Conductance Mechanisms of Rapidly Desensitizing Cation Channelrhodopsins from Cryptophyte Algae

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ABSTRACT

Channelrhodopsins guide algal phototaxis and are widely used as optogenetic probes for control of membrane potential with light. "Bacteriorhodopsin-like" cation channelrhodopsins (BCCRs) from cryptophytes differ in primary structure from other CCRs, lacking usual residues important for their cation conductance. Instead, the sequences of BCCR match more closely those of rhodopsin proton pumps, containing residues responsible for critical proton transfer reactions. We report 19 new BCCRs which, together with the earlier 6 known members of this family, form three branches (subfamilies) of a phylogenetic tree. Here, we show that the conductance mechanisms in two subfamilies differ with respect to involvement of the homolog of the proton donor in rhodopsin pumps. Two BCCRs from the genus Rhodomonas generate photocurrents that rapidly desensitize under continuous illumination. Using a combination of patch clamp electrophysiology, absorption, Raman spectroscopy, and flash photolysis, we found that the desensitization is due to rapid accumulation of a long-lived nonconducting intermediate of the photocycle with unusually blue-shifted absorption with a maximum at 330 nm. These observations reveal diversity within the BCCR family and contribute to deeper understanding of their independently evolved cation channel function.

IMPORTANCE

Cation channelrhodopsins, light-gated channels from flagellate green algae, are extensively used as optogenetic photoactivators of neurons in research and recently have progressed to clinical trials for vision restoration. However, the molecular mechanisms of their photoactivation remain poorly understood. We recently identified cryptophyte cation channelrhodopsins, structurally different from those of green algae, which have separately evolved to converge on light-gated cation conductance. This study reveals diversity within this new protein family and describes a subclade with unusually rapid desensitization that results in short transient photocurrents in continuous light. Such transient currents have not been observed in the green algae channelrhodopsins and are potentially useful in optogenetic protocols. Kinetic UV-visible (UV-vis) spectroscopy and photoelectrophysiology reveal that the desensitization is caused by rapid accumulation of a nonconductive photo-intermediate in the photochemical reaction cycle. The absorption maximum of the intermediate is 330 nm, the shortest wavelength reported in any rhodopsin, indicating a novel chromophore structure.

KEYWORDS

channelrhodopsins, ion channels, optogenetics, patch clamp, photobiology

Citation

Sineshchekov OA, Govorunova EG, Li H, Wang Y, Melkonian M, Wong G-K-S, Brown LS, Spudich JL. 2020. Conductance mechanisms of rapidly desensitizing cation channelrhodopsins from cryptophyte algae. mBio 11:e00657-20. https://doi.org/10.1128/mBio.00657-20.

Editor

Steven J. Norris, McGovern Medical School

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This article is a direct contribution from John L. Spudich, a Fellow of the American Academy of Microbiology, who arranged for and secured reviews by Sergei Balashov, University of California, Irvine, and Kevin Gardner, CUNY.

Received 19 March 2020
Accepted 26 March 2020
Published 21 April 2020
Channelrhodopsins are light-gated channels first discovered in green (chlorophyte) flagellate algae, in which they serve as photoreceptors mediating phototaxis by depolarization of the cell membrane (1–3). Currently, channelrhodopsins are widely used for control of neurons and other excitable cells with light (“optogenetics”) (4) for research and also in clinical trials to restore vision to the blind (5). Channelrhodopsins from chlorophyte algae conduct cations and are therefore referred to as cation channelrhodopsins (CCRs). Anion-conducting channelrhodopsins (ACRs) have been found in the phylogenetically distant cryptophyte algae (6), and a second family has been found more recently in environmental DNA samples of unidentified origin (7). These three channelrhodopsin families share ~50% of overall sequence homology, including several key residues shown to be required for their channel activity.

However, cryptophyte genomes also encode a family of microbial rhodopsins that show a higher sequence homology to haloarchaeal proton-pumping rhodopsins than to any known channelrhodopsins, and yet exhibit cation channel activity, apparently a product of convergent evolution (8–10). In particular, these proteins contain homologs of the two carboxylate residues that serve as the Schiff base proton acceptor and donor in Halobacterium salinarum bacteriorhodopsin (Asp85 and Asp96, respectively), which together with the Thr89 homolog form the “DTD” motif characteristic of proton pumps. In contrast, in all other known channelrhodopsins, one or both of these positions are occupied by noncarboxylate residues.

