Recoverin depletion accelerates cone photoresponse recovery

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The neuronal Ca\(^{2+}\)-binding protein Recoverin has been shown to regulate phototransduction termination in mammalian rods. Here we identify four recoverin genes in the zebrafish genome, rcv1a, rcv1b, rcv2a and rcv2b, and investigate their role in modulating the cone phototransduction cascade. While Recoverin-1b is only found in the adult retina, the other Recoverins are expressed throughout development in all four cone types, except Recoverin-1a, which is expressed only in rods and UV cones. Applying a double flash electroretinogram (ERG) paradigm, downregulation of Recoverin-2a or 2b accelerates cone photoresponse recovery, albeit at different light intensities. Exclusive recording from UV cones via spectral ERG reveals that knockdown of Recoverin-1a alone has no effect, but Recoverin-1a/2a double-knockdowns showed an even shorter recovery time than Recoverin-2a-deficient larvae. We also showed that UV cone photoresponse kinetics depend on Recoverin-2a function via cone-specific kinase Grk7a. This is the first in vivo study demonstrating that cone opsin deactivation kinetics determine overall photoresponse shut off kinetics.

1. Introduction

The vertebrate retina contains two classes of photoreceptors, rods and cones, which function at low and bright light conditions, respectively. Although both share a similar G-protein-coupled phototransduction pathway, the cone phototransduction is characterized by lower sensitivity and faster kinetics, which allows cones to function over almost 9 orders of illumination magnitude [1]. The rate-limiting step of photoresponse recovery is also different between these two cell types [2–4]. Rods and cones use cell-type-specific molecules in the phototransduction cascade, which may account for these differences.

The cascade is initiated by light-activated rhodopsin (Rh*), which induces the detachment of trimeric G-protein Transducin α-subunit. The Transducin α-subunit in turn binds to phosphodiesterase (PDE), causing a decrease in the second messenger cGMP. This drop in cGMP levels leads to the closure of CNG cation channels, hyperpolarizing the photoreceptor and lowering \([Ca^{2+}]_i\) [5]. The deactivation of both Rh* and the PDE–Transducin complex is required to terminate the phototransduction cascade. Rh* in rods is initially phosphorylated by Rhodopsin kinase Grk1 [6] in a Ca\(^{2+}\)-dependent manner via the small Ca\(^{2+}\)-binding protein Recoverin [7,8]. Rod Recoverin is proposed to inhibit Grk1 in darkness when outer segment \([Ca^{2+}]_i\) is high and is released from Grk1 during light response when \([Ca^{2+}]_i\) is decreasing [9,10]. Modulation of Rhodopsin lifetime during light adaption is abolished in Recoverin-deficient mice [11]. The final deactivation of Rh* is achieved by the binding of Arrestin [12], while intrinsic GTPase activity ends the ability of PDE–Transducin complex to hydrolyse cGMP [13].

Cones-specific opsin kinase Grk7 has similar functions as rod Grk1 [14,15]. Interestingly, cone Arrestin is not essential to deactivate short-wavelength opsin in mice [16,17], but its functional loss induces delayed cone photoresponse recovery in cone-dominant zebrafish [18]. A Ca\(^{2+}\)-sensitive phosphorylation of cone...
visual pigment in vivo in zebrafish [19] indicates a possible role of Recoverin in regulating cone opsin quenching via direct control of [Ca2+], during light response.

In this study, we take advantage of the cone-dominant retina of zebrafish to study Recoverin function in cone vision. The zebrafish genome harbours four rcv genes. While Recoverin-1b (Rcv1b) is only expressed in the adult retina the other Recoverins are present throughout development in all cone types, with the exception of Rcv1a, which is expressed in rods and UV cones only.

We determined photoresponse recovery with a double flash electroretinogram (ERG) paradigm [14] analysing the role of Recoverin proteins. Our results establish that the cone photoresponse kinetics are shaped by photopigment quenching and a loss of Recoverin leads to an accelerated photoresponse recovery.

