $pK_a$ change of D92 ARII at N, the rates of photocycle at acidic pH where
protons are in abundance should be the same between the wild-type and mutants. However, it was not the case (Fig. 4). Hence, this implies that not only the change of $pK_{a,N}$ but also change of $\tau_N$ (pH-independent factor) contributes to the delay of N-decay with the replacement of C218^{ARII} even under lower pH conditions. Thus, the delayed N-decay of mutants compared to that of wild-type resulted in the slower turnover rates of mutants in all pH range.

As described above, the replacement of C218^{ARII} changed $\tau_N$ composed of both $\tau_s$ and $\tau_i$. The change of $\tau_i$ may imply the modulation of the rate of the conformational change leading to the delay of the configuration change of retinal.

Although the detailed molecular mechanism for switching of the proton donor between two N-substates has not yet been clearly defined even in the most well-known BR, the presence of a hydrophobic gate formed in the CP domain may be one of key components. D96^{BR} in the CP-side of the channel is surrounded by hydrophobic residues, such as F42^{BR}, I45^{BR}, L92^{BR}, L95^{BR}, L97^{BR}, L99^{BR}, L100^{BR}, and L223^{BR} [43]. A hydrophobic barrier composed of these residues shields the inner space between SB and D96^{BR} from the external CP aqueous phase. The opening of the intracellular segment with the outward tilt of helix F at the M-state leads to the internal hydration of the CP domain [44]. However, a proton uptake pathway remains functionally closed even in this state, limiting the solvent accessibility of D96^{BR}. Wang et al. performed an in silico study using the X-ray crystal structure of a N-state-mimic D96G/F171C/F219L^{BR} triple mutant [43]. Based on their computational results, they proposed that the deprotonation of D96^{BR} along with the M-to-N transition catalyzes the further opening of the proton uptake pathway in the CP-side of the channel at the following N-state [43]. Thus, it may be predictable that the deprotonation of D96^{BR} acts
as a switch to open the hydrophobic gate and change its own accessibility.

Most of the above described residues in BR are conserved in ARII. However, one of the residues in the intracellular hydrophobic cavity (L223\textsubscript{BR}) is replaced by C218\textsubscript{ARII}. Thus, C218\textsubscript{ARII} may not be irrelevant to the function of the hydrophobic CP gate as an accessibility switch of D92\textsubscript{ARII}. If it is the case, it would be unclear why C218\textsubscript{ARII}, which is missing in BR, is utilized for the switching of D92\textsubscript{ARII}. The precise cause for this is unknown, but one of the conceivable reasons may be the lack of an amino-acid residue with a bulky group on its side chain at the end of the cavity. In BR, an aromatic ring of F42\textsubscript{BR} is located between D96\textsubscript{BR} and the CP surface that functions as a hydrophobic lid to separate D96\textsubscript{BR} from the CP media. Therefore, it has been hypothesized that F42\textsubscript{BR} is important for regulating the accessibility to the CP aqueous space [45,46]. According to the model by Wang \emph{et al.}, it is likely that the phenyl ring of F42\textsubscript{BR} is displaced along with the deprotonation of D96\textsubscript{BR}, facilitating the opening of the gate and the influx of water into the CP domain upon N-decay [43]. However, the corresponding residue in ARII is A41\textsubscript{ARII}, whose side chain’s volume is small (Fig. 1B); consequently, the gating control in ARII may be insufficient, unlike BR. Thus, the H-bond between D92\textsubscript{ARII} and C218\textsubscript{ARII} may serve as an alternative switch for compensating for the absence of the gating function by F42\textsubscript{BR}. From this viewpoint, it may be noteworthy that this unique cysteine residue is not conserved in another homologue, ARI, which shows a slower photocycle [21]. The pH-dependent second slower phase of M-decay in ARI, mainly reflecting the conversion from the MN-quasi-equilibrium to O, was even slower than the N-decay rate of ARII [21]. ARI also lacks a residue corresponding to F42\textsubscript{BR} (replaced by T44\textsubscript{ARI}). Therefore, together with the absence of phenolic lid in the CP region, the lack of the Asp-Cys interhelical
interaction may further decelerate the photocycle of ARI.