Earlier we have shown that channel activity in CCR1 and CCR2 from the cryptophyte alga Guillardia theta (GtCCR1 and GtCCR2) is mechanistically distinct from that in chlorophyte CCRs (9). According to our model, channel opening in these proteins requires deprotonation of the Asp96 homolog and occurs >10-fold faster than reprotonation of the retinyliden Schiff base. The latter is achieved by return of the proton from the earlier protonated acceptor, thus preventing vectorial proton translocation across the membrane. To emphasize their distinction from other known CCRs, we named these proteins “bacteriorhodopsin-like cation channelrhodopsins” (BCCRs) (9).

Besides their fundamental importance as independently evolved light-gated cation channels, BCCRs have attracted attention as optogenetic tools, because some of them exhibit more red-shifted absorption, enabling use of deeper-penetrating long-wavelength light and a higher Na⁺/H⁺ permeability ratio favorable for neuron depolarization with minimal acidification (8, 11) compared to blue-light-activated channelrhodopsin 2 from Chlamydomonas reinhardtii (ChR2), the molecule that so far has been most popular in optogenetic studies (12). Recently, ChRmine, a BCCR, has been used successfully to activate mouse neocortical neurons with orange light (13).

Here, we describe 19 BCCRs from nine cryptophyte species, the protein sequences of which form two separate branches (subfamilies) of the phylogenetic tree (Fig. 1A) in addition to that comprising the previously characterized GtCCR1 and GtCCR2. Two of these proteins derived from two Rhodomonas species utilize a different activation mechanism and exhibit rapid desensitization of photocurrent under continuous illumination. We show that the photochemical cycle of these channelrhodopsins involves accumulation of an extremely short-wavelength-absorbing and long-living intermediate responsible for fast inactivation of their photocurrents. These observations reveal diversity within the BCCR family and contribute to deeper understanding of their cation channel function independently evolved from chlorophyte CCRs.

RESULTS
Identification and electrophysiological screening of BCCR homologs. Using probabilistic inference methods based on profile hidden Markov models (14) built on previously known BCCR sequences from G. theta, we identified 19 new BCCR homologs from nine marine cryptophyte strains included in the ongoing algal transcriptome sequencing projects (15, 16). The majority were cold-water species (from the Arctic or Antarctic), but Rhodomonas lens was from the Gulf of Mexico and Rhodomonas salina was from Milford, Connecticut. The previously unclassified strain CCMP 2293 has recently been allocated to the new genus Baffinella (as Baffinella frigidus) (17).
Table 1 lists GenBank accession numbers, source organisms, transcript names, and abbreviated protein names of the BCCR homologs identified in this study. In the abbreviated protein names, the first two or three letters stand for the beginning letters of the genus and species name. One of the sequences derived from *Rhodomonas lens* (*Rl* CCR1) exactly matched the sequence recently reported under the name “ChRmine” and attributed to the marine ciliate *Tiarina fusus* (13). As *T. fusus* culture used for RNA isolation was fed on *R. lens* (BioSample accession number SAMN02740485), the presence of this sequence in the *T. fusus* transcriptome can be explained by insufficient starvation of the organisms prior to RNA extraction.

Figure S1 in the supplemental material shows protein alignment of the opsin domains of BCCRs identified in this study. Asp85 and Thr89 (bacteriorhodopsin numbering) are conserved in all sequences, whereas Asp96 is replaced with Glu and Thr in *Baffinella frigidus* CCR2 (*Bf* CCR2) and *Geminigera* sp. (Caron lab isolate) CCR (*G1* CCR), respectively. However, neither of these sequences were electrogenic upon expression in mammalian cells (see below), and therefore, the functional importance of these substitutions could not be assessed. In many BCCR homologs, the position of bacteriorhodopsin’s Arg82 is occupied by other residues (Lys, Ala, Pro, Gln, or even Glu [in *Rhodomonas abbreviata* CCR2 (*RaCCR2*)]), which is unusual among microbial rhodopsins. Figure 1A shows a phylogenetic tree of the transmembrane domains of BCCRs identified so far. The previously characterized *GtCCR1* and *GtCCR2*, together with *HpCCR*, a closely related sequence from *Hanusia phi*, form a separate branch of this tree.

We synthesized human codon-optimized polynucleotides encoding the opsin domains of 19 newly identified BCCRs, fused them to an in-frame C-terminal EYFP (enhanced yellow fluorescent protein) tag and expressed in HEK293 (human embryonic kidney) cells. Ten of the encoded proteins generated photocurrents, the largest of
which were the peak currents from RaCCR1 and RsCCR1 (R. salina CCR1) (Fig. S2A). The action spectra were determined by measuring the initial slopes of photocurrent in the linear range of the light intensity. The spectrum of RaCCR1 closely matched that of RicCR1 (also called ChRmine [13]) and peaked at ~530 nm; that of RsCCR1 was ~5 nm blue-shifted (Fig. S2B). The spectral maxima of other tested BCCRs are listed in Table 1. The current kinetics was very diverse in the tested BCCRs. In particular, currents recorded from RaCCR1 and RsCCR1 exhibited extremely rapid desensitization during continuous illumination (Fig. 1B, red and blue lines, respectively). Representative photocurrent traces from other tested BCCRs are shown in Fig. S3.

