1 Spectral sensitivity in Onychophora (velvet worms)

² revealed by electroretinograms, phototactic

behaviour and opsin gene expression

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23 ABSTRACT

24 Onychophorans typically possess a pair of simple eyes, inherited from the last common ancestor of Panarthropoda (Onychophora + Tardigrada + Arthropoda). These visual 25 26 organs are thought to be homologous to the arthropod median ocelli, whereas the compound eyes most likely evolved in the arthropod lineage. To gain insights into the 27 28 ancestral function and evolution of the visual system in panarthropods, we investigated phototactic behaviour, opsin gene expression and the spectral sensitivity of the eyes in 29 two representative species of Onychophora: Euperipatoides rowelli (Peripatopsidae), 30 and Principapillatus hitoyensis (Peripatidae). Our behavioural analyses, in conjunction 31 32 with previous data, demonstrate that both species exhibit photonegative responses to wavelengths ranging from ultraviolet to green light (370-530 nm), while 33 electroretinograms reveal that the onychophoran eye is maximally sensitive to blue light 34 (peak sensitivity ~480 nm). Template fits to the obtained sensitivities suggest that the 35 onychophoran eye is monochromatic. To clarify on which type of opsin the single 36 visual pigment is based, we localised the corresponding mRNA in the onychophoran 37 eye and brain using in situ hybridization. Our data show that the r-opsin gene 38 (onychopsin) is expressed exclusively in the photoreceptor cells of the eye, whereas the 39 *c-opsin* mRNA is confined to optic ganglion cells and the brain. Together, our findings 40 suggest that the onychopsin is involved in vision, whereas the c-opsin might have a 41 42 photoreceptive, non-visual function in onychophorans. 43

44 KEY WORDS: Arthropod, Eye, Light response, Vision, Opsins, Phototaxis, Evolution

45 INTRODUCTION

46 Onychophorans (velvet worms) typically bear a pair of simple, ocellus-like eyes (Fig. 1A,B), which are thought to be homologous with the median ocelli of arthropods 47 (Mayer, 2006), one of their closest relatives (Giribet and Edgecombe, 2012). 48 Accordingly, the last common ancestor of Panarthropoda (Onychophora + Tardigrada + 49 Arthropoda) most likely possessed a pair of ocellus-like visual organs, whereas the 50 compound eyes evolved within the arthropod lineage (Mayer, 2006; Hering et al., 2012; 51 Hering and Mayer, 2014). While all arthropod species studied thus far have multiple 52 rhabdomeric opsins (r-opsins) as visual pigments, their presence being a prerequisite for 53 54 colour vision (reviewed by Briscoe and Chittka, 2001), transcriptomic analyses of the opsin repertoire revealed only one r-opsin gene (onychopsin) in five distantly related 55 onychophoran species (Hering et al., 2012). In phylogenetic analyses, onychopsin forms 56 the sister group to the visual r-opsins of arthropods, suggesting that this gene functions 57 in onychophoran vision. However, a ciliary-type opsin (c-opsin, to which type also the 58 visual pigments of vertebrates belong; reviewed by Porter et al., 2012), has also been 59 reported to occur in the onychophoran eye (Eriksson et al., 2013). Hence, a detailed 60 expression study at the cellular level seems necessary to clarify whether r- or c-type 61 opsins, or both, are involved in onychophoran vision. 62

Behavioural studies revealed negative phototactic behaviour in two species of 63 64 Peripatidae: Epiperipatus biollevi (see Monge-Nájera et al., 1993), and Principapillatus hitoyensis (referred to as "Epiperipatus cf. isthmicola" in Hering et al., 2012). 65 Specimens of *P. hitoyensis* showed a photonegative reaction to wavelengths ranging 66 67 from 363 nm (ultraviolet, UV) to 586 nm (yellow). The sensitivity maximum (α -peak) of the visual pigment in this species was therefore estimated to be in the blue-green 68 69 range of the spectrum (Hering et al., 2012). However, neither the specific wavelength of 70 the α -peak nor the actual spectral sensitivity curve of the onychophoran photoreceptors 71 is known. Moreover, it is unclear whether a photonegative reaction to the same 72 wavelengths occurs in representatives of the second major onychophoran subgroup, the 73 Peripatopsidae, for which quantitative data are still missing.

We therefore analysed the behavioural response to light and localised the
expression of the r- and c-type opsins in the peripatopsid *Euperipatoides rowelli*. To
identify the sensitivity maximum (α-peak) of the visual pigment and to complement

recordings from the eye in both *E. rowelli* and *P. hitoyensis*. The obtained data allow

79 conclusions regarding the physiological properties and function of the visual system in

the last common ancestor of Onychophora and Panarthropoda, respectively.

81

82 **RESULTS**

83 Spectral sensitivity of the onychophoran eye

We recorded electroretinograms (ERGs), extracellular, light-induced potential changes 84 in the retina, from the eyes of six specimens of E. rowelli of both sexes and one male of 85 P. hitoyensis (Fig. 2A,B). Responses to light flashes of 40 or 100 ms duration consisted 86 of an initial hyperpolarization truncated by a depolarization (Fig. 3A,C). The half width 87 of the response (width at half maximal hyperpolarization) exceeded 140 ms and the time 88 to peak (time interval from stimulus onset to maximal hyperpolarization) exceeded 95 89 ms for all tested intensities. Both values increased considerably with decreasing 90 stimulus intensity (Fig. 3A,C). Spectral flashes with equal photon flux revealed that the 91 dark-adapted retina was most sensitive to wavelengths around 480 nm, i.e. to the region 92 of the spectrum perceived as blue by humans. We fitted template formulae 93 94 (Govardovskii et al., 2000), which approximate the absorbance spectra of opsin-based visual pigments in invertebrates (Stavenga, 2010), to the data from each individual (Fig. 95 3B,D) and averaged the results for different individuals of the same species. Optimal 96 97 fits were obtained assuming a visual pigment with an absorbance peak at 474 ± 6.5 nm (wavelength λ_{max} : average \pm standard deviation; coefficient of determination: $R^2 = 0.88$ 98 \pm 0.03) for the six specimens of *E. rowelli* and at 484 nm (R² = 0.93) for one specimen 99 of P. hitoyensis. When the eye was adapted to green light, responses were 100 101 indistinguishable from noise level, and no other sensitivity peaks were apparent at 102 shorter wavelengths. During recovery from green adaptation, the shape of the spectral 103 curve continued to be similar to the curve of the dark-adapted retina (supplementary 104 material Fig. S1).

