

Deactivation Mechanisms of Rod Phototransduction

The Cogan Lecture

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The absorption of photons in rods and cones of the retina activate homologous biochemical signaling cascades that lead to the electrical changes that subserve the first steps in vision. Persistent activity of the cascade interferes with the ability of the photoreceptor to signal the absorption of subsequent photons, ultimately limiting the photoreceptor's sensitivity and temporal resolution. This article summarizes recent work on transgenic and knockout mouse rods that has revealed the deactivation mechanisms essential for normal response recovery and how each of these processes contributes to the overall time course of the flash response of rods. (*Invest Ophthalmol Vis Sci.* 2010;51:1283-1288) DOI:10.1167/iovs.09-4366

Vision begins when a photon of light is absorbed in the retinal photoreceptor cells and activates a series of biochemical reactions known as the phototransduction cascade. Photon absorption occurs within the photoreceptor outer segments, cylindrical subcellular compartments containing a stack of intracellular membranes called discs, that house the molecular machinery of phototransduction. The most abundant protein of the disc membranes is the light receptor itself, the G protein-coupled receptor, rhodopsin. A photon of sufficient energy photoisomerizes rhodopsin's covalently attached chromophore, 11-*cis* retinal, to its all-*trans* form, causing the protein to undergo a conformational change to an active state, metarhodopsin II (R^{*}).¹ R^{*} binds and activates the heterotrimeric G protein, transducin (G_t) by catalyzing GDP-GTP exchange on the α subunit at a rate of several hundred per second.² Each of the activated transducin α subunits (G_t α^*) binds the γ -subunit of the phosphodiesterase³ (PDE6), relieving PDE6's inhibition of PDE $\alpha\beta$ catalytic subunits,⁴ producing an active effector complex (G_t α -PDE^{*}) with a greatly increased rate of hydrolysis of cyclic GMP. The decrease in cGMP concentration due to G_t α -PDE^{*} activity rapidly leads to closure of cGMP-gated cation channels in the plasma membrane.⁵ The consequent decrease in inward cation current hyperpolarizes the cell, thereby reducing the rate of glutamate released from the photoreceptor terminal.

The light-evoked decrease in outer segment current, or photoresponse, persists until R^{*} and G_t α -PDE^{*} have deactivated and the cGMP levels have been restored. This review will discuss the molecular events that are essential for the recovery of the light-evoked changes in membrane current.

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KEY KINETIC FEATURES OF RESPONSE RECOVERY

The phototransduction cascade has long been recognized to produce a photoresponse with remarkably short latency, while having a slower offset that is approximately exponential in nature.⁶ The time course of the mammalian rod photoresponse is roughly 10-fold faster than that of amphibians. For suction electrode recordings made from small pieces of isolated mouse retina, like those described here, the time to peak of the dim flash response of a healthy rod is \sim 100 ms and the recovery time constant, \sim 200 ms. In vivo electroretinogram (ERG) recordings in mice have revealed nearly identical time to peak⁷ and recovery time constant.⁸

The kinetics of the photoresponse are remarkably consistent within a given rod from trial to trial and across a wide range of flash strengths. In a mouse rod, the response to a single photon typically reaches a peak amplitude of approximately 0.5 pA. Brighter flashes produce responses that are larger in amplitude, until all the cGMP channels are closed, and the response reaches a maximum, or saturating, amplitude. Further increases in flash strength produce more cascade activity, but no additional increase in amplitude. Rather, the responses remain in saturation for longer times. Plotting the time that a bright-flash response remains in saturation as a function of the natural log of the flash strength (the so-called Pepperberg plot)⁹ yields a linear relation for up to \sim 3000 photoisomerized rhodopsin molecules in mouse rods.^{8,10} The slope of this linear relation is the dominant recovery time constant, τ_D , which is remarkably similar (\sim 200 ms) to the time constant fitted empirically to the final falling phase of the response to dim flashes (so-called τ_{rec}). The correspondence of τ_{rec} and τ_D suggests that the same first-order deactivation step rate-limits recovery from both dim and bright flashes.¹⁰ The molecular identity of this slowest deactivation step was the subject of much study and debate for more than 15 years. Identification of the biochemical steps that are essential for recovery was necessary before it could be determined which step was the slowest and rate-limiting one.

ESSENTIAL DEACTIVATION STEPS FOR PHOTORESPONSE RECOVERY

cGMP Synthesis and the Role of Calcium Feedback to GCAPs/GCs in Mouse Rods

In order for the electrical response to recover, the cGMP-dependent channels must reopen, and for this to occur, the cGMP concentration must be restored. This requires that the rate of cGMP hydrolysis by PDE must decrease, and thus that R^{*}, G_t α -PDE^{*} all turn off. In addition, cGMP must be resynthesized by guanylate cyclase (GC-1 and GC-2 or GC-E and GC-F in mouse).¹¹ The rate of cGMP synthesis increases during the photoresponse, as the accompanying fall in intracellular calcium activates GC-1 and GC-2 through the concerted actions of guanylate cyclase activating proteins, or GCAPs.¹²⁻¹⁴ In normal

rods, calcium feedback to guanylate cyclase sufficiently speeds the rate of cGMP synthesis so that the flash response at late times well-approximates the time course of decline of the overall PDE activity, rather than being limited by the rate of cGMP synthesis. Evidence for this conclusion stems from experiments performed on mouse rods lacking GCAPs: Without calcium feedback to guanylate cyclase, the dim flash response is much larger and longer lasting than normal ($\tau_{\text{rec}} = 313$ ms), though the dominant time constant of recovery ($\tau_D = 240$ ms) is unaffected. These results indicate that the rate-limiting step in deactivation is normal in GCAP-knockout rods and that the dim flash response is larger and longer lasting than normal because of the slow rate of cGMP synthesis in the absence of calcium feedback.¹⁵

