

Guanylate cyclase–activating protein 2 contributes to phototransduction and light adaptation in mouse cone photoreceptors

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Light adaptation of photoreceptor cells is mediated by Ca^{2+} dependent mechanisms. In darkness, Ca²⁺ influx through cGMP-gated channels into the outer segment of photoreceptors is balanced by Ca²⁺ extrusion via Na⁺/Ca²⁺, K⁺ exchangers (NCKXs). Light activates a G protein signaling cascade, which closes cGMP-gated channels and decreases Ca²⁺ levels in photoreceptor outer segment because of continuing Ca²⁺ extrusion by NCKXs. Guanylate cyclase-activating proteins (GCAPs) then up-regulate cGMP synthesis by activating retinal membrane guanylate cyclases (RetGCs) in low Ca²⁺. This activation of RetGC accelerates photoresponse recovery and critically contributes to light adaptation of the nighttime rod and daytime cone photoreceptors. In mouse rod photoreceptors, GCAP1 and GCAP2 both contribute to the Ca²⁺-feedback mechanism. In contrast, only GCAP1 appears to modulate RetGC activity in mouse cones because evidence of GCAP2 expression in cones is lacking. Surprisingly, we found that GCAP2 is expressed in cones and can regulate light sensitivity and response kinetics as well as light adaptation of GCAP1-deficient mouse cones. Furthermore, we show that GCAP2 promotes cGMP synthesis and cGMP-gated channel opening in mouse cones exposed to low Ca²⁺. Our biochemical model and experiments indicate that GCAP2 significantly contributes to the activation of RetGC1 at low Ca²⁺ when GCAP1 is not present. Of note, in WT mouse cones, GCAP1 dominates the regulation of cGMP synthesis. We conclude that, under normal physiological conditions, GCAP1 dominates the regulation of cGMP synthesis in mouse cones, but if its function becomes compromised, GCAP2 contributes to the regulation of phototransduction and light adaptation of cones.

Guanylate cyclase-activating proteins (GCAPs)³ are EF-hand proteins that regulate cGMP synthesis by retinal membrane guanylate (guanylyl) cyclases (RetGCs) in a Ca²⁺-dependent manner (1-6). In low Ca²⁺, when the active EF-hand sites of the GCAP protein are not occupied by Ca²⁺, GCAPs activate Ret-GCs and promote the synthesis of cGMP. High Ca^{2+} blocks the activation of GC by GCAPs, and only a low basal level of cGMP synthesis is maintained in the cells. The presence of several GCAP isoforms in photoreceptor cells has been well-established (7–10). The diversity of GCAPs is particularly apparent in fish photoreceptors where at least seven different GCAP genes are expressed (7). Human photoreceptors express GCAP1-3, whereas only GCAP1 and GCAP2 are present in mouse photoreceptor cells (8, 10). Several mutations in the GUCA1A gene encoding for GCAP1 cause severe hereditary blinding diseases, including Leber congenital amaurosis, macular dystrophy, and cone-rod dystrophies (11-18). Although significant advances have been made in understanding the etiology of these diseases, it is still not clear why mutations in GUCA1A preferentially lead to cone, rather than rod, dystrophies and loss of daytime vision.

GCAP-mediated regulation of cGMP synthesis in the photoreceptors has been shown to be the single most important Ca^{2+} -mediated pathway of light adaptation (19, 20). In darkness, steady-state cGMP concentration in photoreceptor outer segments is maintained by a low basal synthesis of cGMP by RetGCs and its hydrolysis by phosphodiesterase 6 (PDE6). Light activates a G protein signaling cascade, leading to the increased hydrolysis rate of cGMP by PDE6 and a decline of the rod and cone outer segment cGMP concentration. Consequently, cGMP-gated channels in the outer segment plasma membrane close, leading to a decreased inflow of Na⁺ and Ca²⁺ into the outer segments (for a review, see Ref. 21). As Ca²⁺ ions are continuously extruded from outer segments by Na⁺/Ca²⁺, K^+ exchangers (22–25), the Ca^{2+} level drops, and Mg^{2+} replaces Ca^{2+} in the Ca^{2+}/Mg^{2+} -binding sites of GCAPs (26). The Mg²⁺-GCAPs activate RetGCs to accelerate cGMP synthesis, promoting the recovery of the photoreceptor cell to its

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³ The abbreviations used are: GCAP, guanylate cyclase–activating protein; RetGC, retinal membrane guanylate (guanylyl) cyclase; PDE, phosphodiesterase; mCAR, mouse cone arrestin; ERG, electroretinography; LED, lightemitting diode.

dark-adapted state after a transient light stimulus or preventing a closure of all cGMP-gated channels during continuous illumination.

It is believed that mouse rods express both GCAP1 and GCAP2, whereas mouse cones express only GCAP1 in their outer segments (9, 10). In rods, GCAP1 and GCAP2 regulate the cGMP synthesis in a relay fashion. Early in the photoresponse or at dim background light, when the Ca²⁺ level is only slightly lower than in darkness, GCAP1-mediated feedback dominates. Later in the photoresponse or at brighter background light, when Ca²⁺ drops to lower levels, GCAP2-mediated feedback is also engaged (19, 27). This model is consistent with the higher Ca²⁺ affinity of GCAP2 compared with GCAP1 (18, 28, 29). Although previous studies have suggested that GCAP2 may not be substantially present in normal or $Nrl^{-/-}$ mouse cone outer segments (10, 30, 31), direct genetic and functional approaches have not been used to test whether GCAP2 has any physiological role in mouse cones. Here, we aimed to determine the contribution of GCAP1 and GCAP2 in mouse cone phototransduction and light adaptation by using a comprehensive electrophysiology, genetic, biochemistry, and single-cell immunohistochemistry study.