2. Material and methods

2.1. Zebrafish care

Zebrafish (Danio rerio) were kept at a 14 L : 10 D cycle at 28°C [20]. Embryos of WIK wild-type fish were raised in E3 medium containing 0.01% methylene blue or with 0.2 mM PTU (1-phenyl-2-thioaura; Sigma-Aldrich) to avoid pigmentations. Adult zebrafish were sacrificed by fish system water containing 0.4% 3-aminobenzoic acid methyl ester (MESAB, Sigma-Aldrich) and 4.6 mM NaHCO3.

2.2. Cloning of recoverin genes and in situ hybridization

Cloning was performed as described in [21] using oligonucleotide primers listed in electronic supplementary material S1. Digoxigenin-labelled in situ hybridization (ISH) RNA probes were generated according to supplier’s instructions (DIG RNA Labelling Mix, Roche) and used on zebrafish larvae as previously described [22].

2.3. Generation of antibodies

Custom polyclonal antibody peptides were raised by Eurogentec (Seraing, Belgium). Rabbits were immunized with the Rcv1a peptide CIQYDEPKKIQEKLEKKKH or Rcv2b peptide CKLIPKDKQSLPND. Guinea pigs were immunized with the Rcv1b peptide CIQFDKPKQVKQEKLEKTKQ or Rcv2a peptide CKLIPKEDQESLPAD. Antibodies were affinity-purified.

2.4. Immunohistochemistry

Section immunohistochemistry was carried out as described previously [14], except for the following modifications. Adult zebrafish eyes were fixed in Zebrafish (Danio rerio) were kept at a 14 L : 10 D cycle at 28°C (Seraing, Belgium), Rabbits were immunized with the Rcv1a peptide CIQYDEPKKIQEKLEKKKH or Rcv2b peptide CKLIPKDKQSLPND. Guinea pigs were immunized with the Rcv1b peptide CIQFDKPKQVKQEKLEKTKQ or Rcv2a peptide CKLIPKEDQESLPAD. Antibodies were affinity-purified.

2.5. Microscopy

ISH images were taken by a light microscope (BX61, Olympus) with a CCD camera (ColorView IIIu, Soft Imaging System, Olympus) and processed by Adobe PHOTOSHOP CS3. Fluorescence Z-stacks photos were taken by a confocal laser scanning microscope (Leica SP5, Leica Microsystems) and processed by IMARS 7.6.3 (Bitplane, Zurich, Switzerland).

2.6. Targeted gene knockdown

Antisense morpholino oligonucleotides (Gene Tools, Philomath, OR, USA) were designed against translational start sides and injected into one cell-stage embryos (electronic supplementary material S1).

Amounts of morpholino per embryo were rcv1a (20 ng), rcv2a (7.4 ng or 3.6 ng), grk7a (2.4 ng), rcv2b (1.2 ng) and control (7.4 ng). For double knockdowns, rcv2a (3.6 ng)/grk7a (2.4 ng) or rcv1a (11 ng)/rcv2a (7.4 ng) were injected.