Rhodopsin Phosphorylation and Arrestin Binding

Since the early experiments of Bownds et al.¹⁶ and others,¹⁷⁻¹⁹ it has been known that after photoisomerization, rhodopsin becomes phosphorylated, and that after this phosphorylation, the protein arrestin (ARR1) binds with high affinity. Evidence that these deactivation steps must occur on the time scale of the flash response in the mouse was found in experiments on transgenic and knockout rods, which showed that either the absence of rhodopsin's C-terminal phosphorylation sites²⁰⁻²² or the absence of rhodopsin kinase (GRK1)²³ led to single-photon responses that were larger than normal, peeling away from the responses of wild-type rods along the rising phase. This finding indicated that R* activity was normally reduced by GRK1 within 70 ms.²³ Single-photon responses generated by unphosphorylated R*s typically maintained this larger amplitude for several seconds before abruptly turning off. On average, dim flash responses of rods lacking R* phosphorylation showed a τ_{rec} of 2 to 5 seconds.²⁰⁻²³ In response to bright flashes, final recovery was slower still, with a time constant of approximately 40 seconds.²⁴ Together, these results indicate that phosphorylation of R*s C-terminal residues are absolutely essential for normal response recovery, and that this phosphorylation must be mediated by GRK1 within 100 ms of the flash.²³ Whether other kinases contribute to R* phosphorylation on other time scales or illumination conditions remains unknown.

After phosphorylation by GRK1, ARR1 binds to R* with high affinity,²⁵ completely inhibiting its ability to bind and activate additional G_t molecules. Earlier experiments had shown that ARR1 was essential for the final quench in R*s activity: ARR1 knockout rods initially began to recover (presumably because of the effect of phosphorylation alone in reducing R*s catalytic activity^{26,27}), but then in the final phase recovered extremely slowly ($\tau_{\text{rec}} \sim 40$ seconds).^{24,26,28} This slow recovery time constant is probably due to the thermal decay of metarhodopsin II.²⁹ Together, these experiments indicate that both phosphorylation by GRK1 and the binding of ARR1 are essential for normal recovery of the rod flash response.

There are many unanswered questions about the role of phosphorylation and arrestin binding in controlling rhodopsin activity and how these processes may be altered during light adaptation. For example, GRK1 activity is inhibited by calcium-bound recoverin.^{30,31} This inhibition is relieved when calcium levels fall during steady illumination, resulting in more rapid rhodopsin deactivation.³² Phosphorylation of GRK1 by PKA³³ or by autophosphorylation^{34,35} alters GRK1 activity in vitro and could likewise modulate the rate of R* deactivation in vivo. Similarly, understanding the role of ARR1 binding in R* deactivation is made more complex by the expression of different ARR1 splice variants^{36,37} that have different binding properties and selectivity for R* in vitro.²⁵ Surprisingly, ARR1 splice variants²⁴ and an ideally engineered ARR1 mutant³⁸ are equally

efficient at deactivating unphosphorylated rhodopsin at the single-photon level, measured with suction electrodes. In contrast, the splice variants and the enhanced ARR1 mutant show functional rescue by ERG recordings³⁸ and retinal histology,^{24,38} suggesting that at higher light intensities or under in vivo conditions, there is an additional functional role for ARR1.

