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Modulation of spectral properties and pump activity of proteorhodopsins by retinal analogues

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Proteorhodopsins are heptahelical membrane proteins which function as light-driven proton pumps. They use all-*trans*-retinal A1 as a ligand and chromophore and absorb visible light (520–540 nm). In the present paper, we describe modulation of the absorbance band of the proteorhodopsin from Monterey Bay SAR 86 gammaproteobacteria (PR), its red-shifted double mutant PR-D212N/F234S (PR-DNFS) and *Gloeobacter* rhodopsin (GR). This was approached using three analogues of all-*trans*-retinal A1, which differ in their electronic and conformational properties: all-*trans*-6,7-*s-trans*-locked retinal A1, all-*trans*-phenyl-retinal A1 and all-*trans*-retinal A2. We further probed the effect of these retinal analogues on the proton pump activity of the proteorhodopsins. Our results indicate that, whereas the constraints of the retinal-binding pocket differ for the

proteorhodopsins, at least two of the retinal analogues are capable of shifting the absorbance bands of the pigments either bathochromically or hypsochromically, while maintaining their proton pump activity. Furthermore, the shifts implemented by the analogues add up to the shift induced by the double mutation in PR-DNFS. This type of chromophore substitution may present attractive applications in the field of optogenetics, towards increasing the flexibility of optogenetic tools or for membrane potential probes.

Key words: chromophore, microbial rhodopsins, retinal analogues, spectral sensitivity.

INTRODUCTION

Microbial rhodopsins (also called type-I rhodopsins) are a large family of light-sensitive heptahelical transmembrane proteins found in archaea (e.g. bacteriorhodopsin, sensory rhodopsin II), bacteria (e.g. proteorhodopsins, *Anabaena* sensory rhodopsin) and eukaryotes (e.g. algal and fungal rhodopsins) [1–3]. These proteins mediate a variety of functions in their hosts, including phototactic signalling and active transport of ions. The archaeal proton pump bacteriorhodopsin (BR) is the first identified of these rhodopsins and the best studied over the last four decades [4–6]. The first proteorhodopsin was discovered in gammaproteobacteria of the SAR 86 group, from Monterey Bay in California, U.S.A. (PR) [7]. PR contains the transmembrane motif with seven α -helices and the chromophoric ligand all-*trans*-retinal A1, common to all rhodopsins, and shares approximately 20% sequence identity with BR [7].

Many different variants of PR have been found since, widely distributed in marine, freshwater and terrestrial biotopes [8]. PR could successfully be heterologously expressed in *Escherichia coli* [9–14] and was shown to function as a light-driven proton pump after incubation with retinal [7]. The phototrophic potential of PR can be exploited to drive physiological activities in a heterologous environment such as flagellar motility or ATP synthesis in *E. coli* [10,11]. In native organisms, PR has been shown to enhance survival or growth rate under energy-limiting conditions [15,16]. This potential and their wide distribution

suggests that PRs are a key player in the phototrophic energy balance of the biosphere.

PR covalently binds all-*trans*-retinal A1 (Figure 1) through a protonated Schiff base with Lys²³¹ of helix G, yielding a new absorbance band peaking at ~525 nm [7,17,18]. Absorption of a single photon leads to the isomerization of the all-*trans* chromophore to the 13-*cis* configuration, which in turn induces a sequence of specific structural changes in the protein moiety (opsin) [7,19]. As a consequence, a proton is transferred from the cytoplasmic to the extracellular side of the membrane, creating a protonmotive force.

More recently, a type I rhodopsin (GR) was discovered in the genome of the cyanobacterium *Gloeobacter violaceus* PCC 7421 [20]. Heterologous expression in *E. coli* and reconstitution with retinal generated an absorbance band peaking at ~540 nm [21]. GR also exhibits proton pump activity [21,22]. The photocycles of PR and GR are similar to that of BR [19], but their absorbance bands are strongly blue-shifted from BR (~570 nm).

The length of the π -conjugated polyene chain in the chromophore, along with its interactions with protein residues lining the binding pocket, in combination with the distance between the protonated Schiff base and the counterion charge will determine the energy gap between the ground and excited state of the retinal chromophore. These tuning processes in concert shape and position the absorbance band of the proteorhodopsins.

Abbreviations: BR, bacteriorhodopsin; CCCP, carbonyl cyanide *m*-chlorophenylhydrazide; ChR, channel rhodopsin; DDM, 1-*n*-dodecyl- β -D-maltopyranoside; GR, *Gloeobacter* rhodopsin; Ni-NTA, Ni²⁺-nitrilotriacetate; PR, proteorhodopsin from Monterey Bay SAR 86 gammaproteobacteria; PR-DNFS, PR-D212N/F234S; SRII, sensory rhodopsin II; WT, wild-type; XR, xanthorhodopsin.

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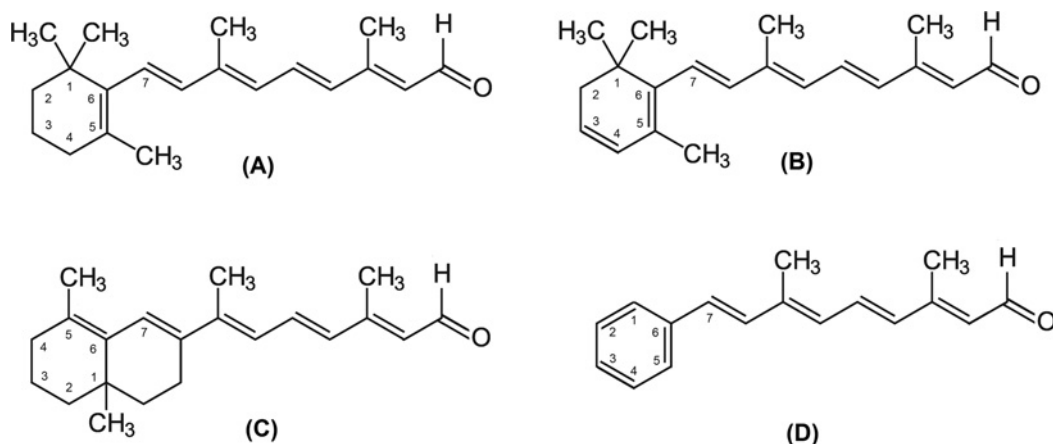


Figure 1 Chemical structures of the all-*trans* isomers of retinal A1 and the analogues used in the present study

(A) Retinal A1, the native ligand of microbial rhodopsins. (B) Retinal A2, which contains an additional double bond between C3 and C4. (C) All-*E*-locked retinal A1, locked in a 6-*s-trans* configuration. (D) Phenyl-retinal A1, containing an aromatized ring lacking the methyl groups.

