Have the eyes of bioluminescent scale worms adapted to see their own light? A comparative study of eyes and vision in Harmothoe imbricata and Lepidonotus squamatus

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ABSTRACT
Annelids constitute a diverse phylum with more than 19,000 species, which exhibit greatly varying morphologies and lifestyles ranging from sessile detritivores to fast swimming active predators. The lifestyle of an animal is closely linked to its sensory systems, not least the visual equipment. Interestingly, many errantian annelid species from different families, such as the scale worms (Polynoidae), have two pairs of eyes on their prostomium. These eyes are typically 100–200 µm in diameter and structurally similar judged from their gross morphology. The polynoids Harmothoe imbricata and Lepidonotus squamatus from the North Atlantic are both benthic predators preying on small invertebrates but only H. imbricata can produce bioluminescence in its scales. Here, we examined the eye morphology, photoreceptor physiology and light-guided behaviour in these two scale worms to assess their visual capacity and visual ecology. The structure and physiology of the two pairs of eyes are remarkably similar within each species, with the only difference being the gaze direction. The photoreceptor physiology, however, differs between species. Both species express a single opsin in their eyes, but in H. imbricata the peak sensitivity is green shifted and the temporal resolution is lower, suggesting that the eyes of H. imbricata are adapted to detect their own bioluminescence. The behavioural experiments showed that both species are strictly night active but yielded no support for the hypothesis that H. imbricata is repelled by its own bioluminescence.

KEY WORDS: Annelida, Polynoidae, Vision, Bioluminescence, Eye physiology, Night active

INTRODUCTION
Light reception takes place in almost all animals and ocelli and eyes are found in about half of the animal phyla (Land and Nilsson, 2012). Within annelids ocelli and eyes are present in a large number of species and the majority of larger clades. Still, in the derived lineage of clitellates (earth worms and leeches) only a few species have small pigment cup eyes and light guided behaviours seem to be limited to simple phototaxis in most of them (Verger-Bocquet, 1992). The tube dwelling or burrowing species of the large clade Sedentaria, are also often eyeless, but some species have highly sophisticated compound eyes like the Christmas tree worm, Spirobranchus giganteus, and the fan worm Acromegalomma vesiculosum (Bok et al., 2017, 2019). These eyes can have more than 1000 ommatidia each, putatively having spatial resolution rivaling many insect eyes, but it remains unknown if and how this image information is used and thus what functional significance the eyes have.

In the large sister group of Sedentaria, the Errantia, eyes are common. They vary in size and complexity from pigment cups with only a few receptors to highly advanced camera-type eyes with large spherical lenses and 1000s of photoreceptors (Hermans and Eakin, 1974; Wald and Rayport, 1977). An often-found arrangement of the visual system in Errantia is two pairs of eyes on the prostomium, as seen in many nereids and syllids, for example, Alitta (=Nerics) virens and Odontosyllis enopla (Dorsett and Hyde, 1968; Wolken and Florida, 1984). These eyes often appear similar in structure though there might be a size difference between the two pairs. They are typically made of only two cell types: pigmented cells forming the pigment screen and rhabdomeric photoreceptors (Suschenko and Purschke, 2009). In some species the pigment cells send in processes between the photoreceptors, which expand distally and form a lens-like structure. Whether this structure does have any optical functions is questionable since it is sometimes irregularly shaped and often lies directly next to the photoreceptor outer segments without a vitreous space in between (Verger-Bocquet, 1992; Purschke et al., 2006).

Eyes have been examined in a large number of errantian species (Purschke, 2010) but little is known about the functional significance of the eyes. Perhaps the most extreme case are the pelagic alciopids, which have a single pair of huge camera type eyes, tripling the width of the prostomium (Hermans and Eakin, 1974). They are assumed to be visual predators, but it has so far not been possible to test this experimentally. The limited available electrophysiological data indicates that the complex morphology is backed by complex physiology, possibly supporting colour vision (Wald and Rayport, 1977). In the bioluminescent syllid, O. enopla, there is a striking match between the spectral sensitivity of one of their photoreceptor populations and the peak wavelength of the emitted light, suggesting visual communication during mating (Nicol, 1978). Based on the general eye structure in Errantia other possible light-guided behaviours could be phototaxis, diurnal activity patterns, depth gauge and predator detection (Nilsson, 2009; Purschke, 2010), but experimental evidence is most often lacking.

