# Melanopsin-Expressing Amphioxus Photoreceptors Transduce Light via a Phospholipase C Signaling Cascade

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## Abstract

Melanopsin, the receptor molecule that underlies light sensitivity in mammalian 'circadian' receptors, is homologous to invertebrate rhodopsins and has been proposed to operate via a similar signaling pathway. Its downstream effectors, however, remain elusive. Melanopsin also expresses in two distinct light-sensitive cell types in the neural tube of amphioxus. This organism is the most basal extant chordate and can help outline the evolutionary history of different photoreceptor lineages and their transduction mechanisms; moreover, isolated amphioxus photoreceptors offer unique advantages, because they are unambiguously identifiable and amenable to single-cell physiological assays. In the present study wholecell patch clamp recording, pharmacological manipulations, and immunodetection were utilized to investigate light transduction in amphioxus photoreceptors. A G<sub>q</sub> was identified and selectively localized to the photosensitive microvillar membrane, while the pivotal role of phospholipase C was established pharmacologically. The photocurrent was profoundly depressed by IP<sub>3</sub> receptor antagonists, highlighting the importance of IP<sub>3</sub> receptors in light signaling. By contrast, surrogates of diacylglycerol (DAG), as well as poly-unsaturated fatty acids failed to activate a membrane conductance or to alter the light response. The results strengthen the notion that calcium released from the ER via IP<sub>3</sub>-sensitive channels may fulfill a key role in conveying - directly or indirectly - the melanopsin-initiated light signal to the photoconductance; moreover, they challenge the dogma that microvillar photoreceptors and phoshoinositide-based light transduction are a prerogative of invertebrate eyes.

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## Introduction

A long-held belief, based on detailed morphological observations of eyes and photoreceptors across a wide variety of animal species, maintained that vision has evolved independently in different phyla [1]. In particular, microvillar photoreceptors, on one hand, and rods and cones on the other - the two canonical classes of light-sensing cells - have been thought to represent an instance of convergent evolution, and to be exclusively confined to invertebrates and vertebrates, respectively. Contrary to such view, recent lines of evidence derived from the molecular analysis of the genes that specify eye development and of the protein families that underlie light transduction, are gradually converging on the conclusion that visual mechanisms in metazoa may share a monophyletic origin [2]. In fact, microvillar photoreceptors are the likely descendants of the light sensor of the most primitive protoeye already present in pre-bilateria [3,4]; as such, one could expect to find them across both protostomia and deuterostomia. Representation of this ancient line of visual cells amongst the vertebrates had been traditionally dismissed, but has strongly reemerged in recent times with the identification of unconventional photoreceptors in the ganglion layer of the mammalian retina [5,6], dubbed intrinsically photosensitive retinal ganglion cells (ipRGCs).

These cells subserve non-visual light-dependent functions, such as controlling the pupillary reflex and photo-entraining the circadian rhythms. Like invertebrate rhabdomeric visual cells, these 'circadian' photoreceptors depolarize to light, and their photopigment, melanopsin, has a significant similarity to the rhodopsin of invertebrates [7]; furthermore, clues have been garnered that the transduction cascade is PLC-dependent [8,9]. Nonetheless, the characteristic microvillar cellular architecture is conspicuously absent in ipRGCs, and a detailed investigation of the effector mechanisms in native cells proved challenging, owing to their extreme scarcity and difficult identification. In the present report an alternative model system was utilized to address the nature of the light-signaling pathway that couples melanopsin photostimulation to the ion channels underlying the receptor potential. The approach adopted was to exploit an early chordate in which melanopsin-expressing cells are not as under-represented as in the retina of mammals, and have retained the distinguishing microvillar traits, making their classification unambiguous and facilitating their targeting in physiological studies. According to molecular phylogeny, amphioxus is the most basal living chordate [10]. Melanopsin has been detected in its neural tube [11], and its pattern of expression coincides with the distribution of two groups of cells, known as Joseph cells and organs of Hesse, respectively,

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which display distinct microvilli [12,13,14]. Our recent patch clamp measurements in enzymatically isolated cells of both types established that they are indeed primary photoreceptors, and their light response has an action spectrum similar to the absorption spectrum reported for amphioxus melanopsin in vitro [15]. The receptor potential of both Joseph and Hesse cells is depolarizing, and is accompanied by an increase in a cationic membrane conductance; furthermore, imaging with fluorescent indicators reveals that light mobilizes calcium from intracellular stores. In the present work, we utilized a multi-pronged approach to examine the nature of the enzymatic cascade that couples light absorption to ion channel activation. The significance of this endeavor is twofold: on the one hand it can help elucidate the evolutionary history of microvillar photoreceptors and their representation amongst chordates; on the other hand, it provides clues on the lightsignalling mechanisms of melanopsin in a native system.

