A comprehensive model of the phototransduction cascade in mouse rod cells†

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Vertebrate visual phototransduction is perhaps the most well-studied G-protein signaling pathway. A wealth of available biochemical and electrophysiological data has resulted in a rich history of mathematical modeling of the system. However, while the most comprehensive models have relied upon amphibian biochemical and electrophysiological data, modern research typically employs mammalian species, particularly mice, which exhibit significantly faster signaling dynamics. In this work, we present an adaptation of a previously published, comprehensive model of amphibian phototransduction that can produce quantitatively accurate simulations of the murine photoresponse. We demonstrate the ability of the model to predict responses to a wide range of stimuli and under a variety of mutant conditions. Finally, we employ the model to highlight a likely unknown mechanism related to the interaction between rhodopsin and rhodopsin kinase.

1 Introduction

Visual phototransduction is the biochemical process by which a light stimulus is translated into a neuronal signal. The response is triggered by the absorption of photons of light by visual pigments, which then activate a prototypical G-protein signaling cascade.1,2 The first steps of phototransduction involve the binding of the heterotrimeric G-protein transducin (Gt) to the activated receptor, rhodopsin (R*). R* catalyzes the exchange of GTP for GDP bound to Gt, which results in the dissociation of the R*·Gt complex and the further dissociation of the Gt, subunit from the Gt heterotrimer.3 Gt is then free to bind and activate a phosphodiesterase (PDE), resulting in the hydrolysis of intracellular cGMP.4,5 This leads to the closure of cGMP-gated ion channels, a subsequent drop in the intracellular Ca2+ concentration due to its continued extrusion via Na+/Ca2+, K+ ion exchangers,6,7 and a consequent hyper-polarization of the cell membrane.

Deactivation of the pathway consists of several concurrent mechanisms. Decreasing Ca2+ concentrations induce a change in conformation of the protein recoverin (Rec),8 causing it to dissociate from rhodopsin kinase (RK).9,10 The kinase is then free to bind and multiply phosphorylate R*,11–13 Increasing phosphorylation levels of R* lead to decreased binding affinities of both RK14 and Gt15 for it, while the affinity of Arrestin (Arr) for R* increases.15,16 Arr dissociates from its homo-dimeric and homo-tetrameric storage forms to bind R*, preventing further activation of the pathway and effecting the release and recycling of the light-sensitive chromophore from the receptor.17–19 Meanwhile, the regulating protein RGS9-1 binds the activated Gt, PDE complex and stimulates the innate GTPase activity of Gt, resulting in the deactivation and dissociation of the complex.20 Finally, decreasing Ca2+ concentrations trigger the activity of two guanylate cyclase activating proteins (GCAPs), which cause guanylate cyclases (GCs) to synthesize cGMP at higher rates.21 This leads to the re-opening of the cGMP-gated ion channels and a return to the dark circulating current.

The rod photoreceptor exhibits the ability to respond to stimuli across several orders of magnitude of intensity, including single photons. At low light intensities, activation requires efficient and effective signal amplification.22 Saturating stimuli, on the other hand, necessitate rapid recovery to allow the continued detection of light. Overly effective recovery mechanisms, however, would threaten to quench dim-light responses before they are sufficiently amplified. Thus, recovery is tightly controlled by parallel negative-feedback mechanisms, including Ca2+-mediated feedback on GCAPs activity21,23–25 and on Rec regulation of RK.9,10,26–29 These serve to ramp up signal...
recovery to more intense stimuli without extinguishing dim-light responses. The result is an exquisitely balanced, but dynamically complex system.

In the past decade, an ongoing effort has been made to produce a comprehensive model of visual phototransduction in rod photoreceptor cells. The model to-date has been built to include nearly all of the known mechanisms involved in phototransduction. It has been constructed according to a bottom-up strategy, in which detailed representations of the underlying reactions are implemented according to the law of mass-action, eschewing high-level, empirical kinetic models. This model has proven to be very successful in reproducing a variety of mutant conditions in many of the underlying proteins. Furthermore, it has been used to make novel predictions of the dynamical role of a homo-oligomerization mechanism of Arr.