A major group within Errantia are the scale worms, Aphroditiformia. A few deep sea and cave-living species are continuous swimmers in the water column, but most scale worms are benthic crawlers found in most marine habitats (Rouse and Pleijel, 2001; Gonzalez et al., 2018). They are characterized by two rows of dorsal scales, elytra, and many of them have two pairs of prostomial eyes (Suschenko and Purschke, 2009). They occupy several different...
niches and trophic levels, but a large number of them are predators feeding on a broad variety of benthic invertebrates (Fauclair and Jumars, 1979; Jumars et al., 2015). In the shallow waters of the North-East Atlantic, two species of polynoid scale worms are commonly found, Lepidonotus squamatus and Harmothoe imbricata. They typically inhabit rocky shores and stony reefs (with H. imbricata preferring blue mussel beds) where they feed on small benthic invertebrates (Fauclair and Jumars, 1979). Their two pairs of eyes appear similar, and from examinations of other species of Harmothoe, it was found that they consist of pigmented photoreceptors and pigmented supporting cells also forming a lens-like structure (Suschenko and Purschke, 2009). An interesting difference between the two species is that in H. imbricata the scales emit light when disturbed and become strongly bioluminescent when autotomized, which is not the case for L. squamatus (Nicol, 1953). This is a defence mechanism allowing H. imbricata to escape, while an attacking predator chases the autotomized scales (Livermore et al., 2018). The two species each belong to their clade or subfamily within Polynoidae: L. squamatus belongs to Lepidonotinae, whereas H. imbricata is a member of the Polynoinae (Zhang et al., 2018). Within the larger Polynoidea clade, several examples of bioluminescent species have been reported (Moraes et al., 2021). Although these subfamilies both occupy derived positions within Aphroditiformia (scale worms), scale worms have been proposed to date back to Devonian times and the split of the Polynoidea subfamilies may also date back >100 million years (Rouse and Pleijel, 2001).

Here, we use H. imbricata and L. squamatus to obtain experimental data on the visual capacity of the typical eyes of Errantia. Through behavioural, morphological, and physiological examinations, we seek to test the following two hypotheses. (1) Each pair of eyes serve different purposes as seen in some other multi-eyed visual systems (Garm and Mori, 2009; Menda et al., 2014), which is reflected in differences in physiology and/or morphology. (2) The eyes of the bioluminescent H. imbricata will be optimized to detect the light emitted by their autotomized scales. Our results provide some support for the second hypothesis but support the first hypothesis to a lesser degree.

MATERIALS AND METHODS

Animals

Fifteen specimens of Lepidonotus squamatus (Linnaeus 1758), 2–5 cm long, were collected at ~35 m in the northern part of Øresund, Denmark, using a standard triangular dredge. The specimens were brought back directly to the Marine Biological Section in Copenhagen. About 50 specimens of Harmothoe imbricata (Linnaeus 1767), 1–3 cm long, were hand collected from mussel beds attached to ropes at a pier in the Kaldbak Fjord, Faroe Islands. They were transported back alive to Copenhagen in pairs in 50 ml Falcon tubes with seawater and a small piece of rope. In Copenhagen, the two species were kept in a 35 or 150 litre tank, respectively, with sea water of 10°C and 32–33 psu. About half of the water was exchanged every 2–3 weeks. Large stones were scattered on the bottom of the tanks and served as hiding places. Worms were fed a variety of local amphipods and isopods along with small benthic anelids from aquarium cultures (Ophryotrocha spp.), but only the anelids were found to be consumed.

Eye morphology and visual fields

Three specimens of each species were anesthetized in sea water mixed 1:1 with 7% MgCl₂ for 10 min. Afterwards they were decapitated and had the prostomia with their two sets of eyes photographed under a standard dissection microscope equipped with a c-mount camera (Evolution MP 5.0). In order to visualize the gaze direction of the pupils, pictures were taken dorsally, laterally and in the case of H. imbricata, also anteriorly.