To test relative permeability for Na\(^+\), we partially replaced this ion in the bath with nonpermeable N-methyl-\(\alpha\)-glucamine (NMG\(^-\)) and determined the reversal potentials (E_{rev}) by measuring the current-voltage relationships in four BCCR variants that generated the largest photocurrents. RaCCR1 and RsCCR1 showed large E_{rev} shifts toward the new equilibrium potential for Na\(^+\) (Fig. S4A), similar to the earlier reported GtCCRs (9, 10). However, RaCCR2 and RsCCR2 showed smaller shifts, similar to that in CrChR2 (18). When we reduced the bath pH without changing its Na\(^+\) concentration, E_{rev} shifts were correspondingly smaller in RaCCR1 and RsCCR1 compared to RaCCR2 and RsCCR2 (Fig. S4B), indicating a higher Na\(^+\)/H\(^+\) permeability ratio of the former two CCRs, compared to the latter. No change in E_{rev} was detected upon partial replacement of Cl\(^-\) in the bath with bulky aspartate, indicating that neither of the tested BCCRs conducted Cl\(^-\) (Fig. S4C).

**Absorption spectroscopy of RaCCR1 and RsCCR1.** To gain more insight into mechanisms of their photoactivation, we expressed and detergent purified RaCCR1 and RsCCR1 from the methylotrophic yeast *Pichia pastoris*. Their absorption spectra in the visible range closely matched the action spectra of photocurrents with main peaks at 530 and 524 nm, respectively (Fig. 2A). In addition, the absorption spectra of dark-adapted RaCCR1 and RsCCR1 showed structured absorption in the near-UV range (with peaks at ~307, 321, and 337 nm), in contrast to that of previously characterized GtCCR2 purified from the same expression host.

Similar UV bands had been reported in GtCCR4 and tentatively attributed to impurities of the sample (10). However, incubation of RaCCR1 and RsCCR1 with hydroxylamine, an agent known to cleave the retinal chromophore from the bacterio-

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**TABLE 1 BCCR homologs tested in this study**

| GenBank accession no. | Abbreviated protein name | Source organism | Transcript no. | Spectral peak (nm) |
|-----------------------|--------------------------|-----------------|----------------|-------------------|
| MNS85290              | *B*CCR1                  | *Baffinella frigidis* (CCMP 2293) | 0987_20121128_7073* | 500               |
| MNS85291              | *B*CCR2                  | *Baffinella frigidis* (CCMP 2293) | 0987_20121128_39196* | NA                |
| MNS85292              | GcCCR1                   | *Geminiger a cryophila* (CCMP 2564) | 0799_20121207_13742* | NA                |
| MNS85293              | GcCCR2                   | *Geminiger a cryophila* (CCMP 2564) | 0799_20121207_42824* | 460               |
| MNS85294              | GCCR1                    | *Geminiger sp.* (Caron lab isolate) | 1102_20130122_14880* | NA                |
| MNS85295              | HpcCCR                   | *Hausia phi* (CCMP 325) | 1048_20121227_6498* | 500               |
| MNS85296              | HrCCR                    | *Hemiselmis rufescens* (PCC 563) | 1357_20121228_3699* | NA                |
| MNS85298              | *P*uCCR2                 | Proteomonas sulcata (CCMP 704) | IRZA_2004242* | 480               |
| MNS85297              | *P*uCCR3                 | Proteomonas sulcata (CCMP 704) | IRZA_2061044* | NA                |
| MNS85299              | *P*uCCR4                 | Proteomonas sulcata (CCMP 704) | IRZA_2001844* | NA                |
| MNS85300              | RaCCR1                   | *Rhodomonas abbreviata* (Caron lab isolate) | 1101_20121128_4039* | 530               |
| MNS85303              | RaCCR2                   | *Rhodomonas abbreviata* (Caron lab isolate) | 1101_20121128_2696* | 470               |
| MNS85301              | RaCCR3                   | *Rhodomonas abbreviata* (Caron lab isolate) | 1101_20121128_32053* | 520               |
| MNS85302              | RaCCR4                   | *Rhodomonas abbreviata* (Caron lab isolate) | 1101_20121128_22530* | NA                |
| MNS85304              | RicCR1                   | *Rhodomonas lens* (CCMP 739) | 0484_2_20121128_11058* | 520               |
| MNS85305              | RicCR2                   | *Rhodomonas lens* (CCMP 739) | 0484_2_20121128_23336* | NA                |
| MNS85307              | RsCCR1                   | *Rhodomonas salina* (CCMP 1319) | 1047_20130122_18677* | 524               |
| MNS85308              | RsCCR2                   | *Rhodomonas salina* (CCMP 1319) | 1047_20130122_17846* | 470               |
| MNS85306              | RsCCR3                   | *Rhodomonas salina* (CCMP 1319) | 1047_20130122_11358* | NA                |