The *trans*→*cis* photo-isomerization in the chromophore induces proton transfer from the Schiff base to the proton acceptor (Asp⁹⁷ in PR, Asp¹²¹ in GR), thereby triggering a complex mechanism involving proton release at the extracellular surface, proton uptake from the cytoplasmic surface and thermal re-isomerization of the chromophore. The latter processes largely determine proton pump activity and depend not only on the protein machinery, but also on the conformational properties of the ligand. For instance, the electronic distribution in the chromophore will affect the p*K*_a of the Schiff base [23–25], and its structure and fit into the binding pocket will determine the distance to the counterion and to the proton-accepting residue and may have long-range effects on the protein structure.

In the present study, we investigate the effects of three retinal analogues, which differ in electronic and conformational properties, on functional properties of wild-type (WT) PR and GR, expressed in *E. coli*. We further include the red-shifted double mutant PR-D212N/F234S (PR-DNFS) [13] to test whether any spectral shift introduced by an analogue is additive to the mutation-induced shift. The three all-*trans* analogues of the native retinal A1 are retinal A2, phenyl-retinal A1 and 6-*s-trans* locked retinal A1 (Figure 1). Retinal A2 has the same structural make-up as retinal A1, but contains an elongated π -conjugated polyene chain. This analogue has been shown to red-shift the absorbance band of archaeal rhodopsins [26–29] and xanthorhodopsin (XR) [30]. We anticipated that this analogue would have a similar effect on the proteorhodopsins. Phenyl-retinal A1 contains an even more complex conjugated system, but lacks the methyl groups in the ring element. This analogue blue-shifts the absorbance band of BR [31,32] and probably of XR as well [30,33]. In 6-*s-trans*-locked retinal A1, the C6–C7 bond is locked in an *s-trans* configuration. The use of this analogue corroborated evidence that the chromophore in BR and XR contains a 6-*s-trans* configuration [34].

Our results show that the binding pocket constraints differ for PR and GR, as well as that retinal analogues can significantly modulate the spectral properties of proteorhodopsins with at least partial preservation of proton pump activity. Furthermore, we show that in PR-DNFS, the analogue-induced spectral shifts are additive to the mutation-induced shift.

MATERIALS AND METHODS

Materials

Escherichia coli strain UT5600 and the pKJ900 plasmids encoding PR or GR with a C-terminal His₆ tag were a gift from Dr K. Jung (University of Seoul, Seoul, South Korea) [13]. UT5600 was used to express the recombinant proteins. All-*trans*-retinal A1 was obtained from Sigma–Aldrich. The all-*trans* analogues retinal A2 and all-*E*-6-*s-trans* locked retinal A1 were prepared as described previously [34,35]. All-*trans*-phenyl-retinal A1 was synthesized in analogy to retinal A1 and A2, but starting from benzaldehyde, and the structure was confirmed by ¹H-NMR [36]. Sources of special chemicals include: IPTG from Promega, 1-*n*-dodecyl- β -D-maltopyranoside (DDM) from Protein Labelling Innovation, benzonase from Novagen, lysozyme, ampicillin, valinomycin and carbonyl cyanide *m*-chlorophenylhydrazine (CCCP) from Sigma, Ni²⁺-nitrilotriacetate (Ni-NTA) columns, Coomassie Brilliant Blue G-250 protein-staining solution, restriction enzymes, protein ladder and Pfu DNA polymerase from Thermo Scientific, and EDTA-free protease inhibitor tablets from Roche. All other chemicals were of analytical grade.

Site-directed mutagenesis

Site-directed mutagenesis was performed on the PR gene using mismatch PCR. In brief, the pKJ900 plasmid containing PR was isolated from a 5 ml overnight culture using the QIAprep Spin Miniprep Kit (Qiagen). The plasmid was linearized by restriction with Esp3I and subjected to mismatch PCR using overlapping primers containing the corresponding mutation sites for the D212N (DN-fwd, pKJ-rev; pKJ-fwd, DN-rev) and the F234S (FS-fwd, pKJ-rev; pKJ-fwd, FS-rev) mutations. Then, 25 cycles of PCR were run at 95 °C for 30 s, 55 °C for 30 s and 68 °C for 1 min. The mutant gene was further amplified using outside vector primers (pKJ-fwd, pKJ-rev) using the same PCR programme. The sequences of the primers is presented in the Supplementary Online Data. The amplified mutant gene and vector were restricted at HindIII and XbaI sites and run on an agarose gel with 0.5 μ g/ml ethidium bromide. The bands corresponding to the restricted mutant gene and the empty vector were excised, extracted using a

Qiagen gel extraction kit and ligated overnight at 4°C. The ligated plasmid was then transformed into *E. coli* UT5600.

Cell transformation

A single colony of *E. coli* UT5600 was inoculated into 25 ml of LB medium and allowed to grow at 37°C for 4–6 h. The cells were made competent using CaCl_2 as described previously [37]. Plasmid DNA (40 ng), isolated as described above, was added to 200 μl of competent cells and incubated on ice for 30 min. The mixture was heat-shocked for 2 min at 42°C. The cells were allowed to recover after addition of 1 ml of SOC (Life technologies) medium for 30 min at 37°C, to express ampicillin resistance. The cells were then plated on to LB agar plates containing ampicillin (50 $\mu\text{g}/\text{ml}$) and incubated overnight at 37°C.

Bacterial cell culture

The cells were grown in LB medium with ampicillin selection at 30°C in an orbital shaker at 180 rev/min. Overnight cultures were grown from frozen glycerol stocks of transformed cells, which were diluted 1:100 to obtain the working culture. At a cell density corresponding to a OD_{600} of 0.3–0.4, expression of the proteorhodopsin apoprotein (opsin) was induced by the addition of IPTG to a final concentration of 1 mM. The cells were allowed to grow for a further 24 h at 30°C, and then harvested.

Regeneration of proteo-opsin with retinal

Retinal stocks were stored at -80°C in hexane solution, with the exception of phenyl-retinal A1, which was stored in methanol. Spectra are presented in Supplementary Figure S1. The molar absorbance values of all-*trans*-retinal A1 (λ_{max} 368 nm in hexane) and of all-*trans*-retinal A2 (385 nm) were taken as 49000 and 44000 $\text{M}^{-1}\cdot\text{cm}^{-1}$ respectively [38,39]. The molar absorbance values of all-*E*-locked retinal A1 (395 nm) and of all-*trans*-phenyl-retinal A1 (360 nm in methanol) were assumed to be similar to that of all-*trans*-retinal A1. At the time of use, the required aliquot of stock solution was evaporated and the residue redissolved in dimethylformamide (DMF) to obtain a concentration of 1 mM. This solution was then added to a crude cellular lysate or to isolated membrane vesicles containing the opsin to achieve a final retinal concentration of 5–10 μM , and incubated under dim light for up to 60 min at room temperature or, if necessary, subsequently overnight at 4°C.

