Ex vivo ERG analysis of photoreceptors using an in vivo ERG system

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The Function of the retina and effects of drugs on it can be assessed by recording transretinal voltage across isolated retina that is perfused with physiological medium. However, building ex vivo ERG apparatus requires substantial amount of time, resources and expertise. Here we adapted a commercial in vivo ERG system for transretinal ERG recordings from rod and cone photoreceptors and compared rod and cone signaling between ex vivo and in vivo environments. We found that the rod and cone a- and b-waves recorded with the transretinal ERG adapter and a standard in vivo ERG system are comparable to those obtained from live anesthetized animals. However, ex vivo responses are somewhat slower and their oscillatory potentials are suppressed as compared to those recorded in vivo. We found that rod amplification constant \( A \) was comparable between ex vivo and in vivo conditions, \( A = 10^{-30} \) s, depending on the choice of response normalization. We estimate that the light funneling factor of 3 for cones in the mouse retina. The ex vivo ERG adapter provides a simple and affordable alternative to designing a custom-built transretinal recordings setup for the study of photoreceptors. Our results provide a roadmap to the rigorous quantitative analysis of rod and cone responses made possible with such a system.

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1. Introduction

Photoreceptors in the retina transform incoming photons into an electrical signal. Rod photoreceptors mediate night vision while cone photoreceptors enable us to see fine details and colors in daylight. Many retinal diseases that lead to blindness are caused by photoreceptor dysfunction. With the advancement of molecular tools over the last decades, genetically modified animals have become widely used to study the physiology and pathology of the retina in an effort to understand how photoreceptor function is affected in visual disorders and for the development of therapeutic treatments for degenerative and blinding diseases (Boye et al., 2013; Dalke & Graw, 2005; Peachey & Ball, 2003). Electroretinography (ERG) is an established technique used to measure light-induced electrical activity of the retina (Armington & Bloom, 1974). ERG can be measured in vivo from humans or anesthetized animals and commercial in vivo systems are routinely used in many laboratories and clinical settings. The in vivo ERG response to a flash of light consists of an initial negative a-wave followed by positive b-wave that can be roughly related to the activity of photoreceptors and ON bipolar cells, respectively (Green & Kapousta-Bruneau, 1999a; Gurevich & Slaughter, 1993; Penn & Hagins, 1969; Robson & Frishman, 1995; Stockton & Slaughter, 1989). However, the initial a-wave is quickly overlapped with the b-wave preventing accurate studies of photoreceptor function. This is especially problematic in photopic ERG recordings, where the a-wave produced by the small percentage of cone photoreceptors is detectable only with very bright test flashes. Furthermore, mechanical movement of the animal during recordings (caused by breathing and heart beat) complicates the quantitative analysis of the in vivo ERG data.

Ex vivo ERG is another well-established method which measures the light-induced voltage change across the retina isolated from the eye and pigment epithelium. Ex vivo signal can be recorded across the whole retina (transretinal ERG, Donner, Hemila, & Koskelainen, 1988; Frank & Dowling, 1968; Granit, 1933; Nymark et al., 2005) or across different retinal layers with microelectrodes (Arden, 1976; Green & Kapousta-Bruneau, 1999b; Penn & Hagins, 1969). In optimal perfusion conditions, the scotopic a- and b-waves of transretinal ERG are comparable to the corresponding in vivo waves (Green & Kapousta-Bruneau, 2014 Elsevier Ltd. All rights reserved.)
transducin. However, it is not clear how well cone signaling is preserved under ex vivo conditions. Recently, transretinal ERG has proved to be a very powerful tool to study mouse rod and cone phototransduction and the visual cycle (Brockway et al., 2005; Heikkinen, Nymark, & Koskelainen, 2008; Heikkinen et al., 2012; Kolesnikov et al., 2011; Sakurai, Chen, & Kefalov, 2011; Wang et al., 2009). This method provides easy means to isolate the photoreceptor signal component by using chemical agents to block synaptic transmission. It also significantly improves the signal-to-noise ratio (SNR) compared to in vivo ERG because of the lack of mechanical movement of the specimen. Overall, transretinal ERG facilitates quantitative studies of photoreceptors while still in their natural environment. In addition, it represents an easy way to test the effect of various drugs on photoreceptor function and retinal physiology. However, ex vivo ERG commercial equipment has not been available until now and construction of a custom-built setup is not trivial. Here we demonstrate that transretinal ERG recordings can be obtained with a commercial ex vivo ERG adapter (Ocuscience.us) in combination with a standard in vivo ERG system such as the one made by LKC (originally designed for human studies but also widely used for rodents) or the one from Ocuscience (made specifically for small animals). The adapter is a cost-effective way for studying mammalian rod and cone photoreceptor function and allows several hours of stable recordings simultaneously from two isolated retinas with very good SNR and minimal rundown. To establish the method and demonstrate the type of physiological analysis possible with it, we compared the kinetics and sensitivity of rods and cones between in vivo recordings (using an LKC ERG system) and ex vivo recordings (using the LKC system with the ex vivo ERG adapter). Consistent with previous studies, we found that rod sensitivity and the magnitude of phototransduction amplification are preserved in the isolated retina. Quantitative analysis of rod and cone ERG a-waves showed that mouse rods have higher molecular amplification constant compared to cones. In addition, comparison of phototransduction amplification between in vivo and ex vivo conditions yielded an estimate of ~3 for the cone light funneling factor in vivo. Finally, we compared rod and cone synaptic transmission between in vivo and ex vivo conditions. Analysis of rod and cone b-waves revealed that their amplitude and kinetics were largely preserved ex vivo. However, we found that in contrast to in vivo recordings, the rod b-wave amplitude increased supralinearly with increasing flash energy in ex vivo conditions.

2. Materials and methods

2.1. Animals

All rod ERG recordings were obtained from wild-type (WT) C57Bl/6 mice, and all cone ERG recordings were done from rod transducin x knock-out (Gnt1<−/−> mice whose rods do not produce light responses (Calvert et al., 2000). Animals were kept in 12/12 h dark/light cycle and dark-adapted 6-12 h before the experiment. All experimental protocols were in accordance with the Guide for the Care and Use of Laboratory Animals and were approved by the institutional animal care and use committee at Washington University and adhered to the EU Directive 2010/63/EU for animal experiments.