Although the model parameters were fit using biochemical and electrophysiological data culled from experiments on amphibians, it was found to produce qualitatively accurate simulations of experiments originally performed in mice, despite the approximately ten-fold slower dynamics in amphibians. Because modern vision research most commonly employs the mouse visual system, it would be valuable to have a model that can also produce quantitatively accurate simulations of it. To this end, we have adapted the most recent version of the amphibian model by Invergo et al. in order to simulate murine electrophysiological data. This was accomplished both through the integration of previously published parameter values and through the use of informed tuning and estimation techniques for unknown values but, importantly, without modification of the underlying reaction network. The resulting model provides accurate reproductions of the mouse visual response under a range of conditions and stimuli, while pointing to potential gaps in our knowledge of the phototransduction system.

2 Methods

2.1 Model implementation

The unaltered reaction network of Invergo et al. was used to construct the model (Table S1, ES†). This reaction network is the most recent in a series of iterative improvements to a comprehensive model of amphibian phototransduction. Each successive iteration of the model has added to or improved the reactions of the previous version according to the latest biochemical knowledge. The current network consists of a system of ordinary differential equations for 96 reactions, deterministically tracking the time evolution of 76 molecular species using 62 parameters. It was implemented using SBTOOLBOX2 for MATLAB (http://www.sbtoolbox2.org). Model files in SBTOOLBOX2 or SBML formats are available upon request. All numerical simulations and parameter estimation were carried out in this framework. Deterministic simulations were run from automatically generated and compiled C-code models, based on the CVODE integrator from SUNDIALS. Simulated light stimulus intensities in units of $R^+/s$ were approximated from published values described in units of photons $\mu m^{-2}$ using a collecting area of 0.43 $\mu m^2$.2,38

2.2 Parameter determination

Parameters values were retrieved or approximated from the literature when possible. In cases where the true parameter values were unknown, they were either manually tuned to meet experimental expectations or they were estimated through parameter optimization techniques. Manual tuning was performed on individual parameters to reproduce expected behaviors for which experimental data is not available (e.g. changes in concentration of a given protein during a photosresponse). Parameter estimation was done using a combination of the Nelder–Mead Simplex method for local optimization and a particle swarm method for global optimization. Optimization was performed against published electrophysiological data of the responses of a three-month-old mouse of the B6D2F1/J strain, which is a F1 hybrid between the C57BL/6J and DBA/2J strains. After estimation, parameter values were held fixed in all subsequent simulation experiments.

2.2.1 Amplification parameters. The model parameters related to the $Gt-R^*$ interaction required significant modification to adapt the model to mammalian data (Table S2, ES†). The parameters controlling the $Gt-PDE$ interaction, on the other hand, were left unchanged due to their insignificant effect on the model dynamics (see Dell’Orco et al. for a global parameter sensitivity analysis). Because the true values of the $Gt-R^*$ rate parameters are unknown, they were manually tuned to produce the expected rate of $Gt_{cat}$ production per $R^*$ per second, as was done in the original formulation of the amphibian iteration of this model. This rate can be estimated according to the following relationship:

$$A = \nu_{Gt} cG \beta_{sub} n_{cG}$$

where $A$ is the amplification constant, estimated to be 5 to 10 s$^{-1}$; $\nu_{Gt}$ is the rate of $Gt_{cat}$ production per $R^*$ per second, to be estimated; $cG$ is the coupling efficiency of $Gt$ to PDE, which, due to the relative insensitivity of the PDE-activation-related parameters, is approximately 1; $n_{cG}$ is the Hill coefficient of the cGMP channel activation, which was manually tuned to 3.8; and $\beta_{sub}$ is the rate constant of cGMP hydrolysis per activated PDE subunit. $\beta_{sub}$ may be approximated as follows:\n
$$\beta_{sub} = \frac{1}{N_{AV} V_{cyste} B_{cG}}$$