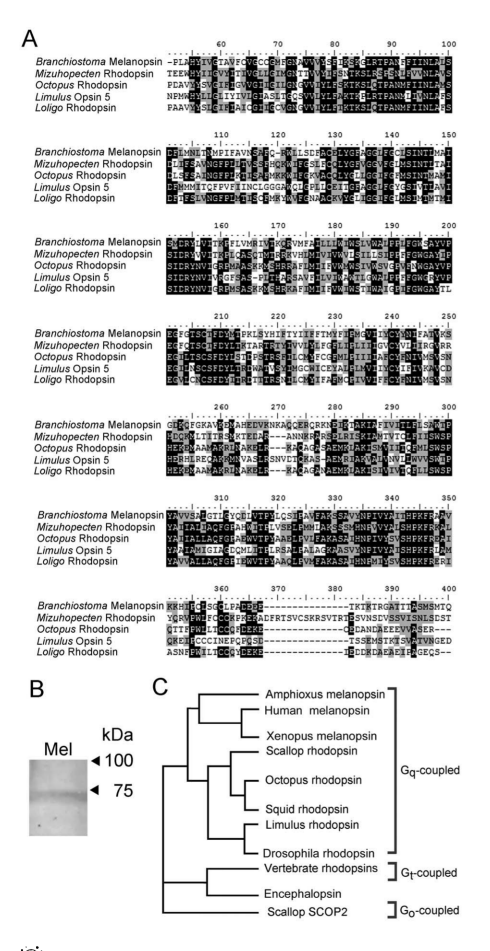
## Results

The microvillar architecture of Joseph and Hesse cells of amphioxus is reminiscent of the rhabdomeric photoreceptors of invertebrates, and so are the functional properties of their photoresponse. Moreover, the predicted aminoacid sequence of amphioxus melanopsin resembles most closely, amongst nonmelanopsin photopigments, the rhodopsin of microvillar visual cells of the scallop (Mizuhopecten yessoensis), octopus (Octopus dofleini), horseshoe crab (Limulus polyphemus), and squid (Loligo forbesi). In all cases the overall identity is 21-22%; by contrast, the similarity to the closest opsins from ciliary-type photoreceptors is substantially lower (15%-17% overall, far below all other reported microvillar photoreceptor rhodopsins). Figure 1A shows a multi-sequence alignment restricted to the core region of the amphioxus melanopsin (encompassing the 7 trans-membrane helices but excluding part of the N-terminus and the C-terminal region, where the greatest degree of divergence appears) against the aforementioned species, revealing a striking similarity. Panel B of the same figure shows a Western blot of amphioxus neural tube lysate, probed with antibodies raised against an extracellular domain of human melanopsin, that cross-react with other vertebrate forms (V-20, Santa Cruz Biotechnology). A single band was detected, with an apparent molecular weight of  $\approx$ 71 kDa; this is somewhat higher than that of mammals (typically  $\approx$ 49–62 kDa, depending on glycosylation state) [16], but compares favorably with the figure of  $\approx 67$  kDa from the predicted aminoacid sequence of Branchiostoma floridae melanopsin. Panel C of Figure 1 shows a simplified phylogenetic tree of representative animal photopigments, illustrating how amphioxus and vertebrate melanopsins group with the rhodopsins that are known to signal via G<sub>q</sub>. It is therefore plausible that the coupling of photon absorption to channel gating in amphioxus may also be based on the G<sub>q</sub>-triggered PLC signaling cascade.