2.2. Ex vivo adapter design

The ex vivo ERG adapter illustrated in Supplementary Fig. S1 is built around the commercial in vivo ERG system from LKC Technologies. The main components of the adapter are (1) the specimen holder, (2) the perfusion system, and (3) the heat exchanger. Light stimulation and data collection are performed by using LKC LED ganzfeld stimulator, universal biomedical amplifier (UBA4200), drivers, and EM for Windows 9.4.0 software.

2.2.1. Specimen holder

The specimen holder design is based on a previously published type of closed perfusion chamber (Nymark et al., 2006; Fig. 1 and Supplementary Fig. S2). This closed design is optimal since it minimizes the risk of perfusion solution spills on the expensive light-reflective coating of the ganzfeld sphere. In addition, the chamber is simple to use and does not require additional stands or micromanipulators for electrode positioning. To further improve the ease of use and the efficiency of data collection, the chamber is machined from two separate pieces of polycarbonate (top and bottom parts) with threaded female connectors for custom-built electrode mounts, designed to host pellet electrodes with gold-plated 2 mm pin connectors (RC2, World Precision Instrument (WPI)). A polished transparent 0.5 mm thick polycarbonate window over the retinas allows efficient light stimulation. The double-specimen holder design enables simultaneous recordings from two isolated retinas.

The bottom part of the specimen holder contains two domes with 3 mm flat top surfaces on which round filter papers (HABG01300, Millipore) are attached. This filter paper is recommended to use because it is gray/black, and thus should not reflect significant amounts of light, facilitating the conversion of light energy into pigment isomerizations (see Section 2.5). Two 1 mm diameter channels filled with physiological solution provide the connection between the proximal side of each retina and pellet electrodes that are connected to the amplifier’s positive pole (see Fig. 1). The domes are surrounded by O-ring seals to prevent leakage of the perfusion solution which would potentially increase the noise, shunting, and mixing the signals between the two retinas. Retinas (red ovals in Fig. 1) are placed photoreceptor side up on the filter paper on top of the domes. The top chamber piece, with grooves matching the domes, is clamped tightly onto the bottom piece with screws and wing nuts. The top piece includes two separate parallel channels to perfuse the two retinas at their photoreceptor side. Thin 0.63 mm channels connected to the perfusion channels provide the electrical interface between the distal sides of the retinas and electrodes that are connected to the amplifier’s negative pole (ERG signal reference). In addition, the top part houses the adapter for the mouse rectal temperature probe (503524, WPI) that monitors the temperature very near the retina, just downstream of it.

2.2.2. Perfusion system

The perfusion solution is fed by gravity from a bottle warmed to 37 °C in a water bath (e.g. 180 Series Water Bath, Thermo Scientific) placed on the top of the LKC LED ganzfeld stimulator. For safety, the water bath was filled with metallic beads instead of water (Lab Armor Bath Beads, Fisher Scientific) that surrounded the perfusion bottle to maintain steady and even temperature of the perfusion solution. The perfusion solution is bubbled in the bottle with 95% O2/5% CO2 gas mixture and guided through the flow-rate valve (2C8891, Baxter) and heat exchanger (see Supplementary Fig. S1) to the perfusion channels of the specimen holder and eventually to the waste bin placed just outside the ganzfeld sphere. The electromagnetic noise introduced through the perfusion system is reduced by a Faraday cage that is attached on the top of the LKC ganzfeld stimulator around the water bath and by shielding the perfusion tubing with aluminum foil. The Faraday cage and tubing shield are electrically connected to the amplifier’s ground with a grounding cable.
Fig. 1. Double-retina specimen holder. The bottom part has two domes topped with filter paper supporting the retinas (red ovals). Electrode channels filled with physiological solution allow electrical contact to the ganglion cell side of the retinas. Two pellet electrodes with custom-build housings (see text) are thread-connected. The top part has individual perfusion channels for each retina. Electrical connection to photoreceptor side of the retinas is achieved through narrow channels that are connected to the perfusion channels. Similar electrodes as in the bottom part are thread-connectable. After placing the dissected retinas on the domes, the top part is clamped tightly to the bottom part with permanent screws in the bottom part and nuts on the top. O-rings (not shown in the figure) around the domes prevent perfusion solution leaks. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

2.2.3. Temperature control

The temperature of the retina is controlled by using the temperature controller (ATC1000, WPI), mouse heating pad (502195, WPI), and mouse rectal temperature probe (503524, WPI) provided with the in vivo ERG system, combined with a custom-built brass heat exchanger. The rectal probe is mounted permanently to the top part of the specimen holder close to the retina, and is connected to ATC1000 temperature controller with the temperature set to 37 °C. The specimen holder is placed on the heating pad. Warming of the retina is further facilitated by circulating the perfusion solution tubing through the brass heat exchanger that is also mounted on the mouse heating pad (see Supplementary Fig. S1).

2.3. Ex vivo ERG recordings

Dark-adapted animals were sacrificed by CO₂ inhalation, the eyes were enucleated and bisected along the equator, and the retinas without retinal pigment epithelium (RPE) were detached in Ames solution under infrared (IR) light with IR converters fit to the dissecting microscope. Both retinas of the mouse were mounted on the domes of the specimen holder as described above and the specimen holder was transferred to the LKC system under dim red light. Ames medium (A1420, Sigma–Aldrich) was used as the perfusion solution, with 50 μM BaCl₂ added to remove the strong negative glial PIII component. DL-AP4 (50 μM) was used to isolate the photoreceptor component. Photoreceptors were isolated with the LKC system’s green (530 nm) LED light which was calibrated as described below. All chemicals except DL-AP4 (Tocris Bioscience) were purchased from Sigma–Aldrich. For all recordings, the responses were low-pass filtered at 300 Hz (8-pole Bessel filter built in the LKC amplifier) and data was acquired at 1000 Hz.