Preparation and analysis of membrane vesicles

The cells were harvested by centrifugation at 3200 g for 20 min at room temperature, and the pellet was resuspended in an ice-cold solution of 50 mM Tris/HCl and 150 mM NaCl, pH 7 (10 ml per 50 ml of culture). The suspended cells were lysed by sonication at 4°C using a Sonics Vibra-Cell™ sonicator for 10 min (4 s pulses, 5 s pauses, 25 % amplitude) and centrifuged at 4000 g for 15 min at 4°C to remove insoluble material and cellular debris. The resulting supernatant with membrane vesicles was incubated with the selected retinal for 1 h at room temperature and the vesicles were then pelleted by high-speed centrifugation (147 000 g) for 1 h at 4°C. The pellet was resuspended in 150 mM NaCl (2 ml per 50 ml culture). For solubilization of membrane proteins, DDM was added to a final concentration of 2 % (w/v), and the suspension was incubated with shaking for 1 h at 4°C, followed by overnight incubation at 4°C. The insoluble material

was removed by centrifugation at 16 000 g for 20 min at 4°C. The supernatant was used for spectral analysis.

Proteorhodopsin purification

The cell pellet was resuspended in ice-cold lysis buffer (5 ml per 100 ml of culture) containing 20 mM Tris/HCl, 50 mM NaCl, 20 mM imidazole and 0.1 % DDM, pH 7, supplemented with an EDTA-free protease inhibitor tablet, benzonase (4 units/100 ml culture) and lysozyme (4 mg per 100 ml of culture). The suspension was sonicated at 4°C and centrifuged to remove cellular debris as described in the previous section. At this stage, the crude mixture was incubated with the selected retinal for 1 h at room temperature. DDM was then added to a final concentration of 1.5 % (w/v) and the sample was kept rotating overnight at 4°C. The insoluble material was removed by centrifugation at 4000 g for 25 min at 4°C and the resulting supernatant was utilized as a crude extract. For purification of the His₆-tagged proteorhodopsins, immobilized metal-ion-affinity chromatography (IMAC) was exploited using 0.4 ml of Ni-NTA resin per 100 ml of original culture volume. The resin was contained in a spin column and first equilibrated with buffer A (20 mM Bis-Tris propane, 0.5 M NaCl and 0.1 % DDM, pH 8) containing 20 mM imidazole. The crude extract was then allowed to equilibrate with the column for 15 min at room temperature. The column was washed five times with five column volumes of buffer A containing 50 mM imidazole at room temperature. Finally, strongly bound protein was eluted using buffer A containing 250 mM imidazole and 0.01 % DDM at room temperature. Fractions of 0.3 ml were collected. Fractions containing the purified proteorhodopsin were combined and analysed by spectroscopy and SDS/PAGE. Thus purified proteorhodopsin could be stored at 4°C for several weeks, but was kept at -80°C for long-term storage.

Spectroscopy

The spectral properties of all samples were measured using a Shimadzu UV-Vis spectrophotometer (UV-1601). In order to test the pH-dependence of the main absorbance band of the proteorhodopsins [9,13], solubilized membrane vesicles were analysed at different pH values by diluting the samples 1:1 with buffers containing either 100 mM Bis-Tris propane at pH 6.5 or 9.5, or 20 mM MES at pH 5. To isolate the major absorbance band of the proteorhodopsin out of the composite spectrum of membrane vesicles, hydroxylamine was added from a 1 M stock solution, pH 7, to a final concentration of 50 mM, followed by incubation at room temperature under ambient light. Hydroxylamine attacks the Schiff base and releases the retinal from the opsin-binding pocket as retinaloxime. A difference spectrum then reveals the major absorbance band of the proteorhodopsin present. Absorbance maxima were determined using the internal peak-pick function of the software UVProbe.

SDS/PAGE

The crude and purified protein fractions were analysed by SDS/PAGE [40]. In brief, aliquots of the samples were diluted with SDS sample buffer, incubated for 30 min at 37°C, and run on a 12.5 % polyacrylamide gel at 20 mA for 2 h. The gel was stained using Coomassie Brilliant Blue G for 2 h according to the manufacturer's instructions and destained overnight using MilliQ. Gel images were obtained using the Bio-Rad GS-800 gel imaging dock.

Starvation of cells

Overnight cultures were diluted 1:100 to obtain a working culture of 25 ml. At a cell density corresponding to an OD_{600} of 0.3–0.4, production of the proteorhodopsin was induced by the addition of IPTG to a final concentration of 1 mM and the selected retinal to a final concentration of 5 μ M. The cultures were allowed to grow for a further 24 h at 30 °C in the dark, and then harvested. The cells were washed twice in starvation buffer (250 mM KCl, 10 mM NaCl, 10 mM $MgSO_4$, 100 μ M $CaCl_2$ and 10 mM Tris/HCl, pH 7) and were starved by incubation with continuous mixing for 4 days at room temperature. The cells were washed another three times and resuspended in 5 ml of starvation buffer supplemented with a 40 μ M final concentration of valinomycin. The cell suspension was incubated for 30 min in the dark at room temperature.

Proton-pumping assay

The cell suspension was illuminated through a bandpass filter (BG-18, Schott) using a halogen light source with fibre optics (Euromex, LE.5211). An illumination level of 40 μ einstein·m⁻²·s⁻¹ was used throughout. Light-induced pH changes were measured with a pH microelectrode (SenTix[®] MIC, WTW) and the readout was monitored by a pH meter (Inolab pH 7310, WTW) fed into a computer. The following light regime was used: 1 min dark, 1 min light, 2 min dark, 1 min light, 2 min dark. Pumping rates were calculated for 5 ml of the cell suspension using two independent trials. Finally, CCCP was added to a final concentration of 40 μ M and the suspension was incubated at room temperature for 30 min in the dark. Light-induced pH changes were then measured again using the same light regime. Pumping rates were calculated as protons/s from the initial rate of the light-induced pH change, corrected if required for baseline drift in controls (starved cells without expression of proteo-opsin or without retinal). Molecular pumping rates could subsequently be calculated after assay of the proteorhodopsin level (see below).

Determination of proteorhodopsin levels for the proton-pumping assay

The above cell suspension from the proton-pumping assay was rinsed with starvation buffer and the pellet was resuspended in 10 ml of buffer B (10 mM Tris/HCl and 150 mM NaCl, pH 7). The cell suspension was sonicated as described above and the membrane vesicles and cell debris were pelleted together at 147 000 *g* for 1 h at 4 °C. The pellet was resuspended in 1.5 ml of 150 mM NaCl. DDM was added to a final concentration of 2.5 % and incubated at room temperature with mixing overnight. Under these conditions, maximal extraction of proteorhodopsin was achieved without significant losses. The following day, the insoluble material was removed by centrifugation at 16 000 *g* for 20 min at 4 °C. The supernatant was used to measure an absorbance spectrum before and after bleaching with hydroxylamine as described above. The optical density value at the absorbance maximum was used to calculate the original proteorhodopsin level in the cell suspension using a molar absorbance of 45 000 M⁻¹·cm⁻¹ [12].