Results

GCAP1 and GCAP2 are expressed in mouse cones

GCAP1 is expressed in outer segments of vertebrate rods and cones from zebrafish to human (8, 9). However, the expression pattern of GCAP2 varies among different species (9). Previous studies have shown contradicting results regarding its presence in mouse photoreceptors (9, 10, 32). Thus, we sought to determine the expression pattern of GCAP2 in mouse cones by single-cell immunohistochemistry in retinas from wildtype (WT) control, $Gcap1^{-/-}$, $Gcap2^{-/-}$, and GCAP1/2 double knockout ($Gcaps^{-/-}$) mice. The *top two panels* of Fig. 1 demonstrate expression of GCAP2 in WT control and Gcap1^{-/-} cones based on the colocalization of mouse cone arrestin (mCAR; green) and GCAP2 (red) antibodies. Additional GCAP2 signal around the cones is from rod photoreceptors that sometimes surrounded the cones even after the mechanical cell isolation (see "Experimental procedures"). We observed overlap between the cone arrestin and GCAP2 signals in both WT control and $Gcap1^{-/-}$ cones, suggesting that GCAP2 is expressed in mouse cones. As expected, the GCAP2 signal was not observed in $Gcap2^{-/-}$ or $Gcaps^{-/-}$ cones (Fig. 1, bottom two panels), thereby confirming the specificity of the GCAP2 antibody (33). Together, these results clearly demonstrate that GCAP2 is expressed in mouse cones.

GCAP1 and GCAP2 regulate the kinetics and sensitivity of mouse cone phototransduction

To determine the specific roles of GCAP2 and GCAP1 in mouse cone phototransduction, we compared light responses of dark-adapted cones from WT control, $Gcap1^{-/-}$, $Gcap2^{-/-}$, and $Gcaps^{-/-}$ mice using *ex vivo* electroretinography (ERG) recordings. To isolate the cone photoreceptor component of the *ex vivo* ERG signal, we used synaptic blockers and Ba²⁺ to remove b- and c-waves and rod-saturating background light.

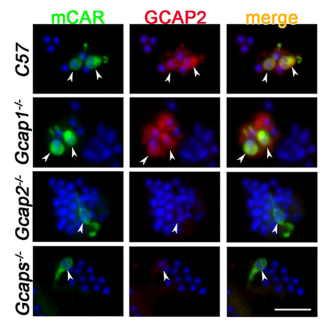


Figure 1. GCAP2 is expressed in the mouse cones. Dissociated retinal cells from mice of the indicated genotypes were incubated with the mCAR antibody (green) to label cones followed by incubation with GCAP2 antibody (red). Arrowheads point to the positions of the cones in each field. Nuclei were stained with DAPI (blue). Images shown are representative of 38 cones from seven fields (Gcap1^{-/-}), 10 cones from seven fields (Gcap2^{-/-}), and nine cones from three fields (Gcaps^{-/-}). Scale bar, 20 μ m.

Key experiments and the light adaptation studies were also done in a $Gnat1^{-/-}$ genetic background to remove the rod component of the ERG signal. As has been shown previously (19), simultaneous deletion of GCAP1 and GCAP2 slowed down light response recovery and increased the sensitivity of cones to light flashes (see Fig. 2d and Table 1). Removal of GCAP1 alone increased time to peak (t_n) of the responses elicited by dim light (Fig. 2, d and h, and Table 1) and increased the sensitivity of cones almost as much as the deletion of both GCAP1 and GCAP2 (Fig. 2d and Table 1). However, the recovery kinetics of the late tail phase of the responses in $Gcap1^{-/-}$ cones was not decelerated for both dim flashes (Fig. 2d) and bright saturating flashes (Fig. 2e). Afterdepolarization, or response recovery overshoot, which was often present both in control and GCAP-deficient cones, prevented us from fitting an exponential function to the late tail phase of the responses to estimate the response recovery time constant (τ_{rec}). However, the faster overall kinetics of $Gcap1^{-/-}$ cone dim flash responses as compared with that of $Gcaps^{-/-}$ cones was demonstrated by their shorter integration time when compared with Gcaps mice (Table 1). Isolating the cone component of the response by using $Gnat1^{-/-}$ mice confirmed that the responses from GCAP1-deficient cones are still substantially faster than these of cones lacking both GCAP1 and GCAP2 (Fig. 2, f and g). These results suggest that GCAP2 can shape the light response kinetics specifically in brighter light, at least in the absence of GCAP1. In contrast, the sensitivity of dark-adapted cones to dim light flashes appears to be mediated mainly by GCAP1 (Fig. 2h and Table 1). We also recorded light responses from $Gcap2^{-/-}$ mice (Fig. 2c) but did not find any significant changes of response kinetics or light sensitivity of GCAP2-deficient cones compared with WT controls (Fig. 2, d, e, and h,



