ABSTRACT: Attractant and repellent signaling conformers of the dual-signaling phototaxis receptor sensory rhodopsin I and its transducer subunit (SRI–HtrI) have recently been distinguished experimentally by the opposite connection of their retinylidene protonated Schiff bases to the outwardly located periplasmic side and inwardly located cytoplasmic side. Here we show that the $pK_a$ of the outwardly located Asp76 counterion in the outwardly connected conformer is lowered by $\sim 1.5$ units from that of the inwardly connected conformer. The $pK_a$ difference enables quantitative determination of the relative amounts of the two conformers in wild-type cells and behavioral mutants prior to photoexcitation, comparison of their absorption spectra, and determination of their relative signaling efficiency. We have shown that the one-photon excitation of the SRI–HtrI attractant conformer causes a Schiff base connectivity switch from inwardly connected to outwardly connected states in the attractant signaling photoreaction. Conversely, a second near-UV photon drives the complex back to the inwardly connected conformer in the repellent signaling photoreaction. The results suggest a model of the color-discriminating dual-signaling mechanism in which phototaxis responses (his-kinase modulation) result from the photointerconversion of the two oppositely connected SRI–HtrI conformers by one-photon and two-photon activation. Furthermore, we find that the related repellent phototaxis SRI–HtrII receptor complex has an outwardly connected retinylidene Schiff base like the repellent signaling forms of the SRI–HtrI complex, indicating the general applicability of macro conformational changes, which can be detected by the connectivity switch, to phototaxis signaling by sensory rhodopsin–transducer complexes.

Many microorganisms use photoreceptors to modulate their motility to seek or avoid regions of illumination depending on light intensity and color. In halophilic archaea, visual pigment-like photoreceptors, sensory rhodopsins, mediate attractant and repellent phototaxis responses by controlling through transducer proteins a his-kinase that modulates the swimming behavior of the cell (1–3). Sensory rhodopsin I (SRI) is of special interest because it mediates opposite behavior depending on light color: attractant responses (due to transient his-kinase inhibition) upon absorption of a long-wavelength photon and repellent responses (due to transient his-kinase activation) when a second short-wavelength photon is absorbed (4). Single mutations have been found in SRI and also in the second subunit of the receptor–transducer complex, the transducer HtrI, which invert the sign of the single-photon response to repellent, and second-site (suppressor) mutations in both subunits cancel the effect of inverter mutations restoring wild-type behavior (5–7).

We developed an electrophysiological method for measuring the vectoriality and kinetics of light-induced charge movements within rhodopsin molecules in suspensions of Escherichia coli cells expressing various microbial rhodopsins (8). Using this method, we found that the SRI–HtrI complex exists in two conformational states with opposite connectivity of the retinylidene Schiff base in the photoactive site (9). By connectivity of the Schiff base, we mean the direction, inward toward the cytoplasm or outward toward the extracellular medium, of proton release following single-photon absorption. In the conformation that mediates attractant photoresponses, the Schiff base deprotonates toward the cytoplasmic side of the membrane to the inner, cytoplasmic half-channel of SRI. In the repellent-mediating conformation of the complex, the proton is transferred in the opposite direction to the acceptor Asp76 localized in the extracellular half-channel. It should be emphasized that the light-induced proton transfer to Asp76 itself does not play a significant functional role in signaling and reflects only the outwardly connected state of the complex, since the repellent receptor conformer of SRI_D76N–HtrI_E56Q mediates repellent responses and does not deprotonate the Schiff base. Inversion of Schiff base connectivity during the photocycle (often called the connectivity or accessibility switch) is the key event enabling vectorial proton transport and light energy conversion by light-driven proton pumps such as bacteriorhodopsin (10). Our data
show that this feature of energy-converting microbial rhodopsins is crucial for the sensory pigments as well.