Construction of homology models

A homology model for PR was built using the structure of sensory rhodopsin II (SRII) as a template (PDB code 1H2S) [41] with which it has the highest sequence identity out of the retinal proteins for which structural information was available at

that time (24 % sequence identity). For GR, we built a model using the structure of XR as a template (PDB code 3DDL, 56 % sequence identity) [42]. Model building and subsequent analysis was performed using the WHAT IF [43] and YASARA [44] twinset with standard parameters.

RESULTS

Reconstitution of proteorhodopsins with retinal analogues

Production of proteorhodopsins following induction with IPTG and simultaneous addition of retinal A1 to the correspondingly transformed UT5600 cells could be easily verified by the red coloration of the pelleted cells or membrane vesicle preparation. However, this requires substantial amounts of retinal (up to 0.5 mg per 100 ml of culture) and in view of the limited quantities of retinal analogues available, we investigated other options. It eventually turned out that the apoproteins (proteo-opsins) produced *in vivo* would still readily bind retinal A1 after harvesting the cells and preparation of membrane vesicles, thereby generating the corresponding holoproteins (proteorhodopsins). We observed that incubation of these vesicles with retinal A1 resulted in at least the same yield of proteorhodopsin as when retinal A1 was supplied to the cell culture, but required 5–10-fold less retinal. Incorporation of retinal A1 as well as the analogues could be detected visually from the development of a red or purple colour (Supplementary Figure S2), and was subsequently characterized by spectroscopy after solubilization in 2 % (w/v) DDM (Figure 2A). Since the spectra of the vesicles also have contributions from other membrane components (cytochromes, excess retinal), the specific absorbance band of the proteorhodopsin was isolated by incubation with 50 mM hydroxylamine (Figure 2B), as detailed in the Supplementary Online Data.

This procedure works well for all proteorhodopsins tested (WT PR, its double mutant PR-DNFS and GR). We observed that the PRs were much more sensitive to this ‘bleaching’ with hydroxylamine than GR, which may be related to the slower photocycle kinetics of PR [19,21,45]. The thus obtained absorbance maxima with retinal A1 at neutral pH for PR, PR-DNFS and GR lie at 526 nm, 545 nm and 540 nm respectively, with an accuracy of ± 3 nm ($n=3$) (Figure 2B). These values are in good agreement with those reported in the literature [7,13,17,46].

Upon incubation of the three proteo-opsin species with the retinal analogues, we observed that all three analogues regenerate the PR opsins with distinct absorbance maxima (Figure 2), whereas only two of the analogues regenerate the GR opsin (see Figure 4). Rapid incorporation of retinal A2 and all-*E*-locked retinal A1 was noticed, being completed within 30–60 min at room temperature. Incorporation of phenyl-retinal A1 was much slower, requiring additional overnight incubation at 4 °C. Incorporation of retinal A2 induces an appreciable red-shift of approximately 30 nm relative to retinal A1 in the visible absorbance band of all three proteorhodopsins. Incorporation of phenyl-retinal A1 on the other hand causes an appreciable blue-shift of approximately 30 nm in the PR species, whereas it does not seem to generate a stable pigment with GR. This was further corroborated by the lack of a distinct negative peak after treatment of GR-opsin vesicles, incubated with phenyl-retinal A1, with hydroxylamine (results not shown). All-*E*-locked retinal A1 can be incorporated in all three opsins and produces only a small red-shift in the absorbance band, relative to retinal A1.

The position of the absorbance band of PR and reported mutants is pH-dependent [9,13,47] probably because of protonation of

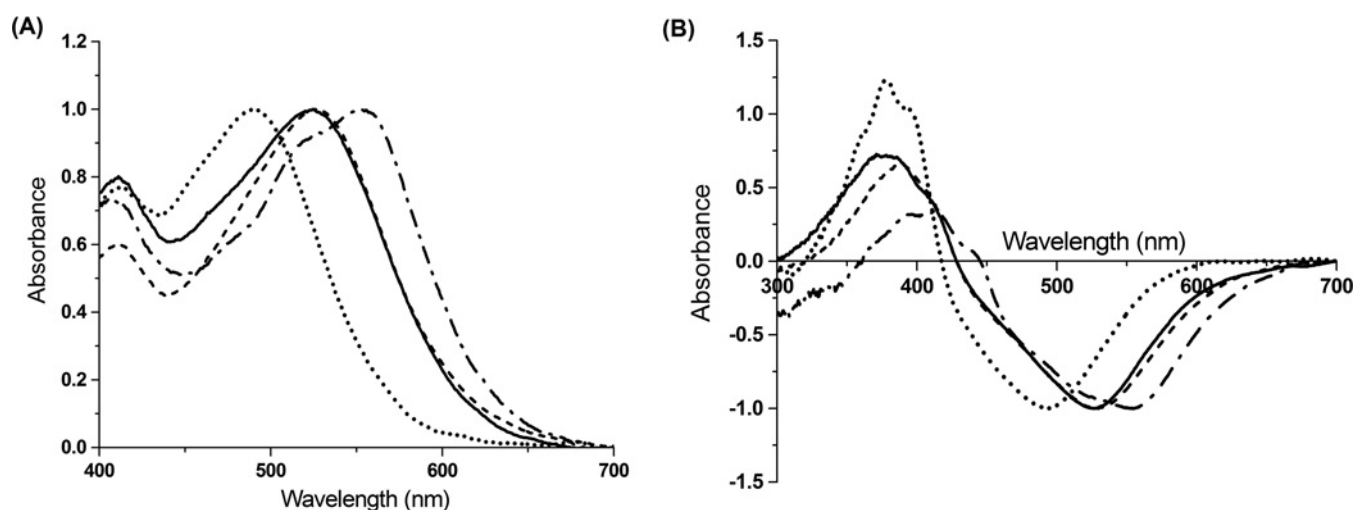


Figure 2 Spectral analysis of solubilized membrane vesicles containing PR reconstituted with the various retinals

(A) Absorbance spectra. Accuracy of the $\lambda_{\max}$ values ± 5 nm. (B) Difference spectra obtained after incubation of the solubilized membrane vesicles with 50 mM hydroxylamine. The absorbance bands of the proteorhodopsins are seen here as negative peaks, whereas the positive peaks between 360 and 400 nm are due to the formation of the retinaloximes. Accuracy of the $\lambda_{\max}$ values was ± 3 nm. —, PR with retinal A1, absorbance maximum at ~ 520 nm (A) and 526 nm (B); - - -, PR with all-*E*-locked retinal A1, absorbance maximum at ~ 520 nm (A) and 529 nm (B). PR with phenyl-retinal A1, absorbance maximum at ~ 490 nm (A) and 494 nm (B).