 $G_q$  has been previously identified in another species of amphioxus, *B. belcheri*, where its distribution matches the expression pattern of melanopsin [11]. We sought to obtain more detailed information on its cellular localization, to buttress the proposed role in light transduction. We found a commercial polyclonal antibody (Millipore anti-Gq/11CT) raised against a synthetic peptide that is 100% identical to the predicted Cterminal aminoacid sequence of the putative  $G_q$  found in the amphioxus genome. This motif is widely conserved across species, and Figure 2A shows an alignment of the immunogenic peptide sequence (QLNLKEYNLV) against the last 50 aminoacids of  $G_q$ of *Branchiostoma* and other organisms. In a Western blot of neural tube this antibody identified a single band (Figure 2B), with the expected molecular weight (≈42 kDa). The same antibody was then utilized in immunohistochemistry. We focused on Hesse cells, because the presence of the companion pigmented screening cell makes their identification unambiguous even in tissue sections. Figure 2C illustrates the main features of a dissociated Hesse cell: the accessory screening cell engulfs the microvilli-covered region of the sensory cell [12]; therefore, in a slice that cuts through the middle of the ocellus, the profile of the villous region (red drawing in Fig. 2C) would be revealed. The left panel of Figure 2D shows a Nomarski micrograph of a 10 µm section of fixed neural tube containing two Hesse cells; the one on the left was sliced near the middle. The right panel shows the corresponding fluorescence image, stained with anti-G<sub>a</sub> antibodies and Alexa Fluo 488conjugated secondaries. The 'crown' of microvilli is distinctly and selectively decorated, abutting a crescent-shaped slice of the dark accessory cell. It can be concluded that Gq expresses in the photosensitive membrane of identified Hesse cells. We sought therefore functional evidence for the participation of the Gq-triggered cascade in light transduction.

Phospholipase C (PLC) is the prime target of  $G_q$ . The compound U-73122 has been shown to inhibit the activation of PLC in a variety of experimental systems [17,18], including microvillar invertebrate photoreceptors [19]. The effects have been reported to be irreversible. We tested this drug on the lightevoked current in voltage-clamped Joseph and Hesse cells. U-73122 was initially dissolved in DMSO at a 1 mM concentration, and an aliquot of this stock solution diluted 1:100 in ASW to a final concentration of 10 µM. Repetitive flashes of light of subsaturating intensity were applied every 60 sec. After attaining stability, two control responses were recorded and then the drug was rapidly introduced by local superfusion via a puffer pipette. Figure 3A illustrates the progressively decreasing photocurrents in representative cells of the two types, while Figure 3B shows a plot of the time course of changes in the peak amplitude of the lightevoked current, averaged separately for 3 Joseph cells and 4 Hesse cells. For each cell, the data were normalized with respect to the size of the first response. In both classes of cells, U-73122 caused a dramatic suppression of the photocurrent, and within  $\approx$ 3–4 minutes the light response was virtually abolished. By contrast, voltage-activated currents were unaffected by the drug. Control application of ASW containing 1% DMSO produced no effect.

Our previous results demonstrated that light mobilizes calcium from internal stores, and that blunting Ca elevation by chelators severely impairs the photoresponse [15]. Together with the effects of U-73122 described above, these observations likely implicate  $IP_3$  in the Ca-dependent regulation of the light response. To corroborate such notion, we examined the impact of antagonists of the IP<sub>3</sub> receptor on the photocurrent. We first tested 2aminoethoxydiphenyl borate (2-APB), which has been widely utilized as an inhibitor of the  $IP_3$  receptor [20]. The advantages of this drug include its membrane-permeability and reversibility, thus allowing extracellular application and within-cell comparisons. The effect of 2-APB on the light response was assessed by superfusion at a concentration of 100 µM. Figure 4A shows photocurrent traces recorded in a Hesse cell and in a Joseph cell in control solution, during exposure to 2-APB, and after washing the drug. In both cases, a rapid, reversible reduction of the peak amplitude of the photocurrent occurred. Panel B of the same figure summarizes the results: the inhibition of the light response, calculated as 1-R\_{2-APB}/R\_{\rm Control}, was 77.6\%  $\pm 11.6\%$  SD for Hesse cells (n = 6), and  $89.9\% \pm 7\%$  in Joseph cells (n = 5). The somewhat lower apparent effectiveness of 2-APB in Hesse cells may hint at the greater difficulty of penetration of the drug into the microvillar