The excised prostomia were fixed in seawater mixed 1:1 with 3% glutaraldehyde and 3.7% paraformaldehyde in 0.1 mol l⁻¹ sodium cacodylate buffer for 3 days at 4°C and post-fixed in 2% osmium tetroxide in 0.1 mol l⁻¹ cacodylate buffer for 2 h at room temperature. Afterwards they were dehydrated in a standard series of ethanol, transferred to acetone, and embedded in Epon 812 resin. The diurnal activity patterns of the scale worms were examined in their holding tanks, which had a light scheme of 16 h light:8 h dark, mimicking local summertime. They were filmed using a standard Handycam (Panasonic HC-VX980) in nightshot mode for 15 h, starting at 16:00 h with 3 h room light followed by 8 h in darkness and ending with another 4 h of room light. During the 8 h of darkness, the tank was illuminated by an array of infrared (IR, 940 nm) LEDs. This was repeated three times for both species. To check for an endogenous diurnal rhythm, two trials were done where the light scheme in the holding tanks was changed such that the 8 h of darkness occurred between 11:00 h and 19:00 h.

The response to flashes of light mimicking the bioluminescent flashes of H. imbricata were examined with the animals placed one at a time in a small container (1.5 l) filled with water from the holding tank and a single stone to hide under. They were transferred to the experimental tank under room light conditions but once in the tank light was turned off and they were left to dark adapt for 15–20 min. Afterwards the setup was illuminated by IR light only and filmed using a Handycam in night shot mode. At the end of the experiment the stone was a 1 mm light guide was placed 0.5 cm above the bottom pointing at the stone. The light guide was illuminated by a computer-controlled optical bench, which supplied flashes of dim green light (520 nm, half width=12 nm) of 0.8 W sr⁻¹ m⁻². Two different 30 min long experiments were conducted: (1) flashes were given at random intervals (1–8 min between flashes) and with varying duration between 5 and 60 s (average of 30 s); (2) flashes of 30 s were given every time the animals were within 10 cm of the
light guide and faced it. This resulted in 4–10 and 2–8 stimulations in each trial, respectively. The intensity, peak wavelength and duration of the flashes were chosen to mimic the bioluminescence of a single scale of *H. imbricata* ( Plyuscheva and Martin, 2009). The light-emitting surface of the light guide was also matched to the scale size of a midsized *H. imbricata*. In total 7 *H. imbricata* and 5 *L. squamatus* were used in the behavioural tests.

Because *L. squamatus* would occasionally walk around with the anteriormost two scales elevated (or laterally retracted) while the rest were laying down flat across the dorsal body, an additional experiment was conducted for this species. When the two anteriormost scales were lifted, the animal would be approached slowly with a histology needle to test if they would lower the scale to protect the prostomium sacrificing vision. This was repeated twice with two specimens.

**Electrophysiology**

The prostomium was dissected from each animal and the tentacles and palps removed. Then, it was transferred to a Petri dish in the electrophysiological setup containing about 5 ml of seawater from the holding tank. A custom-made glass suction electrode (outer diameter ~50 µm, pore size 5–10 µm) was placed at the edge of either one of the posterior or anterior eyes and a 1 mm light guide was placed immediately in front of the pupil. The light guide was several times larger than the diameter of the pupil ensuring a close to even illumination of the retina. The light stimuli were produced by an optical bench with an ultra-bright white LED (Luminus CBT 90), a series of neutral density filters and a spectral filter wheel. The LED was controlled by a custom-made program for LabView (National Instruments, TX, USA).

Initially, 25 ms white flashes were presented to the eye and the electrode was moved around until an impulse response was obtained and then the preparation was dark adapted for 30 min. The duration of dark adaptation was determined by applying the same stimuli every 5 min and waiting for two consecutive stimulations to give the same response magnitude. In the two preparations tested in this way, it occurred after 20 min and 25 min, respectively, thus ensuring a safety margin. In a similar way, the longevity of the preparation was tested, and it was found that no significant change in responsiveness occurred within the first 2.5 h after dark adaptation.