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106 Localisation of the r- and c-type opsins in the onychophoran eye and brain

107 To determine whether the signal obtained from the electrophysiological recordings is

108 related to the onychophoran r- or c-type opsin, we performed fluorescence in situ

109 hybridization experiments on cryosections of the E. rowelli heads (Fig. 4A-F). Our data 110 show that Er-onychopsin is expressed in the photoreceptor layer of each eye, and lacking in other tissues, including the brain (N=6; Fig. 4B–D; supplementary material 111 112 Fig. S2). In contrast, *Er-c-opsin* mRNA is localised both in optic ganglion cells within the eye and in numerous neuronal somata within the brain, in particular in the ventral 113 114 perikaryal layer of the protocerebrum (Fig. 4E,F; supplementary material Fig. S2). A few additional *Er-c-opsin* expressing cells are seen in the deutocerebrum and in the 115 116 medullary cords (=connecting cords) linking the brain to the ventral nerve cords (Fig. 4E,F). Most importantly, and in contrast to optic ganglion cells, *Er-c-opsin* mRNA is 117 118 not expressed in the photoreceptor layer of the eye (Fig. 4E).

Our control experiments using the sense probe for *Er-onychopsin* revealed no 119 signal within the eye, indicating that the obtained labelling using the antisense probe for 120 Er-onychopsin is specific (supplementary material Fig. S2). The unspecific labelling in 121 the cuticle lining the epidermis and pharyngeal lumen, which occurs in all our 122 preparations, is due to autofluorescence. In contrast, applying the *Er-c-opsin* sense 123 probe revealed a similar pattern to the *Er-c-opsin* antisense probe (supplementary 124 material Fig. S2). The same result was obtained repeatedly in all experiments using 125 different sense probes on cryosections of different individuals (N=4), whereas an 126 increase of the hybridization temperature to 58 °C completely abolished the labelling in 127 128 all reactions, irrespective of whether the sense or antisense probes were used.

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130 Behavioural response of onychophorans to light of different wavelengths

To clarify whether *E. rowelli* shows negative phototaxis, single animals were released in a dark arena and left free to move in any direction. On directing a bright white light at their heads, the animals immediately changed walking direction away from the light source (Fig. 5A; Wilcoxon-signed-rank test: p<0.001), but did not veer from course in control experiments without the light stimulus (Fig. 5B).

To determine the sensitivity threshold of *E. rowelli*'s negative phototaxis, we used a blue light-emitting diode (λ max = 465 nm, i.e., close to the sensitivity maximum obtained from the electroretinograms, cf. Fig. 3B). Up to 6 animals were grouped (N=7 groups) and released simultaneously in one half of a dark arena, after which this half was illuminated with blue light of four different intensities (3×10¹¹, 6×10¹¹, 9×10¹¹, 141 12×10^{11} photons cm⁻² s⁻¹, measured at the bottom of the arena) and the behaviour of the 142 animals was then recorded for 5 minutes. In our setup, no significant reaction to blue 143 light was evident after an illumination with 3×10^{11} photons cm⁻² s⁻¹ and avoidance 144 behaviour first occurred at 6×10^{11} photons cm⁻² s⁻¹ (Fig. 5C). Hence, to evaluate 145 reactions to wavelengths outside the sensitivity maximum, the stimulus was delivered at 146 twice the intensity of this value (i.e., 12×10^{11} photons cm⁻² s⁻¹) in subsequent 147 experiments.

To determine whether or not a shorter exposure to light would affect the significance of our results, we compared the animal's reaction to a 5, 3, 2, and 1 minute long illumination, respectively (supplementary material Fig. S3). The data show that illumination for 1 minute is sufficient to elicit a highly significant avoidance reaction. However, since about one third of all our animals did not respond to the 1-minute stimulus, we selected an illumination duration of 2 min for the remaining experiments (Fig. 5C).

For our major experiments, we again released groups of up to 6 animals (N=15 155 groups) in one half of the dark arena, which was then illuminated with quasi-156 monochromatic light of six different wavelengths of the same intensity (12×10^{11}) 157 photons $\text{cm}^{-2} \text{ s}^{-1}$). In these experiments, E. rowelli specimens significantly evaded 158 wavelengths ranging from UV to green light (p<0.001; Friedman-test with Dunn's post-159 test) but showed no evasive reaction to light of longer wavelengths ($p_{591}=0.343$, 160 $p_{631}=0.650$; Friedman-test with Dunn's post-test) (Fig. 6A). In a final set of 161 experiments, we tested our specimens for potentially positive phototaxis, as it is 162 163 unknown whether or not they show preference for a particular wavelength. For these experiments, we used the same setting and released the animals in one half of the dark 164 165 arena, but illuminated the other half with light of each of the six wavelengths, 166 respectively. The obtained data gave no indication that the animals exhibit positive 167 phototactic behaviour (Fig 6B; Friedman-test with Dunn's post-test: p₄₆₅ and p₅₉₁=0.769, p_{rest}>0.999). 168