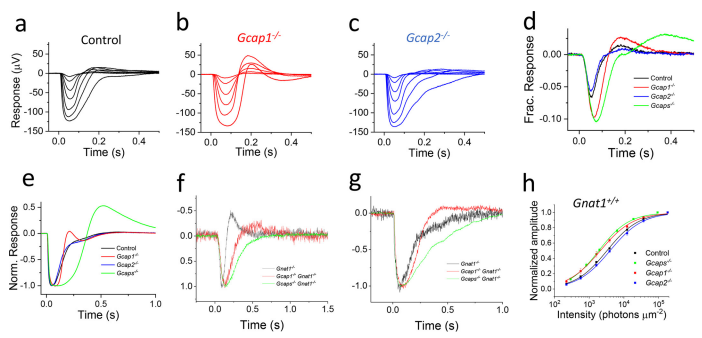


Figure 2. GCAP1 and GCAP2 regulate mouse cone phototransduction. a-c, responses of dark-adapted cones to 1-ms flashes of light with intensity, I_{Fr} , ranging from 220 to 183,000 photons (530 nm) μ m⁻² in the presence of rod-saturating background light from isolated WT control (*a*), $Gcap1^{-/-}$ (*b*), and $Gcap2^{-/-}$ (*c*) mouse retinas. *d*, averaged responses of control (*black*), $Gcap1^{-/-}$ (*red*), $Gcap2^{-/-}$ (*blue*), and $Gcaps^{-/-}$ (*green*) mouse cones to a 220 photons μ m⁻² flash normalized with r_{max} . *e*, saturated responses of control (*black*), $Gcap1^{-/-}$ (*red*), $Gcap2^{-/-}$ (*blue*), and $Gcaps^{-/-}$ (*green*) mouse cones to the 183,000 photons μ m⁻² flash normalized (*Norm.*) with r_{max} . *f* and *g*, normalized dim flash (*f*) and saturated (*g*) light responses recorded from dark-adapted retinas of control (*black*), $Gcap1^{-/-}$ (*green*) mice that were bred on a $Gnat1^{-/-}$ background are shown. *h*, the *smooth* traces plot Equation 1 with $I_{1/2}$ of 3,200, 1,900, 2,100, and 4,000 photons μ m⁻² fitted to the average response amplitude data (r/r_{max} as a function of I_F) of each genotype. *Error bars* give S.E. n = 3 mice (six retinas) for each genotype.

Table 1

Light (flash) response parameters from WT, Gcaps^{-/-}, Gcap1^{-/-}, and Gcap2^{-/-} mouse cones

All recordings were from $Gnat1^{+/+}$ mice except for the light adaptation parameters I_0 and n, which were obtained from $Gnat1^{-/-}$ mice. Retinas were exposed to constant 70,000 photons (530 nm) $\mu m^{-2} s^{-1}$ background light to suppress the rod component of the response except in the light adaptation experiments that were from $Gnat1^{-/-}$ mice. r_{max} , saturated photoresponse amplitude ($I_F = 183,000$ photons $\mu m^{-2} a t 530$ nm); t_p , time to peak ($I_F = 220$ photons $\mu m^{-2} a t 530$ nm); t_p , integration time defined as an area under a dim flash response divided by the amplitude of the response; I_{ip} , light flash intensity eliciting a response with peak amplitude $r = 0.5r_{max}$ determined by fitting Equation 1 to the response flactor determined by fitting Equation data; n, steepness factor determined by fitting Equation data.* and + indicate statistically significant difference as compared with the WT control and $Gcaps^{-/-}$ mouse cones, respectively (p < 0.5, two-tailed Student's t test). NA, not available.

Genotype	r _{max}	t_p	t_i	$I_{1/2}$	I_0	п
	μV	ms	ms	photons μm^{-2}	photons μm^{-2}	
WT	144 ± 15	53 ± 1	51 ± 0.1	$3,200 \pm 130$	$14,500 \pm 3,000$	1.0 ± 0.1
$GCAPs^{-/-}$	156 ± 26	$76 \pm 3^{*}$	90 ± 9*	$1,900 \pm 150^{*}$	$3,600 \pm 800^{*}$	$1.5 \pm 0.03^{*}$
GCAP1 ^{-/-}	135 ± 6	$66 \pm 3^{*}$	$64 \pm 4^{*+}$	$2,100 \pm 320^{*}$	$6,900 \pm 900^*$	$1.0 \pm 0.02 \dagger$
$GCAP2^{-/-}$	125 ± 18	$52\pm2^+$	49 ± 21	4,000 ± 450†	ŃA	NA

and Table 1). Thus, we conclude that GCAP1 can support normal cone photoresponses in the absence of GCAP2.

GCAP2 promotes cGMP synthesis in low Ca^{2+} in mouse cones

Biochemical experiments have demonstrated that GCAP proteins activate cGMP synthesis of RetGCs in low Ca²⁺ (1, 6). Here, we asked whether GCAP2 could promote cGMP synthesis in intact mouse cones. To assess the Ca²⁺-mediated acceleration of cGMP synthesis in cones, we determined the change of the maximal saturated cone photoresponse amplitude (r_{max}) when the retinas were switched from normal perfusion solution with 1.2 mM [Ca²⁺]_o to low ~30 nM [Ca²⁺]_o in *ex vivo* ERG experiments. Such a treatment causes rapid reduction in the level of Ca²⁺ in photoreceptor outer segments and the subsequent GCAP-mediated up-regulation of cGMP synthesis (41). The r_{max} is proportional to the cGMP-gated channel current, and thus, increased cGMP concentration caused by accelerated

cGMP synthesis rate is expected to increase r_{max} . We determined r_{max} from saturated cone responses elicited by periodic bright test flashes in dark-adapted mouse retinas before and after low-Ca²⁺ exposure. In control $Gnat1^{-/-}$ retinas, r_{max} increased about 4-fold after a low-Ca²⁺ exposure (Fig. 3, black squares), demonstrating the up-regulation of cGMP synthesis and subsequent opening of the cGMP-gated channels. However, the cells could not maintain such a high cGMP-gated (CNG) channel channel current for long, and eventually r_{max} declined under low Ca^{2+} . When cones lacking both GCAP1 and GCAP2 (from $Gcaps^{-/-} Gnat1^{-/-}$ retinas) were exposed to low Ca²⁺, a much more subtle increase of r_{max} was observed (Fig. 3, green squares), consistent with the lack of up-regulation of cGMP synthesis in low Ca^{2+} in the absence of both GCAPs. Notably, when we exposed $Gcap1^{-/-} Gnat1^{-/-}$ retinas to low ${\rm Ca}^{2+}$, we observed substantial increase in $r_{\rm max}$ that was comparable with that in control $Gnat1^{-/-}$ mice (Fig. 3, red squares).