Each protocol started with an intensity series covering 5 log units of white light, 2.5×10⁻³ W sr⁻¹ m⁻² to 2.5×10² W sr⁻¹ m⁻² in steps of 0.3 or 0.7 log units using neutral density filter (Linos, Goettingen, Germany), starting at the low intensity end. This was followed by an equal quanta stimulation, 1.7×10⁶ photons s⁻¹ sr⁻¹ m⁻², with spectral filters (half width=12 nm, CVI Laser, Bendheim, Germany) covering 420–680 nm in steps of 10 or 20 nm. The spectral data were transformed by the V-log J curves (Coates et al., 2006) and compared with the absorption spectrum of a theoretical opsin (Govardovskii et al., 2000) using the least-squares method. Following the spectral series, a second intensity series was conducted in steps of 1 log unit to ensure that the sensitivity of the preparation had not changed during the protocol. All stimuli were 25 ms flashes with an interstimulus interval of 1.5 min and the entire protocol lasted approximately 65 min excluding dark adaptation. For each of the pro stomia used, tests were performed on either one eye only, on both an anterior and a posterior eye or on one eye only but followed by a second series also testing the temporal resolution of that eye.

The temporal resolution was tested by flicker fusion frequency (FFF) experiments. Because of differences in the dynamic range between the two species, with *H. imbricata* being more sensitive, the protocols differed. For *H. imbricata*, the eyes were initially light adapted for 5 min to 0.51 W sr⁻¹ m⁻². This was followed by a series of sinusoidal stimuli with an amplitude of ±0.5 W sr⁻¹ m⁻² starting at the intensity of the adaptation light. The frequency span tested was 0.5–8 Hz in steps of 0.5 Hz. For *L. squamatus*, the eyes were light adapted for 5 min to 5.1 W sr⁻¹ m⁻², which was followed by a series of sinusoidal stimuli with an amplitude of ±5 W sr⁻¹ m⁻². The frequency span tested for this species was 0.2–3.8 Hz in steps of 0.4 Hz. The stimulations for *H. imbricata* lasted 10 s with interstimulus intervals of 30 s and stimulations for *L. squamatus* lasted 60 s and had interstimulus intervals of 30 s. FFF curves were constructed through Fourier transformations of the recordings using the normalized power of the principal frequency (stimulus

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**Fig. 1. Arrangement of the eyes of Harmothoe imbricata and Lepidonotus squamatus.** Both species of scale worms have the classical set of two eyes on their prostomium as seen on many species of errant marine annelids. (A) In *H. imbricata* the posterior pair of eyes (PE) are situated on the back of the prostomium on the dorsolateral side. The anterior eyes (AE) are situated on the ventral side just next to the lateral tentacles on the frontal part of the prostomium. (B) Anterolateral view of the prostomium of *H. imbricata* showing the left anterior eye. Note the pupil pointing anterolaterally. (C) In *L. squamatus* both pairs of eyes are clearly visible on the dorsal side of the prostomium. Compared with *H. imbricata*, the anterior eyes are moved backwards and are situated midway on the prostomium. (D) In lateral view, the pupils in *L. squamatus* are pointing dorsolaterally for the posterior eyes and anterolaterally for the anterior eyes.
frequency) of each recording. In all cases a time period matching 10 stimulation cycles was analyzed.

For *H. imbricata*, the intensity and spectral series were obtained for 7 anterior and 7 posterior eyes and FFF series was obtained from 5 anterior and 5 posterior eyes. For *L. squamatus* the intensity and spectral series were obtained for 8 anterior and 8 posterior eyes and FFF series was obtained from 6 anterior and 6 posterior eyes.

The signals from the electrode were amplified 1000 times using a differential AC Amplifier (1700, A-MSystems, USA) and filtered through a 50 Hz notch filter, and 0.1 and 1000 Hz high and low pass filters, respectively. The recordings were stored and processed on a laptop using a custom-made program for LabView (National Instruments, TX).

**RESULTS**

**Eye arrangement and gaze direction**

The posterior eyes are similar in the two species and situated on the dorsolateral part of the prostomium close to the peristomium. They are oval and 100–150 µm along the largest diameter in adult specimens (Fig. 1). In both worm species, the posterior eyes gaze dorsolaterally and their pupils are round and ~50 µm in diameter (Figs 1 and 2). The anterior eyes are also oval but slightly larger, spanning 150–200 µm along the largest diameter in both species. In *H. imbricata*, they are situated on the anteriormost part of the prostomium under the cephalic peaks, whereas in *L. squamatus* they are placed midways on the prostomium on the dorsolateral side (Fig. 1). Similarly to the posterior eyes, the pupils are ~50 µm in diameter and in *H. imbricata* the gaze is directed anteriorly (Fig. 2B, D) whereas in *L. squamatus* the gaze is anterolateral (Fig. 2C, E).