2.4. In vivo ERG recordings

Dark-adapted mice were anesthetized by intraperitoneal injection of ketamine (80 mg/kg) and xylazine (15 mg/kg) cocktail. Pupils were dilated with 1% atropine sulfate. During testing, a heating pad controlled by a rectangular temperature probe maintained mouse body temperature at 37 °C. Full-field ERGs were recorded using a UTAS BigShot apparatus (LKC Technologies) and mouse corneal electrodes (STelesSR, LKC Technologies). The reference electrode needle was inserted under the skin at the skull. Test flashes of calibrated green (530 nm) LED light were applied in darkness. For all recordings, the filtering and acquisition parameters were identical to ex vivo recordings.

2.5. Light calibration

The photopic luminous energy \( [E] = \text{Cd s m}^{-2} \) of green (530 nm) flashes produced by the LKC stimulator from –35 to 10 dB (the nominal flash energies set from the software) was measured with calibrated integrating flexOptometer™ (Gamma Scientific/UDT Instruments). For dimmer flashes \( E \) was calculated from the measured reference point at –35 dB using the dB scale of the LKC stimulus unit. In order to quantify the amplification and sensitivity of the mouse rod and cone phototransduction (see Section 2.6) the luminous energy in Cd s m⁻² had to be converted into flash energy \( Q \) expressed in photons per unit area. Green 530 nm light flashes from the LKC LED ganzfeld stimulator produced up to (our own calibration) 1.5 log (Cd s m⁻²). In addition, very bright broadband spectrum Xenon flash tube, producing up to nominal 25 dB of white light, was used to find the maximal a- and b-wave amplitudes at the end of in vivo experiments. For our green light stimuli, the photon flux \( Q \) can be expressed (see Wyszecki & Stiles, 1982) as:

\[
Q(530 \text{ nm}) = \pi E \times 1.5 \times 10^{-15} \times k(530 \text{ nm})
\] (1)

where the factor \( k \) converts photopic lumens to scotopic lumens and \( 1.5 \times 10^{-15} \) (in units of photons s⁻¹ lumen⁻¹) converts scotopic lumens into photon flux. The value for \( k \) at 530 nm is 2.34, based on the comparison of the standard CIE photopic and scotopic luminosity functions at 530 nm.

The comparison of in vivo and ex vivo data was facilitated by the use of the same LKC system when recording either from live
mouse or isolated retina. On the other hand, light was delivered at wide angle and scattered in our polycarbonate specimen holder affecting the amount of light ultimately absorbed by visual pigments in the photoreceptor outer segments. We measured the effect of stimulation geometry by covering the specimen holder with aluminum foil that had a $\geq 3$ mm aperture above the retina which decreased the linear response by 1.7-fold. In our standard electrophysiology rig (Sundermeier et al., 2014) where a narrow-angle beam was used, the same aluminum sheet did not affect the rod sensitivity. We also compared the sensitivity of rods under identical perfusion conditions between our electrophysiology rig and ex vivo adapter system. After correction for the scattered light, the calculated sensitivity of WT rods was comparable (within 15%) between the two setups. In order to compare the cellular and molecular properties of the retinas between in vivo and ex vivo conditions, we estimated how the optics of the mouse eye affected the photon flux at the level of the photoreceptor outer segments. For this, the following expression was used (Lyubarsky, Daniele, & Pugh, 2004; Lyubarsky et al., 1999)

$$\Phi = Q(\lambda)\tau(\lambda)\frac{S_{\text{pupil}}}{S_{\text{retina}}}k_{\text{shadow}}a_c(\lambda) = \omega_c(\lambda)Q(\lambda)$$

where $\Phi$ is the number of photopigment isomerizations in rods or cones, $Q(\lambda)$ is defined in Eq. (1), $\tau(\lambda)$ is the light transmission of the pre-photoreceptor media of the eye (0.7 for green light), $S_{\text{pupil}}$ is the area of a fully dilated pupil (3.2 mm$^2$ for adult mouse eye), $S_{\text{retina}}$ is the total area of the retina (18 mm$^2$), and $f$ is the light funneling factor by photoreceptor inner segments (~1.3 for rods and 7 for cones, Lyubarsky, Daniele, & Pugh, 2004; Lyubarsky et al., 1999). In transretinal recordings, light was applied from the photoreceptor side so $f = 1$. On the other hand, in this stimulation geometry the amount of light reaching the cone outer segments was affected by the rod shadowing effect (Heikkinen, Nymark, & Koskelainen, 2008). For cone ex vivo recordings this was corrected by setting $k_{\text{shadow}}$ to 0.7 (while for in vivo recordings it was set to 1) in Eq. (2) (Heikkinen, Nymark, & Koskelainen, 2008; Sakurai, Chen, & Kefalov, 2011). The light collecting area ($a_c$) converts the number of photons hitting the outer segments of rods or cones into the number of pigment isomerizations and it is identical for in vivo and ex vivo recordings. We adopted $a_c$ of 0.57 and 0.07 mm$^2$ at 530 nm for rods and cones, respectively (Govardovskii et al., 2000; Lyubarsky, Daniele, & Pugh, 2004; Lyubarsky et al., 1999; Sakurai, Chen, & Kefalov, 2011). The parameters that are different between in vivo and ex vivo in Eq. (2) are $\tau(\lambda)$, $S_{\text{pupil}}/S_{\text{retina}}$ for $f$ and $k_{\text{shadow}}$. For in vivo recordings the term $\tau(\lambda)$ for ex vivo recordings the term $\tau(\lambda) \times (S_{\text{pupil}}/S_{\text{retina}}) \times f \times k_{\text{shadow}}$ is 0.16 for rods and 0.87 for cones whereas in ex vivo case this term equals to 1 for rods and 0.75 for cones. This means that rod outer segments receive about ten times more photons with equal stimuli in ex vivo compared to in vivo recording geometry (including the correction factor of 1.7 for the scattered light, see above) while cone outer segments should receive more photons in vivo (2-fold more if scattered light is included). When calculating this term for cones in vivo, we adapted a funneling factor of 7 which is based on the assumption that all of the light “collected” by cone inner segments is funneled to the outer segment (see Lyubarsky et al., 2002). The validity of this assumption will be discussed below.