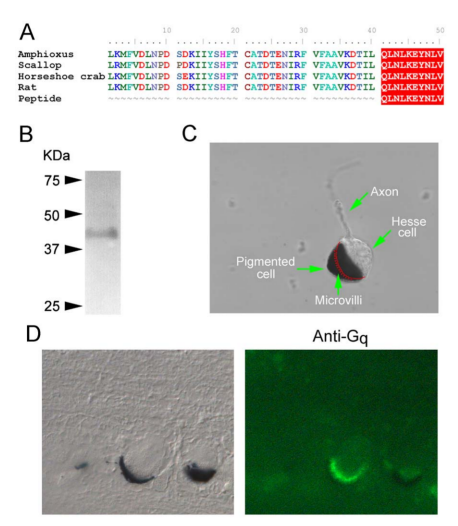


**Figure 1. Amphioxus melanopsin groups with**  $G_q$ **-coupled rhodopsins.** (*A*) The translated sequence of *Branchiostoma* was subjected to a BLAST search, and subsequently aligned with the highest-ranking hits among *non*-melanopsin photopigments. All belong to microvillar photoreceptors from eyes of invertebrates. The Clustal W alignment shows the core region of the proteins (omitting the carboxy-terminus, and part of the amino-terminus, which are generally divergent); black shading indicates identity, whereas grey shading indicates conservative aminoacid substitutions. Accession numbers: amphioxus (*Branchiostoma*) Q4R114; scallop (*Mizuhopecten*), O15973.1, octopus (*Octopus*). P09241.1; horseshoe crab (*Limulus*), ACO05013.1; and squid (*Loligo*) P24603.1. (*B*) Western blot with anti-melanopsin antibodies. A single band of  $\approx$ 71 kDa was detected, closely approaching the size expected from the predicted aminoacid sequence. This confirms that the melanopsin protein expresses in the neural tube of *B. floridae*. (*C*) Phylogenetic tree illustrating the grouping of the main classes of photopigments found in the animal kingdom. Amphioxus melanopsin is closely related to the G<sub>q</sub>-coupled rhodopsin of mollusks and arthropods. doi:10.1371/journal.pone.0029813.q001

region, where access is hindered by the abutting pigmented cell; however, this trend fell short of statistical significance.

Although 2-APB was initially thought to act selectively on the  $IP_3$  receptor, subsequent reports have questioned its alleged high specificity, showing more widespread effects on calcium-signaling, including an antagonistic action on store-operated calcium influx [21] and on SERCA pumps [22]. While 2-APB remains a useful tool for examining phosphoinositide-dependent Ca mobilization we sought additional supporting evidence for the involvement of the  $IP_3R$  in the light response. We examined the effects of low-

molecular weight heparin (<6 KDa), which has long been known to reduce the IP<sub>3</sub>-triggered release of calcium from the endoplasmic reticulum [23]. The effects of heparin on the light response were assessed by dialyzing it into amphioxus photoreceptors at a concentration of 2 mg/ml (dissolved in standard intracellular solution), and administering repetitive flashes of light (1/min), starting immediately after attaining the whole-cell configuration. In Figure 4C the average peak amplitude of the photocurrent, normalized by the largest response, is separately plotted as a function of time for Hesse and Joseph cells, under



**Figure 2.**  $G_q$  expresses in the microvillar membrane of Hesse cells. (*A*) The sequence of the immunogenic peptide used to raise anti- $G_q$  antibodies was aligned with the predicted C-terminal region of  $G_q$  of amphioxus, and those of other organisms. (*B*) Western blot of neural tube using anti  $G_{qr}$  showing the detection of a single band of  $\approx 42$  kDa. (*C*) Morphological characteristics of the Hesse cell: the accessory pigmented cell engulfs the microvilli-covered region of the clear sensory cell; the position of the villous membrane within the occluded region is drawn in red. (*D*) *Left*: Nomarski micrograph of a 10 µm section of fixed neural tube containing two Hesse cells. *Right*: fluorescence image of the same section incubated with anti- $G_q$  antibodies and Alexa Fluo 488-conjugated secondary antibodies. The immunostaining is confined to the region of the microvilli. doi:10.1371/journal.pone.0029813.g002

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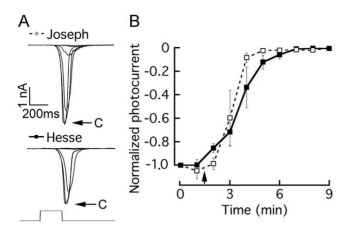
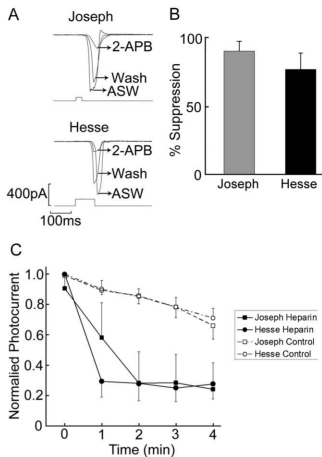