*Functional homologs are shown in bold type.

*Transcripts from the MMEETS project are indicated by an asterisk. Transcripts from the 1KP project are indicated by a pound sign.

*NA, not applicable.
Rhodopsin apoprotein in a light-dependent manner (19), decreased absorption in the UV region with the difference spectrum exhibiting the same triple-peak structure characteristic of protein-bound retinal (Fig. S5A and B). In both proteins, the rate of hydroxylamine bleaching of the UV bands was at least twice as fast as that of the main band (Fig. S5C and D), indicating that the UV-absorbing fractions were more accessible to hydroxylamine than the fractions absorbing in the visible range. Illumination accelerated bleaching in the visible range, as expected for retinylidene proteins, but did not influence the rate of bleaching at 321 nm (Fig. S5C and D). These results suggest that the structured UV absorbance in the dark-adapted sample is attributable to retinal binding to partially misfolded RaCCR1 and RsCCR1. The ratio of the UV absorption to the main peak absorption varied from 0.4 to 1.5 in different preparations, did not depend on the length of the expression construct, purification procedure, or storage conditions, and may reflect the relative amount of misfolded protein.

Continuous illumination of detergent-purified RaCCR1 with visible light decreased absorption at the main band (P530) and led to formation of a product (P330) with structured absorption in the UV region with three peaks at 318, 330, and 346 nm (Fig. 2B), red-shifted from those observed in the dark. Dissipation of P330 occurred on the time scale of seconds in parallel with recovery of the unphotolyzed state P530 (Fig. 2C). The recovery of the unphotolyzed state was ~3-fold slower in Pichia membranes than in detergent-purified protein (Fig. S6A). A very similar product with structured UV absorption was also formed upon illumination of purified RsCCR1 (Fig. S6B), with a rate of dissipation in the dark ~2-fold faster than that in RaCCR1 (Fig. 2D).

**Mechanism of photocurrent desensitization.** As described above, photocurrents from RaCCR1 and RsCCR1 exhibited rapid desensitization under continuous light (Fig. 1B). Desensitization was also observed under stimulation with 6-ns laser flashes at 0.1-Hz frequency even at 10% power (Fig. S6C), which argues against its origin from a secondary photochemical process. An alternative explanation for photocurrent desen-
sitization is the existence of a long-lived nonconductive state in the single-turnover photocycle. To determine the rate of peak current recovery, a second flash was applied after a variable time delay. The rate of restoration of the ability to generate electric current closely matched that of P330 dissipation in both purified proteins (Fig. 2D), strongly suggesting that accumulation of P330 is responsible for the rapid desensitization of photocurrents generated by RaCCR1 and RsCCR1.

Figure S7A shows a series of photocurrent traces generated by RaCCR1 in response to 1-s light pulses of different intensities. The peak photocurrent increased over the entire tested intensity range, whereas the degree of desensitization reached saturation 2 orders of magnitude earlier (Fig. S7B), and similar results were obtained with RsCCR1 (Fig. S7C). These observations show that the long-lived nonconductive P330 is not in equilibrium with the unphotolyzed state of the protein.

Alkalization caused formation of a UV-absorbing species of RaCCR1 with a structured spectrum closely matching that of the form obtained by illumination (Fig. 3A). The pKₐ of this process was identical to that of decrease of absorbance at 530 nm, which showed that the UV-absorbing species was produced from P530 (Fig. 3B). The alkali-induced conversion of the unphotolyzed form absorbing at 530 nm to P330 decreased the amplitude of the photoinduced conversion. When illuminated RaCCR1 samples were incubated in the dark at high pH, only a small decrease in absorbance at 330 nm was observed, compared to neutral pH conditions (Fig. 3C). The pKₐ of this decrease in amplitude was identical to that of conversion of P530 to the UV-absorbing species (Fig. 3D). We conclude from these observations that the same species accumulated at high pH as that obtained by illumination (i.e., P330). To the best of our knowledge, P330 is the shortest wavelength intermediate observed in the photocycle of any microbial rhodopsin. In addition to formation of P330, alkalization caused accumulation of an M-like intermediate absorbing at ~390 nm with a pKₐ of ~9.0,
although its concentration (assuming approximately equal extinction coefficients) was 10-fold smaller than that of P330 (Fig. 3E).