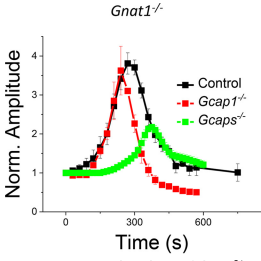


Figure 3. GCAP2 promotes CNG channel current in low Ca²⁺. Normalized (*Norm.*) r_{max} , the saturated photoresponse amplitude of dark-adapted cones, of control *Gnat1^{-/-}* (*black*), *Gcaps^{-/-} Gnat1^{-/-}* (*green*), and *Gcap1^{-/-} Gnat1^{-/-}* (*red*) mice in normal Ca²⁺ (at t = 0 s) and during low-Ca²⁺ exposure (t > 0 s) is shown. The values for r_{max} were normalized to their respective value in normal Ca²⁺ just before the switch to low Ca²⁺ at t = 0 s. n = 3 mice (six retinas) for each genotype. *Error bars* give S.E.

These results demonstrate that Ca^{2+} feedback mediated by GCAP2 can promote acceleration of the cGMP synthesis in intact mouse cones in the absence of GCAP1.

GCAP2 contributes to mouse cone light adaptation in bright background light

GCAP-mediated Ca²⁺ feedback dominates the regulation of rod and cone photoreceptor sensitivity in response to fast increments or decrements of background light (19, 34). However, the distinct contributions of GCAP1 and GCAP2 to the light adaptation capacity of mouse cones is not known. To address this question, we determined how the sensitivity of cones is regulated by background light in isolated retinas from control mice expressing both GCAPs and from mice lacking either both GCAPs or only GCAP1. All mice were on a $Gnat1^{-/-}$ background to eliminate rod signaling and facilitate the quantification of cone light adaptation. When mouse cones are exposed to a step of light, they produce an initial hyperpolarizing response peak followed by partial relaxation to a plateau (Fig. 4, a-c). This relaxation was attenuated after removal of both GCAPs (Fig. 4b), consistent with the dominant role of the GCAP-mediated feedback in cone light adaptation. Notably, GCAP1-deficient cones exhibited prominent relaxation after the peak of the response comparable with that in control cones, indicative of efficient light adaptation. We quantified the relaxation magnitude and kinetics by fitting a sum of two exponential functions from the peak to the plateau of the step responses using Equation 3 (see Fig. 4, a-c). Although we used a two-exponential function, the relaxation was dominated by the exponential term with the faster of the two time constants (τ_1) . Thus, we used τ_1 to assess the kinetics of relaxation and the amplitude from peak to the plateau (*A*) normalized by the peak amplitude (r_0) to assess the magnitude of relaxation (see Equation 3). The A/r_0 was similar between control (76 \pm 1%) and $Gcap1^{-/-}$ (80 ± 1%) mice but significantly smaller in the

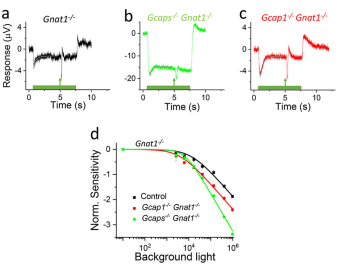


Figure 4. GCAP1 and GCAP2 contribute to the light adaptation capacity of cones. *a*–*c*, responses of cones to 7-s steps of light with a 1-ms flash delivered 4.5 s after the step onset from isolated retinas of control *Gnat1^{-/-}*(*a*), *Gcaps^{-/-}Gnat1^{-/-}*(*b*), and *Gcap1^{-/-}Gnat1^{-/-}*(*c*) mice obtained using *ex vivo* ERG recordings. *Smooth gray* traces plot Equation 3 with best fitting parameters *A*₁, *τ*₁, and *τ*₂. See "Results" for numerical values and statistical analysis. *d*, sensitivity (*S_F*) normalized (*Norm.*) with the dark-adapted sensitivity (*S_{F,D}*) of cones as a function of background light intensity (*I*) in control *Gnat1^{-/-}*(*black*), *Gcaps^{-/-}Gnat1^{-/-}*(*green*), and *Gcap1^{-/-}Gnat1^{-/-}*(*red*). *Smooth* lines plot Equation 2 with *l*₀ of 13,500 photons μ m⁻² s⁻¹ (*n* = 1), 6,700 photons μ m⁻² s⁻¹ (*n* = 1.4), and 4,100 photons μ m⁻² s⁻¹ (*n* = 1) for control *Gnat1^{-/-}*(*black*), *Gcaps^{-/-}Gnat1^{-/-}*(*green*), and *Gcap1^{-/-}Gnat*.

absence of both GCAP1 and GCAP2 (27 \pm 5%). This result demonstrates that the expression of GCAP2 in GCAP1-deficient cones was sufficient to promote robust light adaptation as demonstrated by the substantial relaxation of their response in steady background light. However, the kinetics of the relaxation was decelerated significantly by the deletion of GCAP1 alone (from 165 \pm 30 ms in control to 495 \pm 20 ms in *Gcap1^{-/-}* mice), whereas the value for τ_1 was not statistically significantly different in *Gcaps^{-/-}* cones (423 \pm 30 ms) as compared with that in *Gcap1^{-/-}* cones. This result is consistent with the dominant role of GCAP1 in driving the rapid light adaptation of mouse cones.