The FTIR studies by Heberle and colleagues implied the existence of a functional Asp-Cys (D156\textsuperscript{ChR2}-C128\textsuperscript{ChR2}) interaction in ChR2 [24], although the presence of a direct Asp-Cys H-bond was not observed in the crystal structure of a channelrhodopsin C1C2 chimera [47]. Most recently, Volkov et al. reported the crystal structures of the wild-type ChR2 and a C128T\textsuperscript{ChR2} mutant [48]. According to these structures, the Asp-Cys interaction in the wild-type ChR2 was formed through a water molecule, whereas the Asp-Thr interaction in C128T\textsuperscript{ChR2} was directly formed. Not only disabling mutations of the Asp-Cys interaction (D156A\textsuperscript{ChR2} and C128A\textsuperscript{ChR2}) but also potentially conservative mutations (C128S\textsuperscript{ChR2} and C128T\textsuperscript{ChR2}) commonly caused the prolongation of the lifetime of the photolyzed conductive state [49]. Together with our present study’s findings, this demonstrates that the directly formed stronger interaction makes the gating control cease to function properly, as well as disruptions of the H-bond. Through the FTIR measurements by Heberle et al., the interaction between D156\textsuperscript{ChR2} and C128\textsuperscript{ChR2} was observed at several photoproduct states, although the H-bonding strength was different from that of the dark state [24]. Unfortunately, it has not been yet determined how the H-bond changes at the photolyzed conductive states. Therefore, the detailed mechanism of gating control by the Asp-Cys interaction remains to be clarified with further spectroscopic or structural studies. Nevertheless, the present work implies that a weak interaction between Asp and Cys residues, which could be formed transiently, may function as the optimal switch for control of the gating in the CP-side of the channel.

5. Conclusions
Based on the solved X-ray crystal structure of ARII in the dark state [18], we previously identified the presence of a unique H-bond between D92^ARII and C218^ARII on different helices. In the present article, we examined how this H-bonding alteration with the replacement of C218^ARII affected the photocycle and coupled proton transfer. Together with several kinetic changes of the photoproducts, both C-to-A replacement for abolishment of the H-bond and C-to-S replacement for conservation of the H-bond (but likely establishment of a stronger interaction) resulted in a delay of the turnover of the photocycle. The most crucial effect was observed in the N-to-O transition along with the reprotonation of D92^ARII from the CP bulk, which was slowed down by these substitutions. As the putative role of the native weak Asp-Cys interhelical H-bond in ARII, we inferred that this interaction may be associated with i) the proper modulation of the pK_a of D92^ARII and ii) the mechanism for switching of the accessibility of D92^ARII and/or iii) reisomerization of retinal upon the N-O transition. However, the determination of the authentic role of this interaction and its detailed molecular mechanism awaits further studies.

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Figure legends

**Figure 1:** Structural aspect in the CP-side channel of ARII. (A) Overall structure of ARII monomer based on the previously solved X-ray crystal structure at 3.2-Å resolution (PDB code: 3AM6) [18]. (B) Close-up view of the CP transmembrane region
surrounded by a blue broken line in panel (A). The residues forming two types of interhelical interactions with D92^ARI are shown together with several neighboring residues. The corresponding residues in BR are in parentheses. (C) Formation of hydrogen bond between D92^ARI and C218^ARI.

**Figure 2:** Time-dependent flash-induced absorbance changes in the wild-type (A), C218A^ARI (B), and C218S^ARI (C). The data at three selected wavelengths (400, 530, and 610 nm) are shown. The observed and regression curves by global fitting analysis are represented in gray and black lines, respectively. Experimental conditions were as follows: the media included 400 mM NaCl, 0.05% DDM, and 6 mixed buffers (citrate, MES, HEPES, MOPS, CHES, CAPS, at 10 mM each). The pH was 7.0. The temperature was set to 20°C. The corresponding figure for the wild-type was drawn from the obtained dataset in a previous publication [22].

**Figure 3:** Absorption spectra of photointermediates in kinetic photoproduct P_{P_{i}}-states. Respective spectra of P_{i} (i=1-4 (mutants) or -5 (wild-type)) calculated by global fitting for the wild-type (A), C218A^ARI (B), and C218S^ARI (C) are shown separately in panels (a)-(e) in the order of their appearance in the scheme. The observed data of respective P_{i}-spectra are shown in filled circles, while those of the dark-state (P_{0})-spectra are shown in gray continuous lines. The bold continuous and thin broken lines represent the fitting curves generated from skewed Gaussian functions for the observed data and putative spectra of the photocycle intermediates in each P_{i}-state estimated from the fitting, respectively.
**Figure 4:** Flash-induced absorbance changes for C218A<sub>ARII</sub> (A) and C218S<sub>ARII</sub> (B) at different pH values. The pH of the medium was titrated from 4.0 to 9.0 every 0.5 units. The full data are shown in Fig. S3 (A, C218A<sup>ARII</sup> and B, C218S<sup>ARII</sup>). Of these data, representative data at four pH values, pH 4.0 (a), 6.0 (b), 8.0 (c), and 9.0 (d), are shown. The noisy gray and smooth black lines represent the observed and fitting curves, respectively. The simultaneous multiexponential fitting for three absorption changes was performed using a function with four exponential terms, with the exception of three datasets for C218A<sup>ARII</sup> at pH 4.5, 5.0, and 5.5, which required five terms. For comparison, the corresponding data of the wild-type, which were quoted from our previous article [22], were shown as dotted (400 nm), broken (520 nm), and chain (610 nm) lines.