RGS9-Catalyzed GTP Hydrolysis of the G_tα/PDE Complex

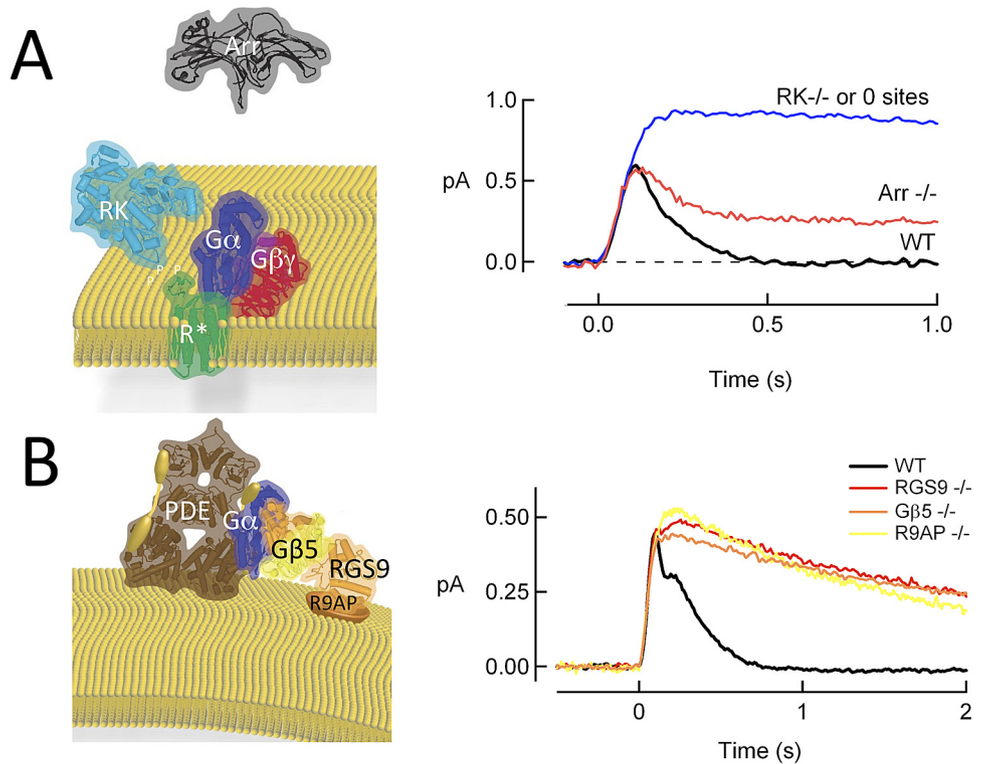
Like all heterotrimeric G proteins, transducin remains active until the α subunit hydrolyzes its bound GTP to GDP. In isolation, this GTP hydrolysis occurs far too slowly to account for the time constant of recovery of the flash response. In the 1990s, Ted Wensel's laboratory^{39,40} discovered that GTP hydrolysis by G_tα* is catalyzed by a photoreceptor-specific protein called RGS9-1 (regulator of G protein signaling, ninth family member, first splice variant). RGS9 also binds to the G protein β subunit Gβ5-L⁴¹ and R9AP (RGS9 anchoring protein),⁴² which holds RGS9/Gβ5-L with high affinity on the disc membrane. Deleting any one of these three genes (RGS9/Gβ5/R9AP) abolishes expression of the entire complex and the GTPase stimulating activity for G_tα* in vitro.^{10,43,44} The single-photon responses of each of these knockout rods are all very similar, recovering roughly 10 times slower than normal (Fig. 1).^{10,44,45} Thus, the RGS9 complex (hereafter, simply "RGS9") is absolutely essential for the normal deactivation of G_tα-PDE* and recovery of the light response in rods.

Despite the requirement for RGS9 in stimulating GTP hydrolysis, the fastest RGS9-hydrolysis occurs specifically when G_tα* is bound to PDEγ.⁴⁶ The requirement of PDEγ for rapid G_tα* deactivation was proposed to increase the gain of transduction by assuring that every G_tα* produced would bind and activate the effector before turning off.⁴⁷ Indeed, mutations in PDEγ that interfered with the ability of PDEγ to stimulate GTP hydrolysis also interfered with the ability of G_tα to bind PDE⁴⁸ and resulted in lower transduction gain and slow photoreceptor recovery.⁴⁹ More recent studies⁵⁰ tested this idea further by replacing the PDEγ-dependent photoreceptor splice variant of RGS9 (RGS9-1) with the more widely expressed neural splice variant RGS9-2, which stimulates GTP hydrolysis of G_tα*, regardless of whether PDEγ is bound.⁵¹ Surprisingly, the gain of transduction was wholly unaffected by expression of the PDEγ-independent RGS-2.⁵⁰ The likely explanation is that the rate by which G_tα* normally binds PDE is extremely high,⁵² whereas the rate of RGS-catalyzed GTP hydrolysis is normally slow (see below), so that virtually none of the G_tα*s hydrolyze GTP before they encounter PDE, even without the specialized co-requirement for PDEγ that seems a unique feature of the photoreceptor-specific RGS9-1.⁵⁰ The evolutionary selection that would seem to have specified the RGS9-1 isoform uniquely for photoreceptors remains unknown.

RATES OF DEACTIVATION STEPS IN INTACT RODS

Although the use of knockouts has proven to be enormously helpful in identifying the biochemical deactivation steps essential for normal recovery of the photoreponse, this approach cannot address the relative rates of these reactions under normal conditions. For example, the 40-second time constant of recovery observed in ARR1 knockout rods does not tell us how rapidly ARR1 normally acts, but rather reveals the time course of R* deactivation in the absence of ARR1, likely the time course of metarhodopsin II decay.²⁶ Likewise, the 10-second dominant time constant of recovery in RGS9-knockout rods does not reveal how RGS9 mediates normal recovery, but rather shows how rapidly G_tα-PDE* deactivation proceeds when the RGS9 complex is missing.¹⁰ To understand the rates

FIGURE 1. Deactivation steps essential for normal photoresponse recovery. (A) R* deactivation requires phosphorylation by GRK1 (RK) and the binding of ARR1. Traces are population-average, single-photon responses adapted or unpublished from previous studies.^{23,24} (B) G_tα-PDE* deactivation requires GTP hydrolysis that is stimulated by the RGS9 complex consisting of RGS9-1 (RGS9), Gβ5-L (Gβ5), and R9AP. Traces are population-average, single-photon responses adapted, or unpublished from previous studies.^{10,44,45} Crystal structures of each protein or enzyme exported from RCSBPDB Protein Data Bank (available at www.pdb.org) using the Protein Workshop viewer (available at www.rcsb.org/pdb/; both are a cooperative venture between Rutgers, Piscataway, NJ, and University of California, San Diego) for illustration. PDB Accession numbers were: Rh, 2L37; RK, 3C50; Arr1, 1CF1; RGS9 and Gβ5, 2PBI; PDEαβ, 1FL4; Gβ1γ1, 1TBG; Gat, 1TAD. The representations provided for PDEγ (*yellow barbell*) and R9AP (*orange disc*) are cartoons, because no crystal structures are yet available.



of R* and G_tα-PDE* deactivation under normal conditions, one needs to make more subtle perturbations in expression level or activity than simply deleting one or the other enzyme altogether. In truncated amphibian rods, this has been achieved by using nucleotide analogues and comparing the response “peel-away” times.⁵³ In intact rods, to determine whether R* deactivation or G_tα-PDE* deactivation is slower under normal conditions, it is easiest to try to speed up one or the other reaction and determine which one decreases the time constant of recovery.