$k_{cat}$ is the turnover rate of a doubly-activated PDE holomer, estimated to be 1200 to 3500 s$^{-1}$; $K_m$ is the Michaelis constant of cGMP hydrolysis by PDE, estimated to be 17 to 23 $\mu$M; $N_{AV}$ is Avogadro’s constant; $V_{cyste}$ is the rod outer segment volume, calculated to be approximately 0.03916 pl for the mouse used in the model fitting; and $B_{cG}$ is the buffering power of the cytoplasm for cGMP, which is approximately 2.$^1$ It was determined that an amplification constant $A$ of 10 s$^{-2}$ was required to fit the electrophysiological data. The ratio $k_{cat}/K_m$ was maximized according to the reported ranges for
the two parameters, resulting in a $\beta_{\text{sub}}$ of $2.1826 \times 10^{-3}$ s$^{-1}$ and a $\nu_{\text{RGO}}$ of 1206 s$^{-1}$. The maximal rate of $G_\alpha$ production per activated receptor has been reported to be approximately 1290/R*$^a$. Therefore, this estimate of $\nu_{\text{RGO}}$ is reasonable. The $G_\alpha$-related parameters in the model were then manually tuned to approximate this rate of $G_\alpha$ production (Table S2, ESI$^\dagger$).

2.2.2 Recovery parameters. The basal binding rates of RK and Arr to R* required re-tuning to fit mammalian recovery dynamics (Table S3, ESI$^\dagger$). It was previously reported that a linear decrease in the affinity of RK for R* due to phosphorylation of the receptor provided a better, less sensitive fit to the amphibian electrophysiological data. In the present model, it was found that such a linear relationship resulted in a poor prediction of the signal responses of a Rec-knockout animal. In particular, it resulted in RK out-competing Gi for binding R* and the subsequent full phosphorylation of the receptor before Gt deactivation was manually tuned to approximate the exponential one and by setting the exponent to a greater value than was previously used (see Dell'Ocro et al.$^{32}$), this problem was alleviated. Note, however, that the true relationship is not known.

The basal rate of RK–R* binding was tuned according to the original estimation by Hamer et al.$^{31}$ such that $RK_{\text{dark}}^1 = 100$ s$^{-1}$, or 100/R*$^a$, where $RK_{\text{dark}}^1$ is the quantity of free RK molecules in the dark. According to a steady-state analysis, $RK_{\text{dark}}^1$ is 580 in the model, thus $RK_{\text{dark}}^1$ was set to 0.1724 s$^{-1}$. The rates of phosphorylation and the following dissociation of the RK–R* complex were required to be approximately ten-fold faster than in the amphibian model (Table S3, ESI$^\dagger$). Parameters determining the rate of the RK–Rec interaction required no changes.

The Arr–R* interaction required significant re-tuning (Table S3, ESI$^\dagger$). In particular, the binding and dissociation rates were required to be several orders of magnitude faster than in the amphibian model. Hamer et al.$^{31}$ predicted the basal rate of Arr–R* binding to be 0.25/R*$^a$ (Arr$^1$ to Arr$^1$ = 25 × 10$^{-3}$ s$^{-1}$). In the present model, with an initial, steady-state Arr monomer quantity of 1 260 760 molecules, the resulting $k_{\text{Arr}}$ of 1.9829 × 10$^{-2}$ s$^{-1}$ was much too slow to fit mammalian recovery times. A value of 9.9147 × 10$^{-6}$ s$^{-1}$ (12.5/R*$^a$) was determined to be sufficient through initial estimation and subsequent manual tuning.

The rate of dissociation of the Arr–R* complex prior to R* deactivation was manually set to 0.1 × 10$^{-3}$ s$^{-1}$ and to 0.1 × 10$^{-5}$ s$^{-1}$ (Arr$^1$ to Arr$^1$ = 173 nM; optimized from ref. 25) and $K_C^1$ (59 nM) are the Ca$^{2+}$ concentrations at which the GCAPs activity is half maximal, for GCAP1 and GCAP2, respectively; and $m_1$ (3; optimized) and $m_2$ (1.5; optimized) are Hill coefficients for GCAP1 and GCAP2, respectively. $\beta_{\text{dark}}$ the dark rate of cGMP hydrolysis, was given a value of 3.19 s$^{-1}$ according to steady-state analysis and cGMP$_{\text{dark}}$, the concentration of cGMP in the dark, was manually tuned to be 6.5 μM (Table S4, ESI$^\dagger$).