the proton-acceptor residue Asp⁹⁷ [12,48] which induces a red-shift of 25–30 nm. The corresponding pK_a values for PR and PR-DNFS are reported to be 7.3 and 8.0 respectively [13]. Such a strong pH-dependence has not yet been observed for GR [21,22,49]. We were interested in whether the analogue pigments would show a similar behaviour, and recorded the absorbance spectra of the corresponding solubilized membrane vesicles at pH 5, 6.5 and 9.5 (Supplementary Table S1). All PR-based analogue pigments showed a clear pH-dependence of their visible absorbance band. The largest red-shift upon going from pH 9.5 to 5 of approximately 25 nm was observed for the all-*E*-locked retinal A1 analogue of PR. The magnitude of the shift between pH 9.5 and 5 will also depend on the pK_a value. An accurate determination of this pK_a in various PR-based analogue pigments is the subject of a follow-up study and outside the scope of the present study. On the other hand, we did not observe a significant pH effect in this range on the absorbance profile of GR or of the GR-based analogue pigments.

Purification of the proteorhodopsins

Following their characterization in membrane vesicles, we aimed for optimal spectral characterization of the native and analogue pigments by purification over an Ni-NTA resin exploiting their C-terminal His₆ tag. According to the ratio A_{280}/A_{5xx} in the absorbance spectra, which varied between 1.5 and 2.5 for all pigments, and the strong band at ~ 25 kDa for PR (Supplementary Figure S3) and at ~ 27 kDa for GR (results not shown) observed upon SDS/PAGE analysis, we conclude that a high degree of purification was achieved for all pigments. The purified fractions remained spectrally stable for several weeks at 4°C.

Purified PR, PR-DNFS and GR show $\lambda_{\max}$ values of 522 nm, 540 nm and 543 nm with retinal A1 respectively (Figures 3 and 4, and Supplementary Figure S4), in excellent agreement with literature data [13,46] and in close agreement with the vesicle data. Incorporation of retinal A2 induces a large red-shift of approximately 30 nm in the specific absorbance band

of all three proteorhodopsins to $\lambda_{\max}$ values of 554 nm, 568 nm and 575 nm for PR, PR-DNFS and GR respectively. In contrast, phenyl-retinal A1 induces a blue-shift of at least 30 nm in PR and PR-DNFS to 493 nm and 501 nm respectively. No specific absorbance band was seen for GR purified after incubation with this analogue. SDS/PAGE analysis revealed that the opsin was purified successfully, which strengthens our conclusion that the GR opsin is not able to stably bind phenyl-retinal A1, or this analogue pigment is not stable under our experimental conditions. Incorporation of all-*E*-locked retinal A1 induces a small red-shift of approximately 5 nm in all analogue pigments to 526 nm, 545 nm and 548 nm for PR, PR-DNFS and GR respectively. These data are compiled in Table 1.

Proton pump activity of reconstituted proteorhodopsins

Although membrane vesicles are more economical with respect to ligand usage, we find them less suitable for a quantitative proton-pumping assay. We were not able to isolate vesicle preparations with largely right-side-out- or inside-out-oriented proteorhodopsin. In addition, isolated vesicles can be more or less leaky. Hence we opted to use intact *E. coli* cells instead. However, the activity of proteorhodopsins is not easily reproducibly assayed in viable *E. coli* cells. The proton electromotive force is used to drive metabolic processes (e.g. ATP synthesis by ATP synthase) and may trigger opening of voltage-gated proton channels in the *E. coli* membrane [9]. To avoid such complications, the cells were starved for a few days, with the most reproducible results obtained after 4–5 days of starvation. The proton pump activity of the native proteorhodopsins and their analogue pigments was measured in starved cell suspensions in the presence of the K⁺ ionophore valinomycin and K⁺ ions, which eliminates the electrical component of the electrochemical proton gradient. This allows quantification of the proton efflux mediated by the proteorhodopsins, without interference by any possible back-pressure effects of the transmembrane electrical potential gradient. For the cells producing proteorhodopsin, this was seen

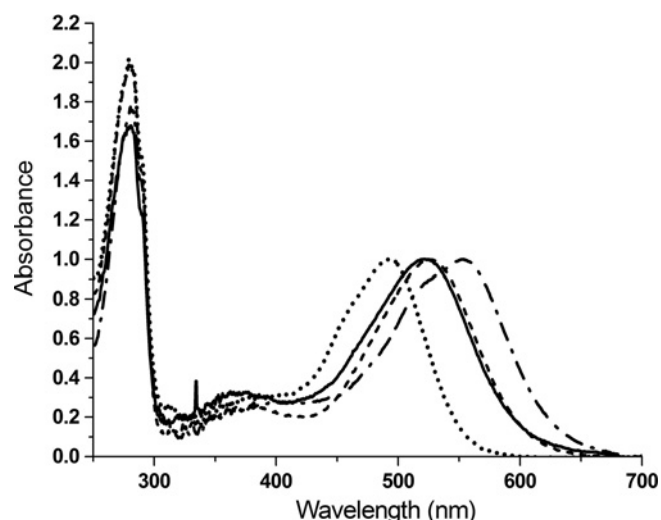


Figure 3 Absorbance spectra of purified PR and its analogues

The absorbance at 280 nm largely arises from the aromatic residues of the protein. The A_{280}/A_{500} ratios are all <2 , indicating a high degree of purity. Spectra were taken at pH 8. Accuracy of the $\lambda_{\max}$ values was ± 2 nm. —, PR with retinal A1, absorbance maximum at 522 nm; ---, PR with retinal A2, absorbance maximum at 554 nm, with a shoulder at ~ 520 nm; PR with all-*E*-locked retinal A1, absorbance maximum at 526 nm; -.-.-, PR with phenyl-retinal A1, absorbance maximum at 493 nm.