Finally, we introduce here a general light collection area $\omega_c(\lambda)$ that combines all the terms needed to convert photon flux into pigment isomerizations in rods or cones in both in vivo and ex vivo conditions.

### 2.6. Data analysis

#### A Hill-type function

$$\frac{r}{R_{\text{max}}} = \frac{E^r}{E^r + E_{1/2}^r}$$

was fitted to ERG a- and b-wave normalized amplitudes ($r/R_{\text{max}}$) to find the flash strength that produces a half-maximal response ($E_{1/2}$).

In the presence of DL-AP4 the maximum rod response was measured at the plateau level after the initial fast negative peak present in bright flash responses.

The gain of phototransduction activation reactions was quantified by determining the amplification constant $A$ as described by Lamb & Pugh, 1992 (LP model). For cone amplification constant the effect of membrane time constant was factored in as described by Smith & Lamb, 1997. Amplification constant ($A$) was fitted individually to each trace of the flash response family and average value reported for each mouse. However, as amplification constant is known to decrease at bright flash energies, values from bright flash responses were excluded from the analysis. In the short delay in LP function was kept between 2 and 3 ms and the membrane time constant of cones was assumed to be 5 ms. Fittings were performed with Origin 9.0.0 (64-bit, SR2, OriginLab) and other analysis was done with Origin and EMMWin 9.4.0 (LKC Technologies). For statistical analysis two-tailed student t-test was used.

### 3. Results

#### 3.1. Comparison of ex vivo and in vivo ERG flash responses from rods and cones

Physiological recordings from isolated retina offer some advantages compared to in vivo experiments (see Sections 1 and 4). However, it is not clear how the photoreceptor responses recorded in ex vivo conditions correspond to those in vivo. Here we compared rod- and cone-driven responses recorded from anesthetized mice with those from isolated mouse retinas obtained with the ex vivo adapter. We suppressed the strong negative glial slow PIII component by barium as described in Section 2.3, because it caused a transient decrease of the ex vivo b-wave amplitude within a certain flash range (see Supplemental Fig. S3). Green and Kapousta-Brunoe observed a similar effect of barium increasing the b-wave amplitudes of rat ERG responses (Green & Kapousta-Brunoe, 1999a). Response families to a wide range of light strengths from representative experiments are illustrated in Fig. 2A–D for both rods (black) and cones (red), for in vivo (A and B) and ex vivo (C and D) conditions. As recently shown by others (Green & Kapousta-Brunoe, 1999b; Heikkinen et al., 2012), we found that rod a- and b-wave amplitudes were not significantly different between in vivo and ex vivo recordings (Table 1). However, the kinetics of rod responses appeared slower and the oscillatory potentials were attenuated in ex vivo compared to in vivo responses. The inset in Fig. 2C plots in vivo and ex vivo responses to flashes estimated to produce identical number of $R^*$ exemplifying about 30% longer $t_p$ for both a-wave and b-wave in the isolated retina as compared to the live animal.

To compare cone function between in vivo and ex vivo conditions, we used Gnat1$^{-/-}$ mice whose rods are morphologically intact but do not respond to light. This simplifies the isolation of small cone responses that would be otherwise masked by large rod responses. However, it is also possible to isolate cone responses in WT mouse by using rod saturating pre-flash or background light (Heikkinen, Nymark, & Koskelainen, 2008; Jaissle et al., 2001; Nikonov et al., 2006). Our cone recordings demonstrate a remarkable resemblance with respect to a- and b-wave amplitudes between ex vivo and in vivo conditions as well as comparable but somewhat slower kinetics ex vivo (Fig. 2B and D, Table 1). Although the maximum ex vivo cone b-wave amplitudes were smaller on average ($p < 0.05$), in our best preparations they were similar to those recorded in vivo (Table 1). As in the rod recordings,
However, based on Eq. (2) with its previously published parameter values, we estimated that rod outer segments ex vivo receive about one log-unit more light compared to in vivo experiments in our recording geometries (see Section 2.5). After correcting for the different light path geometry, the half-saturating flash energies reported in Table 1 were similar between ex vivo and in vivo recordings.

Fig. 2F shows population-averaged b-wave amplitudes as a function of flash strength for rods (black symbols) and cones (red symbols), for in vivo (open symbols) and ex vivo (filled symbols) recordings. The rod data was normalized to the first plateau before b-wave amplitudes started to increase further at brighter light intensities, possibly due to cone contribution (see also Section 4.2). Again, the saturation of b-waves was reached with dimmer light flashes in ex vivo as compared to in vivo conditions. However, this difference vanished at the threshold for rods. Indeed, taking into account the differences in light path geometry, in vivo b-waves were more sensitive across the whole stimuli range. The cone b-wave amplitude data also showed a leftward shift in isolated retina recordings compared to in vivo conditions. This is surprising as cone outer segments should receive more photons in in vivo conditions (see Section 2.5). The discrepancy might be at least partially explained by the overestimation of the cone funneling factor in vivo, as discussed below. Interestingly, the b-wave intensity-response relations were steeper ex vivo compared to those in vivo for both rods and cones. We quantified this by fitting Eq. (3) to the data from individual experiments by assuming that the Hill coefficient $n$ is similar across experiments. The $n$ increased from 0.6 in vivo to 1.2 ex vivo for rods and from 0.7 to 1.1 for cones.