Ca$^{2+}$ regulation-related parameters were either culled from the literature or estimated by parameter estimation. See Table S4 (ESI$^\dagger$).

3 Results

Model fitting was performed against single-cell recordings of photoresponses to stimuli ranging from 1.7 to 4630 photons μm$^{-2}$ in rod photoreceptors of a three-month-old B6D2F1/J mouse. Photoreponses were measured as the suppression of the dark current in the outer segment of the cell. The resulting model successfully reproduces the primary characteristics of the photoresponse to both dim and saturating light flashes and overall closely corresponds to the experimental data (Fig. 1). In order to
test the accuracy of the model in reproducing the mechanisms of light adaptation, we next simulated and compared light flashes in the presence and in the absence of a steady, non-saturating background illumination (Fig. 2A). Consistent with in vivo experiments, we found a shortening of the saturation time in the presence of the steady illumination, however the simulated effect was very moderate compared to experimental observations.46

To further test the performance of the model, we also simulated a variety of mutant conditions that were previously used for model verification.33 Overall, simulations produced with the present model correspond well with the experimental data. Decreasing RK activity via underexpression of the protein results in the expected slowing of recovery to a dim flash stimulus (Fig. 3A') and an increase in saturation time ($T_{\text{sat}}$, the time spent at at least 90% of the peak amplitude) after a bright stimulus (Fig. 3C'). Overexpression of the kinase did result in a constant decrease in $T_{\text{sat}}$ across all stimulus intensities, which does not occur in experimental measurements with mice (compare Fig. 3C', produced by Sakurai et al.,47 with Fig. 3C'), however this phenomenon has previously been observed with this model and the exact reason for this discrepancy remains unclear.33

The model also produces excellent predictions of the values of $\tau_{\text{rec}}$ (the time constant of an exponential fit to the second half of the recovery from a dim flash stimulus) and the dominant time constant of recovery, $\tau_D$, (the slope of the increase in $T_{\text{sat}}$ with flash stimuli of logarithmically increasing intensities) (Fig. 3B' and D'). Both values have been measured to be approximately 250 ms for wild-type mice47 (Fig. 3B and D). Sakurai et al.47 showed that $\tau_{\text{rec}}$ is slightly higher when RK is underexpressed and moderately lower with overexpression, which the model accurately captures (Fig. 3B'). $\tau_D$, on the other hand, is not expected to change with increasing RK expression (however, three-fold overexpression may result in a small decrease in $\tau_{\text{rec}}$, as measured by Sakurai et al.47 and reproduced by our model; Fig. 3D'). In our simulations, RK underexpression resulted in a minor increase in $\tau_D$ (Fig. 3D').

The removal of Ca$^{2+}$ feedback on RK by simulating a Rec knock-out mouse results in lower peak amplitudes after dim flashes and a strongly reduced $T_{\text{sat}}$ (Fig. 4). A slight decrease in $\tau_D$ from 242 in the wild-type to 193 in the simulated knock-out is similar in magnitude to the decrease seen in vivo of 184 ms to 154 ms.28 The model captures well the sharp peak and slowing rate of recovery to non-saturating stimuli exhibited by the knock-out animals (compare Fig. 4B, produced by Makino et al.,28 to Fig. 4E).