169

170 **DISCUSSION**

171 Onychophorans avoid wavelengths ranging from UV to green light

172 Our behavioural data provide evidence for negative phototaxis in the peripatopsid E. 173 rowelli, corresponding to previous results from the peripatid P. hitoyensis (see Hering et al., 2012). In our tests, specimens of E. rowelli were not attracted by light but instead 174 significantly avoided illumination with wavelengths ranging from UV to green light. 175 176 The animals are unlikely to have reacted to heat rather than light, as there was no 177 detectable increase of temperature in the arena even after illumination at maximum light intensity. The observed photonegative reaction is consistent with the nocturnal lifestyle 178 179 and high susceptibility of onychophorans to desiccation (e.g. Manton and Ramsay, 1937; Bursell and Ewer, 1950; Eakin and Westfall, 1965; Read and Hughes, 1987). This 180 might be one of the reasons why these animals generally avoid light, as it is a potential 181 indicator of heat and low humidity. 182

The slow response of the onychophoran photoreceptors, as evidenced by a long 183 time to peak and broad half width of the ERG signal in both species studied, is common 184 in nocturnal invertebrates, which typically show slower reactions to light as compared 185 to diurnal animals. This is regarded as an adaptation to dim light conditions, as the long 186 latency and response duration generally enhance the sensitivity of visual organs 187 (reviewed by Warrant, 2008; Fain et al., 2010; Warrant and Dacke, 2011). According to 188 our behavioural sensitivity tests, the threshold for negative phototaxis in E. rowelli lies 189 at 6×10^{11} photons cm⁻² s⁻¹ for blue light. This value corresponds well to light intensities 190 typically found on the ground of rainforests during the day (e.g. Vazquez-Yanes et al., 191 192 1990), which, again, is in line with the nocturnal lifestyle of velvet worms that usually forage for food at night and seek shelter during the day (Read and Hughes, 1987; 193 194 Mesibov, 1998).

Previous behavioural data from the peripatid P. hitoyensis indicated that the 195 196 maximum sensitivity in this species lies within the blue-green range of the light 197 spectrum (Hering et al., 2012). Our results based on ERGs from the eyes of P. 198 hitoyensis and E. rowelli now provide more precise values, and the estimated maxima at $\lambda_{\text{max}} = 474 \pm 6.5$ nm in *E. rowelli* and $\lambda_{\text{max}} = 484$ nm in *P. hitoyensis* suggest that the 199 200 eyes of both species are most sensitive to blue light, i.e. close to the lower limit of the previously estimated range (cf. fig. 4 in Hering et al., 2012). A maximum sensitivity to 201 blue light is widespread among invertebrates with monochromatic vision, which might 202

be due to the maximum distribution of energy of solar radiation at about 480 nm

204 (Menzel, 1979; Bowmaker and Hunt, 1999; Kelber and Roth, 2006).

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Onychophorans exhibit monochromatic vision with onychopsin as the only visual pigment

208 The electrophysiologically determined spectral sensitivity of the dark-adapted eye could be well approximated assuming the presence of only one opsin-based visual pigment. 209 210 Selectively adapting the retina to green light did not provide evidence for additional visual pigments maximally sensitive to shorter wavelengths. This finding is in line with 211 212 the previous hypothesis of monochromatic vision in Onychophora, which is most likely based on an r-type opsin, the so-called onychopsin (Hering et al., 2012). However, since 213 Eriksson et al. (2013) also detected a c-type opsin in the eye of *E. kanangrensis*, it was 214 still debatable whether a c- or rather an r-type opsin is involved in onychophoran vision. 215 216 Our in situ hybridization experiments in specimens of E. rowelli revealed onychopsin mRNA exclusively in the photoreceptor cell layer of the eye, but no expression in the 217 brain (contrary to the claim of Eriksson et al., 2013) or other tissues. Conversely, we did 218 not detect *Er-c-opsin* mRNA in the photoreceptor cells but rather in optic ganglion cells 219 220 in the proximal portion of the eye as well as within the brain. In the brain, the majority 221 of Er-c-opsin-positive somata occur in the ventromedian portion of the protocerebrum, 222 but a few cell bodies are also found within the deutocerebrum. These findings, in 223 particular the lack of *Er-c-opsin* expression in the photoreceptor cells, speak against an involvement of the c-opsin in onychophoran vision. We therefore conclude that 224 225 onychopsin is most likely the only visual pigment in the onychophoran eye.

Although the function of the onychophoran c-opsin protein is unknown, it might 226 227 be involved in non-visual, extra-ocular photoreception associated with circadian clock 228 mechanisms (Fukada and Okano, 2002; Vigh et al., 2002; Arendt et al., 2004; Velarde 229 et al., 2005; Shichida and Matsuyama, 2009). Notably, our expression data further indicate that the *Er-c-opsin* gene might be transcribed from both the sense and the 230 231 antisense DNA strands, as our in situ hybridization experiments using the sense probe revealed a similar expression pattern of this gene to that using the antisense probe. The 232 putative role of the *Er-c-opsin* antisense transcript in *E. rowelli* is unknown, but it might 233 234 be involved in a self-regulation of the *Er-c-opsin* expression – a function that has been

suggested for other genes exhibiting antisense transcription (reviewed by Pelechano andSteinmetz, 2013).

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238 Conclusions

Our electrophysiological data revealed maximum sensitivity to wavelengths around 480 239 240 nm in representatives of both major onychophoran subgroups, which suggests that the spectral absorption characteristics of the onychophoran photoreceptors in the retina have 241 242 remained nearly unchanged for ~350 million years, i.e., since the divergence of Peripatidae and Peripatopsidae (Murienne et al., 2014). This might be due to the 243 244 generally conserved geographic distribution and lifestyle of onychophorans, which are 245 confined to humid habitats and exhibit cryptic, nocturnal behaviour (Manton and Ramsay, 1937; Bursell and Ewer, 1950; Read and Hughes, 1987; Mesibov, 1998; 246 Oliveira et al., 2012). In conclusion, our electrophysiological recordings, behavioural 247 experiments, and gene expression studies suggest that the onychophoran eye contains 248 only one visual pigment, the r-type opsin onychopsin, which is most sensitive to blue 249 light. The c-opsin is restricted to the brain, and may function in extra-ocular 250 photoreception. Our data thus support the hypothesis of monochromatic vision in the 251 last common ancestor of Panarthropoda (Hering et al., 2012; Hering and Mayer, 2014). 252