3.2. Isolation of rod and cone photoreceptor responses ex vivo and in vivo with synaptic blockers

Transretinal ERG has recently proven its power in studying photoreceptor function in the mouse retina. One of the reasons is that synaptic blockers can easily be applied in the perfusion to isolate the photoreceptor component. It is also possible to inject these blockers intravitreally but that requires expertise and special equipment. The injections also can have potentially harmful effects such as cornea damage and retina detachment. Another approach that would help to dissect the photoreceptor response in vivo is to use mutant mice with defects in synaptic transmission, such as nob mice (Pardue et al., 1998). However, this would require deriving any mutant mouse of interest in the nob background, adding time and expense to this approach. Fig. 3A and B shows flash response families recorded in isolated retinas from rods and cones, respectively, isolated with 50 μM DL-AP4 (APB) and 50 μM BaCl$_2$ in the perfusion. For comparison, we also performed a limited number of in vivo recordings where we intravitreally injected one eye with ~1 μL of 26.7 mM DL-AP4 in PBS (~1.3 mM vitreal concentration) and the other eye with PBS only (compare solid and dashed traces in Fig. 3C and D). DL-AP4 effectively removed the positive b-wave component both in vivo and ex vivo although sometimes a small overshoot persisted in both rod and cone (indicated by red arrow in Fig. 3D) responses. Comparison of the rod responses recorded from PBS- and APB-injected eyes to identical flash strengths revealed that the leading edge of responses remained unaffected by APB treatment (Fig. 3C). The leading edge kinetics, and thus the amplification of rod phototransduction, was also not affected by APB in ex vivo recordings (see also Fig. 4). In response to bright flashes a fast transient negative component persisted in rod responses that could not be explained by cone contribution. Implications of this transient peak (“nose”) typically observed in rod and sometimes in cone responses are discussed in more detail below. Low SNR did not allow us to accurately compare the leading edges of cone responses under similar conditions (Fig. 3D).
The overall kinetics and sensitivity of our pharmacologically isolated transretinal responses were similar to those obtained from single cells (Calvert et al., 2000; Nikonov et al., 2006; Tsang et al., 1998). However, some subtle differences are evident. For instance, the average time-to-peak was ca. 120 ms for WT rods and ca. 80 ms for cones from Gnat1"/" mice, both somewhat faster compared to those in single-cell recordings. On the other hand, in vivo dim flash responses isolated with APB were even faster than ex vivo responses peaking at about 90 ms and 60 ms for rods and cones, respectively. This suggests a general trend of deceleration of physiological processes when the integrity of the system is reduced from the in vivo environment to isolated retina and further to single cells. Fig. 3E shows the ex vivo population-averaged normalized intensity-response data for rods and cones. Half-saturating flash

Table 1
Rod and cone response parameters in vivo and ex vivo. All sensitivity parameters (E1/2) are referenced to luminous energy at the cornea (in vivo) or at the retina (ex vivo). See text for conversion to photosensitizers in rods or cones in both conditions. The effect of scattered light in ex vivo recordings is factored in only for the amplification constant (bottom row). The a-wave amplitudes are the maximum amplitudes of the a-wave to bright flash measured at peak or at plateau after the peak during APB exposure. The b-wave amplitude is the difference of a-wave and b-wave peak amplitudes measured from a response to a bright flash. The amplification constant A (s^-2) is given for rods and A2 (pm^2 s^-2) is used for cones (see Section 2). All values are means ± SEM. Statistically significant differences between in vivo and ex vivo recordings are indicated by * (p < 0.05). Statistical analysis was not performed for cone sensitivities or amplification constants. For rod sensitivities the statistical analysis has been performed after correcting for different light path geometries as described in Section 2.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>In vivo rods (n = 6)</th>
<th>Ex vivo rods (n = 5)</th>
<th>Ex vivo rods, APB (n = 3)</th>
<th>In vivo cones (n = 6)</th>
<th>Ex vivo cones (n = 4)</th>
<th>Ex vivo cones, APB (n = 5)</th>
</tr>
</thead>
<tbody>
<tr>
<td>a-Wave amplitude (µV)</td>
<td>420 ± 45</td>
<td>550 ± 80</td>
<td>350 ± 37</td>
<td>25 ± 2</td>
<td>39 ± 6</td>
<td>92 ± 3</td>
</tr>
<tr>
<td>b-Wave amplitude (µV)</td>
<td>660 ± 80</td>
<td>920 ± 190</td>
<td>–</td>
<td>–</td>
<td>405 ± 20^*</td>
<td>–</td>
</tr>
<tr>
<td>E1/2,a-wave (log [Cd s m^-2])</td>
<td>-0.35 ± 0.1</td>
<td>-1.4 ± 0.09</td>
<td>-2.8 ± 0.03</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>E1/2,b-wave (log [Cd s m^-2])</td>
<td>3.0 ± 0.07</td>
<td>3.7 ± 0.06</td>
<td>-</td>
<td>0.1 ± 0.08</td>
<td>-0.7 ± 0.05</td>
<td>-1.0 ± 0.03</td>
</tr>
<tr>
<td>A or A2 (s^-2 or pm^2 s^-2)</td>
<td>6.8 ± 0.8’</td>
<td>12 ± 1^*</td>
<td>29 ± 2</td>
<td>0.19 ± 0.01</td>
<td>0.42 ± 0.02</td>
<td>0.2 ± 0.01</td>
</tr>
</tbody>
</table>

The overall kinetics and sensitivity of our pharmacologically isolated transretinal responses were similar to those obtained from single cells (Calvert et al., 2000; Nikonov et al., 2006; Tsang et al., 1998). However, some subtle differences are evident. For instance, the average time-to-peak was ca. 120 ms for WT rods and ca. 80 ms for cones from Gnat1"/" mice, both somewhat faster compared to those in single-cell recordings. On the other hand, in vivo dim flash responses isolated with APB were even faster than ex vivo responses peaking at about 90 ms and 60 ms for rods and cones, respectively. This suggests a general trend of deceleration of physiological processes when the integrity of the system is reduced from the in vivo environment to isolated retina and further to single cells. Fig. 3E shows the ex vivo population-averaged normalized intensity-response data for rods and cones. Half-saturating flash

![Fig. 3. Comparison of rod and cone a-waves isolated with DL-AP4 in ex vivo and in vivo conditions.](image)

![Fig. 4. Determination of the rod amplification constant in ex vivo and in vivo conditions.](image)
intensities were -2.8 and -1.0 log(Cd s m⁻²) for rods and cones, respectively. By converting these values to average number of pigment photoisomerizations (see Section 2.5) per rod (R) or cone (C) we estimate that rods needed 17R and cones 112C to produce a half-saturating response.