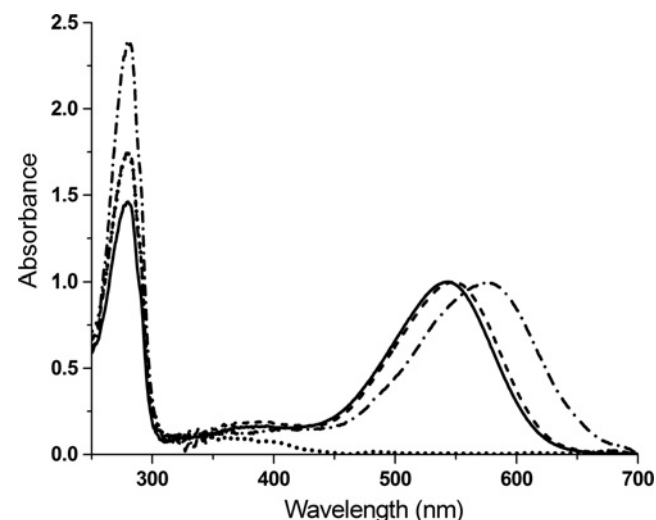


Figure 4 Absorbance spectra of purified GR and its analogues

The A_{280}/A_{500} ratios are all <2.5 . Spectra were taken at pH 8. Accuracy of the $\lambda_{\max}$ values was ± 2 nm. No specific peak around 500 nm was obtained with the phenyl-retinal A1 analogue. —, GR with retinal A1, absorbance maximum at 543 nm; ---, GR with retinal A2, absorbance maximum at 575 nm, with a hint of a shoulder near 540 nm; GR with all-*E*-locked retinal A1, absorbance maximum at 548 nm; -.-.-, GR with phenyl-retinal A1, only a protein peak near 280 nm was observed.

as a decrease in pH upon illumination, which slowly returned to the baseline value in darkness. We did not try to optimize conditions to achieve maximal pumping rates (spectral range, photon intensity, cell density), since this is outside the scope of the present study. Instead, for mutual comparison, the calculated rates were normalized relative to the most active A1 rhodopsin (GR).

Representative traces for PR and its three analogue pigments are shown in Figure 5, and the normalized pumping rates in Table 1. The highest pumping rates obtained with the native

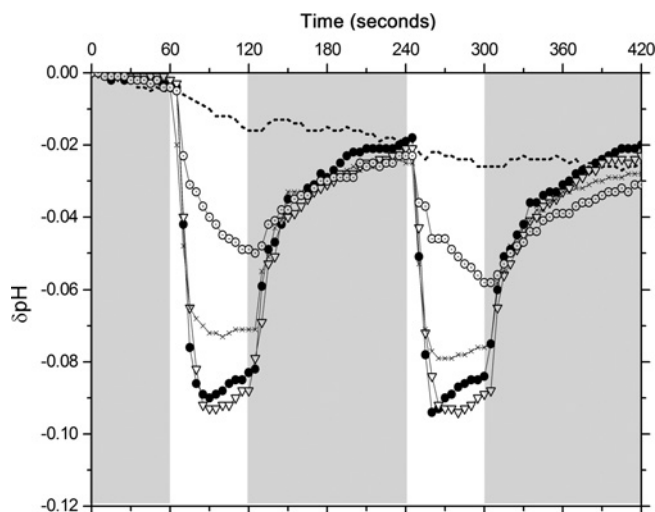


Figure 5 Proton-pumping traces of starved *E. coli* UT5600 cell suspensions containing PR or one of its analogues

pH readings were acquired every 5 s after an equilibration period of 1 min, followed by illumination (white area) for 1 min, and darkness (grey area) for 2 min. The levels of the various pigments varied in the experiments. Hence the calculated initial rates of the traces were divided by the pigment concentration to arrive at molecular pumping rates., control *E. coli* UT5600 cells containing the proteo-opsin without retinal; —●—, *E. coli* UT5600 cells containing PR with retinal A1; ---△---, *E. coli* UT5600 cells containing PR with retinal A2;×....., *E. coli* UT5600 cells containing PR with all-*E*-locked retinal A1; —◇—, *E. coli* UT5600 cells containing PR with phenyl-retinal A1.

ligand retinal A1 were observed for GR (4–5 protons $\cdot$ mol $^{-1}$ $\cdot$ s $^{-1}$), which corroborates findings from previous studies [21,22,50]. The lowest pumping activity of the three proteorhodopsin species was observed for the PR-DNFS mutant, which retains 40–60 % pumping activity of the WT. This is consistent with previous data using sphaeroplast suspensions [13]. Interestingly, with retinal A2 or all-*E*-locked retinal A1 as the chromophore, all three proteorhodopsins presented proton pump activity very similar to that with retinal A1, indicating that these analogues do not perturb the proton translocation mechanism. Also, the phenyl-retinal A1 analogue retains significant proton pump activity for both PR and PR-DNFS.

All light-induced pH changes were largely prevented by the addition of 40 μ M protonophorous uncoupler CCCP. In addition, no light-induced pH changes were observed for the control situation, where the opsin expression was not induced but retinal was present (results not shown), or the converse, i.e. when the opsin was expressed but retinal not added (Figure 5). Taken together, this indicates that the measured pumping rates are a result of the light-induced proton efflux mediated by the respective proteorhodopsins.

Homology models of PR and GR

In order to evaluate the effect of retinal analogues and mutations, a 3D structure of PR and GR would be very useful. However, a crystal structure is not available for these retinal proteins and the PR structure solved using NMR spectroscopy [51] significantly deviates even in the transmembrane domain from the crystal structures of other microbial retinal proteins [18,52] and from earlier PR homology models based on the BR structure [53,54]. Hence we decided to generate homology models for GR and PR, based on the crystal structures of retinal proteins with which they had the highest sequence identity at that time (XR and

Table 1 Pumping scores and absorbance maximum values of PR, PR-DNFS and GR and their analogues

GR does not form a stable pigment with phenyl-retinal A1 (indicated by –). The λ_{max} values of the purified proteins were measured at pH 8. Accuracy of the λ_{max} values was ± 2 nm. The spectral shift relative to A1 is given in wavelength (nm) and energy (cm^{-1}). Proton-pumping rates were normalized with respect to GR with retinal A1. + + +, pumping rate 70–120% of that of A1-GR; + +, pumping rate 40–70% of that of A1-GR; +, pumping rate 20–40% of that of A1-GR.

Retinal analogue	PR			PR-DNFS			GR		
	λ_{max} (nm)	Spectral shift (nm)	Spectral shift (cm^{-1})	Pump rate	λ_{max} (nm)	Spectral shift (nm)	Spectral shift (cm^{-1})	Pump rate	λ_{max} (nm)
A1	522	–	–	+	540	–	–	+	543
A2	554	+ 32	1106	+	568	+ 28	913	+	575
Locked	526	+ 4	146	+	545	+ 5	170	+	548
Phenyl-A1	493	– 29	1127	+	501	– 39	1442	+	–

SRII respectively). The resulting models for PR and GR show high similarity in the structure of the transmembrane domain and the binding pocket, but, as expected, show variation in the loop regions. Recently, a crystal structure was published for a blue-PR having 83 % sequence identity with PR [18] (PDB code 4KNF). A comparison of this structure with our homology model for PR, presented in Figure 6(A), shows excellent conformity in the α -helical domain, and a RMSD of only 1.8 Å (1 Å = 0.1 nm) over all identical residues.

DISCUSSION

Proteorhodopsin selection

PR and, to a lesser extent, GR have been extensively characterized with their native ligand all-*trans*-retinal A1. We obtained good expression levels in the *E. coli* strain UT5600 (at least 10 mg/l for PR and 4 mg/l for GR). The proteorhodopsins produced could be purified to a high degree. Furthermore, we obtained good agreement between the λ_{max} values obtained from the absorbance spectra of the purified protein, and the difference spectrum obtained after bleaching the solubilized membrane vesicle fraction with hydroxylamine.