253

254 MATERIALS AND METHODS

255 Animals

Two species from the two major onychophoran subgroups were studied (Fig. 1A,B): 256 257 Principapillatus hitoyensis Oliveira et al., 2012 (Peripatidae), and Euperipatoides rowelli Reid, 1996 (Peripatopsidae). Specimens of P. hitoyensis were collected from 258 259 leaf litter in the Reserva Biológica Hitoy Cerere (Province of Limón, Region of Talamanca, Costa Rica; 09°40'N, 83°02'W, 300 m). Specimens of E. rowelli were 260 261 obtained from decaying logs in the Tallaganda State Forest (New South Wales, Australia; 35°26'S, 149°33'E, 954 m). Onychophorans were collected and exported 262 263 under the following permits: (1) the Forestry Commission of New South Wales, Australia (permit no. SL100159); and (2) the Gerencia Manejo y Uso Sostenible de RR 264 NN-Ministerio del Ambiente y Energia, Costa Rica (permit numbers 123-2005-SINAC 265 266 and 014950). The animals were kept in plastic boxes with perforated lids as described

267 previously (Baer and Mayer, 2012). Specimens of P. hitoyensis were maintained at 268 room temperature (20–24 °C), whereas those of E. rowelli were kept at 18 °C either in the dark (for behavioural experiments), in the dark and under day/night conditions (for 269 270 gene expression studies), or under a shifted 14/10 hours day/night cycle and tested in their active night period (in electrophysiological experiments). Behavioural experiments 271 272 were carried out at a normal day/night rhythm between 4 and 9 pm at 19 °C. All animal treatments complied with the Principles of Laboratory Animal Care and the German 273 274 Law on the Protection of Animals.

276 Electroretinograms

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To obtain ERGs, we fixed the anterior body portion of each analysed specimen to a 277 halved, tapered pipette tip (Fig. 2A) using dental silicone (polyvinyl siloxane, 278 PRESIDENT light body, Iso 4823; Coltène, Altstätten, Switzerland) or a combination 279 280 of dental silicone and a 1:1 mixture of beeswax and resin, taking care to leave one eye and its surroundings free. In most cases, it was necessary to immobilise the animal 281 temporarily prior to handling by a 2-3 seconds long pulse of carbon dioxide from a soda 282 maker (Genesis, SodaStream®, Tel Aviv, Israel). The reference electrode, a chlorinated 283 284 silver wire, was brought in contact with the trunk of the specimen by conducting electrode paste (Gel101, Biopac Systems Inc., Goleta, California, USA) and kept in 285 286 place with paper tissue wrapped around the trunk and the pipette tip. To prevent 287 dehydration, the tissue was soaked in saline based upon the composition of onychophoran haemolymph (Robson et al., 1966). Just posterior or dorsal to the 288 289 exposed eye, the skin was thinned conically with the sharp tip of a broken razor blade. This allowed us to penetrate the integument with an electrolytically sharpened tungsten 290 291 electrode for recording from the retina, while the skin closed tightly around the 292 electrode and resealed the puncture (Fig. 2B).

A white light stimulus was produced by a 200 W Xenon lamp (Cermax LX175F
ASB-XE-175EX, SP Spectral Products, Putnam, Connecticut, USA) and directed to the
eye via the central, 400 μm-wide fibre of a forked light guide (QR400-7-SR/BX, Ocean
Optics, Dunedin, Florida, USA), when two shutters (VS25S2ZM1R1 and LS6ZM2,
both Uniblitz, Vincent Associates®, Rochester, New York, USA) were opened. The
angular position and distance of the tip of the light guide to the eye were adjusted on a

goniometer such that the entire cornea was illuminated by the light beam of 16°
divergence and ERG responses were maximized. Narrow-band interference filters (10–
12 nm full width at half maximum; Melles Griot, Rochester, New York, USA) and
neutral density filters (fused silica; also Melles Griot) could be inserted into the light
path, providing spectral stimuli with equal photon flux from ultraviolet (330 nm) to red
(700 nm) in 10 or 20 nm steps. All stimuli were delivered as flashes of 40 or 100 ms
duration, separated by pauses of 3 or 5 s.

306 For spectral adaptation, light of a green light-emitting diode (LED) with a dominant wavelength of 521 nm and 34 nm full width at half maximum (LXHL-MM1D 307 308 Green Luxeon Star, Quadica Developments Inc., Brantford, Ontario, Canada) was presented constantly throughout the experiment via the six outer fibres (each 400 µm in 309 diameter) of the forked light guide. The combined light beam of 25° divergence 310 provided between 3×10^{13} and 2×10^{16} photons cm⁻² s⁻¹ at the position of the eye, 311 depending on the operating current of the LED and the distance between the cornea and 312 the tip of the light guide. 313

Responses were amplified by a P15 AC amplifier (Natus Neurology Incorporated - Grass Technologies, Warwick, Rhode Island, USA) and sampled at 2000 Hz, digitized and saved using an NI PCIe-6251 data acquisition board and custom-made LabView scripts (both National Instruments Corporation, Austin, Texas, USA) installed on a conventional computer.