Figure 3. Pharmacological interference with PLC suppresses the photocurrent. The PLC antagonist U-73122 (10  $\mu$ M) was applied to voltage-clamped photoreceptors stimulated every minute with a flash of light of fixed intensity. (A) Superimposed current traces measured in a Joseph cell and in a Hesse cell; after photocurrent stabilization, two control responses were measured ('c' and arrows), and then the drug was introduced by local superfusion, inducing a progressive decline in the peak amplitude of the light-evoked current in both cell types. (B) Time course of the effect of the PLC inhibitor, averaged for 4 Hesse cells (solid lines/filled symbols) and 3 Joseph cells (dashed lines/open symbols); error bars indicate standard error; the starting time of drug application is marked by the arrow. doi:10.1371/journal.pone.0029813.g003

control conditions (standard internal solution) and with heparin dialysis. Heparin-treated photoreceptors of both types exhibited a dramatically enhanced decline of the light response over time, compared to the controls, which only show the normal rundown that typically occurs during the first few minutes of recording, before the photocurrent stabilizes. Because the outcome was similar for Hesse and Joseph cells, the values from the two cell classes were combined and subjected to an analysis of variance with repeated measurements. The difference between the control (n = 10) and heparin-treated cells (n = 7) was statistically significant (F = 29.3, df = 16, p<0.001).

The above results strengthen the contention that the PLC cascade mediates light transduction in amphioxus; moreover, in conjunction with a prior demonstration of the profound inhibitory effect of intracellular BAPTA [15], they suggest that IP<sub>3</sub>-triggered Ca release plays a central role in this process. Hydrolysis of PIP<sub>2</sub> by PLC- $\beta$  generates, in addition to the soluble messenger IP<sub>3</sub>, membrane-bound diacylglicerol (DAG), which is also endowed with signaling properties: beyond its well-known ability to activate protein kinase C [24], DAG or its analogs and metabolites can also directly interact with ionic channels; among the reported effects are the stimulation of a cationic membrane conductance related to the light response in microvillar photoreceptors of Lima [25] and Drosophila [26], as well as the activation of heterologouslyexpressed TRPL channels [27]. We tested several surrogates of DAG, including the phorbol ester PMA  $(1 \mu M, n=6)$ , the structural analog DOG (50–100  $\mu$ M, n=4), OAG (10  $\mu$ M, n=6) and SAG (10  $\mu$ M, n=8). The compounds were either dialyzed intracellularly via the patch electrode, or applied extracellularly by rapid superfusion. The procedure entailed examining possible direct effects on membrane current, as well as alterations of the light-evoked current; to this end,  $I_{\rm m}$  was recorded for 3 minutes in the dark as chemical stimulation was applied, followed by an assessment of photoresponsiveness by a full light-intensity series. Figure 5 illustrates the outcome obtained with

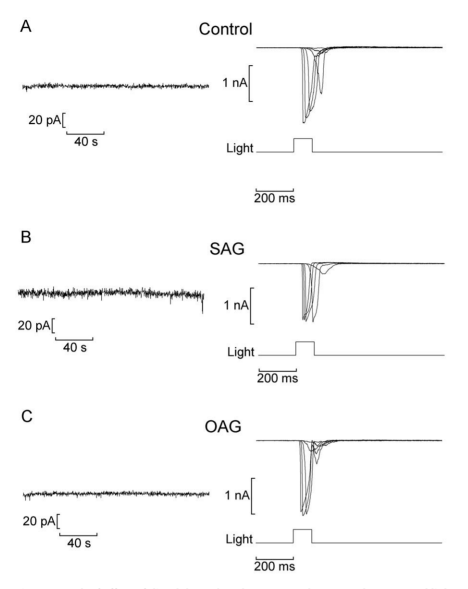


**Figure 4. Inhibition of the light response by antagonists of the IP**<sub>3</sub> **receptor.** (*A*) Effect of 2-APB applied extracellularly at a concentration of 100  $\mu$ M. Examples of photocurrent recordings in Joseph and Hesse cells before, during and after puffer application of 2-APB, highlighting the reversible inhibitory effects. Holding potential -50 mV. (*B*) Average inhibition of the light-evoked current by 2-APB; error bars indicate the standard deviation (n = 5 for Joseph cells, n = 6 for *Hesse cells*). (C) Effect of heparin. Low-molecular weight heparin was dialyzed intracellularly through the patch pipette (2 mg/ml), as voltageclamped photoreceptors were repeatedly stimulated with a 100-ms flash of light. The peak amplitude of the photocurrent is plotted as a function of time, starting immediately after breaking the patch of membrane to access the cell interior (*Control: Joseph n* = 6, *Hesse n* = 4; *Heparin: Joseph n* = 3, *Hesse n* = 4).