3.3. Comparison of rod and cone signal amplification between in vivo and ex vivo conditions

The kinetics of the early rising phase of the ERG signal has been used to derive the molecular gain of rod and cone activation reactions in both ex vivo and in vivo situations. For rods, this amplification constant seems to be invariant across different experimental conditions, from single-cell recordings to in vivo recordings (Breton et al., 1994; Heikkinen, Nymark, & Koskelainen, 2008; Lyubarsky, Daniele, & Pugh, 2004; Nikonov et al., 2005). Fig. 4A and B shows fittings of the model introduced by Lamb and Pugh (LP model, Lamb & Pugh, 1992) to responses recorded from WT mice in vivo and ex vivo, respectively. On average, we found that the amplification constant (A) was 6.8 ± 1 and 12 ± 1 s⁻² as extracted from in vivo and ex vivo recordings, respectively. These amplification values are comparable to previously published results from mouse rods obtained with in vivo, ex vivo, or single cell recordings (Heikkinen, Nymark, & Koskelainen, 2008; Lyubarsky, Daniele, & Pugh, 2004; Nikonov et al., 2005).

Although the analysis of ERG data yields consistent values for A, it still remains unclear if its absolute value represents the true molecular gain of phototransduction activation reactions. Indeed, our analysis yielded somewhat different values for A from in vivo and ex vivo recordings. One complication for this analysis is the choice of maximal a-wave amplitude because it is largely masked by the b-wave. Thus, choosing for instance a 2 times larger rmax would yield 2-fold smaller A. We attempted to determine the proper choice of rmax by comparing the values of A obtained from transretinal recordings with or without APB in the perfusion as well as from in vivo ERG recordings from eyes that where injected with APB-free or APB-containing PBS. Fig. 4D illustrates in vivo responses to bright flashes from a control eye (gray) and APB-injected eye (black). Although the b-wave is removed by APB, it is evident that a substantial early transient peak still remains. In all four experiments performed, the peak amplitude of bright flash response in control and APB-injected eyes matched quite well. This observation suggests that following bright flashes it is not the b-wave that masks the peak of the negative-going a-wave. Analysis of the kinetics of the positive-going part of the ERG signal has led to the same conclusion (Kang Derwent & Linsenmeier, 2001).

This finding might have important implications on the choice of the response normalization when determining the amplification constant. If the plateau level after the peak describes better the saturation level of outer segment current, then the normalization to the noise amplitude would yield substantial underestimation of the amplification constant. Since injections themselves caused desensitization and somewhat reduced amplification in our hands, we analyzed the amplification constant from responses recorded from isolated retina in presence of APB which did not affect the leading edge kinetics in ex vivo conditions (this was found by comparing traces from Figs. 2C and 3A). Fig. 4C illustrates fitting of the LP model with rmax set to the plateau level after the initial peak present in bright flash responses. Average results from 4 retinas yielded over 2-fold increase of rod amplification constant determined in this way compared to the analysis performed as in Fig. 4A (see Table 1). The a-wave appeared to saturate at much lower intensities ex vivo than in vivo where the early peak seems to be more pronounced and probably leads to a substantial overestimation of the rmax that would describe the proper rod photoreceptor saturation (see Robson & Frishman, 2014). The fact that in this study we normalized to the a-wave amplitude at very bright flash in vivo (as is customary, see Fig. 4B) probably explains why we observed a smaller rod amplification constant in vivo as compared to that determined from ex vivo responses.

Determination of cone phototransduction amplification in vivo (and ex vivo) is complicated by several factors. The cone a-wave amplitude is very small, the b-wave intrusion (and phototransduction inactivation) come into play at very early times, and calibration of pigment isomerization is approximate at best. In addition, the extent of light funneling, which cannot be quantified accurately, might affect the estimated value of A for cones by several fold. Recent results from single-cell recordings and ex vivo recordings have suggested that the gain of mouse cone phototransduction activation reactions is quite similar to that in rods (Heikkinen, Nymark, & Koskelainen, 2008; Lyubarsky et al., 2002; Nikonov et al., 2006). In addition in vivo recordings have also suggested that A is similar in rods and cones (Smith & Lamb, 1997). However, the value of cone amplification in vivo is highly dependent on the estimation of the funneling factor (Lyubarsky et al., 1999; Schnapf et al., 1990; Smith & Lamb, 1997). To address this issue, we compared the cone amplification from ex vivo and in vivo ERG flash responses of Gnat1⁻/⁻ mice. Fig. 5A shows a response family recorded from isolated retina to flashes exceeding -2 log(Cd s m⁻²). In this range a small a-wave is evident even without using any blockers in the perfusion. The LP model convolved with the function to account for the rather large membrane time constant of cones were used as described in Section 2.8. The average amplification parameter referenced to the light energy measured at the cornea in vivo or at the retina ex vivo, Aₐ (Aₐ = A₅₃ = A m² s⁻²), was 0.4 m² s⁻² when determined ex vivo in Ames medium and 0.2 m² s⁻² as derived from in vivo recordings (see Fig. 5A, B, and Table 1). Interestingly, we found that Aₐ determined ex vivo in the presence of DL-AP4 and BaCl₂ (Fig. 5C) was over two times smaller compared to that derived from responses recorded in the absence of blockers (Fig. 5A). This may be due to the choice of normalization of the responses. For in vivo and ex vivo recordings, in the absence of synaptic blockers we set the maximal cone amplitude to the peak of a very bright flash response. It is, however, possible that the b-wave interferes at early times and the actual maximal response of the cone photoreceptors would be larger than that observable in the ERG recordings without synaptic blockers. This hypothesis is supported by the doubling of the cone a-wave amplitude in the presence of blockers in ex vivo recordings (see Table 1). Our limited number of in vivo experiments with intravitreal injections of APB also indicated clearly that the maximal cone photoreceptor response (fast PIII) amplitude might be significantly larger than that determined at the peak of a bright flash response in the absence of blockers (see Fig. 3D). The conversion of Aₐ into A is simple ex vivo where the funneling of cone inner segments is absent. Using a cone collection area of 0.07 m², we estimate that the cone amplification constant in our ex vivo conditions is 6 s⁻² and 2.8 s⁻² when determined in the absence or presence of APB, respectively. Assuming a funneling factor of 7 (Lyubarsky et al., 1999) yields A = 1.4 s⁻² when extracted from in vivo recordings with Gnat1⁻/⁻ mice.

