RESEARCH ARTICLE



Seeing Picasso: an investigation into the visual system of the triggerfish *Rhinecanthus aculeatus*

Karen L. Cheney^{1,*}, Jemma Hudson¹, Fanny de Busserolles², Martin Luehrmann², Abigail Shaughnessy^{1,2}, Cedric van den Berg¹, Naomi F. Green¹, N. Justin Marshall² and Fabio Cortesi^{2,*}

ABSTRACT

Vision is used by animals to find food and mates, avoid predators, defend resources and navigate through complex habitats. Behavioural experiments are essential for understanding animals' perception but are often challenging and time-consuming; therefore, using species that can be trained easily for complex tasks is advantageous. Picasso triggerfish, Rhinecanthus aculeatus, have been used in many behavioural studies investigating vision and navigation. However, little is known about the molecular and anatomical basis of their visual system. We addressed this knowledge gap here and behaviourally tested achromatic and chromatic acuity. In terms of visual opsins, R. aculeatus possessed one rod opsin gene (RH1) and at least nine cone opsins: one violetsensitive SWS2B gene, seven duplicates of the blue-green-sensitive RH2 gene (RH2A, RH2B, RH2C1-5) and one red-sensitive LWS gene. However, only five cone opsins were expressed: SWS2B expression was consistent, while RH2A, RH2C-1 and RH2C-2 expression varied depending on whether fish were sampled from the field or aquaria. Levels of LWS expression were very low. Using fluorescence in situ hybridisation, we found SWS2B was expressed exclusively in single cones, whereas RH2A and RH2Cs were expressed in opposite double cone members. Anatomical resolution estimated from ganglion cell densities was 6.8 cycles per degree (cpd), which was significantly higher than values obtained from behavioural testing for black-and-white achromatic stimuli (3.9 cpd) and chromatic stimuli (1.7-1.8 cpd). These measures were twice as high as previously reported. This detailed information on their visual system will help inform future studies with this emerging focal species.

KEY WORDS: Behaviour, Colour vision, Opsin, Gene expression, Visual pigment, Retinal topography, Visual acuity

INTRODUCTION

Behavioural evidence of colour vision in non-human animals was first demonstrated in bees over 100 years ago (von Frisch, 1914). Since then, behavioural experiments have been conducted with a

¹School of Biological Sciences, The University of Queensland, Brisbane, QLD 4072, Australia. ²Queensland Brain Institute, The University of Queensland, Brisbane, QLD 4072, Australia.

*Authors for correspondence (k.cheney@uq.edu.au; f.cortesi@uq.edu.au)

K.L.C., 0000-0001-5622-9494; F.d., 0000-0002-4602-9840; M.L., 0000-0002-4060-4592; A.S., 0000-0001-9330-0093; C.v., 0000-0001-6422-7237; N.F.G., 0000-0001-5199-7857; N.J.M., 0000-0001-9006-6713; F.C., 0000-0002-7518-6159

This is an Open Access article distributed under the terms of the Creative Commons Attribution License (https://creativecommons.org/licenses/by/4.0), which permits unrestricted use, distribution and reproduction in any medium provided that the original work is properly attributed.

Received 21 December 2021; Accepted 25 February 2022

range of animal species to investigate the mechanisms that underly colour and other visual processes (reviewed in Kelber et al., 2003). Furthermore, psychophysical experiments have explored higherorder neural processes such as colour constancy (e.g. Olsson et al., 2016), generalisation (e.g. Baddeley et al., 2001) and categorisation (e.g. Caves et al., 2018; Jones et al., 2001). However, behavioural experiments with animals are often challenging and timeconsuming; therefore, it can be advantageous to use species that can be trained easily and perform complex tasks well. Historically, terrestrial animals such as honeybees, birds (e.g. chicks, finches, budgerigars, blue tits), lizards and mice have performed well in such studies (e.g. de Ibarra et al., 2002; Lind and Kelber, 2011; Silvasti et al., 2021; Stoddard et al., 2020). Focal teleost species have included zebrafish, goldfish, sticklebacks, cichlids, guppies and damselfish (e.g. Escobar-Camacho et al., 2019; Neumeyer, 1985, 1986, 1992; Risner et al., 2006; Sibeaux et al., 2019, 2008).

Over the past decade, a coral reef triggerfish, Rhinecanthus aculeatus (Linnaeus 1758) (commonly known as the Picasso or lagoon triggerfish), has been used in a number of studies on visual processes, including the role of double cones (two single cones joined together) (Pignatelli et al., 2010), colour discrimination thresholds (Champ et al., 2016; Cheney et al., 2019), the perception of visual illusions (Simpson et al., 2016), the impact of caustics on object detection (Matchette et al., 2020), the segregation of objects (Mitchell et al., 2017) and context-dependent luminance perception (van den Berg et al., 2020). Rhinecanthus aculeatus has also been used to understand how animals collect, process and use spatial information to accurately navigate through their environment (Karlsson et al., 2019 preprint), for bioinspired fish robots (Hu et al., 2006) and to provide an example of unusual sound production in teleosts (Parmentier et al., 2017). They are ideal for behavioural studies as they are relatively easy to keep, have a bold personality, and can be trained for complex tasks using operant conditioning. However, a detailed molecular and anatomical investigation of their visual system is needed to help interpret behavioural results, which we present in this study.

In common with other vertebrates, the photoreceptor cells of fish are in a backward-facing layer at the back of the retina, and contain visual pigments, constructed from an opsin protein with a centrally bound light-sensitive retinal chromophore (reviewed in Musilova et al., 2021). The visual pigment absorbs light and initiates the phototransduction cascade, which converts the light input into electric signals. These signals are carried through several neuronal layers before reaching the ganglion cells that converge into the optic nerve. It is the number and spacing of the ganglion cells that provide the upper limit for spatial resolution or visual acuity. Teleosts often show specialisation in ganglion- and photoreceptor-cell densities and distribution that reflect their specific ecologies. Fishes that live in highly complex 3D visual environments such as coral reefs often have concentric areas of higher cell densities that improve visual resolution along their main visual axes. In contrast, teleosts that occupy simpler visual environments, such as at the sand–water interface, have horizontal streaks of higher cell densities that assist in scanning the horizon for food or predators (reviewed in Cortesi et al., 2020).

To perceive colour, animals must have at least two photoreceptor types with different spectral sensitivities, and the relative excitation ratios from each photoreceptor type are compared in a colour opponency process. The peak spectral sensitivity (λ_{max}) of the visual pigment depends both on variability in key opsin amino acids in the retinal binding pocket and the chromophore type used: vitamin A1-based pigment is shorter shifted compared with vitamin A2-based chromophore (reviewed in Carleton et al., 2020). Vertebrate visual opsins are classified into five types based on their photoreceptor specificity, phylogeny and the λ_{max} they confer. These five visual opsins were most likely already present in the vertebrate ancestor (Collin et al., 2003) and encode one rod opsin (rhodopsin, RH1, teleost λ_{max} =447–525 nm) and four cone opsins: a short-wavelength-sensitive protein class 1 opsin (SWS1) maximally sensitive to UV-violet wavelengths (347–383 nm λ_{max}), a second short-wavelength-sensitive class opsin (SWS2) maximally sensitive to the violet–blue part of the spectrum (397–482 nm λ_{max}), a middle-wavelength-sensitive class 2 rhodopsin-like opsin (RH2) maximally sensitive to blue–green wavelengths (452–537 nm λ_{max}) and a long-wavelength-sensitive class opsin (LWS) maximally sensitive to the green-red part of the light spectrum (501-573 nm λ_{max}) (Carleton et al., 2020). SWS proteins are usually found in single cones, whereas RH2 and LWS opsins occur in double cones (Carleton et al., 2020).

Previously, the spectral sensitivities of Rhinecanthus aculeatus photoreceptors were measured using microspectrophotometry (MSP) (Cheney et al., 2013). Rhinecanthus aculeatus has a single cone that houses a short-wavelength pigment (413 nm λ_{max}), whereas medium- (480 nm λ_{max}) and long-wavelength pigments (528 nm λ_{max}) are housed separately in the two members of the double cone. Rhinecanthus aculeatus also use the two members of the double cone independently in colour vision, facilitating trichromatic colour perception (Pignatelli et al., 2010). This suggests that - unlike birds, for example - some fish may not have photoreceptors specialised for either chromatic or achromatic vision (but see the review by Baden, 2021, on visual circuits in zebrafish). However, little is known about the molecular basis of vision in this species. Therefore, we first used whole genome sequencing and high-throughput RNA sequencing (RNAseq) to determine opsin gene repertoire and opsin expression. Second, with each expressed opsin sequence and known data on spectral tuning sites, the spectral sensitivity for each opsin class was predicted to examine whether it compared with previously published MSP data (Cheney et al., 2013). We then used fluorescence in situ hybridisation (FISH) to determine opsin specificity to single and double cone types.

Finally, we used an integrative approach to examine visual acuity in this species using retinal topography of photoreceptor and ganglion cells across the retina, and behavioural acuity experiments of both achromatic and isoluminant chromatic gratings using a paired-choice test with square wave gratings. Previous behavioural testing with this species using achromatic black-and-white gratings suggests that this species has a visual acuity of 1.75 cpd (Champ et al., 2014). However, anatomical investigations of triggerfish suggest higher possible visual acuity of 3.4 cpd for ganglion cells (Champ et al., 2014). Spatial contrast sensitivity may be higher for achromatic gratings than for chromatic gratings, which has been shown in other vertebrates, including humans and birds (Lind and Kelber, 2011; Mullen, 1985). In this study, we provide a deeper investigation into the visual system of this emerging focal species, which we hope will inform future studies of visual perception and navigation.

MATERIALS AND METHODS

Study species and specimen collection

Rhinecanthus aculeatus are common reef fish found on shallow sub-tidal reef flats across the Indo-Pacific region and are generalist omnivores known to feed on a varied diet including algae, detritus, molluscs and crustaceans (Randall et al., 1997). Individuals (*n*=23) were caught using clove oil and hand nets from shallow reefs surrounding Lizard Island, Great Barrier Reef, Australia, or obtained from an aquarium supplier (Cairns Marine Pty Ltd, Cairns, Australia) between 2017 and 2020. Fish ranged in size from 8.1 to 20.6 cm (standard length, SL) and were collected under a Queensland General Fisheries Permit (183990) and Great Barrier Reef Marine Park Authority Permit (G16/38497.1). Sex and age were not determined. Experiments were approved by the University of Queensland Animal Ethics Committee (2017/AEC000077 and QBI/304/16).

For molecular and anatomical techniques, fish (n=15) were anaesthetised with clove oil (10% clove oil; 40% ethanol; 50% seawater) and then euthanised. Eyes were enucleated, then the cornea and lens were removed. This species has a yellow corneal pigment (Siebeck and Marshall, 2001), the density of which increases across the cornea during the day (N.F.G., unpublished observations). Depending on the analysis, the eyes were preserved in different fixative solutions. Eyes that were allocated to retinal mapping were fixed overnight in 4% paraformaldehyde (PFA) and stored at 4°C. Following this, these eyes were stored in 0.1 mol 1^{-1} phosphate-buffered saline (PBS; pH 7.4) until further analysis. For RNA sequencing, retinas were dissected out of the evecup and preserved in RNAlater (Ambion) at -20° C until extraction. For FISH, retinas were prepared following the protocol in Barthel and Raymond (2000) and fixed in 4% PFA at 4°C overnight, washed twice for 5 min in PBS and rinsed briefly in 70% MetOH, before being transferred to 100% methanol and stored at -20° C until use. One individual was fin clipped and the tissue was preserved on 100% ethanol for genome sequencing.