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two of these compounds, SAG and OAG: no change in holding current was detected, and the photocurrent showed no significant differences with respect to controls. All the DAG surrogates tested proved similarly inert in both Hesse and Joseph cells. In *Drosophila* it has been reported that poly-unsaturated fatty acids (PUFAs) such as arachidonic, linolenic and linoleic acids stimulate the light-sensitive channels [26]. Arachidonic acid can be generated from DAG by DAG lipase, and is thus able to serve as a downstream messenger in this branch of the cascade. Because the DAG analogs we utilized are not metabolized to such PUFAs, we examined the effect of direct application of arachidonic acid (5  $\mu$ M; n = 4) and linolenic acid (50  $\mu$ M; n = 4). Again, we failed to observe changes in membrane currents in the dark or in the light-evoked current (*not shown*).

Finally, although the support for  $G_q$ /PLC in light signalling in amphioxus seems compelling, we examined the possible involve-



**Figure 5. Lack of effect of diacylglycerol analogs on membrane conductance and light-evoked current.** (A) Control Hesse cell dialyzed with standard internal solution. (*Left*) Recording of membrane current in the dark immediately after attaining the whole-cell configuration; (*right*) currents evoked by flashes of increasing intensity. (B,C) Similar experiments conducted in Hesse cells dialyzed with SAG (1-Stearoyl-2-arachidonoyl-*sn*-glycerol; 10  $\mu$ M) and OAG (1-Oleoyl-2-acetyl-sn-glycerol; 10  $\mu$ M), respectively. Neither the holding current nor the photocurrents were affected. Membrane potential was clamped at -50 mV throughout. doi:10.1371/journal.pone.0029813.q005

ment of cyclic nucleotides in controlling the light response. Cyclic GMP has been proposed as an effector in the PLC-mediated light transduction of Drosophila [28] and Limulus [29], although no support has been garnered for the required cross-talk between the two pathways [30,31]. However, the issue deserved some scrutiny, because cyclic nucleotides remain the only messenger substances for which the evidence of a *direct* effect on the gating of sensory transduction channels is undisputed, as documented in vertebrate rods [32] and in olfactory neurons [33]. We tested the effects of the poorly hydrolyzable analogs 8-Br-cGMP and 8-Br-cAMP, applied via the patch pipette (20 µM). No change in holding current was observed during intracellular perfusion in the dark while the cell vas maintained under voltage clamp at -50 mV (8-Br-cGMP: 2 Joseph and 2 Hesse cells; 8-Br-cAMP: 3 Joseph and 2 Hesse cells); subsequent tests with flashes of light showed that the photoresponse was equally unaffected (not shown).

## Discussion

The discovery of melanopsin in the vertebrate retina and the realization that it diverges from the rhodopsin of rods and cones prompted a quest for its mechanisms of action. Many of the studies addressing the nature of the biochemical cascade coupling photon absorption to ion channel gating have relied on heterologous expression, sometimes resulting in conflicting observations. In melanopsin-expressing HEK 293 cells [34] and *Xenopus* oocytes [35] it was found that, upon reconstituting a functional photopigment by exogenous retinal, a light-induced electrical response is recorded which is antagonized by inhibitors of  $G_q$  and PLC. By contrast, PLC was apparently not involved in the photosensitivity of Neuro-2A cells expressing melanopsin, and cyclic nucleotides seemingly played a role [36], whereas COS cell-expressed melanopsin activates transducin ( $G_t$ ) *in vitro* [37]. An

important caveat with such an approach is that G protein-mediated cascades can be promiscuous, and implanted receptors often signal through endogenous pathways that differ from those of native cells: a case in point is mammalian rhodopsin, which in rods signals through G<sub>t</sub>/PDE/cGMP, but upon heterologous expression in *Xenopus* oocytes it is capable of mediating light responses by stimulating the  $G_q/PLC-\beta/IP_3$  cascade of the host cell, which normally participates in the fertilization response [38]. It is therefore crucial to conduct functional studies utilizing native melanopsin-expressing cells. The progress in the physiological analysis of mammalian ipRGCs has been painstaking, as their extreme scarcity ( $\approx 1\%$  of the retinal ganglion cell population), together with the lack of morphological landmarks to identify them, proved a formidable hurdle; the recent development of transgenic lines of animals in which such cells express a fluorescent marker [39] is a valuable approach that will help overcome such limitations. A few reports on mice ipRGCs documented phenomena that are consistent with a rhabdomeric-like lightsignaling pathway: for example, the electrical response to light in ipRGCs is disrupted by peptides that preferentially antagonize  $G_{\alpha}$ , and by PLC antagonists [9]. Also, in a cone-less/rod-less background (to eliminate the potential confounding of rod and cone-driven input), a subset of retinal ganglion cells produce a light-stimulated increase in cytosolic calcium [40]; however, this Ca mobilization is only phenomenologically akin to that of rhabdomeric photoreceptors, as it appears to arise from influx (chiefly via voltage-gated Ca channels), rather than phosphoinositide-dependent release [41]. As for downstream effectors, there are suggestions based on microarray analysis that a PKC-z may be implicated [42] and immunohistochemical indications that the light-controlled conductance may be comprised of TRP-class ion channels e.g. TRP6 [43]. Many gaps, therefore, remain to be filled