319 All experiments were carried out in a darkened Faraday cage with either the stimulus or the stimulus and the spectral adaptation light as the only light source. 320 321 Adaptation periods prior to recording had to be kept to a minimum (2 to 5 minutes for spectral adaptation and 5 to 10 minutes for dark adaptation) due to the reduced viability 322 323 of the animals in the setup. The spectral sensitivity under dark adaptation was measured 324 up to 10 times, alternating between series starting with short and proceeding to long 325 wavelengths, and series in the reverse order. Before and after each spectral series, a response-intensity (V-log I) relationship was determined to control for changes in 326 327 recording quality and to establish the saturation level of responses. Initially, V-log I curves were measured with white light. When the peak sensitivity of an individual 328 became evident, we used the available narrow-band spectral light closest to the 329 presumed wavelength of maximal sensitivity for the V-log I. Up to three stable series in 330

331 both directions, i.e. six series altogether, were selected per individual and analysed using custom-made scripts in Matlab (R2012b, The MathWorks, Natick, Massachusetts, 332 USA) as described in detail elsewhere (Telles et al., 2014). In short, the ERG was 333 smoothed by a moving average with a window width of 0.01 s. Response amplitudes were calculated as potential changes from the baseline at stimulus onset to maximal hyperpolarization and converted into sensitivities based on the sigmoidal V-log I relationship obtained before and after each spectral series. We normalized all values to the maximal spectral sensitivity within a series and averaged series from the same individual. An established template for the absorbance of an opsin-based visual pigment (Govardovkii et al., 2000) was fitted to the entire mean spectral sensitivity curve using a non-linear least squares approach. We varied the amplitude and wavelength of the α peak and the amplitude of the β peak independently. Since the sensitivity curve for wavelengths below 390 nm was too flat to determine the wavelength of the β peak, we calculated it as a function of the α peak as suggested elsewhere (Govardovskii et al. 2000). Finally, the wavelengths of the estimated α peak (λ_{max}) and the respective coefficients of determination (R^2) were averaged for different individuals.

48 Fluorescence *in situ* hybridization

Partial sequences spanning most of the transmembrane region of the onychophoran rtype opsin (Er-onychopsin, 736 nt [CDS position 497–1232 of GenBank accession JN661372]) as well as the c-type opsin (Er-c-opsin, 808 nt [KM189804]) were amplified from cDNA, which was obtained by reverse transcription of total RNA 352 (TRIzol[®] extraction protocol) using Superscript II and DNA Pol I polymerase 353 (Invitrogen, Carlsbad, CA, USA) according to the manufacturer's protocols. The 354 fragments were cloned using the pGEM[®]-T Vector System (Promega, Madison, WI, 355 USA) and verified by Sanger sequencing (GATC Biotech, Konstanz, Germany). The 356 357 cDNA clones were amplified by a standard M13 PCR and used directly to transcribe 358 antisense and sense digoxigenin (DIG) labelled RNA probes by using SP6 and T7 RNA 359 Polymerase, respectively, and the DIG RNA labelling mix (Roche Diagnostics, Rotkreuz, Switzerland). Freshly dissected heads of male specimens of Euperipatoides 360 rowelli (N=10) were embedded and immediately frozen in Tissue-Tek[®] O.C.T.[™] 361 Compound (Sakura Finetek, Europe B.V.) and 10-16 µm thick horizontal sections were 362

cut on a Cryostat CM3050 S (Leica Biosystems, Nussloch, Germany). The sections 363 were mounted on SuperFrost[®] plus slides (Menzel GmbH, Braunschweig, Germany), 364 dried for 30 minutes at room temperature, and fixed in 4% paraformaldehyde for 365 another 30 minutes. After several washing steps with PBST (PBS with 0.1% Tween[®] 20) and acetylation, the sections were pre-hybridized in hybridization buffer for 3-4 hours either at 55 °C or 58 °C and then hybridized for 12–16 hours at the same temperature using $\sim 1 \text{ ng/}\mu l \text{ RNA}$ probe. Hybridization was followed by multiple washing steps with saline-sodium citrate buffer (SSC with 0.1% Tween[®] 20). The sections were incubated for 40 minutes in TNT buffer (0.1 M TRIS-HCl, pH 7.5; 0.15 M NaCl; 0.1% Tween[®] 20) containing 0.5% blocking reagent (PerkinElmer, Waltham, MA, USA). Anti-DIG-POD Fab-fragments (Roche Diagnostics), diluted either 1:50 or 1:100 in blocking solution, were then applied to the sections, which were incubated for additional 40-60 minutes at room temperature. After several washing steps with TNT buffer, the sections were incubated for 15 minutes with fluorescein-labelled tyramide (1:50 diluted working solution of the TSA[™] Plus Fluorescein Fluorescence System; PerkinElmer). After counterstaining with propidium iodide for 15 minutes, the slides were mounted in Vectashield® mounting medium (Vector Laboratories, Burlingame, CA, USA) and analysed with a confocal laser-scanning microscope (Leica TCS STED; Leica Microsystems). Confocal image stacks were processed with LAS AF Lite v2.4.1 (Leica Microsystems).

34 Behavioural experiments

The initial light avoidance experiments to test for negative phototaxis were performed with single specimens of *E. rowelli* (N=12) as described previously for spectral experiments on *P. hitoyensis* (see Hering et al., 2012), except that only bright white light was presented. Each animal freely moved in a circular arena and its path was recorded 5 cm before and after stimulus presentation using the freely available video analysis and modelling tool Tracker (Douglas Brown,

- 391 http://www.cabrillo.edu/~dbrown/; v.4.05 [09/04//2013]). The paths of the animals
- tested were then plotted into a single diagram (Fig. 5A,B). The statistical analysis was
- 393 conducted by comparing the turning angles of the animals under illumination to the

control runs without the stimulus by applying the Wilcoxon signed rank test (see Heringet al., 2012).

The remaining behavioural experiments were performed with a modified version 396 of the setup introduced by Monge-Nájera et al. (1993) using a rectangular arena 397 (163×93×70 mm) made of acrylic glass enclosed in black paper (Fig. 2C). A grey 398 399 opaque, removable plastic plate separated the arena in two halves. One half could be illuminated by a cold light source via a double-ended light guide (Fig. 2C,D). For each 400 401 experiment, the bottom of the arena was covered with a sheet of the same white paper 402 towels that were used for maintaining the animals (Baer and Mayer, 2012). The sheet 403 was folded three times, cut to the size of the arena and wetted with 5 ml distilled water, 404 thus ensuring equal humidity in all experiments. No space was left between the paper 405 towel and the border of the arena to prevent the animals from escaping under the paper 406 towel.