**Figure 5:** pH dependence of the decay of the absorbance changes at 400 nm in C218A<sub>ARII</sub> (A) and C218S<sub>ARII</sub> (B). The upper panels (a) show the different decay patterns of the \( \Delta A_{400} \) signals at three pH values. The noisy and smooth lines represent the observed and multiexponential fitting curves, respectively. The lower panels (b) show the plots of the multiple decay time constants against the pH. The inset of the left panel shows the corresponding plots for the wild-type. The plots of \( \tau_1 \), \( \tau_2 \), and \( \tau_3 \) are shown as closed squares, circles, and triangles, respectively. The \( \tau_4 \) and \( \tau_5 \) values stand for the time constants of the rise and decay of a M-like photoproduct (M<sub>a</sub>), which are shown in open and closed diamonds, respectively (also see the Supplementary materials). For comparison, the pH-dependence of \( \tau_3 \) in the wild-type is shown as a dashed line.
**Figure 6:** Photoinduced proton uptake and release in C218A_{ARII} (A) and C218S_{ARII} (B). The scattered signals in the respective panels represent the observed ITO signals at pH 8.5. Since these signals contained significant noise due to their relatively small amplitudes, smoothing lines are shown for the purpose of making traces of the changes easier. Gray solid lines in each panel represent the appropriately multiplied signal of the wild-type at the same pH value. Experiments were performed in the media containing 400 mM NaCl and the aforementioned six buffers at 1 mM each. The temperature was approximately 25°C (room temperature).

**Figure 7:** Estimation of pK_{a} of D92_{ARII} during N-to-O transition from pH dependence of O-yield. The maximum amplitudes of the absorbance changes at 610 nm in the wild-type (filled squares), C218A_{ARII} (filled circles), and C218S_{ARII} (empty circles) were plotted against pH. The plotted values for C218A_{ARII} and C218S_{ARII} were obtained from the data in Fig. S3, whereas those of the wild-type were obtained from the data in ref. [22]. To avoid the change of the fraction of O with the change of the functional pigment, which could be caused by e.g. protonation of a counterion (D81_{ARII}) of PSB at lower pH or deprotonation of PSB at higher pH in the dark, the values of O-yields were normalized to the maximum amplitude of M produced at respective pH values. The unit of the y-axis values is arbitrary. The plotted values were fitted by the Henderson-Hasselbalch equation with a single pK_{a} value:

\[
f_O(pH) = A \left[ \frac{1}{1 + 10^{n_a(pH-pK_{a,N})}} \right],
\]

where \(A\), \(n_a\), and \(pK_{a,N}\) are the scaling constant for the amplitude, Hill coefficient, and pK_{a} of D92_{ARII} during N-decay, respectively. The values of the slopes of the \(n\)-plots in respective proteins (wild-type, 0.55; C218A_{ARII}, 0.60;
C218^{ARII}, 0.64) were used for $n_a$.

Table 1: Comparison of lifetimes of P$_r$-states among ARII and two C218^{ARII} mutants at
### pH 7.0<sup>1)</sup>

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<th>P&lt;sub&gt;2&lt;/sub&gt;</th>
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1) The unit of all time constant values is milliseconds.

2) The corresponding values of the wild-type were cited from a previous paper [22].
Figure 2
Figure 3
Figure 4
Figure 5
Figure 6

(A) C218A

(B) C218S

\[ \text{Voltage (\mu V)} \]

\[ \text{Time (ms)} \]
Figure 7

[Graph showing pH vs. Fraction of O (a.u.) with curves for WT, C218A, and C218S]
Supplementary Materials

Interhelical interactions between D92 and C218 in the cytoplasmic domain regulate proton uptake upon N-decay in the proton transport of *Acetabularia* rhodopsin II