The results described in the present work strongly support the involvement of PLC signaling in the phototransduction cascade of melanopsin-expressing Joseph and Hesse cells of the neural tube of amphioxus. These observations complement the reports describing similarities between these cells and microvillar light-sensing cells of invertebrates, both in terms of morphology [12,44,13,14] and electrophysiology of the photoresponse [15]. Taken together, the evidence strengthens the contention that Joseph and Hesse cells are bona fide members of the family of rhabdomeric photoreceptors. In addition to underscoring the central position of  $G_q$  and PLC for generating the melanopsin-mediated light response in amphioxus, our results provide further clues on the mechanisms implicated. A key role for PLC has recently been demonstrated in the intrinsic light sensitivity of the mammalian iris and ipRGCs [45]. In amphioxus, the robust impact of calcium and IP<sub>3</sub>R manipulations on light responsiveness and the lack of discernible effects by the variety of DAG analogs tested are indicative of a key role of the  $IP_3$  branch of the PLC cascade in the control of the photoconductance. Such a scheme has been proposed to hold for some rhabdomeric photoreceptors such as those of Limulus [46], whereas the importance of signaling by the lipid branch has been emphasized in other species, like Lima [25] and Drosophila [26]; in the latter, in fact, knockout of the  $IP_3$  receptor does not adversely affect the photoresponse [47,48]. Establishing the soundness of such dichotomy will have to await positive confirmation of the identity of the final internal messengers that operate the light-sensitive ion channels in the various cases. Nonetheless, it is noteworthy that in murine ipRGCs, putative descendents of Hesse and Joseph cells, application of OAG reportedly also failed to alter membrane current and photoresponsiveness [9].

On the basis of molecular phylogeny, amphioxus melanopsin has been proposed to constitute a link between  $G_q$ -coupled opsins of invertebrate and those of vertebrate circadian receptors [49,50]. Therefore, amphioxus microvillar photo-sensitive cells provide a glimpse of the rhabdomeric lineage of light detectors in early chordate evolution, and may be regarded as ancestral to the circadian photoreceptors found in present-day vertebrates [51,52].

#### **Materials and Methods**

All animal procedures were conducted according to institutional guidelines.

### Cell isolation

Amphioxus (*Branchiostoma floridae*) were obtained from Gulf Specimens Marine Laboratories (Panacea, FL), and maintained in a sea-water aquarium on a diet of marine phytoplankton, and a 12/12 hr light/dark cycle. Animals were anesthetized by hypothermia, the rostral end was cut and pinned to a Sylgard-coated chamber, and the neural tube was excised. The tissue was then incubated with Pronase (Boehringer; 750 units/ml, 50 min at  $22^{\circ}$ C), followed by extensive washing in ASW supplemented with 4% fetal calf serum, and mechanical trituration with a fine-bore fire-polished Pasteur pipette. The resulting suspension was plated into a perfusion chamber mounted on the stage of an inverted microscope (Zeiss). The coverslip bottom of the chamber was pretreated with Concanavalin-A to promote cell adhesion. Dissociated cells remain physiologically viable for several hours.

## Electrophysiological recording

Patch pipettes were fabricated from borosilicate glass, firepolished, and filled with an intracellular solution (standard composition: 100 mM KCl, 200 K-Aspartate or K-glutamate 5 MgCl<sub>2</sub>, 5 Na<sub>2</sub>ATP, 20 NaCl, 1 EGTA, 300 Sucrose, 10 HEPES, 0.2 GTP, pH 7.3). Electrode resistance in ASW is 2-4 M $\Omega$ ; series resistance was compensated electronically (maximum residual error  $\leq 2$  mV). Data were digitized with an analog-digital interface (Data Translation) which served also to generate stimuli, under the control of software developed in-house. Extracellular ions were exchanged by a system of reservoirs and multi-port valves, perfusing the entire flow-chamber. Alternatively, for rapid application, 'puffer' pipettes were lowered to a pre-set target position near the cell by a programmable positioner (Eppendorf). Pressure-ejection was controlled by solenoid-activated valves. Statistical comparisons of changes in current amplitude over time across conditions were carried out by analysis of variance with repeated measurements (program VassarStats).