**Eye morphology and visual fields**

The overall morphology is similar for the two species and for both pairs of eyes. The outer layer is formed by the nuclei for the two involved cell types: pigment cells forming most of the screening pigment and the pigmented photoreceptors. The pigment is packed in vacuoles, in a 10–15 µm thick layer, and from sub-illuminated...
Diurnal activity patterns
Both species have a distinct diurnal activity pattern and are both strictly night active. During light hours, the worms sat motionless in a dark hiding place, typically under one of the rocks in the tank. When the light was turned off, they started exploring the tank, typically after 5–10 min (Fig. 3). There was no sign of an internal diurnal rhythm since their activity patterns also followed the light in the experiments when the light scheme was shifted. Still, if the worms were stressed in room light with no place to hide, such as a small Petri dish, they would start crawling around. Here, L. squamatus displayed a previously unreported novel behaviour. They would crawl around with only the distalmost pair of elytra lifted, exposing the prostomium and the two pairs of eyes (Fig. 4A). A mechanical threat (needle) made them lower the scales again along with withdrawal of their tentacles (Fig. 4B, Movie 1). After making this discovery, we looked through the recordings of their natural nocturnal activity under IR light, but interestingly, they were never observed lifting their scales in these conditions. This behaviour was not observed for H. imbricata in either situation.

Response to the bioluminescence mimic
After ~15 min of dark adaptation in the experimental tank with only IR light, most specimens of both species would start crawling around, but some moved very little. Overall, the active specimens behaved similarly. In almost all trials they had a strong tendency to walk along the edge of the tank and only rarely crossed the central part (Fig. 5). It did not matter whether the bioluminescence mimic was turned on randomly or only when the animal was facing the light: there was no detectable response and worms did not pause or speed up, nor did they change their walking direction during periods of light on.

Eye physiology
Both species respond normally to short flashes of light with a two-phased potential (Fig. 6). The amplitude of the response is graded following the intensity of the flashes. From the V-log I curves, it is seen that the anterior and posterior eyes of H. imbricata respond very similarly and have a dynamic range of ~2.5 log units from 0.5 W Sr\(^{-1}\) m\(^{-2}\) to 100 W Sr\(^{-1}\) m\(^{-2}\) (Fig. 6C). At maximum intensity, the curves flatten indicating the beginning of photoinhibition. The two pairs of eyes of L. squamatus also respond in the same way but the dynamic range is broader than for H. imbricata, covering at least 4 log units and with no signs of photoinhibition at maximum intensity (Fig. 6F).

All four spectral sensitivity curves have a good match with the theoretical absorption spectrum of opsins (Fig. 7). For H. imbricata, the two pairs of eyes again show similar peak sensitivity of the
matched opsins, being 506±3 and 499±10 nm for the anterior and posterior eyes, respectively. In L. squamatus the peak sensitivities are shifted towards the short wavelength part of the spectrum and the matched opsins show maximum sensitivity at 494±6 and 487±9 nm for the anterior and posterior eyes, respectively. The peak sensitivity in the anterior eyes of H. imbricata was significantly different from both the anterior and posterior eyes of L. squamatus (one way ANOVA, \( F_{2.24} = 7.62, P<0.001 \), followed by Tukey HSD post hoc test, \( P=0.036 \) and 0.0006, respectively). Furthermore, the peak sensitivities of the posterior eyes were also significantly different in the two species (one way ANOVA, \( F_{2.24} = 7.62, P<0.001 \), followed by Tukey HSD post hoc test, \( P=0.036 \)).