Jun Tamogami, Takashi Kikukawa, Keisuke Ohkawa, Noboru Ohsawa, Toshifumi Nara, Makoto Demura, Seiji Miyauchi, Tomomi Kimura-Someya, Mikako Shirouzu, Shigeyuki Yokoyama, Kazumi Shimono and Naoki Kamo
Table S1: Parameters of the skewed Gaussian fit for Pl-spectra in Fig. 3

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Figure S1: Photocycle of D92N^ARH. (A) Flash-induced absorbance changes at three selected wavelengths (400, 530 and 610 nm). The observed signals are represented as gray noisy lines, while the regression curves by the global fitting are represented as black smooth lines. The wild-type data at the same pH are shown as dotted (400 nm), broken (530 nm), and chain (610 nm) lines. Through the global fitting, the lifetimes (decay time constants) of the three P-states were estimated as 0.024 ms for P_1, 238.7 ms for P_2, and 1970.4 ms for P_3, respectively. (B) Absorbance spectra of P_i-states (i=1-3) determined by the global fitting. The spectrum at the unphotolyzed state (P_0) having a λ_{max} at ca. 522 nm is shown as a black broken line. Measurements were performed under the same experimental conditions described in Fig. 2.
Figure S2: Estimation of O- and N-spectra. The presented data are from the previous results of the global fitting in the wild-type at pH 5.0 (A) and 8.6 (B) [S1]. The upper panels (a) show the absorbance changes (gray lines) at three wavelengths (400, 530, and 610 nm) together with the fitting curves (black lines). The vertical red lines in each panel exhibit the time constants of P₄-decay at respective pH values. The lower panels (b) show the results of the fitting analyses with the skewed Gaussian functions for the spectra of P₄. As described previously, the spectrum of P₄ at pH 5.0 contains only an absorption band of O (O₂) as a photoproduct [S2]. Therefore, the analysis for P₄-spectrum yielded an absolute spectrum of O (shown in a broken line). The spectral parameters of O estimated from the fitting were $\lambda_{\text{max}} = 562$ nm, $A_{\text{max}} = 0.84$, $\rho = 1.1$, and $\Delta \nu = 3652$ cm⁻¹. Another small absorbance band at the shorter spectral range (~300-450 nm) may be due to the $\beta$-band. The P₄-spectrum at pH 8.6 is in the
quasi-equilibrium of M, N, and O, where the N/O ratio is >1 because of the slower N-to-O transition at this pH. Thus, the analysis of the P₄-spectrum at pH 8.6 was performed by the function with three Gaussian bands. The calculated spectra of M, N, and O by the fitting are shown as broken lines. During the fitting, the spectral parameters of O, except $A_{\text{max}}$, were fixed to the same values estimated from the analysis for P₄ at pH 5.0. The estimated spectral parameters of N were $\lambda_{\text{max}} = 526$ nm, $A_{\text{max}} = 0.36$, $\rho = 1.8$, and $\Delta \nu = 4165$ cm⁻¹.
Figure S3: Time-dependent absorbance changes at typical three wavelengths in C218A<sup>ARII</sup> (A) and C218S<sup>ARII</sup> (B) at varying pH values.
Figure S4: Formation of $M_a$ in C218SARII under alkaline conditions (pH 8.5 (A) and 9.0 (B)). In two panels, we can observe the increase of the $\Delta A_{400}$ signals after their first decay in the early time region prior to $\sim$0.1 ms, which originates from the decay of a normal $M$. Moreover, these increasing phases appear to agree with the time course of a slower K-decay (monitored at 610 nm) that does not coincide with the rise of normal $M$. The observation of similar absorption changes was also reported for PR at alkaline pH ($\geq \sim$8.5), which was kinetically explained by assuming the existence of another parallel photocycle during which a $M$-like photoproduct ($M_a$) is produced [S3]. Thus, these $\Delta A_{400}$ signals were analyzed by a photocycle scheme via two parallel paths like PR, in which $M$ and $M_a$ are independently produced [S3]. Under the assumption that the precursor intermediates of $M$ and $M_a$ do not affect the formation/decay of these two
M-like states due to their sufficiently faster kinetics, the time-dependent fractional concentration changes of M and Mₐ can be simply represented by the following equations:

\[ \Delta M \approx A_1 \exp\left(-t / \tau_1\right) + A_3 \exp\left(-t / \tau_3\right) \]  

(Eq. S1)

and

\[ \Delta M_a \approx B \frac{\tau_5}{\tau_4 - \tau_3} \left[ \exp\left(-t / \tau_4\right) - \exp\left(-t / \tau_5\right) \right] \]  

(Eq. S2)