To quantify the efficiency of light adaptation, we measured the sensitivity of cones to light flashes at 4.5 s after the step onset at different background light intensities. The sensitivity normalized to the sensitivity in darkness declined more steeply in $Gcaps^{-/-}$ cones than in control or $Gcap1^{-/-}$ cones (Fig. 4d). As expected, based on their higher sensitivity in darkness, the operating range of $Gcap1^{-/-}$ cones was shifted to dimmer background light intensities. However, the slope of the adaptation curve was not changed, and the adaptation capacity was clearly better in $Gcap1^{-/-}$ mice than in $Gcaps^{-/-}$ mice. These results indicate that GCAP2 can contribute to the light adaptation of mouse cones in the absence of GCAP1. We did not have $Gcap2^{-/-}Gnat1^{-/-}$ mice. Thus, in an effort to investigate the role of GCAP2 in light adaptation, we compared light adaptation between $Gcap2^{-/-}$ and WT mice (on $Gnat1^{+/+}$ background). In those experiments, we did not observe any change in light adaptation caused by the deletion of GCAP2 (data not shown), consistent with our flash response data showing only negligible phenotype in GCAP2-deficient cones (Fig. 2, c-e and h).



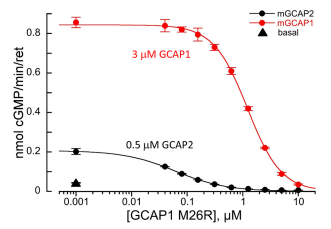


Figure 5. GCAP1 and GCAP2 compete for the activation of RetGC1. The native RetGC1 activity in *Gcaps^{-/-} RetGC2^{-/-}* mouse retinas was assayed as described under "Experimental procedures" in the absence of GCAP5 (\blacktriangle) or in the presence of 3 μ M mouse GCAP1 (*red circles*) or 0.5 μ M GCAP2 (*black circles*). Variable concentrations of the competing bovine M26R GCAP1 were added to the assay as indicated. The data, average \pm S.D. (*error bars*) of two to four independent measurements, were fitted assuming a sigmoidal Hill function.

GCAP1 and GCAP2 compete for activation of RetGC1 in low Ca²⁺

Our results demonstrate that GCAP2 is expressed in mouse cones and that it can contribute to the Ca²⁺-dependent activation of RetGCs and phototransduction feedback in their outer segments. However, it remained unclear whether the expression level of GCAP2 in WT cones is sufficient to contribute to the overall Ca²⁺ feedback. A simple biochemical model (see "Experimental procedures" for details) predicts that a quite small concentration of GCAP2, \sim 0.1–0.5 μ M in the cone outer segment, could explain the ~4-fold increase of $r_{\rm max}$ in low Ca²⁺ observed in $Gcap1^{-/-}$ cones (see Fig. 3). Based on the model prediction, we designed a biochemical experiment to assess the extent of activation of the native RetGC1 (the predominant guanylate cyclase isozyme expressed in the cones (30, 31, 35)) by recombinant GCAP1 and GCAP2. We used photoreceptor membranes from *Gcaps^{-/-} RetGC2^{-/-}* mouse retinas retaining only RetGC1 isozyme to measure cGMP synthesis by RetGC1 in low Ca^{2+} at normal physiological 0.9 mM Mg^{2+} (36). Consistent with our model, the low basal activity of RetGC1 was significantly increased by addition of either 0.5 µM GCAP2 (derived from the biochemical model) or 3 μ M GCAP1 (the estimated GCAP1 concentration in mouse rods (29)) (Fig. 5). Next, we assessed whether GCAP2 can contribute to the regulation of RetGC1 activity in the presence of GCAP1. To test this, we used M26R GCAP1, a mutant form that can bind to RetGC1 like the WT GCAP1 but does not activate it (37, 38). In the presence of 3 µM GCAP1, addition of M26R GCAP1 started to decrease RetGC1 activity at $\sim 0.3 \ \mu\text{M}$ (Fig. 5, red circles), its near-physiological concentration (28, 29), and reached halfmaximal inhibition at 1 μ M. At the same concentration of M26R GCAP1, the activation of RetGC1 by 0.5 µM GCAP2, which in the absence of GCAP1 would be sufficient to effectively accelerate RetGC1 in vivo (Figs. 2 and 3), was almost completely suppressed (Fig. 5, black circles).

Discussion

Ca²⁺-dependent regulation of cGMP synthesis by GCAP2 in mouse cone photoreceptors

Our experiments clearly demonstrate that GCAP2 is expressed in mouse cones (Fig. 1). To address the possible functional role of GCAP2 in cones, we investigated its ability to up-regulate cGMP synthesis in low Ca^{2+} and to mediate light adaptation in cones lacking GCAP1. As previous studies have suggested that GCAP1 and RetGC1 dominate the synthesis of cGMP in the mouse cone outer segments, we expected that $Gcap1^{-/-}$ retinas would respond to low-Ca²⁺ exposure similarly to Gcaps^{-/-} retinas (9, 10, 30, 31, 35). However, Gcap1^{-/-} cones were able to boost their maximal response amplitude in low Ca^{2+} as much as control WT cones (Fig. 3). Based on our model presented under "Experimental procedures," as low a concentration as 0.1 µM GCAP2 in the outer segments of $Gcap1^{-/-}$ cones could explain the ~4-fold increase of their maximal response amplitude in low Ca^{2+} . This concentration is more than 10-fold lower than the known GCAP1 or GCAP2 concentration in mouse rod outer segments (28, 29). The quantitative power of these experiments might be limited due to the cooperativity of the CNG channel for cGMP (39, 40) or the transient nature of the increase in photoreceptor response amplitude in low Ca^{2+} (41–45). However, despite the quantitative limitations of our study, our results clearly demonstrate that GCAP2 can activate RetGC in mouse cone photoreceptor cells when GCAP1 has been deleted.