For behavioural experiments, fish (n=8) were transported to the University of Queensland and housed in individual tanks $(89\times41\times22 \text{ cm})$. Each tank had continuously flowing water supplied by a sump system and air stones to oxygenate the water.

Transcriptome and genome sequencing, quality filtering and *de novo* assembly

The retinal transcriptomes of five *R. aculeatus* were sequenced according to Musilova et al. (2019). One individual was euthanised immediately after capture from the reefs off Lizard Island, Great Barrier Reef, and four individuals were euthanised after being in aquaria at the University of Queensland for between 10 and 18 months. In summary, total RNA was extracted using an RNeasy midi kit (Qiagen), and the quality and concentration of the RNA was checked using a Eukaryotic Total RNA Nanochip on an Agilent 2100 BioAnalyzer. The transcriptome sequencing and RNAseq libraries (paired end, 150 bp insert) were outsourced to Novogene (novogene.com). The transcriptome filtering and *de novo* assembly was performed following the protocol described in de Busserolles et al. (2017). Briefly, the raw reads were uploaded to the Genomics Virtual Laboratory (v.4.0.0) (Afgan et al., 2015) on the

Bioinformatics platform Galaxy Australia (https://usegalaxy.org. au/). Using FastQC (Galaxy v.0.72), the quality of the sequences was assessed. They were then quality filtered using Trimmomatic (Galaxy v. 0.36.6) (Bolger et al., 2014), followed by the *de novo* assembly using Trinity (Galaxy v.2.8.5) (Haas et al., 2013). The newly sequenced samples were subsequently combined with three of our previously sequenced wild-caught samples from Lizard Island (Musilova et al., 2019) to make up our *R. aculeatus* opsin expression dataset.

To identify the visual opsin genes from the transcriptomes, further analyses were performed following the detailed protocol in de Busserolles et al. (2017). The assembled transcripts of R. aculeatus were mapped to the opsin gene sequences of the dusky dottyback (Pseudochromis fuscus; GenBank accession no. KP004335.1) using the medium sensitivity (30% maximum mismatch between transcripts) in Geneious v.2020.1.2 (www. geneious.com). Pseudochromis fuscus was chosen because of its relatively close phylogenetic relationship with R. aculeatus, as well as having representatives from all five visual opsin gene classes. To ascertain that all opsin genes were correctly assembled, and lowly expressed genes were picked up in the assembly, the filtered, unassembled transcriptome reads were then back-mapped against the extracted opsins' sequences using medium-low sensitivity (20% maximum mismatch between reads). For RH2, we detected extra copies that were only partially assembled or that were misassembled. Hence, in this case we used a read mapping approach to disentangle similar gene copies as per de Busserolles et al. (2017) and Musilova et al. (2019). Filtered, unassembled reads were mapped against the P. fuscus RH2 reference using mediumlow sensitivity settings (20% maximum mismatch between reads). We then extracted copy-specific reads by moving along the reference from single-nucleotide polymorphism (SNP) to SNP. Paired-end information allowed us to bridge the gaps between the SNPs. If the coding region was not completely recovered, we remapped unassembled reads using the consensus as a template with low sensitivity settings (0–2% maximum mismatch between reads). This allowed us to reconstruct the whole coding region of all expressed RH2 copies.

To make sure that we did not miss any visual opsin genes that were not expressed in our samples, we sequenced a draft genome for R. aculeatus (ID Olaf) using Illumina short-read technology. Genomic DNA was extracted from a fin clip using the Qiagen DNeasy Blood & Tissue kit according to the manufacturer's protocol (qiagen.com). DNA quality control (Agilent 5400, agilent.com), library preparation and sequencing (paired end 150 bp, 350 bp insert) was outsourced to Novogene (novogene.com). Visual opsin genes were subsequently extracted from the genomic raw reads (212,097,824 paired-end fragments, 31.8 Gb) using the read-mapping approach as outlined above and detailed in Musilova et al. (2019). The only difference was that for the genome, we used the raw reads and mapped them against the single exons of the P. fuscus opsins to avoid long-repetitive intronic sequences. The extracted opsin genes from the genome were then combined with the ones mined from the transcriptomes to generate an R. aculeatus visual opsin gene dataset.

We confirmed opsin gene identity by using BLAST (https://blast. ncbi.nlm.nih.gov/) and by phylogenetic reconstruction. To obtain the opsin gene phylogeny, the opsin gene sequences of *R. aculeatus* were aligned with a reference dataset (obtained from GenBank, https://www.ncbi.nlm.nih.gov/genbank) using MAFFT v.7.450 (Katoh and Standley, 2013) with the L-INS-I settings. On the CIPRES platform, jModeltest v.2.1.6 was used to select the most appropriate model of sequence evolution. Following this, MrBayes v.3.2.7a (Ronquist et al., 2012) was used to infer the phylogenetic relationship between the genes. The following parameters were applied: GTR+I+ γ model, with two independent MCMC searches (four chains each), 10 million generations per run, a tree sampling frequency of 1000 generations, and a burn-in of 25%.

Opsin gene expression

For quantitative gene expression, the filtered reads were mapped to the coding regions of the identified opsin genes with high-specificity settings (98% identity, 80 bp minimum read overlap) as per the protocol in de Busserolles et al. (2017). Only one single cone gene, SWS2B, was expressed. For the other opsins, the gene-specific proportional expression was calculated as the fraction of all expressed for double cone genes *RH2*s and *LWS*, and all opsin genes for *RH1* (see Yourick et al., 2019, for a detailed discussion on opsin gene expression calculations) (Table S2).

For each opsin gene (i), the read count (R_i) was normalised to the length of its coding sequence $(L_i; bp)$:

$$R_{i,\text{normalised}} = (R_i/L_i). \tag{1}$$

The proportion of rod opsin expressed was calculated as the proportion of *RH1* ($p_{i,rod}$) relative to the total normalised opsin expression (T_{opsin}):

$$p_{i,\text{rod}} = (R_{i,\text{normalised}} / T_{\text{opsin}}).$$
⁽²⁾

The proportional expression of double cones $(p_{i,DC})$ was calculated as the proportion of the normalised opsin expression $(R_{i,normalised})$ out of the total normalised expression for double cones (T_{DC}) :

$$p_{i,\text{DC}} = (R_{i,\text{normalised}} / T_{\text{DC}}). \tag{3}$$

Fluorescence in situ hybridisation (FISH)

Dual-labelling FISH was performed on whole-mount retinas of one adult, wild-caught R. aculeatus following standard protocols (Barthel and Raymond, 2000; Dalton et al., 2015). Previously extracted retinal mRNA was reverse transcribed using a High-Capacity RNA-to-cDNA Reverse Transcription Kit (Applied Biosystems). Riboprobe templates were synthesized from cDNA via standard PCR using Q5 High Fidelity DNA polymerase (New England Biolabs) and opsin-specific primers (Table S1). Amplicons were isolated via gel-electrophoresis and gel extraction (Qiagen Gel Extraction Kit), followed by enrichment PCR using gel-extracted amplicons as cDNA template. Primers were designed to bind to the coding sequence of target opsins (RH2A, RH2C, SWS2B), and to contain T3 or T7 RNA polymerase promoter sequences at their 5' ends (T3, reverse primer; T7, forward primer) to allow subsequent strand-specific RNA transcription from cDNA templates for riboprobe synthesis. Antisense riboprobes were synthesised and labelled with digoxigenin-UTP (DIG) or fluorescein-UTP (FL) using DIG/FL RNA labelling mix (Sigma-Aldrich). A single RH2C riboprobe, targeting both expressed RH2C-1 and RH2C-2 paralogues, was synthesised owing to high sequence similarity between these. Hybridised, labelled riboprobes were detected using anti-digoxigenin (Sigma-Aldrich) or anti-fluorescein/Oregon Green (ThermoFisher) antibodies conjugated to horseradish peroxidase. Fluorescent tagging was performed using Alexa Fluor 594 or 488 dyes with tyramide signal amplification (Invitrogen). Finally, retinas were mounted in 70% glycerol in PBS, photoreceptor side up, on microscopy slides with a coverslip.

Dual (*RH2A/RH2C*) or single (*SWS2B*) labelled photoreceptor cells were visualised and imaged using a CFI Apo Lambda S LWD 40X/1.15 NA water immersion objective on a spinning disk confocal microscope (Diskovery, Andor Technology, UK) built around a Nikon Ti-E body (Nikon Corporation, Japan) equipped with two Zyla 4.2 sCMOS cameras (Andor Technology), and controlled by Nikon NIS Elements software. Images were exported in TIF file format and further processed (merging of colour channels, adjusting of brightness, *z*-stack projection) with ImageJ v.1.52p (National Institutes of Health, Bethesda, MD, USA).

Prediction of visual pigment maximal absorbance

Maximal absorbances (λ_{max}) of expressed *R. aculeatus* visual pigments were predicted by comparing opsin amino acid sequences with those of percomorph fish species for which the peak spectral sensitivities of their (A1 chromophore-based) visual pigments were known from in vitro pigment reconstitution (Oryzias latipes, RH1: 502 nm λ_{max} , AB180742.1, Matsumoto et al., 2006; Oreochromis *niloticus*, SWS2B: 425 nm λ_{max} , JF262088.1, RH2B: 472 nm λ_{max} , JF262086.1, RH2Aalpha: 528 nm λ_{max} , JF262086.1, LWS: 560 nm $\lambda_{max},$ JF262088.1, Spady et al., 2006) and by applying tuning effects to their respective peak spectral sensitivities for substitutions at known tuning sites documented in the literature or substitutions at known tuning sites that match the polarity shift of documented substitutions (e.g. for reviews, see Takahashi and Ebrey, 2003; Yokoyama, 2008; Yokoyama and Jia, 2020). For an overview of considered sites and applied effects, see Table S3. We used O. niloticus RH2Aalpha rather than RH2Abeta as the reference owing to its greater amino acid sequence similarity to R. aculeatus RH2A. We used O. niloticus RH2B as the reference for RH2C1 and RH2C2 owing to the lack of in vitro characterised RH2Cs (Musilova and Cortesi, 2021 preprint). Opsin gene sequences were translated into amino acid sequences and aligned using MAFFT (v.7.450) (Katoh and Standley, 2013) in Geneious Prime (v.21.1.1). Bovine rhodopsin (NP_001014890.1) was included in all alignments to identify amino acid residues corresponding to known tuning sites and transmembrane regions according to its crystal structure (Palczewski et al., 2000).

Topographic distribution of ganglion cells and cone photoreceptors

Preparation of retinal whole-mounts

Retinal whole-mounts were prepared according to standard protocols (Coimbra et al., 2006; Stone et al., 1981; Ullmann et al., 2012). The orientation of the retina was kept by noting the position of the falciform process inside the eyecup once the cornea and lens were removed. In all our *R. aculeatus* specimens, the falciform process ended in the ventral margin of the retina. Each retina was bleached overnight at room temperature in a solution of 3% hydrogen peroxide in $0.1 \text{ mol } I^{-1}$ PBS. For photoreceptor analysis, retinas were whole-mounted (photoreceptor layer up) in

100% glycerol on a microscope slide. For ganglion cell analysis, retinas were whole-mounted, ganglion cell layer facing up, on a gelatinised slide, left to dry overnight in formalin vapour to improve fixation and cell differentiation (Coimbra et al., 2006, 2012), stained in 0.1% Cresyl Violet (Coimbra et al., 2006) and cover slipped with Entellan New (Proscitech). Possible shrinkage during staining was considered negligible and, if present, confined to the retinal margins, as the retinal whole-mount was attached to the slide during the entire staining process (Coimbra et al., 2006).