2.7. Western blot

Twenty to forty 5 dpf larvae were homogenized in 150 μl RIPA buffer (150 mM NaCl, 1% Triton-X, 0.5% sodium deoxycholate, 50 mM Tris (pH 8), 1 mM EDTA, 0.1% SDS). β-Actin (approx. 42 kDa) was used as a loading control. Primary antibodies were diluted to the following concentrations: Rcv1a: 1:1000; Rcv2a: 1:2000; Rcv2b: 1:1000 β-Actin: 1:3000. Secondary antibodies were diluted in blocking buffer (goat anti-rabbit: 1:15000; rabbit anti-guinea pig: 1:25000; goat anti-mouse: 1:10000) with a short wavelength absorbing filter and an UV light filter in front of white light source (electronic supplementary material S2B). After calibration, the spectrum graph showed a peak at around 720 nm. To apply together with this UV light filter to receive only the spectrum shown in S2A (SpectraSuite, Ocean Optics). For the same intensity and duration (500 ms) were given [14]. The spectrum shown in S2A (SpectraSuite, Ocean Optics). For the same intensity and duration (500 ms) were given [14]. The interval between two flashes was progressively increasing (100, 200, 300, 500, 1000, 2000, 3500 and 5000 ms). A neutral density filter was applied to have 0.1% double white flash paradigm and the interval between two flashes was increasing (100, 200, 300, 500, 1000, 2000, 3500 and 5000 ms). A neutral density filter was applied to have 0.1% double white flash paradigm and the interval between two flashes was progressively increasing (100, 150, 200, 250, 300, 350, 400, 450 and 500 ms). The interval between two pairs was always 10 s for both bright light and dim light response. Spectrum ERG used the similar set-up but additional background light source (Philips projection lamp type 6958, 20 V, 250 W; housing: Liesegang Diafant 250) with a short wavelength absorbing filter and an UV light filter in front of white light source (electronic supplementary material S2B). After calibration, the spectrum graph showed that most of the light above 385 nm in the visible range was blocked. There was indeed a peak at around 720 nm. To make sure this peak cannot generate any electrical response in the retina, another short wavelength absorbing filter was applied together with this UV light filter to receive only the
genes were expressed in the pineal gland. (the ventral outer retina, with the exception of ing at around 2 dpf. At around 3 dpf, expression is initiated in
are expressed in the pineal gland, a photosensitive organ, start-
Recoverins in larval and adult zebrafish. All four
In order to explore the role of Recoverins (Rcvs) in the termin-
3. Results
In order to explore the role of Recoverins (Rcvs) in the termin-
3.1. Recoverins are expressed in photosensitive organs
We determined both the RNA and protein expression of the
of bipolar cells (electronic supplementary material S3),
light at around 720 nm. But even in darkness this light did not
give any ERG response. The background light was used to
adapt the blue, green and red cones with minimal activation
for UV cones. For paired UV flash recordings, flash interval
duration were the same as for dim flash recordings with
one extra interval of 750 ms. All the experiments were
performed at room temperature (22°C).

3.2. UV cone photoresponse recovery is accelerated
in Rcv-deficient zebrafish larvae
To study the function of Recoverin proteins, we designed
morpholinos against rcs and cone-specific opsin kinase Grk7
[14]. The knockdown efficiency at 5 dpf was evaluated by
Western blot and whole-mount immunohistochemistry (elec-
tronic supplementary material S4). The levels of each Rcv
protein were largely reduced when it was knocked down
while levels of the other Rcv proteins were maintained, indica-
tive of both knockdown efficacy and antibody specificity.
The knockdown efficiency was in the range of 90%
(Rcv1a, 93%; Rcv2a, 97%; Rcv2b, 88%), as semi-quantified by
Western blots using β-actin as loading control (electronic
supplementary material S4B).

In order to study the function of the only mammalian
orthologue Rcv1a in larvae and to investigate how different
Recoverins modulate opsin lifetime in the same cell, we
used spectral ERG to isolate photoresponse, which was lar-
inner retinal staining was neither detected by ISH (figure 1)
where the labelling partially colocalized with the ON-bipolar
cell marker protein kinase C alpha (PKC). Interestingly, this
inner retinal staining was neither detected by ISH (figure 1) nor
immunohistochemical staining in the larval retina (elec-
tronic supplementary material S4A). Rcv1b protein was
found in all adult cone photoreceptors (figure 2), while
being absent in the larval retina (figure 1).

Figure 1. Expression of rcv genes in 3 dpf and 5 dpf zebrafish larvae. (a–h) All the rcv genes except rcv1b showed expression in 3 dpf larvae retina. All the rcv genes were expressed in the pineal gland. (i–l) rcv1b still showed no expression in the 5 dpf larval retina. Scale bar (=50 μm) applies to all panels.
order to recover to 40% of their dark levels, Rcv1a and Rcv2a double knockdown, Rcv2a single knockdown and control morphants required around 200 ms, 275 ms and 350 ms, respectively. Rcv1a knockdown did not induce any phenotype, but Rcv1a and Rcv2a double defect larvae showed significant acceleration compared with Rcv2a single knockdown, indicating that the function of Rcv1a can be replaced by Rcv2a. Concomitant reduction of Rcv2a and Rcv1a cannot be compensated any more. The fact that recovery was faster rather than slower in morphants argues against a toxic side effect of morpholino injection.