The spectral characteristics of a retinylidene chromophore are strongly affected by its interaction with protein residues lining its binding pocket. However, residues can also exert their effect on the chromophore from a distance, via their involvement in the secondary structure of the protein or their contribution to the local electrostatic field. This is exemplified in the D212N/F234S double mutant of PR, which has been described previously [13]. The authors report that mutation of the binding pocket residue Phe²³⁴ to serine causes a red-shift of the main absorbance band and strongly reduces proton pump activity. According to our homology model, Asp²¹² is located in interhelical loop E-3 and it is not surprising that the single mutation D212N does not change the absorbance or pumping activity of PR [13] (results not shown). However, when these mutations are combined, this yields a red-shifted PR-DNFS that retains significant proton pump activity. We could fully reproduce these characteristics and included this double mutant in the present study to test whether spectral shifts induced by a mutation are complementary to those implemented by retinal analogues. With PR-DNFS, expression yields of at least 6 mg/l of UT5600 culture were obtained.

Analogue pigments

To our knowledge, no detailed study has been reported to date on the effect of retinal analogues on molecular properties of PR or GR. In the context of a wide-angle scattering study of PR the 13-desmethyl,13-iodo-retinal A1 analogue was tested and shown to induce a 23 nm red-shift [54]. In the present study, we focus on ring modifications.

Incorporation of retinal A2 is expected to red-shift the absorbance band relative to bound retinal A1 [38,55] (Supplementary Online Data). This has been demonstrated previously in archaeal and channel rhodopsins (ChRs), where red-shifts of approximately 30 nm are reported, with only a small effect on proton pumping or channel functionality [26–29], as well as in XR with a 23 nm red-shift [30]. As anticipated, this analogue reacted rapidly to completeness with all three proteo-opsins within 30 min at room temperature (results not shown), and induced comparable red-shifts of approximately 30 nm corresponding to an energy difference of approximately 1100 cm^{-1} (Table 1). The resulting A2 analogue pigments were quite stable in detergent

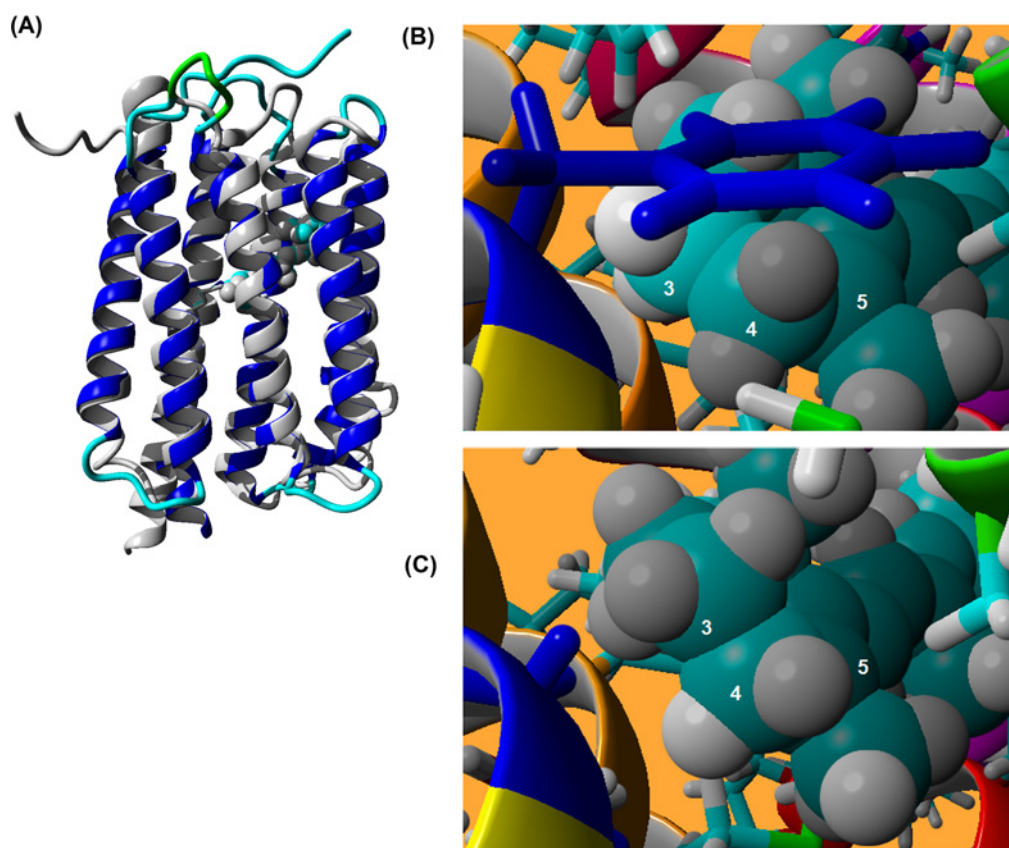


Figure 6 Homology models of PR and GR

(A) Overlay of the backbone of the PR homology model (blue) with that of the recently published blue-light-absorbing PR structure (grey) [23] (PDB code 4KNF). The carbon atoms of the retinal moiety are represented by the small spheres with the ring element at the upper right. The image was generated with YASARA (<http://www.yasara.org>). (B and C) Rear view of the binding pocket in the (B) PR and (C) GR homology models. In PR, the dark blue residue represents Phe¹⁵² which is situated above the ring of retinal A1 (carbons as blue-green spheres; some ring carbons are numbered). In the case of GR, this position contains the smallest possible side chain (Gly¹⁷⁸; dark blue).

solution and could be purified to a high degree (Figures 3 and 4, and Supplementary Figures S3 and S4).

The A2 analogue pigments also present a slightly broadened absorbance band with a small shoulder at the hypsochromic wing of the spectrum (at approximately 510 nm for PR and approximately 530 nm for the other two analogue pigments). Such a shoulder was observed previously in the A2 analogues of BR [26], ChR and archaerhodopsin 3 (AR3) [29]. It is most likely that this shoulder represents fine structure of vibronic origin, possibly due to a more rigid conformation [33], e.g. also evident in the spectrum of all-*E*-locked retinal A1 in Supplementary Figure S1. An important observation is that retinal A2 is fully equivalent to retinal A1 in maintaining proton pump activity in all three proteorhodopsins (Figure 5 and Table 1).