Stereological analyses and topographic map construction

Following the protocols described in de Busserolles et al. (2014a,b), topographic distribution of single cones, double cones, total cones and ganglion cells were assessed using the optical fractionator technique (West, 1991) modified by Coimbra et al. (2009, 2012). Briefly, cone photoreceptors and ganglion cells were randomly and systematically counted using the parameters listed in Table 1 and a 63× oil objective (numerical aperture 1.40) mounted on a compound microscope (Zeiss Imager.Z2) equipped with a motorised stage (MAC 6000 System, Microbrightfield, USA), a digital colour camera (Microbrightfield) and a computer running StereoInvestigator software (Microbrightfield). The counting frame and grid size were chosen carefully to maintain the highest level of sampling and achieve an acceptable Schaeffer coefficient of error (CE<0.1; Glaser and Wilson, 1998). The grid size was adjusted between individuals to take into consideration the variation in size between specimens and allow sampling of around 200 sites per retina (Table 1). To obtain a more accurate estimate of the retinal ganglion cell peak density, sub-sampling was performed in the highest cell density area using the same counting frame but with half the grid size from Table 1.

Single cones and double cones were easily distinguished and counted separately and simultaneously using two different markers to generate data for single cones alone, double cones alone and the two cell types combined (total cones). Ganglion cells were arranged in a single layer within the ganglion cell layer that also comprised displaced amacrine cells and glial cells. Because amacrine cells were not easily distinguished from ganglion cells using cytological criteria alone (Collin and Collin, 1988; Hughes, 1975), especially in high density areas, they were included in the cell counts and only glial cells were excluded. Although the inclusion of amacrine cells in the analysis usually does not influence the overall topography (Collin and Pettigrew, 1989), it may contribute to a slight overestimation of the peak density of ganglion cells and, ultimately, to a slight overestimation of spatial resolving power.

Topographic maps were constructed in R v.2.15.0 (https://www. r-project.org/) with the results exported from the Stereo Investigator Software according to Garza-Gisholt et al. (2014). The Gaussian kernel smoother from the Spatstat package was used (Baddeley and Turner, 2005) and the sigma value was adjusted to the grid size.

Table 1. Summary of the stereology parameters used for the photoreceptor (PR) and ganglion cell (GC) topography analysis

Individual	SL (cm)	N/T eye Ø (mm)	D/V eye Ø (mm)	Cell counted	Counting frame (µm)	Grid (µm)	CE
1	13.3	9.7	7.8	PR	60×60	700×700	0.050
2	12.0	9.0	7.0	PR	60×60	637×637	0.047
3	13.5	8.6	6.3	GC	90×90	720×720	0.056
4	10.4	8.2	6.3	GC	90×90	600×600	0.051

SL, standard length; N, nasal; T, temporal; D, dorsal; V, ventral; Ø, diameter; CE, Schaeffer coefficient of error.

Estimation of spatial resolving power

The upper limit of the spatial resolving power (SRP) in cpd was estimated for each individual using the peak density of ganglion cells (PDG in cells mm^{-1}) as described by Collin and Pettigrew (1989). Briefly, the angle *a*, subtending 1 mm on the retina can be calculated as follows:

$$a = \arctan\left(\frac{1}{f}\right),\tag{4}$$

where f, the focal length or the distance from the centre of the lens to the retina, is 2.55 (Matthiessen's ratio; Matthiessen, 1882) times the radius of the lens. Knowing a, the PDG and the fact that two ganglion cells are needed to distinguish a visual element from its neighbour, the SRP in cpd can be calculated as follows:

$$SRP = (PDG/a)/2.$$
 (5)

Behavioural measurements of achromatic and chromatic acuity Experimental setup

We used eight triggerfish (8.1 to 16.5 cm, SL) to measure achromatic and chromatic acuity using a pairwise discrimination behavioural task with square wave gratings (4×4 cm; Fig. S2). Fish were allowed to acclimatise to their tanks for 2–3 weeks and were fed twice a day with a mixture of blended squid, prawns, fish flakes and peas. Tanks were illuminated with 240 V/50 Hz LED Batten 20 W lights (FL2527; Fuzion Lighting, QLD, Australia), which were hung at 50 cm above the end of the tank at which the target stimuli were placed. The side-welling irradiance of each tank was measured with an Ocean Optics USB2000 spectrometer with a 400 μ m fibre and a cosine corrector (Fig. S2). Measurements were taken in the middle of the water column at 10 cm from the end of the divider pointing towards the stimuli.

Each tank was divided into three sections: a holding area and two passages separated by a 50-cm-long white opaque Corflute board, down which fish could swim to reach the target stimuli (Fig. S2). During testing, a transparent Perspex board was initially placed across the tank at the end of the Corflute divider, which allowed the fish to swim back and forth in front of both stimuli several times before the transparent Perspex board was lifted and they could swim down one passage, which was considered a choice by the fish. This set the point of decision to 50 cm.

We tested black–white square wave achromatic gratings and two combinations of chromatic square wave gratings: green–yellow to mainly stimulate the medium and long cones, and pink–purple to stimulate the short and long cones. Stimuli were between 0.5 to 5 cpd for black–white achromatic stimuli (n=10 stimuli) and 0.5 and 3 cpd for chromatic stimuli increasing in increments of 0.5 cpd (n=6 stimuli). A control ('solid') stimulus was made using a square wave grating of 11 cpd, which was deemed unresolvable by the triggerfish from previous data (Champ et al., 2014). Overall, the total number of fish that were tested for each colour combination was: n=4 for achromatic, n=5 for green–yellow and n=5 for pink–purple.

Square wave gratings were printed on Steinbeis TrendWhite paper using a Canon LaserJet Pro 400 printer and laminated with 80 μ m gloss laminating sheets. We tried to ensure colours within a chromatic combination were as isoluminant as possible (green–yellow and pink–purple) based on quantum catches for an average of double cone members, which are thought to process luminance information (e.g. van den Berg et al., 2020). To calculate quantum catches, the spectral reflectance of printed colours was measured under a black-cloth-covered box with an Ocean Optics USB2000 spectrophotometer (Ocean Optics, FL, USA), a PX-2 pulsed xenon light source and a 200 µm diameter, bifurcated cable held at 45 deg 1 mm above the paper. A Spectralon white standard was used to calibrate the spectrophotometer, and a piece of black velvet covering the end of the fibre was used as the dark standard. Quantum catches of photoreceptors were calculated using photoreceptor spectral sensitivities of triggerfish (Cheney et al., 2013) and illumination and reflectance spectra of printed colours as per eqn 1 in Vorobyev and Osorio (1998). Average normalised quantum catch for the double cone output for the grating colours were: pink 19.3 and purple 18.3, green 27.5 and yellow 27.3, and black 0.8 and white 72.5.

Training

In the first round of testing, each fish was randomly assigned to one of three treatments: black-white (n=2), green-yellow (n=3) or pink-purple (n=3) stimuli (fish were retrained to another treatment in a second round of testing). Within each of these groups, one fish was assigned the control (11 cpd) stimulus as the rewarded (S+) stimulus and the remaining individuals were assigned the striped (0.5 cpd) stimulus as the rewarded (S+) stimulus. This was done to account for any bias in the type of stimulus (control/striped) being learnt. Fish were trained using operant conditioning to approach and peck at a stimulus to receive a food reward. Initially, a small piece (<5 mm) of squid mantle was stuck (using the natural adhesive properties of squid) on the rewarded stimulus, while the unrewarded stimulus was left without. Once fish had associated the S+ stimulus with a reward, the squid was removed from the stimulus and fish were then only presented with a reward from above on forceps once they had pecked the S+ stimulus.

Before commencing each trial, an opaque corflute board was placed in front of the transparent Perspex board to ensure the fish could not see the stimuli being placed at the end of each passage. Stimuli were placed in the left or right passage using a random number generator to ensure that fish did not develop a side bias. Once stimuli were in place, the opaque board was lifted, and the fish were given 20 s to view the stimulus through the transparent Perspex board. The transparent board was then lifted and the side that the fish swam down was recorded. Fish did not peck the stimulus in all cases but were given a food reward for approaching the correct stimulus. In instances when a fish swam down one passage, but then changed direction and then went down the other passage during a trial, only the original choice was recorded. If a fish had not swum down any passage within 1 min of the transparent board being lifted, the fish was deemed unmotivated and trials for that fish were stopped for the rest of that session and continued again in the next session. Once fish successfully chose the S+ stimulus >80% of the time over four sessions, fish moved on to the testing phase. It took 3-4 weeks to train most fish (total of 20–29 sessions for each); however, one fish only took 1 week (nine sessions) to reach the required standard for testing.

Testing

Testing trials were conducted as per training methods, and one to two sessions were conducted each day with a minimum of 2 h between sessions. The training stimulus (0.5 cpd) was used as reinforcement through the testing phase and was presented 20–98 times per fish, depending on the individual. Each grating between 1.0 and 3.0 cpd was presented to each fish 7–20 times for black–white achromatic stimuli and 9–30 times for chromatic stimuli. Once fish had completed testing with their first treatment stimuli (black–white, green–yellow or pink–purple), they were randomly assigned to one of the other two treatments, then trained and tested again as above. However, two fish only completed one treatment owing to the time taken for training and testing. In total, 2418 trials were conducted with each grating presented 110 ± 14 times (mean±s.e.m.) (Table S4).

Data analysis

Data analysis for the behavioural experiment was performed in RStudio v.1.3.1056 (https://www.rstudio.com/). We used the R package quickpsy v.0.1.5.1 (Linares and Lopez-Moliner, 2016) to produce logistic functions for all fish for each target colour set (goodness of fit: deviance <8.6, P>0.47). Our response variable was whether the fish chose (1) or did not choose (0) the positive S+ stimulus. We then used the function 'threshold' to interpolate the 62% threshold, which corresponds to a correct choice frequency significantly different from random behaviour, assuming a binomial distribution of the pooled data per grating (n=110, P<0.01, onetailed binomial test). Once thresholds were calculated for each fish, we tested for significant differences between treatments using a general linear mixed model with the lme4 package v.1.1-27.1 (Bates et al., 2015) and the function lmer. Fish ID was included as a random variable. We produced P-values using the ImerTest package v.3.1-3 (Kuznetsova et al., 2017). There was no difference in whether fish were trained to receive a food reward from the control stimulus (11 cpd) or the grated stimulus ($t_{4,7}=0.33$, P=0.76); fish did not perform differently within a colour treatment between the first and second sessions $(t_{3,0}=1.31, P=0.28)$ and there was no effect of the size of the fish on the visual acuity ($t_{4.6}=0.53$, P=0.62). Therefore, these factors were excluded from the final analysis.