Similarly, when we examined light adaptation in $Gcap1^{-/-}$ cones, we found that the slope of the light adaptation curve was comparable with that in control cones. In addition, the adaptation capacity of GCAP1-deficient cones was substantially better than that of $Gcaps^{-/-}$ cones (Fig. 4). Together, these results demonstrate that GCAP2 is able to up-regulate cGMP synthesis and to mediate light adaptation in cones in the absence of GCAP1.

The role of GCAP1 and GCAP2 in cone phototransduction and light adaptation

The relative contribution of GCAP1 and GCAP2 in rod physiology has been established in mouse rod photoreceptors (27). There, GCAP1 is more important in determining the peak amplitude of the dim flash response, whereas GCAP2 shapes the response recovery kinetics after the peak amplitude. These results are consistent with the known biochemical properties of GCAP1 and GCAP2. Namely, GCAP2 has a higher affinity to Ca^{2+} ($K_{Ca} = 50$ nM) as compared with GCAP1 ($K_{Ca} = 130$ nM) (18, 28, 29). In darkness, Ca^{2+} concentration in mouse rod outer segment is \sim 250 nM, and it declines to \sim 20-50 nM in bright light (46). Hence, after a dim flash, Ca²⁺ dissociates first from GCAP1, and the GCAP1-mediated feedback dominates over the GCAP2-mediated pathway. Later, when Ca²⁺ has dropped to a lower level, it can also dissociate from GCAP2, up-regulating the GCAP2-mediated feedback to contribute to the recovery phase kinetics of the dim flash response. Notably, the primary target for GCAP1 in mouse photoreceptors is RetGC1, whereas regulation of the ancillary isozyme RetGC2 is carried out mostly by GCAP2 (47). Hence, in mouse rods, acti-



vation of the cyclase after the flash of light occurs first as activation of RetGC1 by GCAP1 followed by additional activation of RetGC1 and RetGC2 by GCAP2 (27). Here, we compared dark-adapted cone flash responses from WT, $Gcaps^{-/-}$, $Gcap1^{-/-}$, and $Gcap2^{-/-}$ mice to understand the relative contributions of GCAP1 and GCAP2 in determining the sensitivity and response kinetics of mammalian cones (Fig. 2). We found that deletion of GCAP1 causes a comparable increase of the sensitivity and GCAP2 (Fig. 2, *d* and *h*, and Table 1). Thus, just as in rods, GCAP1 seems to dominate the up-regulation of cGMP synthesis up to the peak of the dim flash response, and the sensitivity of cones is set almost completely by GCAP1.

Comparison of saturated bright flash responses from WT, $Gcaps^{-/-}$, $Gcap1^{-/-}$, and $Gcap2^{-/-}$ mice revealed that deletion of both GCAP1 and GCAP2 significantly delays the escape of cones from saturation, whereas the deletion of GCAP1 had a much less dramatic effect, and the deletion of GCAP2 had almost no effect at all on the recovery kinetics (Fig. 2e). Notably, the recovery kinetics of $Gcap1^{-/-}$ cones were not slower than those of WT cones so that cone responses from $Gcap1^{-/-}$ mice recovered to the baseline level at the same time as those of WT and $Gcap2^{-/-}$ mice (Fig. 2d). Thus, it appears that both GCAP1 and GCAP2 can compensate for the lack of the other isoform in accelerating the recovery of bright flash responses (Fig. 2e). These results are also consistent with the idea that a larger drop in Ca²⁺ caused by brighter light is required to activate the GCAP2 pathway. In support of this notion, we observed deviation between $Gcap1^{-/-}$ and $Gcaps^{-/-}$ mouse light adaptation only at brighter background light. This, again, suggests that GCAP2 is more important under brighter illumination and at lower Ca^{2+} .

Although our data clearly show that GCAP2 contributes significantly to the physiology of mouse cones in $Gcap1^{-/-}$ mice, it is not clear whether GCAP2 plays a role in the phototransduction and/or light adaptation of healthy WT cones. Evidently, GCAP2 is present in native mouse and $Nrl^{-/-}$ cones at much lower levels than in rods, whereas GCAP1 expression in cones is very strong (5, 10, 31). However, our functional data from $Gcap1^{-/-}$ mice could be explained even by a rather low $0.1-0.5 \ \mu\text{M}$ GCAP2 concentration in the absence of GCAP1. In contrast, our biochemical experiments assessing the relative contribution of the two GCAP isoforms show that, even at equal concentrations of the two GCAP isoforms, GCAP1 effectively outcompetes GCAP2 from RetGC1 (see Fig. 5 and Refs. 28 and 38). Assuming further that GCAP2 expression in mouse cones is lower than that of GCAP1, we conclude that under normal physiological conditions GCAP1 would dominate the regulation of cGMP synthesis in mouse cones. However, if the function of GCAP1 becomes compromised, GCAP2 should be able to effectively regulate the phototransduction feedback and light adaptation of cones.

Experimental procedures

Ethical approval

All experimental procedures were in accordance with the Guide for the Care and Use of Laboratory Animals and were

approved by the Institutional Animal Care and Use Committees at Washington University in St. Louis, Salus University, and University of Southern California.