All experiments were carried out in total darkness at 19.3 ± 0.6 °C. The light 407 stimulus was delivered by narrow-banded LEDs (Nichia Corporation, Tokushima, Japan 408 and Avago Technologies, San José, CA, USA). The LEDs emitted no light in the 409 410 infrared range, and thus generated no detectable heat. To control for the possibility of 411 the animals' reaction to heat rather than light, the temperature at the bottom of the arena was measured after illumination for 5 minutes with each LED at maximum intensity 412 413 (i.e. for the 373, 402, 465, 528, 591 and 631 nm LEDs, the maximum intensity was 40, 414 220, 370, 130, 90 and 160 times higher than that used in our behavioural experiments, respectively). These measurements revealed no stimulus-correlated change in 415 416 temperature. The stimulus was generated by the PowerLab 26T data acquisition system (AD Instruments, Dunedin, New Zealand) and equalised by using different output 417 418 voltages and neutral density filters (Tinxi GmbH, Eggenstein-Leopoldshafen, 419 Germany). On- and offset of the stimulus was triggered with the Chart software (v5.5.6; 420 AD Instruments). An infrared-sensitive camera (Sony Handycam HDR-HC7, Tokyo, 421 Japan) was mounted above the arena to automatically record the experiments. To avoid 422 potential bias caused by possible physical or chemical influence, such as smell, noise, heat, or mechanical vibrations, all scoring and analyses were computerised and 423 performed with the experimenter distant from the recording arena, except when 424 425 releasing the animals. Before the next trial, the entire arena was cleaned and the paper

towels renewed to remove any traces left by the animals. In addition, the entire setup
was rotated by 90° every second day.

To determine the optimal duration and illumination intensity for the main 428 429 behavioural experiments, 7 groups of up to 6 specimens of *E. rowelli* (34 individuals in 430 total) were dark-adapted for 20 minutes and placed in the illuminable half of the arena. 431 After leaving them to settle for two minutes, the plastic plate separating the two halves was removed, the light switched on and each group tested for four different intensities 432 $(3 \times 10^{11}, 6 \times 10^{11}, 9 \times 10^{11}, 12 \times 10^{11} \text{ photons cm}^{-2} \text{ s}^{-1})$ of the 465 nm blue light. The 433 movements of the animals were recorded for 1, 2, 3 or 5 minutes (supplementary 434 435 material Fig. S3).

These experiments revealed the optimal duration (2 minutes) and illumination 436 intensity (twice the identified threshold, i.e. 12×10^{11} photons cm⁻² s⁻¹), which were then 437 used for the major behavioural experiments. For these experiments, narrow-band lights 438 of six different wavelengths were used: 373 ± 13 nm, 402 nm ± 10 nm, 465 nm ± 19 439 nm, 528 nm \pm 26 nm, 591 nm \pm 14 nm, and 631 nm \pm 21 nm total width at half-440 maximum, respectively (Fig. 2E). Like in the preliminary tests, up to six specimens 441 were grouped (N=15 groups, 80 animals in total), dark-adapted for 20 minutes and 442 tested in parallel. Each group experienced one run per day. In the tests for negative 443 phototaxis (N=15 groups in total, three trials per wavelength), the animals were placed 444 445 in the illuminable half of the arena, whereas in those for positive phototaxis (N=15446 groups in total, one trial per wavelength) they were set in the dark half of the arena. The behaviour of the animals was recorded for 2 minutes after the light was switched on. 447 448 Experiments, in which animals interacted (e.g. pushed or bit each other, or aggregated), were not analysed, and the animals were retested the following day (30 out of 410 449 450 experiments, 7.3%). To exclude potential biases, the wavelength of the presented light 451 stimulus was changed randomly every day, only precluding the use of the neighbouring 452 wavelengths in subsequent tests.

In all experiments, the time each animal spent in the dark half of the arena was measured, averaged for all specimens of the group and divided by the total experimental time. This resulted in a value ranging from 0 for no avoidance to 1 for total avoidance. The obtained data were analysed using the non-parametric Friedman-test, followed by Dunn's multiple comparison test to compare each wavelength against the control runs.

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These calculations were performed using the statistics program Prism v.6 (GraphPad
Software, La Jolla, CA, USA) and plotted with Adobe Illustrator CS5.1 (Adobe
Systems, San José, CA, USA).

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477 Competing interests

478 The authors declare no competing interests.

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480 Author contributions

H.B., P.A.S. and G.M. designed the behavioural experiments and H.B. carried out these 481 482 experiments and analysed the data. M.H. and A.K. designed the electrophysiological 483 experiments and H.B. and M.H. performed these experiments and analysed the data. 484 L.H. and G.M. designed the gene expression experiments and L.H. carried out these experiments. G.M. and P.A.S. provided the setup and analysing tools for the 485 486 behavioural experiments and A.K. contributed the electrophysiology setup. H.B. wrote the first draft and all authors read, made comments on and approved the final 487 488 manuscript. 489

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583

584 FIGURE LEGENDS

585 Fig. 1. Position of eyes in the two onychophoran species studied, *Euperipatoides*

rowelli and *Principapillatus hitoyensis*. Both species possess a pair of simple lateral
eyes, one at the base of each antenna (arrowheads). Scale bars: 500 μm. (A) The
peripatopsid *E. rowelli*. (B) The peripatid *P. hitoyensis*. Details in A and B are from
specimens preserved in 70% ethanol.