Our electrical data revealed that both conformers are present in the wild-type SRI–HtrI complex, as well as in the inverted (single mutant) and restored (double mutant) complexes (9). However, in the wild-type and restored complexes, both of which mediate attractant responses, the conformational equilibrium is poised strongly in favor of that with an inwardly accessible Schiff base. Signal-inverting mutations shift the equilibrium in favor of the outwardly accessible Schiff base form. The amplitude of charge movement depends on two parameters unknown for SRI: the distance over which the Schiff base proton is displaced and the angle between the direction of its movement and the membrane. Therefore, the electrical data do not allow quantitative determination of the relative sizes of the attractant and repellent conformer pools in the wild-type and mutated complexes.

To test for the existence of the two conformer pools by another method and also to enable quantization of the two pools, we applied absorption spectroscopy of E. coli and native Halobacterium salinarum membranes containing wild-type (attractant), inverted mutant (repellent), and restored (double mutant attractant) SRI–HtrI complexes. This characterization confirms and extends our understanding of the conformational equilibrium and its role in signaling and, moreover, led us to a mechanistic model of SRI–HtrI complex color-discriminating signaling presented here. Predictions from this model regarding Schiff base connectivity switching in the one-photon and two-photon signaling process in the SRI–HtrI complex and the one-photon SRII–HtrII complex are tested and confirmed by electrical measurements. The data provide compelling evidence of the general applicability of the model to phototaxis signaling by SR–Htr complexes.

MATERIALS AND METHODS

Mutagenesis and Expression. Genes encoding SRI–HtrI fusion proteins, in which the C-terminus of SRI is joined through a flexible linker peptide (ASASNGASA) to the N-terminus of HtrI truncated at position 147, were cloned into expression vector pET-21d (Novagen, Merck KgaA, Darmstadt, Germany) under the control of the T7 promoter between NcoI and BamHI sites. The gene encoding the SRI–HtrII fusion protein, in which the C-terminus of NpSRII (SRII from Natronomonas pharaonis) is joined through the same flexible linker peptide to the N-terminus of NpHtrII truncated at position 157, was cloned into pET-21d between NcoI and HindIII. Residue replacement mutations were made using the QuickChange site-directed mutagenesis kits (Stratagene, La Jolla, CA). Expression of the genes in BL21(DE3) was induced by the addition of 1 mM isopropyl β-D-thiogalactopyranoside and 5 μM all-trans-retinal.

Genes encoding SRI joined through the same linker peptide to the N-terminus of full-length HtrI were cloned into a halobacterial expression vector under the control of the bop promoter as described previously (11) between NcoI and XbaI sites. Mutations were introduced by the two-step mega-primer polymerase chain reaction (PCR) method (12) with pfu turbo polymerase (Stratagene) or Phusion polymerase (Finnzyme, Espoo, Finland). H. salinarum strain Pho81Wr (BR HR’ SRI’ HtrI’ SRII’ HtrII’), carotenoid-deficient and restriction-deficient (12) was used as the recipient in plasmid transformations. The transformed H. salinarum cells were grown to early stationary phase in complex medium containing 1 mg/mL mevinolin as described previously (7).

Membrane Preparation. E. coli cells expressing SRI–HtrI47 suspended in 4 M NaCl and 25 mM MES (pH 6.0) were disrupted by a microfluidizer (Microfluidics, Newton, MA). H. salinarum cells expressing SRI–HtrI were disrupted by sonication. Unbroken cells and cell debris were pelleted by low-speed centrifugation (Sorvall RC6 plus, Thermo Fisher Scientific Inc., Asheville, NC). The membrane fraction in the supernatant was then pelleted for 1 h at 147000 g in a Beckman Optima TM L-100 XP ultracentrifuge (Beckman Coulter Inc., Brea, CA) and suspended in 4 M NaCl and 25 mM MES (pH 6.0).