4. Discussion

4.1. Advantages and potential applications for the ex vivo ERG adapter

The ex vivo ERG adapter (Figs. 1, S1 and S2) was designed to be used as easily as possible together with any commercial in vivo ERG apparatus. It makes use of all electronics, software, and the light stimulation of the commercial system, to minimize the cost of the ex vivo system. These features should facilitate the imple-
Fig. 5. Determination of the cone amplification constant under different conditions from Gnat1−/− mice. Ex vivo: (A) The black traces show the recorded flash responses to stimulus energies from −1.0 to 1.5 log (Cd s m−2) of green light. Gray traces are LP model functions with σ[…](mean of the best-fitting value for each trace ± SEM). A convolution to account for the membrane time constant τ = 5 ms was used when the flash energy was between 0.5 and 0.5 log (Cd s m−2) but not for the two dimmest flashes. In vivo: (B) Convoluted version (τ = 5 ms) of the LP activation model with A0 = 0.2 μm2 s−2 (no correction for eye optics, see text) gave the best fit to these representative responses. Flash energies were 1.0 and 1.5 log (Cd s m−2) for green light, and 20 dB for xenon flash. Ex vivo with blockers: (C) Black traces show dark-adapted cone responses recorded from a retina perfused with Ames medium containing DL-AP4 and BaCl2. Green light flash energies ranged from −2.1 to 1.0 log (Cd s m−2). Gray traces show the best-fitting LP activation model functions where convolution has been used for the four brightest flashes. The average best-fitting A0 for the five dimmest flashes was 0.28 ± 0.02 μm2 s−2. Above 0 log (Cd s m−2) A0 started to decrease, and was found to be 0.2 and 0.12 μm2 s−2 for 0.5 and 1.0 log (Cd s m−2), respectively, for this retina.

4.2. Correspondence of the ex vivo environment to physiological in vivo conditions

In our present study we used the same light stimulation and data acquisition system for in vivo and ex vivo ERG recordings to facilitate the comparison between the two conditions. To validate the proper functionality of our easy-to-use ex vivo adapter, we compared mouse rod and cone responses from in vivo and ex vivo recordings. We chose commercially available bicarbonate buffered Ames perfusion medium as it has been shown to support all retinal cells for extended periods. Both retinas were used for simultaneous recordings immediately after their isolation, by using a double-retina specimen holder. Consistent with previous studies (Green & Kapousta-Bruneau, 1999b; Heikkinen et al., 2012), both rod and cone ERG a- and b-waves had amplitudes and kinetics comparable to those determined from in vivo recordings (Fig. 2). However, the kinetics assessed by the τp of either the a-wave or b-wave tended to be slightly slower ex vivo where oscillatory potentials were often attenuated or absent as well (Fig. 2C). Our ex vivo responses remained still faster when compared to single cell recordings suggesting a general trend of response slowdown by isolation process of the retina and cells. It is possible that isolation of the retina from RPE damages rod and cone outer segments and/or that the conditions for the isolated retina are sub-optimal. Indeed, as illustrated in the Supplementary Fig. S3, barium used to remove slow PIII component, appeared to affect the sensitivity of rod signaling pathway. However, it is hard to distinguish whether the effects of barium are due to removing the slow PIII or due to altered rod signaling. In our hands, the intravitreal injection of barium (1 μl of 27 mM BaCl2 in PBS) completely abolished in vivo ERG responses leaving us unable to compare them to those recorded from isolated retina.

A striking difference between the in vivo and ex vivo ERG signals became evident after blocking metabotropic glutamatergic synaptic transmission with DL-AP4. In ex vivo signals this treatment revealed a very large slow negative glial PIII component, especially in response to bright flashes (data not shown in this paper but see the inset of Fig. 3 in Heikkinen et al., 2012). In all of the data shown here (except in Supplementary Fig. S3), the glial component was removed by exogenous application of BaCl2. However, in vivo flash responses lacked this component even when only DL-AP4 (and not BaCl2) was injected intravitreally (Figs. 3C and 4D). There can be at least two factors contributing to this difference. First, the isolation process of the retina from the rest of the eye and pigment epithelium might alter some ERG response components. Second, it is possible that perfusing only the photoreceptor side of the retina is not optimal for maintaining inner retina function.

The determination of sensitivity of individual cells or pathways in the retina is complicated by a number of factors. The in vivo ERG is a non-invasive method to probe retinal function but several overlapping signal components hamper the analysis of individual processes. In addition, it is not straightforward to calculate the amount of photons that are eventually absorbed by photoreceptors.

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First, isolation of the retina leads to removal of most of the physiological noise caused by breathing or cardiac activity of the animal. Second, the ex vivo recordings are more stable than those performed in vivo because they require neither prolonged anesthesia nor repetitive lubrication of the eyes for proper electrode contact. In addition, the ex vivo ERG technique provides an easy and efficient way to test the effects of broad spectrum of drugs on photoreceptors, synaptic transmission, and overall retinal function. Further, the use of the adapter is not restricted to mouse retina but is applicable to other species including amphibians, fish, pri- mates and other mammals. The system also makes possible record-
ings. In addition, the operating range determined from b-wave responses seemed to be suppressed in our ex vivo recordings (Fig. 2F). One possible explanation for this difference might be a glial component opposing the increase of the b-wave in brighter light and thus expanding the apparent range of b-wave amplitudes in vivo where we did not use barium to remove the glial component. However, as shown in Supplementary Fig. S3 removal of BaCl2 from the perfusion ex vivo did not affect significantly the slope of the b-wave amplitudes plotted against flash strength in the range thought to represent rod signaling (the data points before the first plateau in Fig. S3). In addition, it seems that the glial component is not that prominent for in vivo recordings based on the lack of a slow negative wave in the bright flash responses recorded from APB-injected eyes of anesthetized animals (Fig. 3C). The half-saturating b-wave both in our in vivo and ex vivo conditions is about 1R per rod (Fig. 2F and Table 1) which is consistent with direct recordings from mouse rod bipolar cells (Sampath & Rieke, 2004). Interestingly, our ex vivo data showed much steeper increase of b-wave amplitudes as a function of flash energy when compared to in vivo data (Fig. 2F). The reason for this difference is unclear but it is possible that the slower b-wave recovery phase (compare e.g. cone b-waves between Fig. 2B and D) in ex vivo compared to in vivo conditions contributes to this phenomenon.