Effect of RGS9 Complex Overexpression on Recovery of Both Dim and Saturating Flash Responses

For any first-order biochemical reaction scheme, when there is an excess of enzyme over substrate, the overall rate of the reaction varies linearly with the enzyme concentration. For the flashes used to determine τ_{rec} and τ_{D} , there is never more than a single photoisomerization per disc face and thus never more than approximately 20 or so G_tα-PDE*s per disc face, so that there is always excess enzyme (GRK1 or RGS9) available for the respective substrates (R* or G_tα/PDE*). Thus, overexpression of either of these two enzymes should, in principle, accelerate the rate of their reactions, and thus accelerate the rate of R* phosphorylation and RGS9-catalyzed GTP hydrolysis of the G_tα/PDE* complex. (Because recordings from rods expressing lower than normal levels of ARR1 showed responses with normal flash responses, it had been concluded that ARR1 binding does not rate limit recovery of the flash response.²⁶)

In collaboration with Ching-Kang Jason Chen’s group, we set out to test whether R* or G_tα-PDE* deactivation limits the time course of the light response in mouse rods. In a study of more than 20 different transgenic lines that expressed various levels of GRK1 and the RGS9 complex, none of the rods with increased GRK1 expression showed faster flash response kinetics.⁵⁴ Yet quantitative Western blot analysis and immunocytochemistry verified overexpression and outer segment lo-

calization, and, in vitro rhodopsin phosphorylation assays of GRK1-overexpressing rod outer segments, confirmed a greater rate and extent of rhodopsin phosphorylation in response to a full bleach. Furthermore, crossing the GRK1-overexpressing mice with GRK1-knockout mice yielded rods with flash responses that showed normal flash response kinetics. Together, all these results suggest that the elevated quantity of GRK1 in these rods was indeed functional, but did not result in a change in the flash response kinetics, indicating that binding of GRK1 to R* does not rate limit response recovery in normal rods.⁵⁴

In contrast, overexpression of the RGS9 complex resulted in dramatic speeding of recovery from both dim (τ_{rec}) and bright (τ_{D}) flashes.⁵⁴ Analysis of six different lines of mice that expressed the RGS9 complex at different levels over a 20-fold range (0.2× to 4×) showed clear dose-dependency of the response recovery: The greater the expression, the faster the recovery, with τ_{rec} reaching an apparent asymptote of 80 ms at the highest level of expression. Remarkably, for all the lines, recovery from bright, saturating responses (τ_{D}) showed identical concentration dependence, with perfect agreement in the time constant of recovery for a range of flash strengths from a single R* up through several thousand R*/flash.⁵⁴ These results provided unequivocal evidence that the same first-order process—namely, RGS9-catalyzed GTP hydrolysis of G_tα/PDE*, rate limits recovery of responses from the single-photon level, up through flashes that activate approximately 1 R* per disc face.⁵⁴

Exhaustion of Deactivation: The Limited Abundance of PDE

What accounts for the slowing of recovery at flashes that produce more than 1 R* per disc face? Previous work has suggested that the slowing at very bright flash strengths arises from the depletion of some essential deactivation enzyme,⁸ such as GRK1 or PDEγ/RGS9. Recent evidence suggests that indeed the “Pepperberg break” that occurs at ~8100 photons μm^{-2} (or two R*/disc face) arises when G_tα* is produced in

excess of PDE6, resulting in $G_t\alpha^*$ subunits that are uncomplexed with PDE γ , and thus hydrolyze GTP more slowly.⁵⁰

Michaelis Scheme Describing RGS9-Concentration Dependence: $V_{\max}/K_m = 1/\tau_D$