The temporal resolution of the two species appears rather different but again, within each species the anterior and posterior eyes are similar (Fig. 8). Both eyes of H. imbricata have a FFF of 4–5 Hz and interestingly, the power of the Fourier transformation declined rapidly with increasing frequency of stimulation (Fig. 8B). In L. squamatus, the stimulations did not reach FFF and at the highest frequency, 3.8 Hz, a response was still visible (Fig. 8C). Even though unfortunate, we do not consider this a major problem, since the most important for the intra- and interspecific comparison is not the absolute FFF but rather the shape of the initial part of the curve. Still, if the curves for L. squamatus were extended following the shape of the decline between 1.8 and 3.8 Hz the 0.05 cut off level was reached between 5 and 7 Hz for both the anterior and posterior eyes (not shown). The frequency curves are again similar for the two pairs of eyes even though the large standard errors indicate larger variation between preparations in this species. The power of the response in L. squamatus remained high until about 2 Hz, where it started to decline (Fig. 8D).

**DISCUSSION**

Here, we have examined and compared the eye morphology, eye physiology and light-guided behaviours between two scale worm species both having two pairs of prostomial eyes. H. imbricata is found to have more-sensitive photoreceptors with a green shift in the spectral sensitivity matching their bioluminescence, which supports our hypothesis that they may detect their own and conspecifics’ bioluminescence. Furthermore, their low temporal resolution strongly favours detection of slowly changing low intensity signals, which again corresponds with the fact that their light emission lasts several seconds. However, it was not possible to detect a behavioural response to a mimic of their bioluminescence. Despite both species being strictly night active, the results suggest that the less-sensitive eyes in L. squamatus could function during daytime, possibly detecting hiding places or threats such as predators. Except for a difference in gaze direction, the two eyes pairs within each species are morphologically and physiologically close to identical, thus opposing the hypothesis of a functional difference between the eye pairs.

**Seeing your own bioluminescence**

When H. imbricata autotomize their scales, they become bioluminescent, which works as a defence against, at least, crustacean predators (Livermore et al., 2018). While the initial escape is important, it is equally important that the worm keeps following an escape route away from the predator. The obvious way to do this is to see your own glowing scales and crawl/swim in the opposite direction. Whereas our *in vitro* behavioural experiments with a mimic of the glowing scale could not provoke such a behavioural response, the physiological data does support this theory. The intensity of their bioluminescent signal is low (Nicol, 1953; Widder, 2010) and declines with the square of the distance, so the eyes should have enhanced sensitivity in order to detect this signal. We did find that when compared with the closely related non-bioluminescent polynoid species, L. squamatus, the outer segments of the photoreceptors are longer in H. imbricata and so is the integration time as seen by the steep decline on the flicker fusion frequency graphs. This will result in enhanced sensitivity as also seen on the \( V \)-log \( I \) curves, which are shifted 1–1.5 log units to the low intensity side for H. imbricata. Furthermore, it is equally important that the spectral sensitivity of the eye matches the spectral peak of the emitted light (Haddock et al., 2010; Garm et al., 2016). The light emitted from the scales is estimated to peak between 510 and 520 nm (Nicol, 1953; Lecuyer and Arrio, 1975), which has a rather good match with the green shift in sensitivity of especially the anterior eyes in H. imbricata, which peaks at \( \sim 506 \) nm. A green shift in the spectral sensitivity is also often seen as an adaptation to the colour of the water in the habitat with open water species having opsins typically peaking in the deep blue part of the spectrum at \( \sim 470 \) nm and opsins of costal living animals peaking closer to 500 nm (Cherroske and Cronin, 2005). Still, despite the difference in depth distribution, H. imbricata and L. squamatus live in the same coastal greenish water and should not differ in spectral sensitivity if the ambient light was the only determining factor.
In addition to detecting your own light emission, it will also be beneficial to detect light emitted from conspecifics, which warns you about the whereabouts of predators. It would in fact also be advantageous for \textit{L. squamatus} to get this predator warning since the two species co-occur in some habitats, but judging from the physiology of their eyes and their behavioural response to the mimic, this is not supported.

Considering the physiological data, we see two possible explanations for why \textit{H. imbricata} did not respond to a mimic of their own bioluminescence and both might have been at play in our experiments. First, the animals were possibly too stressed to respond after being moved from the holding tank to the experimental set-up. This is indicated by the trajectories showing that they spent most of the time crawling along the sides, which is a typical escape/stress response. Second, the flashes were designed to mimic the bioluminescence of a single scale, which might not be enough to elicit a behavioural response. When under attack, \textit{H. imbricata} will typically shed 4–8 scales (Livermore et al., 2018; our own observations), and even though the shed scales do not stick together, they do stay relatively close to each other and the combined light emission could be needed to trigger an escape response. Additional behavioural experiments are needed to explore these possibilities further.