RESULTS

Opsin gene repertoire

The *R. aculeatus* genome contained one rod opsin gene (*RH1*) and at least nine cone opsins: one violet-sensitive *SWS2B* gene, seven duplicates of the blue-green-sensitive *RH2* gene (*RH2A*, *RH2B* and *RH2C1-5*) and one red-sensitive *LWS* gene. For *RH2B* and *RH2C-4* and *RH2C-5*, only the first two exons could be annotated using the whole-genome read-mapping approach. This is because the intron between exons 2 and 3 of *RH2* is too long to be reconstructed using the short reads alone (for details, see Musilova and Cortesi, 2021 preprint). A high-resolution *R. aculeatus* genome based on longread technology will be needed in the future to resolve these genes and to assess whether even more copies are present. Except for *RH2B* and three *RH2C* copies (*RH2C3-5*), all other opsins were recovered from the transcriptomes. The phylogenetic reconstruction confirmed the class of each opsin gene identified (Fig. 1).

Opsin gene expression

The retina of *R. aculeatus* was rod dominated, where *RH1* was expressed on average $68.9\pm3.75\%$ (*n*=8, mean±s.e.m.). *SWS2B* accounted for 100% of the cone opsin expression in single cones. For the double cone opsins, there was change in expression of the *RH2* and *LWS* genes in relation to whether fish were housed in aquaria or sampled directly from the field. Aquarium fish kept for behavioural trials showed higher *RH2A* expression (*n*=4, $60.30\pm1.65\%$) compared with individuals sampled directly from the field (*n*=4, $35.04\pm4.38\%$). However, *RH2C-2* was expressed at higher levels for individuals collected directly from the field ($45.89\pm2.12\%$) compared with aquarium individuals ($27.38\pm3.81\%$). *RH2C-1* was expressed at similar levels between field site collected ($17.34\pm4.55\%$) and aquarium individuals ($12.14\pm2.17\%$). In addition, *LWS* was expressed at low levels for both aquarium and

field site collected individuals (field site: $1.70\pm1.23\%$, aquarium: $0.18\pm0.09\%$) (Fig. 2).

FISH

Double labelling RNA FISH of expressed opsin mRNAs in *R. aculeatus* whole-mount retinas showed that medium wavelength opsins identified from the retinal transcriptome, *RH2A* and *RH2Cs* (*RH2C* probes did not discriminate between the two identified *RH2C* paralogues), were expressed in opposite members of double cone photoreceptors across the retina (Fig. 3A–D). The sole single cone opsin identified from the retinal transcriptome, *SWS2B*, was expressed exclusively in single cones (Fig. 3E,F).

Prediction of visual pigment maximal absorbance

Amino acid sequence similarity between R. aculeatus opsins and the used reference opsins ranged from 85.5% (51 variable amino acids) for RH2C-1 and RH2C-2 to 94.1% (21 variable amino acids) for RH1 (Table S3). Among variable amino acids, 33% (RH1) to 47% (LWS) (data not shown) lay outside transmembrane regions and were thus unlikely to impact spectral tuning. The numbers of variable amino acids across transmembrane regions ranged from 15 in RH1 to 32 in SWS2B, the majority of which were substitutions that did not incur a physicochemical change. One (RH1), five (SWS2B), nine (RH2A), 12 (RH2C-1) and 11 (RH2C-2) were at known tuning sites. However, many of these were undocumented substitutions that did not incur a polarity shift, resulting in consideration of one (RH1), four (SWS2B), three (RH2C-1), two (RH2C-2) and two (RH2A) sites for λ_{max} prediction (Table S3). Rhinecanthus aculeatus LWS did not differ from the reference LWS at any known tuning sites.

Predicted *R. aculeatus* visual pigment peak sensitivities only in part matched the cone and rod sensitivities obtained via MSP (Cheney et al., 2013). RH1 was predicted to be maximally sensitive at 500 nm owing to S299A (-2 nm; Dungan et al., 2016; Fasick and Robinson, 1998), and closely matched the rod λ_{max} (498 nm) determined using MSP. SWS2B was predicted to be maximally sensitive at 403 nm, primarily tuned to shorter wavelengths compared with O. niloticus SWS2B (425 nm) by W265T (-29 nm) and F46V (+8 nm) (Yokoyama et al., 2007). The predicted λ_{max} differed from single cone λ_{max} of 413 nm obtained via MSP. RH2C visual pigments were predicted to be maximally sensitive at 474 nm (RH2C-1) and 476 nm (RH2C-2). Both RH2Cs were slightly red-shifted compared with O. niloticus RH2B by primarily M88C (+3 nm; Chinen et al., 2005). RH2C-1 and RH2C-2 differed by only six amino acids (98.3% similarity), with one of these, T266V in RH2C-1, predicted to cause a 2 nm reduction of its λ_{max} (Chinen et al., 2005). The predicted λ_{max} are 6 nm (RH2C-1) and 4 nm (RH2C-2) shorter than the peak sensitivity of the shorter medium-wavelength-sensitive (MWS) double cone determined via MSP (480 nm). RH2A was predicted to be maximally sensitive at 526 nm, with only two sites, C88A (-3 nm; Chinen et al., 2005) and I112V (+1 nm; Chinen et al., 2005), considered to cause a small net blue shift compared with O. niloticus RH2Aalpha (528 nm). This predicted λ_{max} almost matched the peak sensitivity of the second, longer MWS double cone in R. aculeatus (528 nm) as determined via MSP. As R. aculeatus LWS did not differ from O. niloticus LWS at any known tuning sites, its predicted λ_{max} was 560 nm.

We identified several substitutions that, although not previously investigated via site-directed mutagenesis (SDM) and *in vitro* pigment regeneration, owing to their incurred polarity shifts and the site of their occurrence, may explain the discrepancies between MSP and predicted λ_{max} (see Table S3). In RH1, S166A incurred a

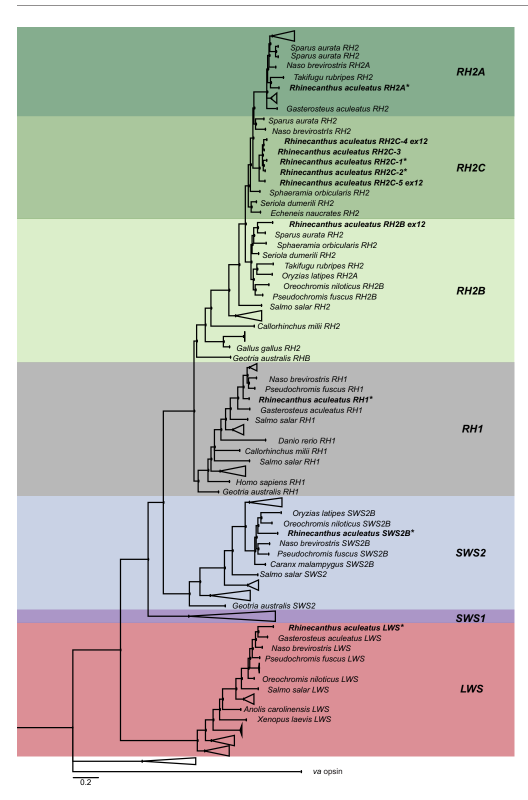


Fig. 1. A phylogenetic tree of the visual opsin genes seen in

vertebrates. The opsin genes belonging to *Rhinecanthus aculeatus* are highlighted in bold. The sequences extracted from the genome and transcriptome (indicated by *) are positioned within their respective opsin class. *RH1*, rhodopsin 1 (rod opsin); *RH2*, rhodopsin-like 2; *SWS2*, shortwavelength sensitive 2; *LWS*, longwavelength sensitive; *va*, vertebrate ancient opsin (outgroup). The black circles represent the Bayesian posterior probabilities >0.8. Scale bar: 0.2 substitutions per site.

polarity shift at a site close to the known RH1 tuning site A164S (Chan et al., 1992) and is also a substitution known to be involved in RH2 tuning (Yokoyama and Jia, 2020). *Rhinecanthus aculeatus* SWS2B showed three polarity changing substitutions, C163F, S166F and S168A, in immediate proximity to the documented tuning site G164A (Yokoyama and Tada, 2003). In both RH2Cs, V185C caused a polarity shift, whereas previously SDM in zebrafish RH2s had shown a -4 nm blue shift for substitution from T to C at this site (Chinen et al., 2005). C98A incurred a

polarity shift at a site adjacent to the powerful RH2 tuning site T97A (-8 nm; Takenaka and Yokoyama, 2007). In *R. aculeatus* RH2A, A151T incurred a polarity shift at a site at which SDM from N to S caused a +4 nm red shift in zebrafish RH2 (Chinen et al., 2005).

Retinal topography and anatomical spatial resolving power

The topographic distribution of ganglion cells and cone photoreceptors was analysed in four individuals (two individuals per cell type). Because intraspecific variability in topography

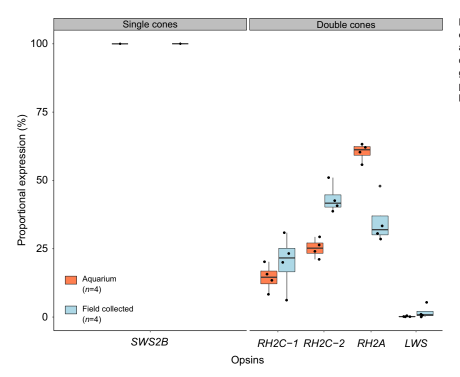


Fig. 2. The proportional cone opsin gene expression of *R. aculeatus* individuals collected from the field and the aquarium. All individuals expressed one single cone opsin gene, *SWS2B*, and four double cone opsin genes, *RH2C-1*, *RH2C-2*, *RH2A* and *LWS*. The proportional expression of the *RH2* and *LWS* differed between aquarium and field-collected individuals.

pattern was very low for both ganglion cells and cone photoreceptors, only one representative map is presented in Fig. 4 and individual maps are provided in Fig. S1.

Ganglion cell topographic distribution revealed the presence of a well-developed horizontal streak with a peak cell density located in the temporal part of the retina (Fig. 4) that ranged from 37,161 to 40,125 cells mm⁻² in the two individuals analysed (Table 2). Based on these peak cell densities, estimated spatial resolving power for *R. aculeatus* ranged from 6.4 to 7.2 cpd (Table 2).

Cone photoreceptors in R. aculeatus were arranged in a regular retinal mosaic where one single cone was surrounded by four double cones, resulting in a double cone to single cone ratio of 2:1. This mosaic pattern was consistent across the entire retina, resulting in similar topographic distributions for each cone type. As a result, we only present and describe the distribution pattern for the total cone population (Fig. 4). Maps for each cone type are provided in Fig. S1. The total cone topography varied slightly from the ganglion cell pattern by having a less developed horizontal streak and a more pronounced area temporalis (Fig. 4). However, the peak cell density of total cones, which ranged from 64,167 to 70,278 cells mm^{-2} (Table 2), was found in the same location (i.e. temporal) as the peak density of ganglion cells. Even though ganglion cells and cone photoreceptor topographies were not analysed in the same retina or individuals, comparison of two individuals of similar size (i.e. \sim 13 cm SL, individuals 1 and 3) suggested a summation ratio of total cones to ganglion cells in the peak density area of around 2:1.

Behavioural measurements of achromatic and chromatic acuity

Behavioural thresholds (62% correct choice) were significantly higher for black–white achromatic stimuli (3.94 cpd, 95% confidence intervals, CI: 3.47–4.49 cpd) than for green–yellow (1.71 cpd, CI: 1.46–1.93 cpd) ($t_{9.31}$ =–7.39, *P*<0.001) or for pink–purple stimuli (1.89 cpd, CI: 1.59–2.38 cpd) ($t_{7.51}$ =–6.00, *P*<0.001). Green–yellow and pink–purple treatments were not significantly different from each other ($t_{6.29}$ =0.58, *P*=0.58) (Fig. 5).