Animals

WT C57Bl/6J control and age-matched adult mice devoid of guanylate cyclase–activating protein 1 ($Gcap1^{-/-}$ (48)), 2 ($Gcap2^{-/-}$ (49)), or both ($Gcaps^{-/-}$ (19)) were used in this study. The mutant strains were bred to the control C57Bl/6J background for several generations but were not siblings of the control mice. For some electrophysiology experiments, $Gcap1^{-/-}$ and $Gcaps^{-/-}$ mice were bred into a $Gnat1^{-/-}$ background to remove the rod-driven light responses (50). Mice were kept under a 12/12-h light/dark cycle and had free access to regular mouse chow and clean water.

Single-cell immunohistochemistry

Freshly dissected retinas from C57, Gcap1^{-/-}, Gcap2^{-/-}, and $Gcaps^{-/-}$ mice were washed in Ames' medium, placed on an ice-cooled glass slide with a few drops of cold Ames' buffer, and chopped with razor blade. Dissociated cells and small cell clumps were collected into 8-chamber slides (Lab-Tek®, catalog number 177445) that were precoated with wheat germ agglutinin (100 μ M wheat germ agglutinin was added to the wells and incubated for 1 h). After cells were collected into wells, equal volumes of formaldehyde (4% in PBS) were added. The slides were centrifuged for 10 min at 168 \times *g* to attach the cells to the glass surface. Cells were washed in PBS; blocked with 5% goat serum, 0.1% Triton X-100 in PBS for 1 h; and incubated overnight with a rabbit polyclonal anti-mCAR antibody (51) (1:700 in blocking buffer). The next day, slides were washed in PBS and incubated with a secondary anti-rabbit antibody to visualize mCAR-labeled cones. Following PBS washes, cells were incubated with biotinylated anti-GCAP2 antibody (33) (1:300 of 1 mg/ml in blocking buffer). The GCAP2 signal was visualized by Texas Red-avidin (1:200; Vector Laboratories). The cells were mounted in Vectashield with DAPI (Vector Laboratories). Fluorescence images were acquired using a Zeiss Axio Scope microscope using the same settings and exposure times for the different genotypes.

Ex vivo electroretinography

We used ex vivo ERG to assess the function of mouse cone phototransduction and light adaptation (52). Either a background light of 70,000 photons (530 nm) μ m⁻² s⁻¹ or $Gnat1^{-/-}$ genetic background (53) was used to remove the rod component of the ERG signal. The $Gnat1^{-/-}$ mouse rods do not respond to light but maintain normal morphology. The background light needed to fully saturate rods was surprisingly high and would have been expected to bleach a significant amount of pigments during our experiments. However, after about 10 min of exposure to the background light, the cone responses remained stable for up to at least 2 h (the longest experiment), potentially due to a balance between pigment bleaching and regeneration via the Müller cell (54) visual cycle pathway. Retinas were dissected from dark-adapted eyes under IR illumination and mounted to a custom-built ERG specimen holder described in Vinberg et al. (52). Flashes and steps of light



were provided by green LEDs (530 nm; Luxeon Rebel LED SR-01-M0090) via an inverted microscope light path where the condenser was replaced by a $10 \times$ objective forming a homogenous 2.35-mm spot of light at the sample. The intensity of the light stimulus was calibrated at the level of the sample by a photometer (Model 211, UDT Instruments). Retinas were perfused at 1 ml/min at 37 °C with bicarbonate-buffered Locke's solution containing 112 mM NaCl, 3.6 mM KCl, 2.4 mM MgCl₂, 1.2 mм CaCl₂, 10 mм HEPES, 20 mм NaHCO₃, 3 mм disodium succinate, 0.5 mM sodium glutamate, and 10 mM glucose. The solution was equilibrated with 95%O₂ and 5%CO₂ at 37 °C. Low-Ca²⁺ solution was prepared by using 0.1 mM CaCl₂ instead of 1.2 mM and adding 0.4 mM EGTA. Addition of EGTA caused acidification of the medium, and we used NaOH to equalize the pH of our normal Locke's and low-Ca²⁺ media. We estimate that the free $[Ca^{2+}]$ of the low- Ca^{2+} medium is ~ 30 nM in the presence of 2.4 mM Mg^{2+} (55).

A differential amplifier (DP-311, Warner Instruments) and Bessel filter (model 3382, Krohn-Hite Corp.) together with a DigiData 1440 digitizer and pCLAMP software (Axon Instruments) were used to acquire data at 10 kHz with a 300-Hz lowpass filter. Clampfit (Axon Instruments), Origin 9.0.0 (Originlab), and Excel (Microsoft) software were used to analyze and graph the data. A Naka-Rushton function was fitted to the response amplitude (r) data.

$$\frac{r}{r_{\max}} = \frac{I_F}{I_{1/2} + I_F}$$
(Eq. 1)

where $r_{\rm max}$ is the maximal saturated response amplitude, I_F is flash intensity, and $I_{1/2}$ is the light intensity (in photons μm^{-2}) required to elicit a half-maximal response. A modified Weber-Fechner function was fitted to light adaptation data.

$$\frac{S_F}{S_{F,D}} = \frac{I_0^n}{I_0^n + I^n}$$
(Eq. 2)

where S_F is the sensitivity of cones to a flash of light (I_F that elicits $r < 0.2r_{\text{max}}$) defined as r/I_F , $S_{F,D}$ is the sensitivity in darkness, I is the background light intensity (in photons $\mu \text{m}^{-2} \text{ s}^{-1}$), I_0 is the background light intensity in which $S_F = 0.5S_{F,D}$, and n is a factor determining the steepness of the adaptation curve.