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591 Fig. 2. Experimental design of the electrophysiological and behavioural

experiments. (A) Diagram of an electrophysiological preparation. The head of a 592 593 specimen is embedded in dental cement (green), so that only the area surrounding the 594 eye is accessible for the electrode and the light guide (light cone indicated in blue). (B) Diagram of a sagittal section of the eye (based on a histological section from Mayer, 595 2006) illustrating the desired position of the electrode. Anterior is left. The electrode 596 does not penetrate the cornea of the eye but is inserted through the adjacent cuticle and 597 tissue. (C) Top view of the arena used for behavioural experiments. Only one half was 598 illuminated (indicated in light-blue). (D) Overview of the entire behavioural setup. (E) 599 Normalised emission spectra of the narrow-banded light-emitting diodes used in the 600 601 behavioural setup. Abbreviations: an, antenna; ar, arena; ca, video camera; co, cornea; 602 ct, connective tissue; da, dark half of the arena; dp, dermal papilla; el, electrode; ey, eye; 603 ia, illuminated half of the arena; le, leg; lg, light guide; og, optic ganglion cells; on, 604 optic neuropil; os, outer segments of photoreceptors; pl, perikaryal layer of photoreceptors; sp, slime papilla; tr, trunk; vb, vitreous body (=lens-like structure). 605 606

Fig. 3. Spectral sensitivity of the eye in the peripatopsid *E. rowelli* and the

608 peripatid *P. hitoyensis* determined by electroretinograms (ERGs). (A,C) ERGs

recorded for flashes of white light with increasing intensity (upper trace) consisted of an
initial hyperpolarization truncated by a depolarization (lower trace). Response
amplitudes (insets) were calculated as potential changes from the baseline at stimulus
onset to maximal hyperpolarization. Dots represent measurements, for which no ERG is
shown. (B,D) Averaged spectral sensitivity of the eye in two individuals (mean ±
standard deviation, six measurements per data point) based on response amplitudes to

615 light stimuli of different wavelengths and equal photon flux. Fitting a template (grey

616 line) for the absorbance of an opsin-based visual pigment to the measurements yielded a 617 wavelength of peak absorbance (λ_{max}) of about 480 nm with a coefficient of 618 determination (R²) over 0.9 for both species.

619

620 Fig. 4. Expression pattern of Er-onychopsin and Er-c-opsin mRNA in the 621 peripatopsid E. rowelli visualised using antisense probes. Horizontal sections of heads. Anterior is up in all images. Light (A) and confocal micrographs (B–F) 622 623 illustrating the results of fluorescence in situ hybridization experiments using antisense RNA probes which were visualized using fluorescein-labelled tyramide. DNA was 624 625 stained using propidium iodide. (A) Cryosection showing the spatial relationship of the eyes to the brain (dashed line). Scale bar: 250 µm. (B) Section at the level of the eyes 626 demonstrating the expression of *Er-onychopsin* in the eyes but not in the brain or other 627 tissues. Scale bar: 250 µm. (C) Section of an eye showing that Er-onychopsin is 628 expressed exclusively in the photoreceptor cell layer (arrows). The cuticle exhibits 629 autofluorescence, which is also evident in sections labelled with the sense probe as a 630 negative control (see supplementary material Fig. S2). Scale bar: 50 µm. (D) Detail of 631 the photoreceptor cell layer. Er-onychopsin expression is restricted to the perikarya 632 633 surrounding each nucleus (arrows). Scale bar: 10 µm. (E) Overview of Er-c-opsin expression in a section of the head at the level of the eyes. Er-c-opsin is expressed in the 634 635 deutocerebrum (arrows) and in the optic ganglion cells (arrowhead) in the proximal 636 portion of the eye (sectioned horizontally on the left side; see inset in the lower right corner for a higher magnification; scale bar: $10 \,\mu$ m). Note the lack of signal in the 637 638 photoreceptor cell layer of the eye on the right side. Scale bar: 250 µm. (F) Overview of *Er-c-opsin* expression in a section through the ventral part of the brain at the level of the 639 640 mushroom bodies. Er-c-opsin is expressed in the median portion of the protocerebrum 641 (arrows) and in the connecting cords linking the brain to the ventral nerve cord 642 (arrowheads). Scale bar: 250 µm. Inset in the lower left corner shows detail of expression in the medioventral perikaryal layer within the protocerebrum. Scale bar: 10 643 644 μm. Abbreviations: at, antennal tract; br, brain; cn, central brain neuropil; cc, connecting cord; co, cornea; de, deutocerebrum; dp, dermal papilla; ep, epidermis; ey, 645 eye; fn, frontal brain neuropil; lu, lumen of the eye vesicle; mb, mushroom body; og, 646

optic ganglion cell layer; on, optic neuropil; pc, photoreceptor cell layer; pe, perikaryallayer of the brain; ph, pharynx; vb, vitreous body.

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650 Fig. 5. Light avoidance behaviour in the peripatopsid *E. rowelli*. (A) White light: 651 Plotted pathways of N=12 E. rowelli (Peripatopsidae), each tested twice (n= 24). Bright 652 white light was presented onto the head of each animal. All animals change their walking direction and turn away from the light stimulus as compared to the control (B). 653 654 The pathways of the same 12 animals are unaffected when no light was presented. The differences in turning are highly significant (p<0.001; Wilcoxon-signed-rank test). (C) 655 656 Sensitivity threshold of *E. rowelli* under 5 and 2 minutes of blue light illumination $(\lambda max = 465 \text{ nm})$. Boxplots (N=7 groups of up to 6 animals, circles give the median, 657 boxes the quartiles and whiskers represent 10/90 percentiles) illustrate the fraction of 658 time spent in the dark half of the arena relative to the total time of the test. Significant 659 avoidance reaction (** p<0.01; Friedman-test with Dunn's post-test) occurs at an 660 intensity of 6×10^{11} photons cm⁻² s⁻¹, whereas no significant reaction is seen at 3×10^{11} 661 $cm^{-2} s^{-1}$ (ns, p=0.570; Friedman-test with Dunn's post-test). 662