One additional complication to the quantitative analysis of the rod ERG signals both in vivo and ex vivo is the fast negative peak (“nose” component) observed in pharmacologically isolated a-wave (Arden, 1976; Robson & Frishman, 2014; Vinberg, Strandman, & Koskelainen, 2009). This component becomes larger upon increasing flash intensity even after rod photocurrent is saturated, as evident by the plateau level after the peak (Figs. 3C and 4C). It is possible that the biphasic nature of dark-adapted a- or b-waves when plotted as a function of flash strength (see Fig. 2F) might not represent simply rod and cone responses but also the rod-originating peak component as suggested recently (Robson & Frishman, 2014). As discussed below, the resulting overestimation of \( r_{\text{max}} \) would also lead to underestimation of the rod amplification constant, \( A \).

The comparison of cone sensitivity and their synaptic transmission between in vivo and ex vivo environments is complicated by the conversion of the luminous energy into cone pigment isomerizations. It has been suggested that cone inner segments and Müller cells act as light guides to increase the amount of photons reaching the cone outer segments in vivo (Agte et al., 2011; Franze et al., 2007; Labin & Ribak, 2010; Lyubarsky et al., 1999; Schnapf et al., 1990). However, the extent of the guiding effect has not been directly measured. In our ex vivo recording geometry, light comes from the photoreceptor side and is exempt from this effect. Based on the geometry of mouse cone photoreceptors, the light funneling by cone inner segments could increase the photon capture by a factor of 7 (Lyubarsky et al., 1999). An additional complication arises from the rod shadowing effect in our ex vivo recording geometry where light is partially absorbed by rhodopsin molecules in rods before reaching the cone outer segments. By taking into account previously suggested magnitudes of light guiding in vivo and light shadowing by rods ex vivo (decrease photon capture of cones by 1.4-fold ex vivo), we conclude that cone outer segments should actually receive more photons in vivo than ex vivo under identical stimulation conditions such as the 530 nm LED ganzfeld used in this study. This means that, differently from rods, where identical stimuli are sensed brighter ex vivo compared to in vivo, cones in the isolated retina should be more sensitive to identical light stimulation in in vivo conditions. As cone amplification constant of phototransduction is expected to be preserved most consistently across different recording conditions, we estimate that identical \( A \) in both in vivo and ex vivo conditions requires a funneling factor of 2.7.

4.3. Comparison of molecular amplification of rod and cone phototransduction activation reactions between in vivo and ex vivo conditions

The nose peak also affects the normalization choice for the rod amplification constant analysis both in vivo and ex vivo, when synaptic blockers are not used. Our results suggest that the maximum response amplitude might contain a substantial contribution from the nose component. When this parameter is set as the peak of the a-wave at the nose level, the determination of rod amplification constant ex vivo yields ~2-fold lower \( A \)-values as compared to those obtained by normalization to the plateau level of the photoreponse. In our conditions, this generates a corrected amplification constant of about 29 s\(^{-2}\) for rods (see Table 1). Thus, it is possible that the real molecular amplification constant of rods in vivo (or ex vivo) is over two times larger than previously thought (see Fig. 4C, D, and Table 1).

Cones are less sensitive and operate under brighter light compared to rods. Correspondingly, it would be logical for cones to have a lower gain of phototransduction. Indeed, it has been shown that cones in eastern chipmunk have significantly lower amplification of phototransduction as compared to their rods (Zhang, Wensel, & Kraft, 2003). However, other studies suggested that the amplification constants of mouse and human rods and cones are comparable (Heikkinen, Nymark, & Koskelainen, 2008; Nikonoruk et al., 2006; Smith & Lamb, 1997). As discussed above, several factors complicate the determination of the cone amplification constant. Here, we used Gnat1\(-/-\) mice whose rods do not respond to light to facilitate the extraction of a truly dark-adapted cone responses in both in vivo and ex vivo settings. Nikonoruk et al., 2006 showed by single-cell recordings that cones of Gnat1\(-/-\) mice have close to normal cone amplification constant (\( A_{\text{WT,cone}} = 3 \times \text{s}^{-2} \), \( A_{\text{Gnat1,-/-,cone}} = 2 \times \text{s}^{-2} \)). Hence, our amplification constant values should be comparable to those of WT cones under the same experimental conditions. Our results indicate that the amplification constant of Gnat1\(-/-\) mouse cones is between 1.4 s\(^{-2}\) (\( f = 7 \)) and 10 s\(^{-2}\) (\( f = 1 \)) in vivo and ca. 6 s\(^{-2}\) to 3 s\(^{-2}\) in ex vivo conditions (in the presence of APB) when the collection area of the cone outer segment was assumed to be 0.07 \( \mu \text{m}^2 \). Overall, our results suggest that the molecular amplification in mouse cones might be indeed lower compared to that in rods. A potential complication could be that part of the negative dim-flash a-wave in Gnat1\(-/-\) ERG recordings might be generated by off-bipolar cells rather than by cone photoreceptors. However, our present results as well as previous study (Lyubarsky et al., 2002) demonstrate that cone amplification constants determined from ERG and single cell recordings (Nikonoruk et al., 2006) are comparable, indicating that the leading edge of the a-wave describes well the activation of the cone phototransduction in Gnat1\(-/-\) mice.

Disclosures

Washington University in St. Louis has a license agreement with Xenotec, Inc. and may receive a royalty from the sale of the ex vivo adapter.

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Appendix A. Supplementary material

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References