**Fig. 6.** Absolute sensitivity of the posterior and anterior eyes. (A,B) Examples of impulse responses from the anterior eyes and posterior eyes of \textit{H. imbricata} covering 5 log units of light intensity. The red curve indicates the 25 ms light stimulation. (C) \(V\text{-log} I\) curves from anterior (blue line) and posterior (red line) eyes of \textit{H. imbricata}. Both curves are close to sigmoidal and have a dynamic range of \(\sim 2.5\) log units. (D,E) Examples of impulse responses from the anterior and posterior eyes of \textit{L. squamatus} covering 5 log units of light intensity. The color-coded intensities follow A and the red curve indicates the 25 ms light stimulation. (F) \(V\text{-log} I\) curves from eyes of \textit{L. squamatus}. No saturation is seen in the high intensity end and both eyes have a dynamic range of at least four log units. Error bars indicate s.e.m.; \(n=7\) in C, and \(n=8\) in F.
Why have two pairs of prostomial eyes?

Eyes are some of the most expensive organs to build and maintain (Moran et al., 2015), but still several animal groups have a large number of eyes. This is seen in box jellyfish (Nilsson et al., 2005), spiders (Harland et al., 2012), chitons (Li et al., 2015), scallops (Speiser and Johnsen, 2008), fan worms (Bok et al., 2017), starfish (Garm and Nilsson, 2014) and many others. One reason for this common pattern in animals with sparse nervous systems is to allow for special purpose eyes, where the visual tasks are divided between eyes, minimizing the need for post-processing (Land and Nilsson, 2006). However, our results do not support such a functional differentiation among the anterior and posterior prostomial eyes, often found in scale worms and other members of Errantia. In both examined species the anterior and posterior eyes are close to identical, only with the anterior eyes being slightly larger. The most conspicuous difference lies in the gaze direction of the two pairs of eyes, suggesting that the benefit of having four prostomial eyes is an extended visual field, as also seen in other animals, such as starfish (Garm and Nilsson, 2014).

Fig. 7. Spectral sensitivity curves. (A,B) Spectral sensitivity curves from the two eyes of H. imbricata (red squares). The two curves are close to identical and the templates for a 506 and 499 nm opsin (black lines) (Govardovskii et al., 2000) are the best matches for the anterior and posterior eye respectively using the least squares method. (C,D) Spectral sensitivity curves from the two eyes of L. squamatus (blue squares). The two curves are also close to identical and a 494 and 487 nm opsin (black lines) are the best matches for the anterior and posterior eye, respectively. Error bars indicate the s.e.m.; n=7 in A,B and n=8 in C,D.

Visual ecology of H. imbricata and L. squamatus

As described above, one of the visual tasks of H. imbricata seems to be detection of their own and possibly conspecific bioluminescence to escape from predators. But there are likely several other behaviours associated with the eyes. Both species are strictly nocturnal and our results also show that this is not controlled by an endogenous rhythm since turning the light off during the day activated them and turning it on at night made them hide and become quiescent. Such a diurnal activity pattern can be achieved by any type of photoreceptor and does not require any spatial information (Nilsson, 2009). It could be controlled by the...
prostomial eyes even when covered by the scales as the relative intensity change throughout the day is not affected by the shading. Their ability to seek out a dark hiding place, such as a rock or mussel shell, at dawn could be guided by low spatial resolution vision, which is supported by the morphology. However, we do not currently have data that allow us to elaborate further on this.

Other behaviours where vision is putatively involved in other species of errantian annelids are prey detection and mating (Hermans and Eakin, 1974; Nicol, 1978). We only observed our animals to be active and feeding in darkness under IR light and under these conditions with higher light intensities. One possibility is that they visually detect predators sneaking up on them in their hiding places during the day, which is supported by the differential scale lifting only occurring in bright light.

It is noteworthy, that the photoreceptors in *L. squamatus* are less sensitive and have somewhat higher temporal resolution than those of *H. imbricata*, which indicates that they are used under conditions with higher light intensities. One possibility is that they visually detect predators sneaking up on them in their hiding places during the day, which is supported by the differential scale lifting only occurring in bright light.

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