Absorption Spectroscopy. Absorption spectra in the UV–visible range were recorded on a Cary 4000 spectrophotometer (Varian, Palo Alto, CA) with an integrating sphere. Control absorption spectra of membranes not expressing receptor–transducer complexes were subtracted from those with expression. Optical densities at 850 nm, reflecting mostly light scattering and proportional to membrane concentration, were set equal in experimental and control samples. Residual scattering effects in corrected spectra of the complexes were eliminated (if necessary) by subtraction of a logarithmic-scale linear function based on the slope of optical density between 750 and 850 nm, where no significant absorption of membrane components exists. Titration of absorption changes was conducted with the same samples by an increase in pH in small steps by addition of microliter amounts of non-neutralized 0.5 M Tris buffer (pH 10) and in later steps by 1 M NaOH. The pH of the sample was measured before and after spectral scans, and in cases of minor differences, the averaged value of the pH was used for plotting the titration curves.

Electrical Measurements. Intramolecular charge movements were measured in suspensions of E. coli cells expressing the indicated proteins by the method described previously (8). The suspension of E. coli cells was flushed along the line between two platinum electrodes by a Vibrant HE 355II Tunable Laser (OPOTEK Inc., Carlsbad, CA) set at desired wavelengths. A macroscopic electrical current in the cuvette appeared due to asymmetric absorption of light by each bacterial cell. The electrode remote from the light source was fed into a model 428 low-noise current amplifier (Keithley) with a 2 μs rise time. The signals were digitized and stored using Digidata 1325A and pCLAMP 9.0 (both from Axon Instruments). Flash intensities were attenuated to be below the saturation of the signals. Twenty to 50 signals in 10 s intervals with a maximum sampling rate of 2 μs/pixel were averaged. If not otherwise indicated, the measuring buffer contained 5 mM Tris-HCl, 1.5 mM NaCl, 0.15 mM CaCl2, and 0.15 mM MgSO4 (pH 7.6).

Cell Motility Measurements. Transformed H. salinarum cells were grown in complex medium containing 1 mg/mL mevinolin as described previously (7). Cultures at the end of their exponential growth phase were diluted 1:10 in fresh CM and incubated for 1 h at 37 °C with agitation. For motility defined as being in the dark, cell trajectories in nonactinic light at wavelengths of >720 nm were captured as real-time AVI files using a Flashbus Spectrum Lite Video Capture PCI Card on a Dell Dimension 8300 personal computer running VirtualDub 1.6.19.0 (http://www.virtualdub.org), AVI encoder software for video capture. VirtualDub was set to record 10 frames per second during 30 s of cell swimming. Means of spontaneous reversals of the cells in the dark were measured by tracking each cell captured in the AVI files for 30 s to count the number of reversals. Means and standard deviations were calculated from 20–30 cells.
RESULTS

The Attractant and Repellent Conformers of the SRI–HtrI Complex Differ in Their $pK_a$ Values of the Schiff Base Counterion. The $p$H titration of the SRI–HtrI complex in *H. salinarum* membranes (Figure 1A) shows difference spectra characteristic of the well-established shift of the absorption maximum from 587 to 552 nm due to deprotonation of the Schiff base counterion Asp76 (1). This deprotonation gives rise to absorption difference spectra with a minimum at $\sim 610$ nm, a smaller maximum at $\sim 520$ nm, and an isosbestic point at 560 nm.

However, for the SRI–HtrI complex in *E. coli* membranes, in addition to these long-wavelength bands, a second $p$H-induced absorption change is observed with a minimum at $\sim 440$ nm, a maximum at $\sim 365$ nm, and an isosbestic point at 400 nm (Figure 1B).