#### Light stimulation

Broad-band light stimuli were generated by a tungsten-halogenquartz (THQ) light source (Oriel); IR was removed by a heatabsorbing filter ( $\lambda$ >800 nm). Solenoid-driven shutters (Uniblitz), calibrated neutral density filters (Melles-Griot, Zevenaar, The Netherlands), and interference filters (Omega Optical, Brattleboro, VT) controlled the duration, intensity, and wavelength of stimulation. A pin-hole restricted the illuminated region to a focused spot ( $\approx$ 150 µm). Alternatively, a blue LED (peak emission 470 nm) was driven by a computer-controlled precision current pump, and delivered a full-field light stimulus via a fiber optics bundle. Light was measured with a radiometer (UDT, Hawthorne, CA), and converted to effective photon flux via an in vivo calibration which made possible to compare across arrangements; the unattenuated beam intensity from the THQ illuminator was  $3.53 \times 10^{15}$  photons×s<sup>-1</sup>cm<sup>-2</sup>, while the maximum intensity of the LED was  $2.2 \times 10^{16}$  photons  $\times s^{-1}$  cm<sup>-2</sup>. During experimental manipulations the cells were illuminated with near-IR light ( $\lambda > 780$  nm; Andover Corporation long-pass filter) and viewed with the aid of a CCD camera (Sony).

#### Chemicals

The phospholipase C antagonist U-73122 (1-[6-[((17 $\beta$ )-3-Methoxyestra-1,3,5[10]-trien-17-yl)amino]hexyl]-1H-pyrrole-2,5dione) was from Calbiochem. Low-molecular weight heparin (<6 KDa), 2-APB (2-aminoethoxydiphenyl borate) and 8-BrcAMP (8-Bromo cyclic adenosine monophosphate) were obtained from Sigma-Aldrich. PMA (phorbol ester Phorbol 12-myristate 13acetate), DOG (1,2 Dioctanoyl-*sn*-glycerol), OAG (1-Oleoyl-2acetyl-sn-glycerol), SAG (1-Stearoyl-2-arachidonoyl-*sn*-glycerol), arachidonic acid, and linolenic acid were purchased from Alexis biochemicals/Enzo Life Sciences. 8-Br-cGMP (8-Bromo cyclic guanosine monophosphate) was from Fluka.

#### Western blots

One isolated neural tube was homogenized (teflon/glass) in the presence of protease inhibitors (100  $\mu$ M PMSF, 1  $\mu$ M pepstatin, and 0.1% Sigma protease inhibitor cocktail), acetone-precipitated for 1 hour at  $-20^{\circ}$ C, and centrifuged 20 min at 10,000×g. The pellet was air-dried, resuspended in sample buffer and separated by SDS PAGE (8–12%). Proteins were electrotransfered (1 hour, 100 V) onto nitrocellulose membrane which was blocked overnight with 3% BSA. The membrane was then sequentially

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incubated with primary antibodies, washed in tris buffer saline (TBS) and incubated in alkaline phosphatase-conjugated secondary ABs. After the final washes, the nitrocellulose membrane was developed in Western Blue (Promega, Madison, WI, USA).

## Immunohistochemistry

After removal of the dorsal fin, the rostral portion of amphioxus was fixed either in 4% paraformaldehyide overnight at 4°C, or by flash-freezing in isopentane chilled in a dry-ice/liquid nitrogen slurry. Subsequently, samples were dehydrated in EtOH at  $-80^{\circ}$ C for 4 days, before impregnating with polyester wax, sectioning at 10 µm and mounting onto glass slides subbed with 0.1% gelatin in 0.05% chromium potassium sulfate. Before use, the wax was dissolved by briefly dipping in 100% EtOH, and sections were rehydrated in 70% EtOH followed by H<sub>2</sub>O, permeabilized in 0.2% Triton-X (5 min), and blocked with 1% goat serum (1 hr). Antibodies were applied in PBS+0.5% BSA; secondary antibodies were conjugated to Alexa Fluo- 488 (Molecular Probes), and the samples were viewed with a Zeiss Axioscope epi-fluorescence microscope.

#### **Author Contributions**

Conceived and designed the experiments: EN MPG. Performed the experiments: JMA CP GM MPG. Analyzed the data: EN MPG JMA CP GM. Contributed reagents/materials/analysis tools: EN. Wrote the paper: EN MPG.

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