DISCUSSION

Previous MSP measurements of R. aculeatus photoreceptors suggested three spectrally distinct cone types: one blue-sensitive single cone (λ_{max} =413 nm), one blue-green-sensitive double cone member (λ_{max} =480 nm) and a second, green-sensitive double cone member (λ_{max} =528 nm), as well as a single spectral type of rod photoreceptor (λ_{max} =498 nm) (Cheney et al., 2013). Our transcriptome analysis largely - but not fully - reflects this by revealing the expression of a single rod opsin (RH1) and for the cone opsins, a short-wavelength-sensitive opsin (SWS2B), yet four medium-long-wavelength-sensitive cone opsins (RH2C-1, RH2C-2, RH2A and LWS). Except for one field caught individual, LWS expression was at levels barely high enough to reconstruct this gene's coding sequence, suggesting that measured double cone λ_{max} were primarily (if not entirely) due to the three identified *RH2* opsins. Amino-acid-based λ_{max} predictions clarified this picture, indicating that the RH2A visual pigment with a λ_{max} of 526 nm was likely to account fully (i.e. not co-expressed) for the green-sensitive double cone member, and this was further supported by our FISH analysis. Predictions also indicated that the two RH2C paralogues were very close in terms of their λ_{max} , and as both showed high expression, either one or both may account for the blue-greensensitive cone member. Unfortunately, owing to high mRNA sequence similarity (97.3% identity) our FISH analysis did not allow a distinction between the two expressed *RH2C* paralogues. Co-expression of either of the two RH2C paralogues with RH2A, in theory, could have explained the discrepancy between predicted RH2C λ_{max} and measured λ_{max} of the blue–green-sensitive MWS double cone, a pattern seen, for example, in some freshwater cichlids (Dalton et al., 2014; Torres-Dowdall et al., 2017). However, our FISH analysis did not show evidence of RH2A/RH2C coexpression. Instead, it showed that these opsins were expressed in opposite double cone members, a pattern also observed in other reef fish such as some anemonefishes (Stieb et al., 2019). Taken together, the transcriptome data, visual pigment λ_{max} predictions and FISH analysis corroborate the existing, MSP-based picture of the Picasso triggerfish photoreceptor spectral sensitivity repertoire.

Ganglion cells

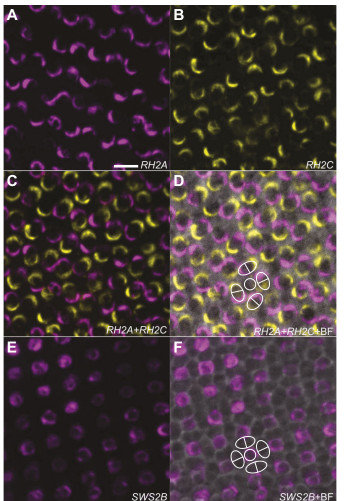


Fig. 3. Double-labelling *in situ* hybridisation of expressed opsin mRNAs in retinal double cone and single cone photoreceptors in *R. aculeatus.* (A–D) *RH2A* (magenta) and *RH2C* (yellow) mRNA were expressed in opposite members of double cones across the retina. (E,F) *SWS2B* (magenta) mRNA was exclusively expressed in single cone photoreceptors across the retina. Representative single and double cones are outlined with white circles and white ovals, respectively. BF: bright field. Scale bar (A–F): 10 µm.

However, although the number of spectrally distinct photoreceptors might be limited, differences in *RH2* expression between individuals that were sampled in the field and the ones that were kept in aquaria shows that opsin gene expression in adult *R. aculeatus* is plastic over time. As for some freshwater fishes (e.g. cichlids; Nandamuri et al., 2017) and reef fishes (e.g. cardinalfishes and damselfishes; Luehrmann et al., 2018), the change in opsin gene expression in *R. aculeatus* was likely caused by differences in the light environment

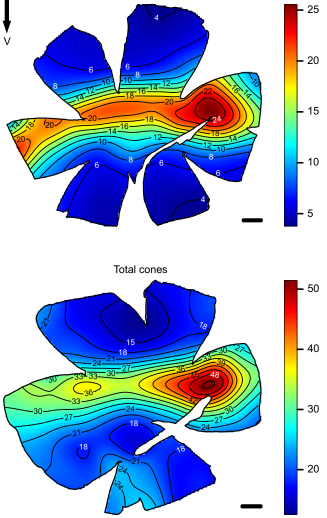


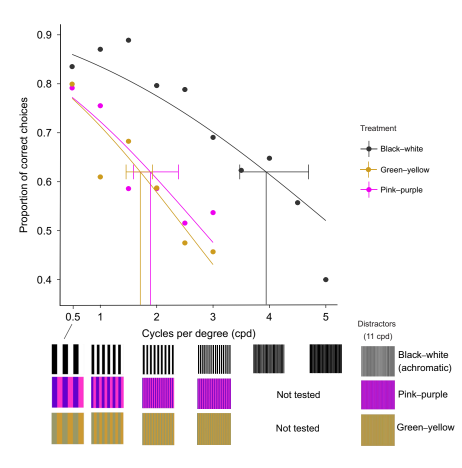
Fig. 4. Topographic distribution of ganglion cells and total cone photoreceptors in *R. aculeatus.* The black lines represent iso-density contours and values are expressed in densities ×10³ cells mm⁻² (see colour scale). The black arrow indicates the orientation of the retinas. T, temporal; V, ventral. Scale bars: 1 mm.

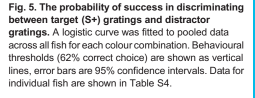
between the reef and the aquarium. The higher expression of RH2A in aquarium-held fishes suggests an overall quantitative shift to longer wavelength sensitivity, i.e. more LWS cones when kept under longer-shifted green/red-dominated LED lights. Although it remains to be investigated whether a change in opsin gene expression truly causes a change in visual discrimination, the findings do highlight the importance of assessing an animal's visual system – from molecule to behaviour – in its specific light environment.

Table 2. Summary of the photoreceptor (Individual 1 and 2) and ganglion cell (Individual 3 and 4) data using the optical fractionator method

Individual	Total	Peak (cells mm ⁻²)	Total DC	Peak DC (cells mm ⁻²)	Total SC	Peak SC (cells mm ⁻²)	Lens Ø (mm)	SRP (cdp)
	Photoreceptors	•						
1	2,034,997	64,167	1,343,688	41,389	691,036	24,722		
2	1,851,208	70,278	1,239,849	48,612	610,231	21,667		
	Ganglion cells							
3	1,074,112	37,161					3.3	7.2
4	784,088	40,125					2.8	6.4

DC, double cones; SC, single cones; Ø, diameter; SRP, spatial resolving power; cpd, cycles per degree.





Prediction of visual pigment maximal absorbance

We do note that the reliability of predicted visual pigment λ_{max} based on amino acid sequences is inherently limited despite tuning sites and their effects having been extensively studied in many taxa, and in fish arguably more so than in any other vertebrate group (for reviews, see Takahashi and Ebrey, 2003; Yokoyama, 2000; Yokoyama and Jia, 2020). Omission of yet unknown tuning sites, effects (if any) of unknown substitutions at known tuning sites, and overestimation or underestimation of tuning effects owing to unknown synergistic effects (including effects that affect visual processing in non-spectral ways, e.g. via modulation of retinal release dynamics; Castiglione and Chang, 2018), are among the shortcomings of this approach (e.g. Chinen et al., 2005; Yokoyama and Jia, 2020). For example, for both RH2Cs the substitution of polar C at site 185, where SDM of polar T to polar C has previously shown to shift λ_{max} by as much as 4 nm (Chinen et al., 2005), could explain some or all of the discrepancy between predictions and MSP. Similarly, the polarity-changing substitutions identified in R. aculeatus SWS2B could, on their own or by reducing the effect of W265T, cause an increase of the pigment's λ_{max} . Consequently, one would consider predictions more robust for more conserved genes (e.g. RH1) or genes in which the extent of λ_{max} variability across many different taxa could be nearly fully described by variations at only a few sites, as is the case for many LWS pigments ('five-sites rule'; Yokoyama and Radlwimmer, 1998). In contrast, λ_{max} of SWS pigments (SWS1, SWS2As, SWS2B) and RH2 pigments are notoriously difficult to predict owing to a plethora of variable sites that have, to some extent, been implicated with spectral tuning, but complicated by apparent yet not fully understood, synergistic effects (Shi et al., 2001; Yokoyama and Jia, 2020; Yokoyama and Tada, 2003). In the future, additional in vitro expression and pigment

regeneration assays, particularly in non-model organism species, as well as further investigation of possible tuning sites via SDM, could improve the reliability of spectral sensitivity predictions.

Retinal topography and anatomical acuity

In this study, ganglion cell topography showed a complete and welldeveloped horizontal streak and no additional high-density areas outside the streak. The total cone topography had a less developed horizontal streak and a more pronounced area temporalis. The peak ganglion cell density was found in the temporal retina with an average of 38,643 cells mm⁻², resulting in an acuity estimate of 6.8 cpd. These estimates fit within the typical range of shallow water reef fishes, which have an anatomical estimate of visual acuity between 4 and 27 cpd (Collin and Pettigrew, 1989).

The well-defined streak pattern of *R. aculeatus* found in this study is very similar to the pattern found for another triggerfish, *Balistoides conspicillum* (Collin and Pettigrew, 1989). According to the 'terrain theory' (Hughes, 1977), the topography of cells across the retina represents the symmetry of the habitat. Triggerfishes live in open flat areas, where they scour the seafloor for potential prey items and have an uninterrupted view of the sand–water horizon; therefore, a horizontal streak may help them scan their environment for potential predators while searching for prey.

For cone densities and distribution, our results were similar to a previous study on *R. aculeatus* (Champ et al., 2014); however, they substantially differed for the ganglion cell analysis, especially in density and acuity estimates. Ganglion cell topography in Champ et al. (2014) revealed the presence of a partial horizontal streak that extends from the nasal to the central retina, as well as several other high-density areas outside the streak. A peak cell density of 12,450 cells mm⁻² was found in the temporal retina, resulting in an

acuity estimate of 3.4 cpd. These differences in topography pattern, ganglion cell density and acuity estimate between the two studies could be due to several factors such as differences in individual size, the methods of analysis used and/or issues encountered during sample preparation.

In teleost fishes, ganglion cell densities and topography patterns usually show very little intraspecific variability (e.g. de Busserolles et al., 2014b), except between individuals of different life stages and therefore sizes (Shand, 1997; Stieb et al., 2019; Tettamanti et al., 2019). In terms of cell densities, while larger individuals with larger eyes usually have a higher total number of ganglion cells in their retina, the number of ganglion cells per retinal area is usually smaller (Stieb et al., 2019; Tettamanti et al., 2019). According to this, the lower cell densities found in the previous study compared with ours could have been explained if they had used larger individuals. However, Champ et al. (2014) used similar sized individuals or slightly smaller ones compared with our study (7–12 cm versus 10–17 cm SL), indicating that size is not a likely explanation for the differences in densities observed between the two studies.