A sum of two exponential functions was used to quantify the kinetics and magnitude of light response relaxation after the initial peak during light steps.

$$r(t) = r_0 + A_1(1 - e^{-\frac{t - t_d}{\tau_1}}) + (A - A_1)(1 - e^{-\frac{t - t_d}{\tau_2}})$$
(Eq. 3)

where r_0 is peak amplitude measured at t_d , A is amplitude measured from the peak to the steady-state plateau of the step response, A_1 is the fraction of recovery covered by the time constant τ_1 , and $(A - A_1)$ is the fraction of the recovery covered by the time constant τ_2 .

Biochemical model of RetGC1 activation by GCAP2

We used the following equations to model binding of Ca^{2+} to GCAP2 and binding/activation of RetGC1 by Ca^{2+} -free

GCAP2. The parameter values were taken from Peshenko *et al.* (28).

$$GCAP2 + 2Ca^{2+} \longleftrightarrow GCAP2-2Ca^{2+}$$
(Eq. 4)

$$\Rightarrow \text{GCAP2} = [\text{GCAP2}] = \frac{K_{\text{Ca}}^2}{K_{\text{Ca}}^2 + [\text{Ca}^{2+}]^2} [\text{GCAP2}]_{\text{total}}$$
(Eq. 5)

where $K_{Ca} = 50 \text{ nM}$ is the apparent dissociation constant of Ca²⁺ from GCAP2. We model the activation of RetGC1 by GCAP2 by assuming that only Ca²⁺-free GCAP2 can activate the RetGC1.

$$\mathsf{GC1} + \mathsf{GCAP2} \longleftrightarrow \mathsf{GC1} - \mathsf{GCAP2} \tag{Eq. 6}$$

$$\Rightarrow \text{GC1-GCAP2} = \frac{\text{GCAP2}}{K_{\text{GC1}} + \text{GCAP2}} \text{GC1}_{\text{total}} \qquad (\text{Eq. 7})$$

where $K_{\rm GC1} = 1.25 \ \mu \text{M}$ and $\rm GC1_{total} = 3.2 \ \mu \text{M}$. Cyclase activity (α ; in $\mu \text{M s}^{-1}$) can be calculated as follows.

$$\alpha = k_{n1} \operatorname{GC1} + k_{s1} \operatorname{GC1-GCAP2}$$
(Eq. 8)

if we assume that GTP (the substrate) $\gg K_{m(\text{GTP-GC})}$ (dissociation constant of the GTP from RetGC1). We assume that the basal RetGC1 activity $k_{n1} = 2.6 \text{ s}^{-1}$ and for the activated RetGC1 $k_{s1} = 33 \text{ s}^{-1}$ (28). Concentrations of GCAP2-free GC1 and GCAP2-bound GC1-GCAP2 in a specific [Ca²⁺] and [GCAP2] can be calculated from Equations 5, 6, and 7.

At steady state,

$$\alpha = \beta \text{ cGMP}$$
(Eq. 9)

$$\Rightarrow cGMP = cG = \frac{\alpha}{\beta}$$
 (Eq. 10)

where $\beta = 4.1 \text{ s}^{-1}$ is the spontaneous cGMP hydrolysis activity of rod PDE in darkness (56). The CNG channel current (57) can be calculated as follows.

$$J_{cG} = J_{max} \frac{cG^3}{cG^3 + (20 \ \mu\text{M})^3}$$
(Eq. 11)

where $J_{\rm max}$ is the CNG channel current at high [cGMP]. Assuming that [Ca²⁺] is 250 nM in a dark-adapted mouse cone outer segment under normal extracellular Ca²⁺ and declines to 25 nM during our low-Ca²⁺ exposure (see above), as low as a 0.1 μ M total concentration of GCAP2 in the cone outer segment is predicted to cause a 4.4-fold increase of $J_{\rm cG}$ when switched from normal (1.2 mM) Ca²⁺ to low Ca²⁺.

Expression and purification of GCAPs

We used recombinant mouse myristoylated GCAP1 (E6S) and GCAP2 expressed from pET11d vector (Novagen/Calbiochem) in BLR(DE3) *Escherichia coli* strain harboring yeast *N*-myristoyltransferase as described previously (28). GCAP2

was purified using urea extraction from the inclusion bodies and size-exclusion chromatography (26, 58). GCAP1 was purified using urea extraction and hydrophobic and size-exclusion column chromatography as described previously to reach a final protein of 95% purity by SDS-PAGE (28, 59). The M26R bovine GCAP1 mutant was produced and purified as described previously (37, 38).

RetGC assays

The native mouse RetGC1 activity was assayed under IR illumination in dark-adapted Gcaps^{-/-} RetGC2^{-/-} triple-knockout mouse retina homogenates isolated as described previously (28). Briefly, the assay mixture (25 μ l) containing retinal homogenate, 30 mM MOPS-KOH (pH 7.2), 60 mM KCl, 4 mM NaCl, 1 mм DTT, 2 mм EGTA, 0.9 mм free Mg^{2+} , 0.3 mм ATP, 4 mм cGMP, 1 mM GTP, 1 μ Ci of [α -³²P]GTP, 100 μ M zaprinast and dipyridamole, 10 mM creatine phosphate, and 0.5 unit of creatine phosphokinase was incubated at 30 °C for 8 min, and the reaction was stopped by heat inactivation at 95° for 2 min. The resultant [³²P]cGMP product was separated by TLC using fluorescently backed polyethyleneimine cellulose plates (Merck) developed in 0.2 M LiCl and eluted with 2 M LiCl, and the radioactivity was counted using ScintiSafe liquid scintillation mixture (Thermo Fisher Scientific) with addition of 20% ethanol.

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