Finally, it is currently understood that RGS9-1-mediated shutdown of the effector PDE is the rate-limiting step in phototransduction recovery, which our previous simulations confirmed,13 and experimental and modeling results further point to the general importance of RGS proteins in speeding up the kinetics of G-protein signaling. This was most convincingly demonstrated by Burns and Pugh,48 who showed that varying the concentration of RGS9-1 has a strong effect on $T_{\text{sat}}$ and $\tau_D$ (Fig. 5A). When the experiments of Burns and Pugh48 are simulated with our model, we find an excellent correspondence with the published results (Fig. 5B). While some quantitative differences exist between simulations produced with the present model and that of Burns and Pugh, it should be pointed out that the latter model’s parameters were specifically fit by maximum likelihood to this data while the simulations were performed with the present model without any specific tuning. In the simulations, underexpression of RGS9-1 leads not only to a strong increase in $T_{\text{sat}}$ but it also results in a notable increase in the slope, $\tau_P$. Conversely, overexpression results in a distinct drop in $T_{\text{sat}}$ and a moderate decrease in $\tau_P$. Thus, RGS9-1 plays a primary role in shaping the dynamics of the murine phototransduction response. Furthermore, the estimated rate of binding of RGS9-1 to the G$\alpha$PDE complex, which is approximately two orders of magnitude faster than that estimated for amphibians, points to
Fig. 3 The effects of varying RK expression on the photoresponse. (A–D) Experimental results of Sakurai et al.47 (A–D) Simulations of the same experiments. (A') Normalized responses to a non-saturating flash (75°). 3× overexpression (dotted-dashed trace) results in a slowed recovery compared to wild-type (solid trace). 3× underexpression (dotted trace) has a marginal effect. (B') The time constant of recovery, \( t_{\text{rec}} \), versus RK expression. (C') Time spent in saturation (\( T_{\text{sat}} \)) as a function of logarithmically increasing stimulus intensities. RK underexpression (open squares) results in an increase in \( T_{\text{sat}} \) compared to wild-type (open circles). RK overexpression (stars) leads to a slight decrease in \( T_{\text{sat}} \). The lines show the least-squares best fit of the first four points, used in the determination of \( t_D \). (D') \( t_D \), the slope of the relationship of \( T_{\text{sat}} \) with logarithmic stimulus intensity, versus RK expression levels. Underexpression results in a small increase in \( t_D \) while overexpression has no effect. Notice that \( t_{\text{rec}} \) and \( t_D \) are approximately equivalent for wild-type mice. (Panels A–D copyright 2011 The Association for Research in Vision and Ophthalmology, Inc.) NB: the authors used “Grk1” and “GRK1” interchangeably.
this mechanism as a major determinant of the relatively rapid recovery dynamics exhibited by mammalian species.

4 Discussion

4.1 Current challenges in modeling the visual photoresponse

The model presented herein accurately reproduces murine photoresponses to a wide range of stimuli and under a variety of mutant conditions. Notably, this could be accomplished by adapting a model created to reproduce the amphibian photoresponse without any alteration to the underlying reaction network. Given the bottom-up approach used to build the model, this success illustrates the degree to which the true biology of the vertebrate phototransduction system is captured in the modeled reactions. Nevertheless, challenges were encountered during the production of this model, which could point to yet unknown or poorly understood mechanisms. We took advantage of the relative ease of computational simulation in order to investigate some of these shortcomings.

In effect, RK would entirely out-compete $G_t$ for binding $R^*$. By reverting to the previously implemented exponential relationship of RK affinity for $R^*$, this binding competition effect was eliminated. The true relationship between RK–$R^*$ affinity and $R^*$ phosphorylation level remains unknown and the sensitivity of the model to this relationship indicates that it is an important gap in the present understanding of phototransduction recovery dynamics.

Despite the overall good performance of the model, a notable shortcoming was observed when simulating light-adaptation. In measuring the effect of light adaptation on the photoresponse, we observed an expected shortening of the saturation time in the presence of a steady background illumination. However, the simulated effect was not as strong as expected (Fig. 2A; see, e.g., Fig. 7C in ref. 46). To determine if cGMP regulation was responsible for this behavior, we performed the same simulations with a model of a GCAPs knock-out animal, implemented by setting the GCAPs Hill coefficients (model parameters $m_1$ and $m_2$) to zero (Fig. 2B). The effect of the steady background was slightly stronger in the mutant, at least partially implicating the cGMP regulatory dynamics in the discrepancy. It was found through parameter manipulation that the $R^*$ deactivation dynamics have a strong effect on this characteristic of light adaptation. Slowing $R^*$ deactivation, through decreased RK or Arr activity, results in a larger gap between dark-adapted and light-adapted saturation times for both wild-type and GCAPs knock-out mutants, in line with experimental expectations (data not shown). However, doing
so also results in these mechanisms having a stronger influence on the recovery parameters $\tau_{\text{rec}}$ and $\tau_{D}$, disrupting the model’s faithful reproduction of the mutant scenarios described in the Results section. On the other hand, simulating the light-adaptation experiments without Ca\textsuperscript{2+} feedback on RK, thereby accelerating R* deactivation, removed any visible effect of light adaptation (data not shown).