At pH 10, essentially all molecules were converted from the unphotolysed form to P330. This allowed us to use Fourier transform (FT) Raman spectroscopy to probe its chromophore structure in the dark. The Raman spectra measured at pH 7.2 and 10 and their difference spectrum are shown in Fig. 3F. The main ethylenic C=C stretch at 1,530 cm\(^{-1}\), which corresponds to the main visible peak at 530 nm (20), and the fingerprint C-C stretches at 1,200 and 1,163 cm\(^{-1}\) showed that at pH 7.2, retinal was predominantly in an all-trans configuration. Upon alkalization, the band at 1,530 cm\(^{-1}\) was strongly reduced, and new bands appeared at 1,590 cm\(^{-1}\) and 1,562 cm\(^{-1}\) which presumably corresponded to P330 and the M-like intermediate absorbing at \(\sim\)390 nm, respectively. The same two bands were clearly resolved in the difference spectrum.

**Fast photochemical conversions.** Fast photochemical conversions in the near-UV and visible range were analyzed by flash photolysis. Figure S8A shows a series of absorption changes in RsCCR1 detected at wavelengths from 390 to 570 nm at 10-nm increments. Only negligible (less than 0.5 milli optical density [mOD] unit) oppositely directed components with the time constant (\(\tau\)) values of \(\sim\)60 to 100 \(\mu s\) were observed at the wavelengths at which maximal absorption of the red-shifted K and blue-shifted L intermediates are expected (480 and 560 nm, respectively) (Fig. S8B). Therefore, we could not follow the K-to-L transition, which occurred on a much faster time scale. To obtain the spectral changes due to L formation, we plotted the mean absorption changes in the time window between 50 and 100 \(\mu s\) after the flash against wavelength (Fig. 4A). The maximum of the L intermediate in this difference spectrum was at \(\sim\)460 nm.

The spectral characteristics of the later transitions were obtained by global fit analysis. The appearance of a typical M intermediate with the absorption maximum at \(\sim\)390 nm (the positive peak in Fig. 4B) was observed within 1 ms. After that, biphasic bleaching at all visible wavelengths took place, which was obviously related to generation of the P330 form. Fast bleaching with \(\tau\) of \(\sim\)4.5 ms reflected the decay of the

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**FIG 4** (A) Mean photoinduced absorbance changes recorded from purified RsCCR1 in the 50- to 100-\(\mu s\) time window. (B to F) Spectral transitions in RsCCR1 derived by global fit analysis.
bulk of the initial form and may involve the appearance of a blue-absorbing (N?)
intermediate (Fig. 4C), which was more obvious during the slow bleaching with
τ~40 ms (Fig. 4D). The recovery of the initial state proceeded in two steps with τ of 1.4
and 3.8 s. At least the fast recovery involved depletion of a blue-absorbing intermediate
(Fig. 4E). The τ of the main slow recovery component was equal to those of P330
dissipation and restoration of electrical sensitivity (Fig. 4F and Fig. 2D). Qualitatively
similar phototransitions were observed in the second pigment RaCCR1 with time
constants of components 0.3, 6, 40, 3.2, and 11.4 ms (Fig. 59). In agreement with slower
dissipation of the P330 intermediate and restoration of light sensitivity in this pigment
compared to RsCCR1 (Fig. 2D), the recovery in the visible range was also slower, and
depletion of the blue-absorbing form was also observed (Fig. S9E and F). However, the
40-ms component which in RsCCR1 corresponded to slow bleaching, in RaCCR1
revealed fast recovery.

We recorded photocurrents in HEK cells upon 6-ns laser flash excitation at 532 nm
as in flash-photolysis measurements for kinetic comparison with absorption changes in
purified proteins. Channel opening and closing in RaCCR1 and RsCCR1 took place in the
same time windows as absorption changes at the wavelengths of M-intermediate
absorption in which proton transfers occur (Fig. 5A and B).

In the current traces generated by GtCCR1 and GtCCR2, a large peak was observed
in the 30- to 100-μs time domain prior to channel opening (9). This peak, also exhibited
by some low-efficiency CCRs from green algae, reflects intramolecular transfer of the
Schiff base proton to an outwardly located acceptor, integrated by the measuring
system (21). This component could also be resolved in the current traces from RaCCR1
and RsCCR1 recorded at the voltages near the reversal potential for Na⁺, but it was
~100-fold smaller than that in GtCCR1 and GtCCR2 (Fig. S10).