In both studies, displaced amacrine cells were included into the ganglion cell counts. Contrary to the present study, Champ et al. (2014) subsequently applied a multiplication factor of 0.76 to all their ganglion cell counts to account for the inclusion of displaced amacrine cells, which could explain some of the discrepancies in densities between the two studies. However, even if we take this correction into consideration in our study, our peak ganglion cell density is still at least twice what was reported in the previous study. We chose not to apply a correction factor in this study as displaced amacrine cells in coral reef fishes are mainly present in peripheral and non-specialised areas of the retina, and their inclusion in ganglion cell retinal topographic analyses usually has very little impact on the general topography pattern or the peak cell density estimation (Collin and Pettigrew, 1988). Consequently, even though the inclusion of amacrine cells in our study might slightly overestimate the peak density of ganglion cells and therefore acuity, we believe it provides a more accurate estimate than applying a general correction factor.

The most likely explanation for the differences in ganglion cell densities and topography pattern observed between the two studies is sample processing. During the preparation of our retinal wholemounts, we found that R. aculeatus had a very thick vitreous that proved challenging to remove, especially in the centre of the retina along the retinal meridian. During several attempts which were not included in this study, leftover vitreous resulted in very weak ganglion cell staining making counting unreliable or even unfeasible in certain areas. Similar challenges might have been encountered in Champ et al. (2014), resulting in an under-sampling of the ganglion cell population along the streak. This could explain why an incomplete streak and patchier topography pattern was found in that study, as well as why relatively similar ganglion cell densities (i.e. ~5000 cells mm⁻²) were found in the dorsal and ventral part of the retina in the two studies, but much lower densities were found along the streak in the previous study.

Behavioural measurements of achromatic and chromatic acuity

For achromatic gratings, we found that *R. aculeatus* had a behavioural acuity of 3.9 cpd, which was significantly lower than anatomical measurements of 6.8 cpd. For fish, anatomical measurements are often higher than those determined using behavioural experiments (e.g. Brokovich et al., 2010; Pankhurst

et al., 1993; Parker et al., 2017). Matthiessen's ratio (2.55), which is an average value from a measured range of 2.4–2.82, is used to estimate focal length instead of using the true focal length of the fish lens, and therefore may cause discrepancies in estimations. Visual acuity estimates based on ganglion cell counts only represent a theoretical upper limit of visual acuity. This is because not all ganglion cells contribute to visual acuity. For example, in primates, spatial resolving power is set by the midget ganglion cells, which only account for 70–80% of the total ganglion cell population (Wässle, 2004). Furthermore, the function of different types of retinal ganglion cells varies both temporally and spatially across the retina in zebrafish (Zhou et al., 2020).

Behavioural studies of visual acuity are likely to represent a more accurate estimate of an animal's functional visual abilities than its anatomical visual acuity. Furthermore, experiments that utilise ecologically relevant paradigms to measure acuity should give closer estimates between behavioural and anatomical measured thresholds. For example, Temple et al. (2013) found that the maximum acuity of archerfish was similar to anatomical measurements when fish spat at prey using the area centralis in the ventro-temporal region of the retina. In contrast, optomotor/ optokinetic tests provide some measure of average retinal acuity but will not capture the acuity in specialised regions of the retina, such as in the area centralis or fovea. Behavioural measurements of acuity may also vary depending on the stimuli used. Triggerfish did not perform as well in acuity tests when trained to circular stimuli as opposed to grating stimuli of horizontal and vertical stripes (Champ et al., 2014). Furthermore, different acuities were measured when honeybees were presented with radial (sectored) compared with linear (square-wave) gratings (Srinivasan and Lehrer, 1988), which could be explained by orientation-specific feature detecting receptive fields (Marr and Hildreth, 1980).

We found that achromatic acuity (3.94 cpd) of triggerfish was significantly higher than chromatic acuity for both the green–yellow (1.71 cpd) and pink–purple (1.89 cpd) stimuli. This supports previous findings in birds, mammals and bees (Giurfa et al., 1997; Lind and Kelber, 2011; Mullen, 1985); however, in humans and birds, differences in acuity were found between different colour channels. In budgerigars, visual acuity was lower for blue–green than for red–green gratings, which may be explained by lower numbers of SWS cones in the retina compared with MWS and LWS cones (Hart et al., 2000). Similarly in humans, acuity was lower for blue–yellow than for red–green contrasts (Mullen, 1985). However, the black–white achromatic stimulus had higher overall luminance compared with the chromatic stimuli based on double cone quantum catch; this may have also improved acuity for achromatic stimuli.

In conclusion, we hope that data from this study will inform future studies with this species investigating a range of topics including, but not limited to, teleost perception, navigation and cognition. We believe that Picasso triggerfish will continue to play a key role in contemporary research in these areas owing to their versatility as a focal study organism. We highlight the need for further information on the specific chromatic opponent mechanisms in animals, which is currently poorly understood (Baden and Osorio, 2019).

Acknowledgements

We would like to acknowledge the Turrbal, Jagera, Dingaal, Ngurrumungu and Thanhil people, the Traditional Owners and custodians of the lands on which the University of Queensland and Lizard Island Research station operate. We pay our respects to their ancestors and their descendants, who continue cultural and spiritual connections to Country and recognize their valuable contributions to Australian and global society. We thank the staff at the Lizard Island Research Station for support during field work and Cairns Marine for supplying fish. We would also like to acknowledge Dr Rumelo Amor from the QBI Advanced Microscopy Facility for technical support, and two anonymous reviewers and the editor for valuable feedback.

Competing interests

The authors declare no competing or financial interests.

Author contributions

Conceptualization: K.L.C., F.C.; Methodology: K.L.C., F.d.B., M.L., F.C.; Formal analysis: K.L.C., J.H., F.d.B., M.L., A.S., C.v.d.B., N.F.G., F.C.; Investigation: K.L.C., J.H., F.d.B., M.L., A.S., C.v.d.B., N.F.G., F.C.; Data curation: K.L.C., J.H., F.d.B., A.S., F.C.; Writing - original draft: K.L.C., J.H., F.d.B., M.L., A.S., F.C.; Writing - review & editing: K.L.C., F.d.B., M.L., A.S., C.v.d.B., N.J.M., F.C.; Visualization: K.L.C., F.d.B., M.L., A.S.; Supervision: K.L.C., N.J.M., F.C.; Project administration: K.L.C.; Funding acquisition: K.L.C., N.J.M., F.C.

Funding

This work was supported by the Australian Research Council (ARC) [Future Fellowship FT190100313 to K.L.C.; Discovery Early Career Researcher Awards: DE180100949 to F.d.B. and DE200100620 to F.C.; and Discovery Projects DP150102710 and DP180102363 to K.L.C., N.J.M. and F.C.]. The stereology analyses were performed at the QBI Advanced Microscopy Facility using Stereo Investigator, which was supported by an ARC LIEF grant (LE10010074). Open Access funding provided by The University of Queensland. Deposited in PMC for immediate release.

Data availability

All data supporting this manuscript including information for spectral sensitivity estimations, transcriptome assemblies, *R. aculeatus* opsin sequences and the raw data and R code for behavioural experiments are available on UQ eSpace: https://doi.org/10.48610/bb0f237. Genomic and transcriptomic raw-reads are available through GenBank (https://www.ncbi.nlm.nih.gov/genbank/) under the bioproject accession number: PRJNA747115 (Biosamples: SAMN26764115–SAMN26764120) and accession numbers

ON060692–ON060701 for single opsin gene sequences.

References

- Afgan, E., Sloggett, C., Goonasekera, N., Makunin, I., Benson, D., Crowe, M., Gladman, S., Kowsar, Y., Pheasant, M. and Horst, R. (2015). Genomics virtual laboratory: a practical bioinformatics workbench for the cloud. *PLoS ONE* 10, e0140829. doi:10.1371/journal.pone.0140829
- Baddeley, A. and Turner, R. (2005). spatstat: An R package for analyzing spatial point patterns. J. Stat. Softw. 12, 1-42. doi:10.18637/jss.v012.i06
- Baddeley, R., Osorio, D. and Jones, C. D. (2001). Colour generalisation by domestic chicks. *Behav. Brain Sci.* 24, 654-654. doi:10.1017/ S0140525X01210085
- Baden, T. (2021). Circuit mechanisms for colour vision in zebrafish. *Curr. Biol.* **31**, R807-R820. doi:10.1016/j.cub.2021.04.053
- Baden, T. and Osorio, D. (2019). The retinal basis of vertebrate color vision. Annu. Rev. Vis. Sci. 5, 177-200. doi:10.1146/annurev-vision-091718-014926
- Barthel, L. K. and Raymond, P. A. (2000). In situ hybridization studies of retinal neurons. *Methods Enzymol.* 316, 579-590. doi:10.1016/S0076-6879(00)16751-5
- Bates, D., Machler, M., Bolker, B. M. and Walker, S. C. (2015). Fitting linear mixedeffects models using Ime4. J. Stat. Softw. 67, 1-48. doi:10.18637/jss.v067.i01
- Bolger, A. M., Lohse, M. and Usadel, B. (2014). Trimmomatic: a flexible trimmer for Illumina sequence data. *Bioinformatics* 30, 2114-2120. doi:10.1093/ bioinformatics/btu170
- Brokovich, E., Ben-Ari, T., Kark, S., Kiflawi, M., Dishon, G., Iluz, D. and Shashar, N. (2010). Functional changes of the visual system of the damselfish *Dascyllus marginatus* along its bathymetric range. *Physiol. Behav.* **101**, 413-421. doi:10.1016/j.physbeh.2010.07.006
- Carleton, K. L., Escobar-Camacho, D., Stieb, S. M., Cortesi, F. and Marshall, N. J. (2020). Seeing the rainbow: mechanisms underlying spectral sensitivity in teleost fishes. J. Exp. Biol. 223, jeb193334. doi:10.1242/jeb.193334
- Castiglione, G. M. and Chang, B. S. W. (2018). Functional trade-offs and environmental variation shaped ancient trajectories in the evolution of dim-light vision. *Elife* 7, e35957. doi:10.7554/eLife.35957
- Caves, E. M., Green, P. A., Zipple, M. N., Peters, S., Johnsen, S. and Nowicki, S. (2018). Categorical perception of colour signals in a songbird. *Nature* 560, 365-367. doi:10.1038/s41586-018-0377-7
- Champ, C., Wallis, G., Vorobyev, M., Siebeck, U. and Marshall, J. (2014). Visual acuity in a species of coral reef fish: *Rhinecanthus aculeatus. Brain Behav. Evol.* 83, 31-42. doi:10.1159/000356977
- Champ, C. M., Vorobyev, M. and Marshall, N. J. (2016). Colour thresholds in a coral reef fish. *R. Soc. Open Sci.* **3**, 160399. doi:10.1098/rsos.160399