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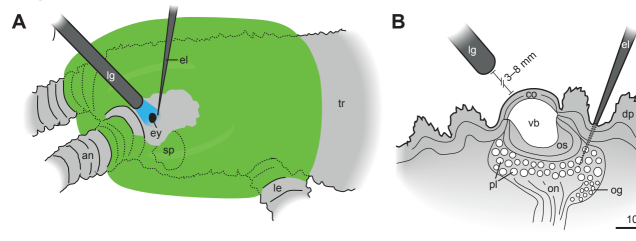
664 Fig. 6. Behavioural response to light stimuli of equal quanta of different

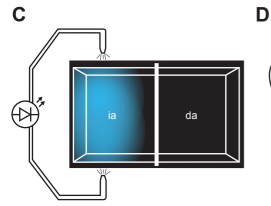
wavelengths in the peripatopsid E. rowelli. Groups (N=15) of up to 6 animals were 665 stimulated for 2 minutes with a spectral light stimulus of 12×10^{11} photons cm⁻² s⁻¹. In 666 667 control experiments, the LEDs remained switched off. Friedman-test with Dunn's posttest was used for data analysis. Boxplots (format as in Fig. 5) illustrating the fraction of 668 669 time spent in the dark half of the arena relative to the total test time. Colours symbolise the wavelengths used. (A) Animals started in the illuminated half and showed negative 670 phototactic behaviour to the light spectrum ranging from UV to green (*** p<0.001), 671 but not to longer wavelengths ($p_{591}=0.343$ and $p_{631}=0.650$, ns=not significant). Every 672 673 group is represented by a mean of three repetitions per wavelength. Dashed line indicates the average spectral sensitivity curve for E. rowelli obtained from 674 675 electrophysiological recordings, with the baseline adjusted to the median of the behavioural control and the maximum standardised to 1. (B) Animals started in the dark 676

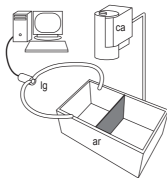
half of the arena, while the other half was illuminated. No significant difference from

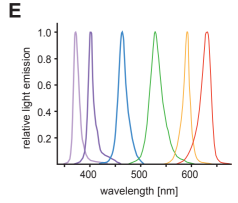
- 678 the control is evident for any of the tested wavelengths (p_{465} and $p_{591}=0.769$,
- $p_{others} > 0.999$). Dashed line indicates the median of the control.





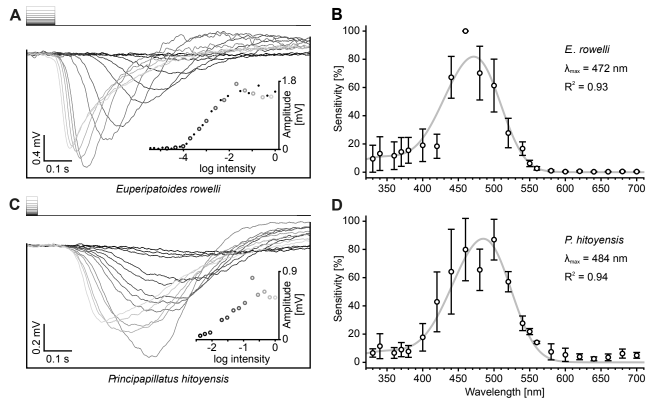


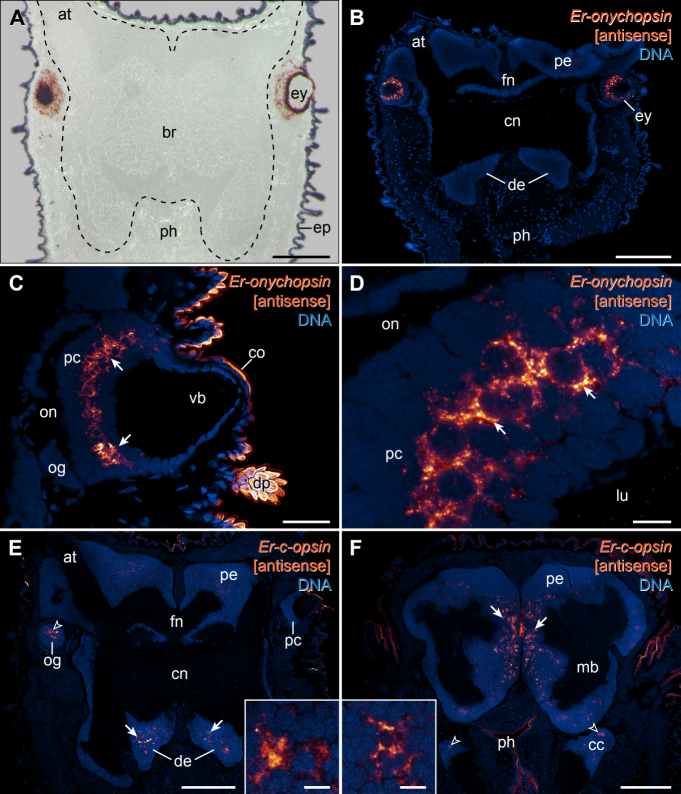


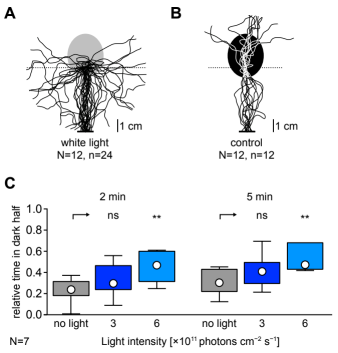


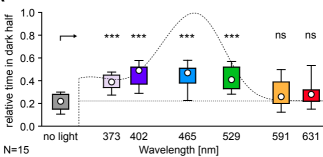
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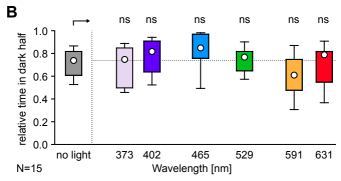
100 µm











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