The short-wavelength titration bands in *E. coli* membranes (Figure 1B) are similar to the titration bands of misfolded microbial rhodopsins [e.g., bacteriorhodopsin and a subcellular sensory rhodopsin-like protein (Figure 1C) and *Chlamydomonas* sensory rhodopsin B (also known as channel rhodopsin-2)], which are expressed at high concentrations according to immunoblots yet produce no light-induced charge movements. These bands are most likely due to deprotonation of the retinylidene Schiff base in the misfolded protein, as observed for model retinylidene Schiff base compounds (1). We commonly observe variable amounts of spectrally similar misfolded fractions in many other microbial rhodopsins. For our purposes here, it is important to carefully monitor and avoid the contribution of the misfolded protein effect to titrate accurately the Asp76 counterion of SRI in *E. coli* membranes. At low to neutral $p$H, the superposition of signals due to the functional and misfolded SRI–HtrI complexes leads to suppression of the positive absorption band of the former and shifts its apparent position to a longer wavelength (Figure 1A,B). On the other hand, negligible absorption changes occur at wavelengths above 600 nm in the misfolded proteins (Figure 1C). Therefore, the absorbance changes in the long-wavelength region of the difference spectra (between 610 and 650 nm) were chosen as a measure of the transitions between acid and alkaline forms of SRI. In addition, special attention was paid to exclude other possible distortions of the titration data by processes other than protonation and deprotonation of the Asp76 counterion. Deprotonation of the Schiff base at extremely high $p$H values was also analyzed, and its perturbing effects were excluded.

A titration curve of wild-type SRI–HtrI complex in native halobacterial membranes obtained with these precautions is fit poorly with a single $pK_a$, whereas the sum of two independent functions $[A_1/(1 + 10^{-pK_{a1}}) + A_2/(1 + 10^{-pK_{a2}})]$ fits well (Figure 2, top). The $pK_a$ values of the two components differ by almost 2 units (7.0 and 8.8), and the relative amplitude of the low-$pK_a$ component is 20–24%, which matches our earlier published observations (11).

We compared the titration of the wild-type complex, which mediates an attractant single-photon response, to that of the SRI–HtrI_E56Q complex, an inverted mutant that produces a strong repellent single-photon response. This mutation is of special interest because it demonstrates that the complex undergoes concerted structural changes since the Glu56 residue in the transducer has a long-distance effect on the photoresponse center in SRI (9).

Initially, we investigated both complexes in *E. coli* membranes at 4 M NaCl because of the higher level of expression and stability.
of the complexes under high-salt conditions. Two components of the counterion titration are present in this system in the wild-type as well as in the mutated SRI–HtrI complexes (Figure 2, bottom). A remarkable result is that the absolute \( pK_a \) values of the two components are not significantly affected by the mutation (7.9 and 9.0 for the wild type and 7.7 and 8.7 for the mutant, respectively), whereas their relative amplitudes change 7.3-fold (Figure 3, left panels). Our interpretation is that the inverting mutation shifts the population of SRI–HtrI complex molecules from the high-\( pK_a \) conformer to the low-\( pK_a \) conformer.

The large increase in the ratio of the low-\( pK_a \) to high-\( pK_a \) components caused by the inverting mutation matches closely the changes in the relative size of the pools of outwardly and inwardly directed Schiff bases found earlier by charge movement measurements (9). Because these measurements were performed at low ionic strengths, for comparison we performed the pH titration of the wild-type and mutated complexes expressed in *E. coli* membranes also under low-salt conditions (0.1 M NaCl). Results similar to those at 4 M NaCl were obtained: the inverting mutation changes the relative sizes of the low-\( pK_a \) and high-\( pK_a \) amplitudes in the same direction as at high salt without significant changes in these \( pK_a \) values (Figure 3, middle panels). Finally, the effect of the inverting mutation on the relative amounts of low- and high-\( pK_a \) conformers in *E. coli* membranes was confirmed to exist in the natural halobacterial membranes (Figure 3, right panels).

Figure 4 illustrates the nearly perfect quantitative correlation of the effects of inverting and suppressor mutations on the ratio of titration components and on the relative fractions of SRI–HtrI complexes with inwardly and outwardly directed Schiff bases. The inverting (repellent) mutation in transducer HtrI_E56Q increases both ratios by 1 order of magnitude in *E. coli* membranes. The second-site suppressor mutation (HtrI_E56Q_N53D), which restores wild-type attractant signaling, restores nearly completely both parameters.