**Mutagenesis analysis.** We found in GtCCR1 and GtCCR2 that a neutralizing muta-
tion of the Asp96 homolog (Asp98) completely suppressed channel activity, so that
only intramolecular transfer of the Schiff base proton could be detected (9). The
_corresponding D125N and D128N mutations in RaCCR1 and RsCCR1, respectively, did
not eliminate channel currents (Fig. 6A). Neutralization of the Asp85 homolog in
RaCCR1 (the D114N mutation) reduced expression of the construct, as judged by the
tag fluorescence, and no photocurrents above the noise level could be detected.
Replacement of Cys119 (corresponding to Thr90 in bacteriorhodopsin) with Ala com-

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**FIG 5** (A and B) Laser flash-evoked photocurrents at ~60 mV (red) and photoinduced absorbance
change (blue) recorded from RaCCR1 (A) and RsCCR1 (B).
pletely abolished photocurrents in RaCCR1, as did also the corresponding mutations in RaCCR2, GtCCR2, and Proteomonas sulcata CCR (PsuCCR). In the RaCCR1_C119T mutant, the photocurrent amplitude was greatly reduced (the mean peak current in response to a first light pulse of maximal intensity was 10 ± 3 pA in 14 cells). These tiny currents, however, exhibited only ~40% desensitization during 1-s continuous illumination (Fig. 6B), i.e., much less than in the wild type. The photocurrent decay after switching the light off was biphasic, with the slow phase on the second time scale. However, the most striking difference of this mutant from the wild type was the absence of the long-living UV-absorbing form with the structured spectrum corresponding to P330 in the wild type (Fig. 6C). Instead, a smooth peak with the maximum at 380 nm (the M state) was produced. The dissipation of this state and the recovery of the unphotolyzed state with the peak absorbance at 506 nm were very slow (Fig. 6D).

**DISCUSSION**

Our results show that BCCRs are widely spread among cryptophyte algae and form three branches (subfamilies) of a phylogenetic tree. BCCRs exhibit diverse current kinetics, spectral sensitivity, and Na⁺/H⁺ permeability ratios, as has also been found in other channelrhodopsin families. Two representatives of the currently studied BCCRs differ in their mechanism of photoactivation from previously described GtCCR1 and GtCCR2, which belong to a different subfamily of cryptophyte CCRs. Most notably, a particular proton transfer essential to trigger channel opening in the earlier reported subfamily is not required in the subfamily described in this study. Photocurrents by RaCCR1 and RsCCR1 exhibit very rapid desensitization under continuous illumination, which we show is related to the formation of a long-living UV-absorbing intermediate in their photocycles. Similar rapid photocurrent desensitization was observed in Mer-
MAIDs (marine anion-conducting and intensely desensitizing channelrhodopsins), explained by accumulation of a long-lived M intermediate with an unusual short-wavelength maximum absorption peak at 364 nm (7). In both RaCCR1 and RsCCR1, two distinct UV-absorbing intermediates were accumulated upon illumination, one at 390 nm, typical of M intermediates, and the other a triple-peak species with a uniquely far-blue-shifted spectrum with a 330-nm maximum.

Flash photolysis measurements revealed an extremely fast (<1 μs after the flash) appearance of the L intermediate that might be in equilibrium with K during the first 100 μs in RaCCR1 and RsCCR1. The typical M intermediate absorbing at 390 nm was formed during 0.1 to 1 ms. We could not monitor the appearance of the second blue-shifted intermediate (P330) because the low signal-to-noise ratio in the near-UV range limited measurements to wavelengths of ≳380 nm. However, we observed bleaching in the entire visible range on the millisecond time scale, which likely indicated accumulation of P330. A decrease of absorption in the blue region was observed during the fast recovery of the unphotolyzed states of both RaCCR1 and RsCCR1. This most probably reflects dissipation of an N-like intermediate that appears earlier in the photocycle.

The short-wavelength absorption of photointermediate P330 is unique among photocycles of microbial rhodopsins. The extremely short wavelength absorption and very high ethylenic stretch wave number of a corresponding band in Raman spectra suggest an extremely hydrophobic environment of the retinal moiety. The chromophore in P330 could, for example, be retroretinal, a derivative in which all double bonds are shifted toward the ring by one position (22), or a free retinal that remains in the binding pocket (23). A linear photocycle involving P330 is the simplest scheme that fits our results; however, we cannot exclude a branched photocycle as was proposed for the CrChR2_C128T mutant (22).