In all-*E*-locked retinal A1, the polyene system is locked in the 6-*s-trans* configuration effectuating optimal conjugation. This analogue was previously tested on bacterio-opsin and xantho-opsin, to study the effect of the configuration about the C6–C7 single bond [35]. It was observed that the 6-*s-trans* form of this compound smoothly reacted with both opsins, but the 6-*s-cis* form did not. The 6-*s-trans* analogue provided spectral properties very similar to those of native BR and XR, and retained 90% of the pumping activity of BR. This corroborated previous solid-state NMR studies indicating that retinal A1 has taken up a 6-*s-trans* configuration upon binding to bacterio-opsin [56]. The transition from the twisted 6-*s-cis* in the free retinal A1 to the planar 6-

s-trans form in the binding pocket of the protein accounts for a part of the spectral shift in the binding site. In all-*E*-locked retinal A1, the 6-*s-trans* form has already been enforced, which explains its red-shifted absorbance band relative to free retinal A1 (Supplementary Figure S1) and its small effect on the absorbance band when bound in BR or XR. The all-*E*-locked retinal A1 was not tested previously on PR or GR. This analogue reacted with all three proteo-opsins, reaching completion within 30–60 min at room temperature. The resulting analogue pigments exhibit only small red-shifts relative to the native pigments (4–5 nm). These results provide strong evidence that, in bacterial rhodopsins, the retinal ligand is bound in the 6-*s-trans* configuration as well. This implies that the bathochromic shift of the absorbance band in XR and archaeal rhodopsins relative to these bacterial rhodopsins is largely induced by the protein environment. All three proteorhodopsin analogues containing all-*E*-locked retinal A1 retained strong proton pump activity. This would render this retinal analogue interesting for optogenetic applications. It can be supplied directly or in precursor form in the diet and will hardly interfere with the visual system. In addition, it may be less prone to generate metabolic products such as retinoic acids with strong signalling function.

Phenyl-retinal A1 was previously reported to react very slowly with bacterio-opsin, yielding a strongly blue-shifted analogue pigment [31,32]. In the case of the bacterial opsins, phenyl-retinal A1 behaves quite differently from the other two analogues.

It forms a stable pigment with the PR and PR-DNFS opsins, but not with the GR opsin. Furthermore, it regenerates the PR opsins slowly, with only partial regeneration after 60 min of incubation at room temperature, requiring additional overnight incubation at 4°C. Finally, it induced a significant blue-shift in absorbance, compared with retinal A1. The slow regeneration is most likely to be due to a poor fit in the opsin-binding pocket because of the absence of the methyl groups on the ring element, which contribute sterically to correct positioning and stabilization of the ring and to its interaction with its protein environment [33,57]. Nevertheless, the resulting analogue pigments maintained significant proton pump activity and could be purified successfully.

The structural models we generated for GR and PR may present a clue to understanding their different reactivity with phenyl-retinal A1. Figure 6(B) shows the models for the rear end of the binding pockets. The most striking difference is seen at the position of Phe¹⁵² in PR, where GR has a glycine residue (Gly¹⁷⁸). This difference is physiologically relevant, since it was demonstrated that mutating Gly¹⁷⁸ in GR to a bulkier tryptophan residue abolished strong binding of a carotenoid (echinenone), which is involved in energy transfer to the retinal [21,58,59], and eliminated this energy transfer. The smaller glycine residue at position 178 apparently is required to allow enough space for immobilizing the carotenoid ring [33,42], thereby allowing energy transfer and increasing the cross-section for photo-activation of retinal. We surmise that the aromatic ring of Phe¹⁵² in PR, which is positioned right above the ring element of retinal, would contribute significant π - π stacking interaction. Together with the methyl groups on the polyene chain, this may sufficiently stabilize this analogue in the binding pocket. GR cannot provide this stacking interaction, and thus may not provide sufficient interaction energy or too much motional freedom for the phenyl-retinal A1 to form a stable protonated Schiff base. We will verify this hypothesis by testing whether the G178F mutant of GR will yield a stable pigment with phenyl-retinal A1.

In this context, a comparison with XR, a distant relative of GR, is appropriate, since XR also binds a carotenoid (salinixanthin, a derivative of echinenone) [60]. The position equivalent to Gly¹⁷⁸ in GR carries the same residue in XR (Gly¹⁵⁶). Xanthopsin is reported to react slowly with phenyl-retinal A1 in the presence of bound salinixanthin, generating a stable pigment [30]. Possibly, the presence of the carotenoid ring sufficiently stabilizes phenyl-retinal A1 in the binding pocket. Since there is no echinenone available in our *E. coli* expression system, it needs to be investigated whether complementation with echinenone would result in a stable analogue pigment of GR with phenyl-retinal A1. It has been suggested that carotenoid fixation is required for pigment formation in XR [33,61]. Our results show that this is not the case for GR, which easily generates analogue pigments that retain full proton-pumping capacity.

Conclusions and future prospects

Our results demonstrate that ring modification can affect the affinity of the analogue for the retinal-binding site and will usually modulate the spectral properties of the respective rhodopsin, but, importantly, can largely maintain proton pump activity. This presents good prospects for further modification trials in the ring and eventual biotechnological applications. The smooth reaction with all-*E*-6-*s-trans*-locked retinal while implementing only a small red-shift corroborates available evidence that the retinal chromophore in bacterial rhodopsins is also bound in the 6-*s-trans* configuration. We show further that the spectral shifts effected

by the analogues can be additive to spectral shifts induced by mutagenesis. Hence combining selected mutagenesis with ligand analogues offers promising prospects for further extending the wavelength range of rhodopsins.

Retinal analogues also present attractive prospects for application in the field of optogenetics. Electrogenic microbial rhodopsins (ion pumps and ChRs) are widely used to modulate the activity of neurons and other cells by light [62,63]. Chromophore substitution provides a complementary strategy to improve the flexibility of these optogenetic tools, e.g. allowing multiphoton excitation or improved light penetration in biological tissues. For instance, supplementation with all-*E*-6-*s-trans*-locked retinal A2 would allow the red-shifting of microbial rhodopsins, probably, as argued above for its A1 counterpart, without significant interference with the visual system and with signalling metabolites of retinal A1.

The non-pumping mutant PR-D97N was recently engineered as a sensor of the membrane potential in bacteria [64]. Spectral modulation of such sensors using chromophore substitution towards specific spectral ranges would have widespread electrophysiological applications in systems that are difficult or too small to study with traditional methods (bacteria, neuronal compartments, mitochondria or other cellular organelles).

AUTHOR CONTRIBUTION

Srividya Ganapathy and Willem de Grip designed the study and wrote the paper. Srividya Ganapathy, Odette Bêcheau and Siebren Frölich tested retinal analogues on the proteorhodopsin species. Hanka Venselaar constructed the homology models. Sarah Radwan contributed to the characterization of proteorhodopsins in *E. coli*. Johan Lugtenburg synthesized the retinal analogues. Jeroen van der Steen, Que Chen, Klaas Hellingwerf and Huub de Groot contributed towards the design of experiments, and the writing and discussion of the paper.

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