Both components of pH-induced spectral changes reflect the protonation state of the Schiff base counterion Asp76, since the spectral shift is typical of counterion protonation in SRI and other microbial rhodopsins (13–15), and no pH dependence of the main absorption maximum was observed when Asp76 was mutated to neutral asparagine (data not shown). In summary, we conclude that the conformer with a low Asp76 \( pK_a \) has an outwardly accessible Schiff base and when photoactivated initiates a repellent response, whereas the conformer with the opposite connectivity of the Schiff base has the higher Asp76 \( pK_a \) and mediates an attractant response.

**The Absorption Spectrum of the Repellent Receptor Conformer Is Blue-Shifted from That of the Attractant Receptor Conformer.** The absorption spectra of the wild-type SRI–HtrI complex and the SRI–HtrI_E56Q complex are identical when the Asp76 counterion is neutralized by acidic pH (Figure 5A) or by the D76N mutation (Figure 5B). On the other hand, when Asp76 is ionized the pH shift-induced differential absorption spectra in the range of low \( pK_a \) are blue-shifted compared to those in the range of high \( pK_a \) (Figure 5C). Hence, the absorption spectrum of the outwardly connected Schiff base (repellent receptor) conformer is blue-shifted compared to that of the inwardly connected (attractant) conformer. The bluer absorption of the low-\( pK_a \) conformer with ionized Asp76 is expected from a stronger interaction of Asp76 with the protonated Schiff base (16).

**Kinase-Activating Activity of Dark-Adapted Attractant and Repellent Receptor Conformers.** The frequency of spontaneous swimming reversals in *H. salinarum* cells reflects the activity of the histidinyl kinase CheA (17, 18). In wild-type cells under natural conditions, alterations of the kinase activity following an increase in light intensity are transient, and the steady-state swimming reversal frequency is the same in the dark and in the light (19). This homeostasis is accomplished by adaptation machinery in the cells that counteracts the effects of light intensity changes and, at the low natural wild-type SR–Htr complex concentrations, resets the kinase activity, and consequently the swimming reversal frequency, to its prestimulus value (I). However, when the receptor–transducer complexes are overexpressed in the cells to >10-fold levels, the adaptation system is not sufficient to reset the kinase activation level (6, 20, 21).
Signal-inverting mutations such as E56Q shift the equilibrium of the two conformers in favor of the outwardly accessible Schiff base form (the repellent receptor conformer). The spontaneous swimming reversal frequency in the dark (i.e., dark kinase activity) of these mutants is ∼10-fold lower as seen for the SRI-HtrI_E56Q complex (Figure 6). Attractant signaling behavior restored to this mutant by the N53D suppressor mutation restores the wild-type ratio of the two conformer pools (Figure 4) and restores the higher kinase activity as shown by the higher dark reversal frequency (Figure 6). Therefore, the repellent receptor outwardly connected conformer exhibits notably less activation of the kinase than the attractant receptor inwardly connected conformer in the dark. Photostimulation of the outwardly connected form induces swimming reversals (i.e., a repellent response) that indicates its photoproduct activates the kinase, and conversely, photostimulation of the inwardly connected form suppresses swimming reversals (i.e., an attractant response) which indicates its photoproduct inhibits the kinase. These expectations are confirmed by steady-state reversal frequency measurements in the light (J. Sasaki and J. L. Spudich, manuscript in preparation). Note that there is apparently a higher molar efficiency of kinase activation than kinase inhibition from the complex since the overall response is strongly repellent in the presence of close to equal fractions of the two conformers in native membranes (Figure 3, right panels).