Rapid desensitization observed in RaCCR1 and RsCCR1 under continuous illumination would potentially allow temporally precise neuronal activation even in the presence of light that can be used for fluorescent imaging. Additional advantages of these two BCCRs are their relatively red-shifted absorption and high Na+/H+ permeability ratio. Better understanding of their molecular mechanisms will facilitate their rational design for optogenetic needs.

MATERIALS AND METHODS

Bioinformatics. Bacteriorhodopsin-like cation channelrhodopsin (BCCR) homologs were identified by searching cryptophyte transcriptomes from the Marine Microbial Eukaryote Transcriptome Sequencing (MMETS) project (15) and the 1,000 Plants (1KP) project (16) using probabilistic inference methods based on profile hidden Markov models (profile HMMs). Profile HMMs were built from previously known BCCR sequences using HMMER software (version 3.1b2) (14) with default parameters and refined upon functional testing of the homologs by patch clamping. Search procedures were automated with Python 2.7 and the Biopython module (24). The protein sequence alignment was created using the MUSCLE algorithm implemented in DNASAR Lasergene (Madison, WI) MegAlign Pro software. The phylogenetic tree was visualized using Dendroscope software (25).

Molecular biology. For expression in HEK293 cells, DNA polynucleotides encoding the BCCR opsin domains optimized for human codon usage were synthesized (GenScript, Piscataway, NJ) and cloned into the mammalian expression vector pcDNA3.1 (Life Technologies, Grand Island, NY) in frame with an EYFP tag. For expression in Pichia, the opsin-encoding constructs were fused in frame with a C-terminal eight-His tag and subcloned into the pPIC9K vector (Invitrogen). Mutants were generated with Quikchange XL kit (Agilent Technologies, Santa Clara, CA) and verified by sequencing.

HEK293 transfection and patch clamp recording. HEK293 cells were transfected using the ScreenFectA transfection reagent (Waco Chemicals USA, Richmond, VA). All-trans retinal (Sigma) was added at the final concentration of 3 μM immediately after transfection. Photocurrents were recorded 48 to 96 h after transfection in the whole-cell voltage clamp mode with an Axopatch 200B amplifier (Molecular Devices, Union City, CA) using the 10-kHz low-pass Bessel filter. The signals were digitized with a Digidata 1440A using pClamp 10 software (both from Molecular Devices). Patch pipettes with resistances of 2 to 4 MΩ were fabricated from borosilicate glass. The standard pipette solution contained 126 mM KCl, 2 mM MgCl₂, 0.5 mM CaCl₂, 5 mM Na-EGTA, and 25 mM HEPES (pH 7.4). The standard bath solution contained 150 mM NaCl, 1.8 mM CaCl₂, 1 mM MgCl₂, 5 mM glucose, and 10 mM HEPES (pH 7.4). A 4 M KCl bridge was used in all experiments, and possible diffusion of Cl⁻ from the bridge to the bath was minimized by frequent replacement of the bath solution with fresh buffer. For measurements of the reversal potential shifts under varied ionic conditions, Na⁺ was substituted for K⁺ in the pipette solution to minimize the number of ionic species in the system. To reduce the Cl⁻ concentration in the bath, NaCl was replaced with Na aspartate. To reduce the Na⁺ concentration, NaCl was replaced with N-methyl-D-glucamine.
chloride. To increase the H+ concentration, the pH was adjusted with H2SO4. The holding voltages were corrected for liquid junction potentials calculated using the Clampex built-in LJP calculator (26). Continuous light pulses were provided by a Polychrome V light source (TILL Photonics GmbH, Graefelfing, Germany) in combination with a mechanical shutter (Uniblitz model LS6; Vincent Associates, Rochester, NY) (half-opening time, 0.5 ms). The maximal quantum density at the focal plane of the 40× objective was 7.7 mW mm−2 at 515 nm. The action spectra were constructed by calculation of the initial slope of photocurrent and corrected for the quantum density measured at each wavelength. Laser excitation was provided by a Minilite Nd:YAG laser (532 nm; pulse width, 6 ns; energy, 12 mJ) (Continuum, San Jose, CA). The current traces were logarithmically filtered using a custom software program, and the laser artifact was digitally subtracted. Curve fitting was performed by Origin Pro software (OriginLab Corporation, Northampton, MA).