The striking concentration-dependence for RGS9-mediated recovery obtained by Krispel et al.⁵⁴ provides a unique opportunity to ask deeper questions about the mechanism of RGS9-catalyzed GTP hydrolysis in the intact rod. For example, does the rate of GTP hydrolysis itself become limiting for recovery when RGS9 expression is sufficiently high, or is the deactivation of R^* then rate limiting? In a recent investigation, the standard theoretical framework for rod phototransduction was expanded to incorporate a "Michaelis module" to describe the RGS9-dependent decay of $G_t\alpha/E^*$ activity (Fig. 2A). Solutions of the differential equations for this augmented scheme were able to precisely account for the dominant recovery rate over the 20-fold range of RGS9 expression levels in the rods of the Krispel study. Screening the parameter space of the augmented scheme with maximum likelihood methodology revealed that the dominant time constant of recovery follows the predicted tail-phase kinetics for the rate of the decline of substrate $G_t\alpha\text{-PDE}^*$ of a standard Michaelis scheme: the rate of recovery, ν ($=1/\tau_D$), was equal to V_{\max}/K_m for the RGS9 reaction.⁵⁵ In other words, the Michaelis module for RGS9-mediated deactivation of $G_t\alpha\text{-PDE}^*$ (Fig. 2A) was able to precisely account for the RGS9-concentration dependence of τ_D that was experimentally observed (Fig. 2B). The analysis also revealed that the value of τ_D (80 ms) for rods of the line with the highest level of RGS9 expression was *not* determined by the RGS9 turnover number, k_{cat} , but rather primarily limited by the RGS9-binding step.⁵⁵ In theory, if R^* 's lifetime is sufficiently short (see below), still higher levels of RGS9 expression could yield still faster photoresponse kinetics, providing the conditions of the Pepperberg analysis are still met (like translation invariance and time required for calcium to equilibrate at its minimum level^{9,56}). The fact that still higher levels of RGS9 expression could accelerate the response recovery further has important implications for the temporal regulation of cone responses, as cones express up to 10-fold higher levels of RGS9 than rods,^{57,58} and have response recovery kinetics that are likewise much faster than those of rods.^{58,59}

A Short R^* Lifetime: Implications for Reproducibility, Efficiency, and Mechanism

The maximum likelihood methodology was also used to test specific hypotheses about the other key rate constant in the theoretical scheme: that of rhodopsin deactivation ($1/\tau_R$; Fig. 2A). The results of these statistical tests showed that it is highly improbable that R^* 's lifetime exceeds 53 ms ($P < 0.05$); values of τ_R longer than this qualitatively failed to account for the vertical separation of the T_{sat} relations among the different RGS9-expressing lines⁵⁵ (Fig. 2B).

Such a short average lifetime of R^* has important implications for single-photon responses, since the amplitude and time course of the single-photon response is highly reproducible from trial-to-trial (coefficient of variation ~ 0.2).^{22,60-63} Some studies of reproducibility have asserted that R^* decay determines the overall time course of the single-photon response and thus that R^* decay is slow and must itself be reproducible,^{22,61,63} an uncommon and complex feat for a single molecule. The results of rods overexpressing the RGS9 complex reveal R^* deactivation to be much more rapid than $G_t\alpha\text{-PDE}^*$ decay,^{54,55} and therefore relatively inconsequential for the overall response time course in normal rods. Instead, other mechanisms most likely contribute to reproducibility, including highly cooperative feedback of calcium-dependent cGMP