A Switch in Schiff Base Connectivity Occurs in the Single-Photon Response of the Wild-Type SRI–HtrI Complex. To test whether the Schiff base switches connectivity after single-photon absorption, we measured the intramolecular charge movement in response to 410 nm laser flashes in the dark-adapted wild-type complex and in the same complex under strong long-wavelength (> 520 nm) background illumination that leads to accumulation of the long-lived unprotonated Schiff base photointermediate M (λ_{max} = 373 nm, also known as S_{373}) in the SRI photocycle (Figure 7). In darkness, only a very small signal with kinetics typical of the photoconversion of the attractant receptor conformer from the dark-adapted SRI-HtrI complex (λ_{max} = 587 nm) was observed, which we attribute to the weak absorption of SRI at 410 nm. Intense orange background illumination converts the dark-adapted form to the M intermediate which absorbs with a λ_{max} of 373 nm and has strong absorption at 410 nm (1). The fraction of molecules converted to M was tested by the decrease in the magnitude of the photoelectric signal produced by 580 nm laser flashes (data not shown). Under our condition, 67% of the dark-adapted state is converted. Hence, under orange background illumination, the 410 nm laser-induced electrical signal caused by excitation of the residual nonconverted SRI–HtrI complex should be 33% of the small signal seen without background illumination (Figure 7, curve 1). Therefore, the charge movement elicited by the 410 nm flash in an orange background can be attributed essentially entirely to photoconversion of the M intermediate. The electrical response
of intermediate M photoexcitation is an inwardly directed charge transfer (Figure 7, curve 2), with kinetics >10-fold faster than those of the inward charge movement from photoexcitation of dark-adapted SRI (Figure 7, curve 3); 410 nm is the shortest output wavelength available from the laser used, and a large decrease in the amplitude of the signal at longer wavelengths, which disappears at 430 nm, corresponds perfectly to the absorption spectrum of SRI M (data not shown).

Photoexcitation of SRI M causes rapid reprotonation of the Schiff base beginning with submicrosecond times (22). As shown in curve 2 of Figure 7, this reprotonation causes inwardly directed charge transfer, which therefore must be from the extracellular side (likely from the protonated Asp76). We conclude that SRI M before the absorption of the second short-wavelength photon is in the conformation with the outwardly connected Schiff base. The accelerated inwardly directed charge movement from light-induced Schiff base reprotonation of M as compared to that of the inwardly directed Schiff base deprotonation in dark SRI is in agreement with previously measured kinetics of these processes (22).

In conclusion, single photon excitation of the SRI–HtrI attractant conformer causes a Schiff base connectivity switch from the inwardly connected kinase-activating conformation to the outwardly connected kinase-inhibiting conformation. A second near-UV photon drives SRI M back to the unphotolyzed Schiff base state (Figure 8). As shown previously (23–25) and in this study, we confirm the existence of two conformer pools not available from photoelectrophysiology, including quantitative determination of the relative amounts of the two conformers in the wild-type and behavioral mutants, assessment of the relative signaling efficiency by the conformers, and measurement of the absorption properties of each conformer.

Two conformers with the oppositely connected Schiff base ensure opposite sides of release and uptake of protons in light-driven ion-pumping rhodopsins (28). Our results show that two similar conformers play key roles also in signaling by sensory rhodopsins. The presence of analogous conformers in bacteriorhodopsin’s photocycle was likely crucial to the conversion of this proton pump with only three mutations to a robust phototaxis receptor mimicking SRII signaling through HtrII (29).