**Expression and purification of BCCRs from Pichia.** The plasmids encoding BCCRs were linearized with SalI and used to transform P. pastoris strain SMD1168 (his4 pep4) by electroporation. Transformants were first screened for their ability to grow on histidine-deficient medium, and second, for their Genetin resistance. Single colonies that grew on 4 mg/ml Genetin were screened by small-scale cultivation, and clones of the brightest color were selected. For protein purification, a starter culture was inoculated into buffered complex glycerol medium until A600 reached 4 to 8, after which the cells were harvested by centrifugation at 5,000 rpm and transferred to buffered complex methanol medium supplemented with 5 μM all-trans retinal (Sigma-Aldrich). Expression was induced by the addition of 0.5% methanol. After 24 to 30 h, the cells were harvested and disrupted in a bead beater (BioSpec Products, Bartlesville, OK) in buffer A (20 mM sodium phosphate [pH 7.4], 100 mM NaCl, 1 mM EDTA, 5% glycerol). After removing cell debris by low-speed centrifugation, membrane fragments were collected by ultracentrifugation at 40,000 rpm in a TLA-100 rotor, resuspended in buffer B (20 mM HEPES [pH 7.4], 300 mM NaCl, 5% glycerol) and solubilized by incubation with 1.5% dodecyl maltoside (DDM) for 1.5 h or overnight at 4°C. Nonsolubilized material was removed by ultracentrifugation at 50,000 rpm in a TLA-100 rotor. The supernatant was mixed with nickel-nitrilotriacetic acid agarose beads (Qiagen, Hilden, Germany) and loaded on a column. The proteins were eluted with buffer C (20 mM HEPES [pH 7.4], 300 mM NaCl, 5% glycerol; 0.02% DDM) containing 300 mM imidazole. The pigments were concentrated, and imidazole was removed by repetitive washing with imidazole-free buffer C using YM-10 centrifugal filters (Amicon, Billerica, MA).

**Absorption spectroscopy and flash photolysis.** Absorption spectra of purified BCCRs were recorded using a Cary 4000 spectrophotometer (Varian, Palo Alto, CA). The pHw was determined by fitting the classical Henderson-Hasselbalch equation in the form y = A/[1 + 10E(pHw − pH)] to experimental data. Light-induced absorption changes were measured with a laboratory-constructed crossbeam apparatus. Excitation flashes (532 nm, 6 ns, 40 μJ) were provided by a Surelile I Nd-YAG laser (Continuum, Santa Clara, CA). Measuring light was from a 250-W incandescent tungsten lamp combined with a McPherson monochromator (model 272; McPherson, Acton, MA). Absorption changes were detected with a Hamamatsu Photonics (Bridgewater, NJ) photomultiplier tube (model R928), protected from excitation laser flashes by a second monochromator of the same type. Signals were amplified by a low-noise current amplifier (model SR445A; Stanford Research Systems, Sunnyvale, CA) and digitized with a GaGe Octopus digitizer board (model CS8327; DynamicSignals LLC, Lockport, IL) at a maximum sampling rate of 50 MHz. Logarithmic filtration of the data was performed using the GageCon program (27).

**Fourier transform Raman spectroscopy.** Fourier transform Raman spectra were collected in 5 μl of a concentrated detergent-solubilized protein in pH-adjusted elution buffer placed in a metallic holder and covered by adhesive tape. The scattering was recorded in 180° backscattering geometry, using FRA106/s accessory to the Bruker IFS66vs spectrometer, with Nd:YAG laser excitation provided at 1064 nm, at a 2-cm−1 resolution, controlled by the OPUS software. At least 10,000 scans were averaged per sample. Raman spectra of the buffers were taken separately in the same geometry and subtracted to obtain pure protein spectra.

**Statistics.** Descriptive statistics was used as implemented in Origin software. The data are presented as means ± standard error of the mean (SEM) values; the data from individual replicates are also shown when appropriate. The sample size was estimated from previous experience and published work on similar subjects, as recommended by the NIH guidelines (28).

**Data availability.** The polynucleotide sequences of BCCR homologs reported in this study have been deposited in GenBank (accession numbers MN585290 to MN585308).

**SUPPLEMENTAL MATERIAL**

Supplemental material is available online only.

**FIG S1**, PDF file, 1.3 MB.

**FIG S2**, TIF file, 0.8 MB.

**FIG S3**, TIF file, 1 MB.

**FIG S4**, TIF file, 1.1 MB.

**FIG S5**, TIF file, 2 MB.

**FIG S6**, TIF file, 1.7 MB.

**FIG S7**, TIF file, 1.5 MB.

**FIG S8**, TIF file, 2.3 MB.

**FIG S9**, TIF file, 2.6 MB.

**FIG S10**, TIF file, 0.9 MB.
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