Here, as defined in the text, \( \tau_1 \) and \( \tau_3 \) in Eq. S1 are time constants of the first and third decay components of M, whereas \( A_1 \) and \( A_3 \) are scaling constants for their amplitude. A second decay component (\( \tau_2 \)) cannot be clearly observed, probably due to overlapping with the decay of Mₐ (\( \tau_5 \)); thus, it was omitted. \( \tau_4 \), \( \tau_5 \), and \( B \) in Eq. S2 represent time constants of the rise and decay of Mₐ and a scaling constant for the amplitude of Mₐ, respectively. A function of the sum of the two equations S1 and S2 noted above (\( \Delta A_{400} \approx \Delta M + \Delta M_a \)) was used for the fitting. The adequate fitting curves are shown in red continuous lines. Respective calculated fractional changes for M and Mₐ are shown in red broken and dotted lines, respectively.
Figure S5: Observation of two O-intermediates in C218A<sup>ARII</sup> (A) and C218S<sup>ARII</sup> (B) under acidic condition (pH 4.0). The decreasing phases of the 610 nm-signals originating from K-decay, which is monitored at the early time range (< ~0.1 ms), are not observed at pH 4.0, suggesting sufficiently fast K-decay to be undetectable in our employed apparatus for measurements. As described in the text, the replacement of C218<sup>ARII</sup> induces the apparent detection of L in the early kinetic photoproducts (P<sub>1</sub> and P<sub>2</sub>) (see Fig. 3), which may not contradict the faster K-decay in these mutants. The latter time region of the 610 nm-signal reflects the formation and decay of O. As described previously [S2], the photocycle of the wild-type required the assumption of two sequential Os (O<sub>1</sub> and O<sub>2</sub>) with different molar extinction coefficient (ε) values under
lower pH conditions ($\leq \sim 5.5$), although $O_1$ cannot be detected at neutral and basic pH values due to slowing of the $N \rightarrow O_1$ reaction as the medium pH increases [S1]. The $O_2$-yield also decreases with further increases in pH. However, in the wild-type, a small amount of $O$ ($O_2$) was observable even at pH 8.8 [S1]. In contrast, a pH-increase above $\sim 7.5$ in C218*ARII* mutants gave rise to no accumulation of $O_2$ during their photocycles (Fig. S3). This may be interpretable as the effects of the significantly slower N-decay in these mutants. The kinetic analyses by three models concerning $O$ were performed for the $\Delta A_{610}$ signals. The principle of these analyses is described in our previous paper [S2]. The panels (a), (b), and (c) show the fitting results by the models containing single $O$, two sequential $Os$ ($O_1$ and $O_2$) with equal $\varepsilon$, and sequential $O_1$ and $O_2$ with different $\varepsilon$ values, respectively. The bold continuous red lines represent the best fit curves in respective analyses. The calculated fractional concentration changes of $O_1$ and $O_2$ from these analyses are shown in red broken and dotted lines, respectively. Among three analyses, the third scheme (panel (C)) gave the most preferable fitting results for both mutants.
Figure S6: Kinetics of photoinduced proton release and uptake in C218A$^{\text{ARII}}$ (A) and C218S$^{\text{ARII}}$ (B). The rates of the first proton release and consecutive uptake were estimated by the fitting with the following equation:

$$\Delta\text{Voltage} \propto \Delta [\text{H}^+] = C\frac{\tau_{\text{uptake}}}{\tau_{\text{release}} - \tau_{\text{uptake}}} \left[ \exp\left(-t / \tau_{\text{release}}\right) - \exp\left(-t / \tau_{\text{uptake}}\right) \right],$$

where $\tau_{\text{release}}$, $\tau_{\text{uptake}}$, and $C$ are the time constants of the first proton release and second uptake, and a constant reflecting the fraction of the subpopulation inducing the initial proton release, respectively [S3]. The continuous red lines in respective panels represent the fit curves. For comparison, the time constant of the H$^+$-uptake in the wild-type ($\tau'_{\text{uptake}}$) was also estimated by the fitting for the decreasing phases in the observed signals (continuous
grey lines) with a single exponential equation (broken red lines).

**Figure S7**: pH dependence of O-decay. (A) Time-dependent absorbance changes at 610 nm in the wild-type (a), C218A<sup>ARII</sup> (b), and C218S<sup>ARII</sup> (c) at three pH values. (B) Plot of time constants of O-decay versus pH. The time constants of O-decay in respective proteins (■, wild-type; ●, C218A<sup>ARII</sup>; ○, C218S<sup>ARII</sup>) were calculated from the regression
for the decaying phases of the 610 nm-signals with a single exponential equation.

References