The pH titration of absorption shows that in the wild-type complex ~80% of the molecules are in the inwardly connected attractant receptor conformation. An inverting mutation does
not change significantly the $pK_a$ values of the attractant and repellent receptor conformers; rather, the relative sizes of the conformer pools is increased 6–10-fold in favor of the repellent conformer with the outwardly connected Schiff base. Suppressor mutations reverse this effect to near the wild-type ratio. The net phototaxis response is strongly repellent in the presence of close to equal fractions of the two conformers in *H. salinarum* cells. Therefore, there appears to be a higher molar efficiency of light-induced kinase activation than kinase inhibition from the complexes. This observation agrees with the finding that in the wild-type SRI–HtrI complex the long-lived unprotonated Schiff base photointermediate M ($\lambda_{\text{max}} = 373$ nm, also known as S373) in the SRI photocycle is the attractant signaling state and that it functions as well as a repellent receptor; i.e., its photoactivation by a second photon leads to a repellent response (1, 4, 30). In view of the results described above, an elegantly simple model results from the generalization of these early findings. Our generalized and more explicit model proposes that concomitant changes between the same two conformers identified here are responsible for the attractant signal from SRI M accumulation, the repellent response from SRI M photoactivation, the repellent response mediated by inverted mutant SRI–HtrI complexes, and the repellent responses mediated by the SRII–HtrII complex (Figure 9). The essence of the model is that the two conformers are photo-interconvertible; i.e., the photointermediate of the kinase-activating inwardly connected conformer is the kinase-inhibiting outwardly connected conformer, which would produce a decrease in kinase activity, i.e., an attractant response. Conversely, the photointermediate of the kinase-inhibiting conformer is the kinase-activating conformer, which produces an increase in kinase activity. A second key feature is that the conformers inhibit or activate the kinase in a manner independent of whether they are photointermediate states, such as SRI M, or thermostable molecules created by mutation (SRI–HtrI vs SRI–HtrI_E56Q) or by evolutionary changes (SRI–HtrI vs SRII–HtrII). It should be emphasized that upon photoexcitation SRI–Htr complexes switch their conformations, which can be distinguished by their opposite Schiff base connectivity, in photointermediate states and the photocycles end with the same thermostable conformation. The return of connectivity to that of the initial state is demonstrated by the observation that the direction of proton movement caused by a second flash after the reappearance of the unphotolyzed spectral state is the same as in response to the first flash.

The switch in Schiff base connectivity from inward to outward in the one-photon attractant signaling photocycle of the wild-type SRI–HtrI complex reported here is a crucial aspect of the model. Note that this conformational change is the reverse of that occurring in the well-studied bacteriorhodopsin photocycle in which the switch is from outward to inward (28). The structural basis of the Schiff base connectivity switch and its coupling to the global conformation of transport rhodopsins has been a subject of great interest, and several models and views have been proposed (reviewed in refs 10, 31, and 32). An atomic-level understanding of this phenomenon is of additional interest given its importance also in sensory rhodopsins. Note that following the finding that HtrI-free SRI carries out light-driven proton transport (33) several authors raised the possibility that the conformational changes of transport rhodopsins are involved in sensory rhodopsin signaling (34–37). Our characterizations of the properties of the signaling conformers here and in our previous study (9) confirm that involvement and further show that specifically the Schiff base connectivity switch, the process that ensures vectoriality to the transporters, plays a key role in sensory rhodopsin function.

We emphasize that the connectivity switch is a measurable aspect of a macroconformational transition of a domain of the SRI–Htr complex that behaves as a single unit undergoing a concerted transition between two states. Characterization of SRI–HtrI mutants indicates this domain includes the SR photoactive site as well as the membrane portion and membrane-proximal (HAMP) domain of the Htr transducer (5, 9). Evidence of such tight coupling within the complex is the fact that mutations in the transducer subunit of the complex cause long-distance effects in the receptor subunit’s photoactive site (9). Whether the connectivity switch is a necessary component or
merely an indicator of the photoinduced transition is not yet known.

It is known that a steric conflict with the isomerizing retinal that requires a Thr-Tyr hydrogen-bonded pair in the photoactive site (38), a process not observed in transport rhodopsins, is required for signaling by the SRII–HtrII complex (39). Introduction of the corresponding Thr-Tyr pair into bacteriorhodopsin by mutation creates the steric conflict (40) and converts bacteriorhodopsin into a functional repellent phototaxis receptor (29). The relationship of the roles of the steric interaction process and the Schiff base connectivity switch in signaling is not clear and merits further investigation.

One of the most intriguing questions is how the opposite direction of switching between the two conformational states is achieved by the same light input. The absorption spectral maxima of the two conformers differ by 5–10 nm. This difference is eliminated by the Asp76Asn mutation or Asp76 neutralization at ionized Asp76, the wild-type and inverted mutant SRI.

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