Because Rcv proteins are highly expressed in the photoreceptor synaptic terminal (figure 2; electronic supplementary material, figure S4) and Rcv1 in mice has been reported to modulate synaptic transmission in the rod pathway [31], it is possible that the faster b-wave recovery in the morphants not

**Figure 2.** Zebrafish Recoverins are expressed in the different cone subtypes. Adult retinal sections from transgenic zebrafish highlighting the different cone subtypes were co-stained with Rcv antibodies. White arrowhead in (a) marked the rod photoreceptors. While Rcv1b, Rcv2a and Rcv2b proteins are present in all cone subtypes, Rcv1a is only expressed in rods and UV cones. In order to highlight also ON-bipolar cells, Rcv2a antibodies were supplemented with PKC antibodies (violet staining). Scale bar (¼ 20 μm) applies to all panels.
only comes from the faster phototransduction decay, but is additionally mediated by an effect on synaptic transmission. The contribution of Rcv downregulation to synaptic transmission can be easily evaluated by quantifying the b-wave amplitude and time-to-peak (table 1). Indeed in this experiment, we did not find any difference between morphants and controls, arguing that synaptic transmission is not affected, which suggested the effect we observed in b-wave recovery is purely contributed by the accelerated phototransduction termination.

3.3. Recoverin affects SWS1 recovery via Grk7a

In rod photoreceptors, it is known that Recoverin regulates the lifetime of activated rhodopsin via Rhodopsin kinase [7]. In order to confirm that cone-specific visual pigment kinase

![Diagram](http://rsob.royalsocietypublishing.org/doi/abs/10.1098/rsob.150086)
Grk7a (electronic supplementary material S5) interacts with Recoverin, we compared the photoresponse recovery in Grk7a single knockdown larvae with the recovery time in double knockdowns of Grk7a and Rcv2a (figure 3b).

We confirmed that the response recovery is significantly delayed in the absence of Grk7a not only under normal ERG [14] but also in UV spectrum ERG (p < 0.05 at interval 250–500 ms), indicating Grk7a is a general cone opsin kinase (figure 3b). Importantly, the photoresponse recovery was not influenced by the additional reduction of Rcv2a (p > 0.5), indicating that Rcv2a acts via Grk7a. Hence, Recoverin regulates SWS1 opsin lifetime via cone opsin kinase.

### 3.4. Rcv2a and Rcv2b depletion accelerate photoresponse recovery under varying light conditions

It was suggested that white light ERG is dominated by the double cone photoresponse [18]. The fact that (unlike UV response) there was no significant difference in b-wave recovery between Rcv1a and Rcv2a double knockdown and Rcv2a single knockdown larvae in the double white flash paradigm proved this notion for the first time (figure 3c). Indeed, there was a significant acceleration of b-wave response recovery in rcv2a morphants with paired saturating flashes compared with control morphants (figure 3c). The rcv2a morphant recovery at an interval of 300 ms was three times faster than in control morphants. However, normal response kinetics were observed in rcv2b morphants (p > 0.1). Nevertheless, under dim flash conditions (0.1% of maximal light intensity) downregulation of rcv2a and rcv2b both accelerated the response decay (p < 0.05 at intervals from 250 ms to 500 ms; figure 3d). It took about 400 ms for rcv2a morphants, 450 ms for rcv2b morphants and 550 ms for the controls to recover about half of the photoresponse, suggesting a role of rcv2b in modulating the photoresponse kinetics only at a smaller Ca$^{2+}$ dynamic range.

### 4. Discussion

Rhodopsin quenching requires the phosphorylation by rhodopsin kinase [6,32] and subsequent binding of Arrestin [33]. Rhodopsin kinase is regulated by Recoverin in a Ca$^{2+}$-dependent manner [7,8]. Exogenous Recoverin prolongs the rod response [34] and genetic deletion does the opposite [35]. Before this study, little was known about the function of Recoverin in cones. But both the Ca$^{2+}$-sensitive cone opsin phosphorylation in zebrafish [19] and the Ca$^{2+}$-sensitive cone opsin quenching step dominating the overall response kinetics in salamander [3,4] imply an important role of Recoverin in shaping the cone photoresponse.