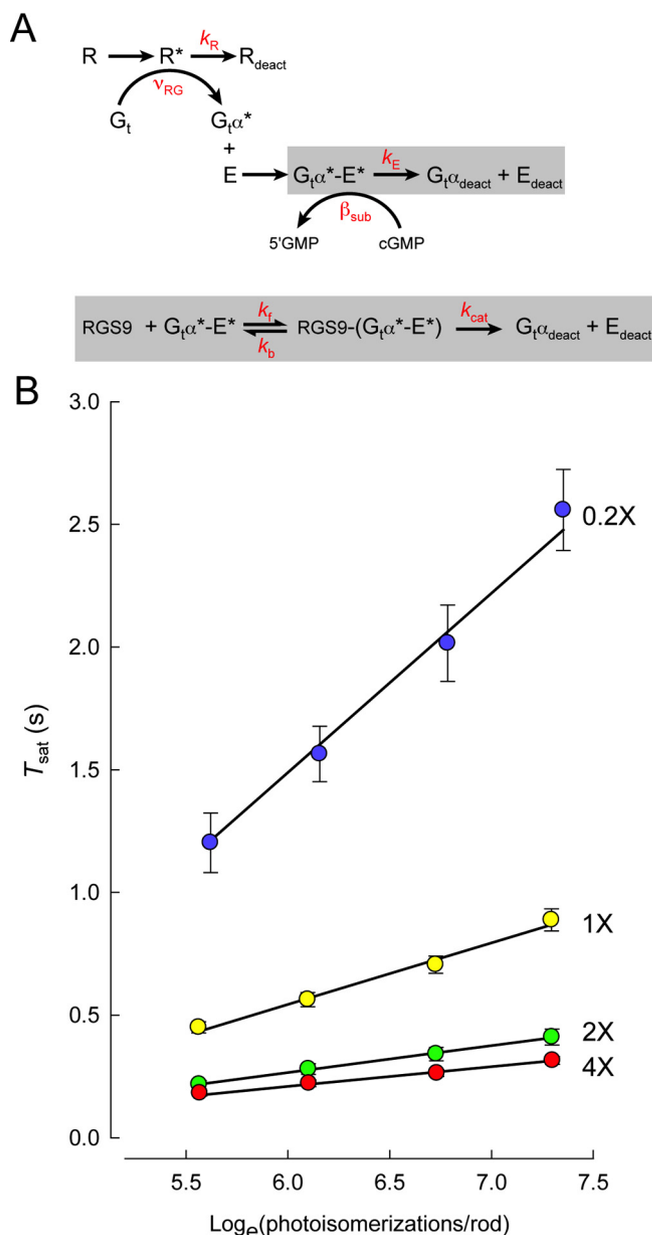


FIGURE 2. Michaelis-module for the RGS9-dependence of $G_t\alpha\text{-PDE}^*$ deactivation reveals the rate constants of RGS9 binding and catalysis, and constrains R^* lifetime. (A) Standard scheme for phototransduction, in which the Michaelis module for RGS9-mediated GTP hydrolysis (gray box below) was substituted for the first order decay of $G_t\alpha\text{-PDE}^*$ (gray box above). (B) The time that flash responses remained in saturation (T_{sat}) as a function of the natural log of the number of R^* (photoisomerizations) produced by each flash for mouse rods expressing a 20-fold range of RGS9 complex.⁵⁴ Error bars represent SEMs. Straight lines are the best-fitting curves produced using simplex searches of the solutions to the differential equations representing the expanded scheme in (A). Parameter values were $k_R = 33 \text{ seconds}^{-1}$, $k_f = 0.051 \mu\text{m}^2 \text{ s}^{-1}$, $k_b = 13.8 \text{ seconds}^{-1}$ and $k_{\text{cat}} = 52.8 \text{ seconds}^{-1}$. Reprinted with permission from Burns ME, Pugh EN Jr. RGS9 concentration matters in rod phototransduction. *Biophys J.* 2009;97:1538-1547. © 2009 Biophysical Society Published by Elsevier Inc.

synthesis,¹⁵ second-messenger diffusion,⁶⁴ and local saturation.^{63,64}

Another apparent consequence of a short R^* lifetime is that the signal transduction from GPCR to effector is nearly perfectly efficient. Because $G_t\alpha\text{-PDE}^*$ deactivation is normally

~five-fold slower than that of R* deactivation, nearly all the G_tα-PDE*s produced after photon absorption are for a time simultaneously active, so that their signal is maximally efficient. If G_tα-PDE* lifetime was shorter, or R* lifetime longer, a significant fraction of the G_tα-PDE* molecules would turn off during the activation phase, resulting in a net loss of signal.

Although it has long been accepted that R*phosphorylation and ARR1 binding must occur on the time scale of the flash response,⁶⁵ it is only since the experiments on genetically targeted mouse rods that the time scale for these reactions in intact photoreceptors has begun to be uncovered. Initial estimates concluded that it must occur within 100 ms of photon absorption,^{20,23} then complete within 80 ms,⁵⁴ and now the upper limit has been refined to 50 ms or less.⁵⁵ No in vitro studies of rhodopsin phosphorylation have yet measured phosphorylation on this time scale (all have sampled phosphorylation on a 30- to 10,000-fold slower time scale than the electrophysiological recordings of mammalian rods). However, it is known that GRK1 can bind R* within 10 ms⁵⁵ and that many other serine/threonine kinases have turnover numbers exceeding 30 seconds⁻¹, and even 500 seconds⁻¹.⁶⁶ Thus, there is a great deal more biochemical work to be done to directly measure the kinetics of the interactions of rhodopsin with its kinase and ARR1; more physiology experiments are needed to understand how these interactions shape R* activity and affect the time course of the light response, and many unanswered questions yet remain about how all these interactions might be altered under light-adapted conditions.