- Chan, T., Lee, M. and Sakmar, T. P. (1992). Introduction of hydroxyl-bearing amino acids causes bathochromic spectral shifts in rhodopsin. Amino acid substitutions responsible for red-green color pigment spectral tuning. J. Biol. Chem. 267, 9478-9480.
- Cheney, K. L., Newport, C., McClure, E. C. and Marshall, N. J. (2013). Colour vision and response bias in a coral reef fish. J. Exp. Biol. 216, 2967-2973.
- Cheney, K. L., Green, N. F., Vibert, A. P., Vorobyev, M., Marshall, N. J., Osorio, D. C. and Endler, J. A. (2019). An Ishihara-style test of animal colour vision. J. Exp. Biol. 222, jeb189787. doi:10.1242/jeb.189787
- Chinen, A., Matsumoto, Y. and Kawamura, S. (2005). Reconstitution of ancestral green visual pigments of zebrafish and molecular mechanism of their spectral differentiation. *Mol. Biol. Evol.* 22, 1001-1010. doi:10.1093/molbev/msi086
- Coimbra, J. P., Marceliano, M. L., Andrade-da-Costa, B. L. and Yamada, E. S. (2006). The retina of tyrant flycatchers: topographic organization of neuronal density and size in the ganglion cell layer of the great kiskadee *Pitangus sulphuratus* and the rusty margined flycatcher *Myiozetetes cayanensis* (Aves: Tyrannidae). *Brain Behav. Evol.* 68, 15-25. doi:10.1159/000092310
- Coimbra, J. P., Trevia, N., Marceliano, M. L., da Silveira Andrade-Da-Costa, B. L., Picanco-Diniz, C. W. and Yamada, E. S. (2009). Number and distribution of neurons in the retinal ganglion cell layer in relation to foraging behaviors of tyrant flycatchers. *J. Comp. Neurol.* **514**, 66-73. doi:10.1002/cne. 21992
- Coimbra, J. P., Nolan, P. M., Collin, S. P. and Hart, N. S. (2012). Retinal ganglion cell topography and spatial resolving power in penguins. *Brain Behav. Evol.* 80, 254-268. doi:10.1159/000341901
- Collin, S. P. and Collin, H. B. (1988). Topographic analysis of the retinal ganglion cell layer and optic nerve in the sandlance *Limnichthyes fasciatus* (Creeiidae, Perciformes). J. Comp. Neurol. 278, 226-241. doi:10.1002/cne.902780206
- Collin, S. P. and Pettigrew, J. D. (1988). Retinal ganglion cell topography in teleosts: a comparison between nissl-stained material and retrograde labelling from the optic nerve. J. Comp. Neurol. 276, 412-422. doi:10.1002/cne.902760306
- Collin, S. P. and Pettigrew, J. D. (1989). Quantitative comparison of the limits on visual spatial resolution set by the ganglion cell layer in twelve species of reef teleosts. *Brain Behav. Evol.* 34, 184-192. doi:10.1159/000116504
- Collin, S. P., Knight, M. A., Davies, W. L., Potter, I. C., Hunt, D. M. and Trezise, A. E. (2003). Ancient colour vision: multiple opsin genes in the ancestral vertebrates. *Curr. Biol.* **13**, R864-R865. doi:10.1016/j.cub.2003.10.044
- Cortesi, F., Mitchell, L. J., Tettamanti, V., Fogg, L. G., de Busserolles, F., Cheney, K. L. and Marshall, N. J. (2020). Visual system diversity in coral reef fishes. Semin. Cell Dev. Biol. 106, 31-42. doi:10.1016/j.semcdb.2020.06.007
- Dalton, B. E., Loew, E. R., Cronin, T. W. and Carleton, K. L. (2014). Spectral tuning by opsin coexpression in retinal regions that view different parts of the visual field. *Proc. R. Soc. B Biol. Sci.* 281, 20141980.
- Dalton, B. E., Lu, J., Leips, J., Cronin, T. W. and Carleton, K. L. (2015). Variable light environments induce plastic spectral tuning by regional opsin coexpression in the African cichlid fish, *Metriaclima zebra. Mol. Ecol.* 24, 4193-4204. doi:10.1111/ mec.13312
- de Busserolles, F., Marshall, N. J. and Collin, S. P. (2014a). The eyes of lanternfishes (Myctophidae, Teleostei): novel ocular specializations for vision in dim light. J. Comp. Neurol. 522, 1618-1640. doi:10.1002/cne.23495
- de Busserolles, F., Marshall, N. J. and Collin, S. P. (2014b). Retinal ganglion cell distribution and spatial resolving power in deep-sea lanternfishes (Myctophidae). *Brain Behav. Evol.* 84, 262-276. doi:10.1159/000365960
- de Busserolles, F., Cortesi, F., Helvik, J. V., Davies, W. I. L., Templin, R. M., Sullivan, R. K. P., Michell, C. T., Mountford, J. K., Collin, S. P., Irigoien, X. et al. (2017). Pushing the limits of photoreception in twilight conditions: The rod-like cone retina of the deep-sea pearlsides. *Sci. Adv.* **3**, eaao4709. doi:10.1126/ sciadv.aao4709
- de Ibarra, N. H., Giurfa, M., Vorobyev, M. (2002). Discrimination of coloured patterns by honeybees through chromatic and achromatic cues. J. Comp. Physiol. A 188, 503-512. doi:10.1007/s00359-002-0322-x
- Dungan, S. Z., Kosyakov, A. and Chang, B. S. W. (2016). Spectral tuning of killer whale (Orcinus orca) rhodopsin: evidence for positive selection and functional adaptation in a cetacean visual pigment. *Mol. Biol. Evol.* 33, 323-336. doi:10.1093/ molbev/msv217
- Escobar-Camacho, D., Taylor, M. A., Cheney, K. L., Green, N. F., Marshall, N. J. and Carleton, K. L. (2019). Color discrimination thresholds in a cichlid fish: *Metriaclima benetos. J. Exp. Biol.* 222, jeb201160. doi:10.1242/jeb.201160
- Fasick, J. I. and Robinson, P. R. (1998). Mechanism of spectral tuning in the dolphin visual pigments. *Biochemistry* 37, 433-438. doi:10.1021/bi972500j
- Garza-Gisholt, E., Hemmi, J. M., Hart, N. S. and Collin, S. P. (2014). A comparison of spatial analysis methods for the construction of topographic maps of retinal cell density. *PLoS One* 9, e93485. doi:10.1371/journal.pone.0093485
- Giurfa, M., Vorobyev, M., Brandt, R., Posner, B. and Menzel, R. (1997). Discrimination of coloured stimuli by honeybees: Alternative use of achromatic and chromatic signals. J. Comp. Physiol. A 180, 235-243. doi:10.1007/ s003590050044
- Glaser, E. M. and Wilson, P. D. (1998). The coefficient of error of optical fractionator population size estimates: a computer simulation comparing three estimators. *J. Microsc. Oxford* **192**, 163-171. doi:10.1046/j.1365-2818.1998.00417.x

- Haas, B. J., Papanicolaou, A., Yassour, M., Grabherr, M., Blood, P. D., Bowden, J., Couger, M. B., Eccles, D., Li, B. and Lieber, M. (2013). De novo transcript sequence reconstruction from RNA-seq using the Trinity platform for reference generation and analysis. *Nat. Protoc.* 8, 1494. doi:10.1038/nprot.2013. 084
- Hart, N. S., Partridge, J. C., Cuthill, I. C. and Bennett, A. T. D. (2000). Visual pigments, oil droplets, ocular media and cone photoreceptor distribution in two species of passerine bird: the blue tit (*Parus caeruleus* L.) and the blackbird (*Turdus merula* L.). J. Comp. Physiol. A Neuroethol. Sens. Neural Behav. Physiol. 186, 375-387. doi:10.1007/s003590050437
- Hu, T. J., Wang, G. M., Shen, L. C. and Li, F. (2006). Bionic inspirations of fish-like robots from *Rhinecanthus aculeatus*. IEEE ICMA 2006: Proceedings of the 2006 IEEE International Conference on Mechatronics and Automation, Vol. 1-3, 639. IEEE.
- Hughes, A. (1975). A quantitative analysis of the cat retinal ganglion cell topography. J. Comp. Neurol. 163, 107-128. doi:10.1002/cne.901630107
- Hughes, A. (1977). The topography of vision in mammals. In *The Visual System in Vertebrates: Handbook of Sensory Physiology*, 7/5 (ed. F. Crescitelli), pp. 613-756. Berlin: Springer.
- Jones, C. D., Osorio, D. and Baddeley, R. J. (2001). Colour categorization by domestic chicks. P Roy Soc B-Biol Sci 268, 2077-2084. doi:10.1098/rspb.2001. 1734
- Karlsson, C., Williis, J. K., Patel, M. and Perera, B. d. (2019). Teleost fish can accurately estimate distance travelled. *bioRxiv*.
- Katoh, K. and Standley, D. M. (2013). MAFFT multiple sequence alignment software version 7: improvements in performance and usability. *Mol. Biol. Evol.* 30, 772-780.
- Kelber, A., Vorobyev, M. and Osorio, D. (2003). Animal colour vision behavioural tests and physiological concepts. *Biol. Rev.* 78, 81-118. doi:10.1017/ S1464793102005985
- Kuznetsova, A., Brockhoff, P. B. and Christensen, R. H. B. (2017). ImerTest package: tests in linear mixed effects models. J. Stat. Softw. 82, 1-26. doi:10.18637/jss.v082.i13
- Linares, D. and Lopez-Moliner, J. (2016). quickpsy: An R package to fit psychometric functions for multiple groups. *R J.* 8, 122-131. doi:10.32614/RJ-2016-008
- Lind, O. and Kelber, A. (2011). The spatial tuning of achromatic and chromatic vision in budgerigars. *J. Vis.* **11**, 2. doi:10.1167/11.7.2
- Luehrmann, M., Stieb, S. M., Carleton, K. L., Pietzker, A., Cheney, K. L. and Marshall, N. J. (2018). Short-term colour vision plasticity on the reef: changes in opsin expression under varying light conditions differ between ecologically distinct fish species. J. Exp. Biol. 221, jeb175281. doi:10.1242/jeb.175281
- Marr, D. and Hildreth, E. (1980). Theory of edge detection. *Proc. R. Soc. B* 207, 187-217. doi:10.1098/rspb.1980.0020
- Matchette, S. R., Cuthill, I. C., Cheney, K. L., Marshall, N. J. and Scott-Samuel, N. E. (2020). Underwater caustics disrupt prey detection by a reef fish. *Proc. R. Soc. B Biol. Sci.* 287, 20192453. doi:10.1098/rspb.2019.2453
- Matsumoto, Y., Fukamachi, S., Mitani, H. and Kawamura, S. (2006). Functional characterization of visual opsin repertoire in Medaka (*Oryzias latipes*). Gene 371, 268-278. doi:10.1016/j.gene.2005.12.005
- Matthiessen, L. (1882). Ueber die beziehungen, welche zwischen dem brechungsindex des kerncentrums der krystalllinse und den dimensionen des auges bestehen. *Pflugers Arch.* 27, 510-523. doi:10.1007/BF01802978
- Mitchell, L., Cheney, K. L., Cortesi, F., Marshall, N. J. and Vorobyev, M. (2017). Triggerfish uses chromaticity and lightness for object segregation. *R. Soc. Open Sci.* 4, 171440. doi:10.1098/rsos.171440
- Mullen, K. T. (1985). The contrast sensitivity of human color vision to red green and blue yellow chromatic gratings. J. Physiol. Lond. 359, 381-400. doi:10.1113/ jphysiol.1985.sp015591
- Musilova, Z. and Cortesi, F. (2021). Multiple ancestral and a plethora of recent gene duplications during the evolution of the green sensitive opsin genes (RH2) in teleost fishes. *bioRxiv*.
- Musilova, Z., Cortesi, F., Matschiner, M., Davies, W. I. L., Patel, J. S., Stieb, S. M., de Busserolles, F., Malmstrom, M., Torresen, O. K., Brown, C. J. et al. (2019). Vision using multiple distinct rod opsins in deep-sea fishes. *Science* 364, 588-592. doi:10.1126/science.aav4632
- Musilova, Z., Salzburger, W. and Cortesi, F. (2021). The visual opsin gene repertoires of teleost fishes: evolution, ecology, and function. *Annu. Rev. Cell Dev. Biol.* 37, 21.1-21.28. doi:10.1146/annurev-cellbio-120219-024915
- Nandamuri, S. P., Yourick, M. R. and Carleton, K. L. (2017). Adult plasticity in African cichlids: rapid changes in opsin expression in response to environmental light differences. *Mol. Ecol.* 26, 6036-6052. doi:10.1111/mec.14357
- Neumeyer, C. (1985). An ultraviolet receptor as a 4th receptor type in goldfish color vision. *Naturwissenschaften* 72, 162-163. doi:10.1007/BF00490410
- Neumeyer, C. (1986). Wavelength discrimination in the goldfish. J. Comp. Physiol. A 158, 203-213. doi:10.1007/BF01338563
- Neumeyer, C. (1992). Tetrachromatic color vision in goldfish evidence from color mixture experiments. J. Comp. Physiol. A 171, 639-649. doi:10.1007/ BF00194111