Thus, we have found that the RK activity required to fit dark-adapted mammalian flash responses in this model may be too fast to accommodate expected light-adaptation behaviors. Additionally, as previously described, all iterations of this model have also shown inaccurate results of simulating RK overexpression, such that it incorrectly results in a decrease in saturation times.\textsuperscript{32,33} The incorporation of a dynamic Arr-oligomerization mechanism partially eliminated this effect by delaying the shutdown of the activated receptor,\textsuperscript{33} however some discrepancy with the \textit{in vivo} results remains (Fig. 3C'). Furthermore, as described above, the model has been also found to be particularly sensitive to the affinity relationship of RK for phosphorylated R*.\textsuperscript{33} No realistic, satisfactory combination of parameters could resolve this suite of problems. Because all well-established mechanisms related to receptor deactivation are included in the model, one would speculate that a yet unknown or poorly understood mechanism related to RK activity exists but is absent in the model. It is possible that further clarifying the effects of R* phosphorylation and RK autophosphorylation on the R*–RK interaction may resolve these discrepancies. In particular, our knowledge of the effect of RK autophosphorylation on this interaction is currently limited to only the kinase’s unphosphorylated and fully phosphorylated states\textsuperscript{14}, leaving the true relationship uncertain in a multi-step phosphorylation paradigm.\textsuperscript{31,30}

Additionally, it is probable that the cGMP-regulatory mechanisms in the model are insufficient. Only one of the two GC isozymes is present in the model, when in fact both exhibit different properties.\textsuperscript{25} Unfortunately, we lack mechanistic information on the putative switch between GC\textsubscript{1} and GC\textsubscript{2} functionality in order to properly implement both cyclases in the model.\textsuperscript{21} It is also worth noting that cGMP regulation remains implemented according to an empirical Hill equation rather than the desired, bottom-up representation of the underlying reactions. While a recent detailed model of cGMP kinetics in phototransduction using a similar Hill representation showed excellent performance in predicting local spatio-temporal kinetics, it was only used to simulate single-photon responses\textsuperscript{49} and not a full range of stimuli as we present here. A more complete implementation, capturing the complexities of the interactions of the two GCs and their Ca\textsuperscript{2+}-sensitive regulation by the GCAPs, is necessary to improve the dynamics of this important signal recovery mechanism and would likely provide more accurate simulations of light adaptation.

4.2 Future directions

At very dim stimulus intensities, down to a single photon, the probability of two proteins encountering each other as well as the general supramolecular organization become important factors in the phototransduction response,\textsuperscript{22} as does the local saturation of second messengers such as cGMP.\textsuperscript{49,50} Due to its
deterministic nature, our model cannot accurately simulate the single photon response (SPR). Furthermore, the representation of the photoreceptor outer segment as a well-mixed volume may omit important localized depletion of the second messengers. A series of spatially accurate models of mouse phototransduction have been produced in order to more accurately simulate this scenario. These models reproduce quite well the SPR and have revealed insights into the mechanisms underlying its variability, however they have not been demonstrated to also faithfully simulate responses to a wider range of stimuli. Additionally, they primarily give focus to the second messengers, while reducing protein activity to high-level, empirical parameters; thus, the intricacies of the inter-protein dynamics are lost. Ideally, the future will see a merging of the two techniques, allowing a detailed, spatially accurate model that can reproduce the full range of responses exhibited by a rod photoreceptor.

4.3 Conclusions

One of the chief aims of systems biology research is the production of quantitatively predictive models from which experimental hypotheses can be derived. It is hoped that our model will be a useful tool in guiding future research on phototransduction. By offering quantitatively accurate predictions across a range of conditions for the primary species used in vision research, it may be used to avoid costly lines of unproductive investigation. Meanwhile, its shortcomings may raise interesting questions regarding gaps in our knowledge of the phototransduction process. Lastly, its modular structure allows for the easy integration of novel features without disrupting the existing network, allowing it to be expanded as research progresses and obviating the need for constructing “one-off” models to explore novel mechanisms.

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