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References

- Wald G, Durell J, St George CC. The light reaction in the bleaching of rhodopsin. *Science*. 1950;111:179-181.
- Arshavsky VY, Lamb TD, Pugh EN Jr. G proteins and phototransduction. *Annu Rev Physiol*. 2002;64:153-187.
- Wensel TG, Stryer L. Activation mechanism of retinal rod cyclic GMP phosphodiesterase probed by fluorescein-labeled inhibitory subunit. *Biochemistry*. 1990;29:2155-2161.
- Hurley JB, Stryer L. Purification and characterization of the gamma regulatory subunit of the cyclic GMP phosphodiesterase from retinal rod outer segments. *J Biol Chem*. 1982;257:11094-11099.
- Karpen JW, Zimmerman AL, Stryer L, Baylor DA. Gating kinetics of the cyclic-GMP-activated channel of retinal rods: flash photolysis and voltage-jump studies. *Proc Natl Acad Sci U S A*. 1988;85:1287-1291.
- Hodgkin AL, Nunn BJ. Control of light-sensitive current in salamander rods. *J Physiol*. 1988;403:439-471.
- Hetling JR, Pepperberg DR. Sensitivity and kinetics of mouse rod flash responses determined in vivo from paired-flash electroretinograms. *J Physiol*. 1999;516:593-609.
- Lyubarsky A, Nikonov S, Pugh EN Jr. The kinetics of inactivation of the rod phototransduction cascade with constant Ca²⁺. *J Gen Physiol*. 1996;107:19-34.
- Pepperberg DR, Cornwall MC, Kahlert M, et al. Light-dependent delay in the falling phase of the retinal rod photoresponse. *Vis Neurosci*. 1992;8:9-18.
- Chen CK, Burns ME, He W, Wensel TG, Baylor DA, Simon MI. Slowed recovery of rod photoresponse in mice lacking the GTPase accelerating protein RGS9-1. *Nature*. 2000;403:557-560.
- Sokal I, Alekseev A, Palczewski K. Photoreceptor guanylate cyclase variants: cGMP production under control. *Acta Biochim Pol*. 2003; 50:1075-1095.
- Gorczyca WA, Gray-Keller MP, Detwiler PB, Palczewski K. Purification and physiological evaluation of a guanylate cyclase activating protein from retinal rods. *Proc Natl Acad Sci U S A*. 1994;91: 4014-4018.
- Dizhoor AM, Olshevskaya EV, Henzel WJ, et al. Cloning, sequencing, and expression of a 24-kDa Ca(2+)-binding protein activating photoreceptor guanylyl cyclase. *J Biol Chem*. 1995;270:25200-25206.
- Haeseleer F, Sokal I, Li N, et al. Molecular characterization of a third member of the guanylyl cyclase-activating protein subfamily. *J Biol Chem*. 1999;274:6526-6535.
- Burns ME, Mendez A, Chen J, Baylor DA. Dynamics of cyclic GMP synthesis in retinal rods. *Neuron*. 2002;36:81-91.
- Bownds D, Dawes J, Miller J, Stahlman M. Phosphorylation of frog photoreceptor membranes induced by light. *Nat New Biol*. 1972; 237:125-127.
- Kuhn H, Dreyer WJ. Light dependent phosphorylation of rhodopsin by ATP. *FEBS Lett*. 1972;20:1-6.
- Kuhn H, Hall SW, Wilden U. Light-induced binding of 48-kDa protein to photoreceptor membranes is highly enhanced by phosphorylation of rhodopsin. *FEBS Lett*. 1984;176:473-478.
- Wilden U, Hall SW, Kuhn H. Phosphodiesterase activation by photoexcited rhodopsin is quenched when rhodopsin is phosphorylated and binds the intrinsic 48-kDa protein of rod outer segments. *Proc Natl Acad Sci U S A*. 1986;83:1174-1178.
- Chen J, Makino CL, Peachey NS, Baylor DA, Simon MI. Mechanisms of rhodopsin inactivation in vivo as revealed by a COOH-terminal truncation mutant. *Science*. 1995;267:374-377.
- Mendez A, Burns ME, Roca A, et al. Rapid and reproducible deactivation of rhodopsin requires multiple phosphorylation sites. *Neuron*. 2000;28:153-164.
- Doan T, Mendez A, Detwiler PB, Chen J, Rieke F. Multiple phosphorylation sites confer reproducibility of the rod's single-photon responses. *Science*. 2006;313:530-533.
- Chen CK, Burns ME, Spencer M, et al. Abnormal photoresponses and light-induced apoptosis in rods lacking rhodopsin kinase. *Proc Natl Acad Sci U S A*. 1999;96:3718-3722.
- Burns ME, Mendez A, Chen CK, et al. Deactivation of phosphorylated and nonphosphorylated rhodopsin by arrestin splice variants. *J Neurosci*. 2006;26:1036-1044.
- Gurevich VV, Hanson SM, Gurevich EV, Vishnivetskiy SA, How rod arrestin achieved perfection: regulation of its availability and binding selectivity. In: Fliesler SJ, Kisselev O, eds. *Signal Transduction in the Retina*. CRC Press: Boca Raton, FL; 2007:55-88.
- Xu J, Dodd RL, Makino CL, Simon MI, Baylor DA, Chen J. Prolonged photoresponses in transgenic mouse rods lacking arrestin. *Nature*. 1997;389:505-509.
- Palczewski K, Rispoli G, Detwiler PB. The influence of arrestin (48K protein) and rhodopsin kinase on visual transduction. *Neuron*. 1992;8:117-126.
- Chan S, Rubin WW, Mendez A, et al. Functional comparisons of visual arrestins in rod photoreceptors of transgenic mice. *Invest Ophthalmol Vis Sci*. 2007;48:1968-1975.
- Cone RA, Cobbs WH, 3rd. Rhodopsin cycle in the living eye of the rat. *Nature*. 1969;221:820-822.
- Kawamura S. Rhodopsin phosphorylation as a mechanism of cyclic GMP phosphodiesterase regulation by S-modulin. *Nature*. 1993; 362:855-857.
- Chen CK, Inglese J, Lefkowitz RJ, Hurley JB. Ca(2+)-dependent interaction of recoverin with rhodopsin kinase. *J Biol Chem*. 1995;270:18060-18066.
- Makino CL, Dodd RL, Chen J, et al. Recoverin regulates light-dependent phosphodiesterase activity in retinal rods. *J Gen Physiol*. 2004;123:729-741.
- Horner TJ, Osawa S, Schaller MD, Weiss ER. Phosphorylation of GRK1 and GRK7 by cAMP-dependent protein kinase attenuates their enzymatic activities. *J Biol Chem*. 2005;280:28241-28250.
- Buczylko J, Gutmann C, Palczewski K. Regulation of rhodopsin kinase by autophosphorylation. *Proc Natl Acad Sci U S A*. 1991; 88:2568-2572.
- Pulvermuller A, Palczewski K, Hofmann KP. Interaction between photoactivated rhodopsin and its kinase: stability and kinetics of complex formation. *Biochemistry*. 1993;32:14082-14088.
- Palczewski K, Buczylko J, Ohguro H, et al. Characterization of a truncated form of arrestin isolated from bovine rod outer segments. *Protein Sci*. 1994;3:314-324.