Four recoverins were found in the zebrafish genome. Larval zebrafish double cones exclusively express rcv2a and rcv2b. During bright light conditions, Rcv2a works to delay red and green opsin quenching, while both Rcv2a and Rcv2b work under dim light conditions. Hence Rcv2b contributes only when Ca$^{2+}$ dynamic range is small [36,37] and less opsin is bleached, suggesting that Rcv2a may be the primary Recoverin and works over broader light conditions. This suggests variations in Ca$^{2+}$ sensitivity among Recoverin isoforms, which would correlate with different numbers of lysine residues at the C-terminus of various Revs. The number of positively charged lysine residues indirectly determines the differences in membrane affinity and efficiency of rhodopsin kinase inhibition between salamander rod and cone Recoverin [38,39]. A reduced number of lysine residues results in decreased Ca$^{2+}$ sensitivity and requires higher Ca$^{2+}$ concentrations to inhibit rhodopsin kinase [40]. Rcv2a and Rcv2b possess 2 and 1 lysine residues at the C-terminus, respectively. Another consistent mechanism for Rcv2b working in dim light conditions could be different functional pairs of Grk and Recoverin. Two paralogous genes of grk7 (grk7a and grk7b) and one paralogue for grk1 (grk1b) were found to be expressed in zebrafish cones, while grk1a is restricted to rods [14,41]. The detailed cellular distribution of Grks in cones is not known, but Grk7a is clearly expressed in all cone subtypes (electronic supplementary material S5). According to Revs expression pattern and functional analysis, there are at least two functional pairs in zebrafish: Grk1a-Rcv1a exclusively in rods and Grk7a-Rcv2a in UV cones. The specific activity of Grk7a for rhodopsin phosphorylation is around 30 times higher than that of all the other Grks [41]. Rcv2b may regulate Grk1b or Grk7b instead of Grk7a, which would be consistent with Rcv2b working under different conditions than Rcv2a. Different Ca$^{2+}$ sensitivity and Grk affinity may both contribute to different working ranges between Rcv2a and Rcv2b.

In rods, it is believed that Ca$^{2+}$ regulates light response and light adaptation via three mechanisms: the dynamic drop in intracellular Ca$^{2+}$ concentration accelerates rhodopsin phosphorylation [7], speeds up the synthesis of cGMP through guanylyl cyclase [42] and enhances the cGMP affinity of CNG-channel [43]. In cone photoreceptors there is a larger fraction of dark current carried by Ca$^{2+}$ [44], a faster light-induced Ca$^{2+}$ dynamic decline and a bigger Ca$^{2+}$ dynamic range than in rods [37,45]. Furthermore, cones show lower sensitivity, faster response kinetics and adaption to a much wider range of light intensity than rods. It is expected that there is a more powerful Ca$^{2+}$ negative feedback in cones. The Ca$^{2+}$-dependent regulation on CNG channel ligand sensitivity is significantly more potent in cones with CNG-modulin as a modulator than in rods using calmodulin [46,47]. However, the Ca$^{2+}$-sensitive guanylyl cyclase activating proteins (GCAPs) show weaker modulation in mammalian cones than in rods [48]. Ca$^{2+}$ probably mediates a more powerful feedback via visual pigment phosphorylation in cones than in rods because the rate-limiting step of bright light response recovery in cones has been found to be opsin quenching [3,4], instead of Ca$^{2+}$ insensitive PDE–Transducin deactivation in rods [2].

Our study is the first in vivo demonstration that cone opsin phosphorylation is indeed regulated by the Ca$^{2+}$-binding protein Recoverin, which allows Ca$^{2+}$ to directly control cone response kinetics and modulate the time scale during light response and adaptation.

**Data accessibility.** Genbank accession numbers: KT325590–KT325593. Authors’ contributions. J.Z. carried out the molecular, imaging and electrophysiological work, data analysis and design of the study, and drafted the manuscript; J.K. participated in imaging and electrophysiological work; E.K. contributed to cloning; M.G. contributed to molecular analysis, antibody design and drafted the manuscript; S.C.F.N. conceived, designed and coordinated the study, and drafted the manuscript. All authors gave final approval for publication.

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