37. Smith WC, Milam AH, Dugger D, Arendt A, Hargrave PA, Palczewski K. A splice variant of arrestin. molecular cloning and localization in bovine retina. *J Biol Chem.* 1994;269:15407-15410.
38. Song X, Vishnivetskiy SA, Gross OP, et al. Enhanced arrestin facilitates recovery and protects rods lacking rhodopsin phosphorylation. *Curr Biol.* 2009;19:700-705.
39. Angleson JK, Wensel TG. A GTPase-accelerating factor for transducin, distinct from its effector cGMP phosphodiesterase, in rod outer segment membranes. *Neuron.* 1993;11:939-949.
40. He W, Cowan CW, Wensel TG. RGS9, a GTPase accelerator for phototransduction. *Neuron.* 1998;20:95-102.
41. Makino ER, Handy JW, Li T, Arshavsky VY. The GTPase activating factor for transducin in rod photoreceptors is the complex between RGS9 and type 5 G protein beta subunit. *Proc Natl Acad Sci U S A.* 1999;96:1947-1952.
42. Hu G, Wensel TG. R9AP, a membrane anchor for the photoreceptor GTPase accelerating protein, RGS9-1. *Proc Natl Acad Sci U S A.* 2002;99:9755-9760.
43. Chen CK, Eversole-Cire P, Zhang H, et al. Instability of GGL domain-containing RGS proteins in mice lacking the G protein beta-subunit Gbeta5. *Proc Natl Acad Sci U S A.* 2003;100:6604-6609.
44. Keresztes G, Martemyanov KA, Krispel CM, et al. Absence of the RGS9.Gbeta5 GTPase-activating complex in photoreceptors of the R9AP knockout mouse. *J Biol Chem.* 2004;279:1581-1584.
45. Krispel CM, Chen CK, Simon MI, Burns ME. Prolonged photoreponses and defective adaptation in rods of Gbeta5^{-/-} mice. *J Neurosci.* 2003;23:6965-6971.
46. Skiba NP, Hopp JA, Arshavsky VY. The effector enzyme regulates the duration of G protein signaling in vertebrate photoreceptors by increasing the affinity between transducin and RGS protein. *J Biol Chem.* 2000;275:32716-32720.
47. Arshavsky V, Bownds MD. Regulation of deactivation of photoreceptor G protein by its target enzyme and cGMP. *Nature.* 1992;357:416-417.
48. Slepak VZ, Artemyev NO, Zhu Y, et al. An effector site that stimulates G-protein GTPase in photoreceptors. *J Biol Chem.* 1995;270:14319-14324.
49. Tsang SH, et al. Role for the target enzyme in deactivation of photoreceptor G protein in vivo. *Science.* 1998;282:117-121.
50. Martemyanov KA, Krispel CM, Lishko PV, Burns ME, Arshavsky VY. Functional comparison of RGS9 splice isoforms in a living cell. *Proc Natl Acad Sci U S A.* 2008;105:20988-20993.
51. Martemyanov KA, Hopp JA, Arshavsky VY. Specificity of G protein-RGS protein recognition is regulated by affinity adapters. *Neuron.* 2003;38:857-862.
52. Leskov IB, Klenchin VA, Handy JW, et al. The gain of rod phototransduction: reconciliation of biochemical and electrophysiological measurements. *Neuron.* 2000;27:525-537.
53. Sagoo MS, Lagnado L. G-protein deactivation is rate-limiting for shut-off of the phototransduction cascade. *Nature.* 1997;389:392-395.
54. Krispel CM, Chen D, Melling N, et al. RGS expression rate-limits recovery of rod photoresponses. *Neuron.* 2006;51:409-416.
55. Burns ME, Pugh EN Jr. RGS9 concentration matters in rod phototransduction. *Biophys J.* 2009;97:1538-1547.
56. Nikonov S, Engheta N, Pugh EN Jr. Kinetics of recovery of the dark-adapted salamander rod photoresponse. *J Gen Physiol.* 1998;111:7-37.
57. Cowan CW, Fariss RN, Sokal I, Palczewski K, Wensel TG. High expression levels in cones of RGS9, the predominant GTPase accelerating protein of rods. *Proc Natl Acad Sci U S A.* 1998;95:5351-5356.
58. Zhang X, Wensel TG, Kraft TW. GTPase regulators and photoreponses in cones of the eastern chipmunk. *J Neurosci.* 2003;23:1287-1297.
59. Nikonov SS, Kholodenko R, Lem J, Pugh EN Jr. Physiological features of the S- and M-cone photoreceptors of wild-type mice from single-cell recordings. *J Gen Physiol.* 2006;127:359-374.
60. Baylor DA, Lamb TD, Yau KW. Responses of retinal rods to single photons. *J Physiol.* 1979;288:613-634.
61. Rieke F, Baylor DA. Origin of reproducibility in the responses of retinal rods to single photons. *Biophys J.* 1998;75:1836-1857.
62. Whitlock GG, Lamb TD. Variability in the time course of single photon responses from toad rods: termination of rhodopsin's activity. *Neuron.* 1999;23:337-351.
63. Field GD, Rieke F. Mechanisms regulating variability of the single photon responses of mammalian rod photoreceptors. *Neuron.* 2002;35:733-747.
64. Bisegna P, Caruso G, Andreucci D, et al. Diffusion of the second messengers in the cytoplasm acts as a variability suppressor of the single photon response in vertebrate phototransduction. *Biophys J.* 2008;94:3363-3383.
65. Sather WA, Detwiler PB. Intracellular biochemical manipulation of phototransduction in detached rod outer segments. *Proc Natl Acad Sci U S A.* 1987;84:9290-9294.
66. Boyer PD, Krebs EG. Control by Phosphorylation. Vol. 17. *The Enzymes.* 3 ed. Academic Press; 1986.