- Olsson, P., Wilby, D. and Kelber, A. (2016). Quantitative studies of animal colour constancy: using the chicken as model. Proc. R. Soc. B Biol. Sci. 283, 20160411.
- Palczewski, K., Kumasaka, T., Hori, T., Behnke, C. A., Motoshima, H., Fox, B. A., Trong, I. L., Teller, D. C., Okada, T., Stenkamp, R. E. et al. (2000). Crystal structure of rhodopsin: a G protein-coupled receptor. *Science* 289, 739-745. doi:10.1126/science.289.5480.739
- Pankhurst, P. M., Pankhurst, N. W. and Montgomery, J. C. (1993). Comparison of behavioral and morphological measures of visual acuity during ontogeny in a teleost fish, *Forsterygion varium*, Tripterygiidae (Forster, 1801). *Brain Behav. Evol.* 42, 178-188. doi:10.1159/000114151
- Parker, A. N., Fritsches, K. A., Newport, C., Wallis, G. and Siebeck, U. E. (2017). Comparison of functional and anatomical estimations of visual acuity in two species of coral reef fish. J. Exp. Biol. 220, 2387-2396.
- Parmentier, E., Raick, X., Lecchini, D., Boyle, K., Van Wassenbergh, S., Bertucci, F. and Kever, L. (2017). Unusual sound production mechanism in the triggerfish *Rhinecanthus aculeatus* (Balistidae). *J. Exp. Biol.* 220, 186-193. doi:10.1242/jeb.157263
- Pignatelli, V., Champ, C., Marshall, J. and Vorobyev, M. (2010). Double cones are used for colour discrimination in the reef fish, *Rhinecanthus aculeatus*. *Biol. Letters* 6, 537-539. doi:10.1098/rsbl.2009.1010
- Randall, J. E., Allen, G. R. and Steene, R. C. (1997). Fishes of the Great Barrier Reef and Coral Sea. University of Hawaii Press.
- Risner, M. L., Lemerise, E., Vukmanic, E. V. and Moore, A. (2006). Behavioral spectral sensitivity of the zebrafish (*Danio rerio*). Vision Res. 46, 2625-2635. doi:10.1016/j.visres.2005.12.014
- Ronquist, F., Teslenko, M., van der Mark, P., Ayres, D. L., Darling, A., Höhna, S., Larget, B., Liu, L., Suchard, M. A. and Huelsenbeck, J. P. (2012). MrBayes 3.2: efficient Bayesian phylogenetic inference and model choice across a large model space. Syst. Biol. 61, 539-542. doi:10.1093/sysbio/sys029
- Shand, J. (1997). Ontogenetic changes in retinal structure and visual acuity: a comparative study of coral-reef teleosts with differing post-settlement lifestyles. *Environ. Biol. Fish.* 49, 307-322. doi:10.1023/A:1007353003066
- Shi, Y. S., Radlwimmer, F. B. and Yokoyama, S. (2001). Molecular genetics and the evolution of ultraviolet vision in vertebrates. *Proc. Natl. Acad. Sci. USA* 98, 11731-11736. doi:10.1073/pnas.201257398
- Sibeaux, A., Cole, G. L. and Endler, J. A. (2019). Success of the receptor noise model in predicting colour discrimination in guppies depends upon the colours tested. *Vision Res.* 159, 86-95. doi:10.1016/j.visres.2019.04.002
- Siebeck, U. and Marshall, N. J. (2001). Ocular media transmission of coral reef fish - Can coral reef fish see ultraviolet light? Vision Res. 41, 133-149. doi:10.1016/ S0042-6989(00)00240-6
- Siebeck, U. E., Wallis, G. M. and Litherland, L. (2008). Colour vision in coral reef fish. *J. Exp. Biol.* 211, 354-360. doi:10.1242/jeb.012880
- Silvasti, S. A., Valkonen, J. K. and Nokelainen, O. (2021). Behavioural thresholds of blue tit colour vision and the effect of background chromatic complexity. *Vision Res.* **182**, 46-57. doi:10.1016/j.visres.2020.11.013
- Simpson, E. E., Marshall, N. J. and Cheney, K. L. (2016). Coral reef fish perceive lightness illusions. *Sci. Rep.* 6, 35335. doi:10.1038/srep35335
- Spady, T. C., Parry, J. W., Robinson, P. R., Hunt, D. M., Bowmaker, J. K. and Carleton, K. L. (2006). Evolution of the cichlid visual palette through ontogenetic subfunctionalization of the opsin gene arrays. *Mol. Biol. Evol.* 23, 1538-1547. doi:10.1093/molbev/msl014
- Srinivasan, M. V. and Lehrer, M. (1988). Spatial acuity of honeybee vision and its spectral properties. J. Comp. Physiol. A 162, 159-172. doi:10.1007/BF00606081
- Stieb, S. M., de Busserolles, F., Carleton, K. L., Cortesi, F., Chung, W.-S., Dalton, B. E., Hammond, L. A. and Marshall, N. J. (2019). A detailed investigation of the visual system and visual ecology of the Barrier Reef anemonefish, *Amphiprion akindynos. Sci. Rep.* 9, 16459. doi:10.1038/s41598-019-52297-0
- Stoddard, M. C., Eyster, H. N., Hogan, B. G., Morris, D. H., Soucy, E. R. and Inouye, D. W. (2020). Wild hummingbirds discriminate nonspectral colors. *Proc. Natl. Acad. Sci. USA* **117**, 15112-15122. doi:10.1073/pnas.1919377117
- Stone, J., Rapaport, D. H., Williams, R. W. and Chalupa, L. (1981). Uniformity of cell distribution in the ganglion cell layer of prenatal cat retina: implications for mechanisms of retinal development. *Brain Res.* 254, 231-242. doi:10.1016/0165-3806(81)90034-1
- Takahashi, Y. and Ebrey, T. G. (2003). Molecular basis of spectral tuning in the newt short wavelength sensitive visual pigment. *Biochemistry* 42, 6025-6034. doi:10.1021/bi020629+
- Takenaka, N. and Yokoyama, S. (2007). Mechanisms of spectral tuning in the RH2 pigments of Tokay gecko and American chameleon. *Gene* **399**, 26-32. doi:10.1016/j.gene.2007.04.036
- Temple, S. E., Manietta, D. and Collin, S. P. (2013). A comparison of behavioural (Landolt C) and anatomical estimates of visual acuity in archerfish (*Toxotes chatareus*). Vision Res. 83, 1-8. doi:10.1016/j.visres.2013.02.014
- Tettamanti, V., de Busserolles, F., Lecchini, D., Marshall, N. J. and Cortesi, F. (2019). Visual system development of the spotted unicomfish, *Naso brevirostris* (Acanthuridae). *J. Exp. Biol.* **222**, jeb209916. doi:10.1242/jeb.209916
- Torres-Dowdall, J., Pierotti, M. E. R., Harer, A., Karagic, N., Woltering, J. M., Henning, F., Elmer, K. R. and Meyer, A. (2017). Rapid and parallel adaptive

evolution of the visual system of neotropical Midas cichlid fishes. *Mol. Biol. Evol.* **34**, 2469-2485. doi:10.1093/molbev/msx143

- Ullmann, J. F., Moore, B. A., Temple, S. E., Fernandez-Juricic, E. and Collin, S. P. (2012). The retinal wholemount technique: a window to understanding the brain and behaviour. *Brain Behav. Evol.* **79**, 26-44. doi:10.1159/000332802
- van den Berg, C. P., Hollenkamp, M., Mitchell, L. J., Watson, E. J., Green, N. F., Marshall, N. J. and Cheney, K. L. (2020). More than noise: context-dependent luminance contrast discrimination in a coral reef fish (*Rhinecanthus aculeatus*). *J. Exp. Biol.* 223, jeb232090. doi:10.1242/jeb.232090
- von Frisch, K. (1914). Der Farbensinn und Formensinn der Biene. Zool. Jb. Physiol. 37, 1-238.
- Vorobyev, M. and Osorio, D. (1998). Receptor noise as a determinant of colour thresholds. *Proc. R. Soc. B Biol. Sci.* 265, 351-358. doi:10.1098/rspb.1998.0302
 Wässle, H. (2004). Parallel processing in the mammalian retina. *Nat. Rev. Neurosci.* 5, 747-757. doi:10.1038/nrn1497
- West, R. A. (1991). Optical properties of aggregate particles whose outer diameter Is comparable to the wavelength. *Appl. Optics* 30, 5316-5324. doi:10.1364/AO.30. 005316
- Yokoyama, S. (2000). Molecular evolution of vertebrate visual pigments. Prog. Retin. Eve Res. 19, 385-419. doi:10.1016/S1350-9462(00)00002-1
- Yokoyama, S. (2008). Evolution of dim-light and color vision pigments. Annu. Rev. Genomics Hum. Genet. 9, 259-282. doi:10.1146/annurev.genom.9.081307.164228

- Yokoyama, S. and Jia, H. (2020). Origin and adaptation of green-sensitive (RH2) pigments in vertebrates. *FEBS Open Biol.* **10**, 873-882. doi:10.1002/2211-5463. 12843
- Yokoyama, S. and Radlwimmer, B. (1998). The "five-sites" rule and the evolution of red and green color vision in mammals. *Mol. Biol. Evol.* **15**, 560-567. doi:10.1093/oxfordjournals.molbev.a025956
- Yokoyama, S. and Tada, T. (2003). The spectral tuning in the short wavelength-sensitive type 2 pigments. *Gene* **306**, 91-98. doi:10.1016/S0378-1119(03)00424-4
- Yokoyama, S., Takenaka, N. and Blow, N. (2007). A novel spectral tuning in the short wavelength-sensitive (SWS1 and SWS2) pigments of bluefin killifish (*Lucania goodei*). *Gene* **396**, 196-202. doi:10.1016/j.gene.2007.03.019
- Yourick, M. R., Sandkam, B. A., Gammerdinger, W. J., Escobar–Camacho, D., Nandamuri, S. P., Clark, F. E., Joyce, B., Conte, M. A., Kocher, T. D. and Carleton, K. L. (2019). Diurnal variation in opsin expression and common housekeeping genes necessitates comprehensive normalization methods for quantitative real-time PCR analyses. *Molecular Ecology Resources* 19, 1447-1460. doi:10.1111/1755-0998.13062
- Zhou, M., Bear, J., Roberts, P. A., Janiak, F. K., Semmelhack, J., Yoshimatsu, T. and Baden, T. (2020). Zebrafish retinal ganglion cells asymmetrically encode spectral and temporal information across visual space. *Curr. Biol.* 30, 2927-2942.e7. doi:10.1016/j.cub.2020.05.055

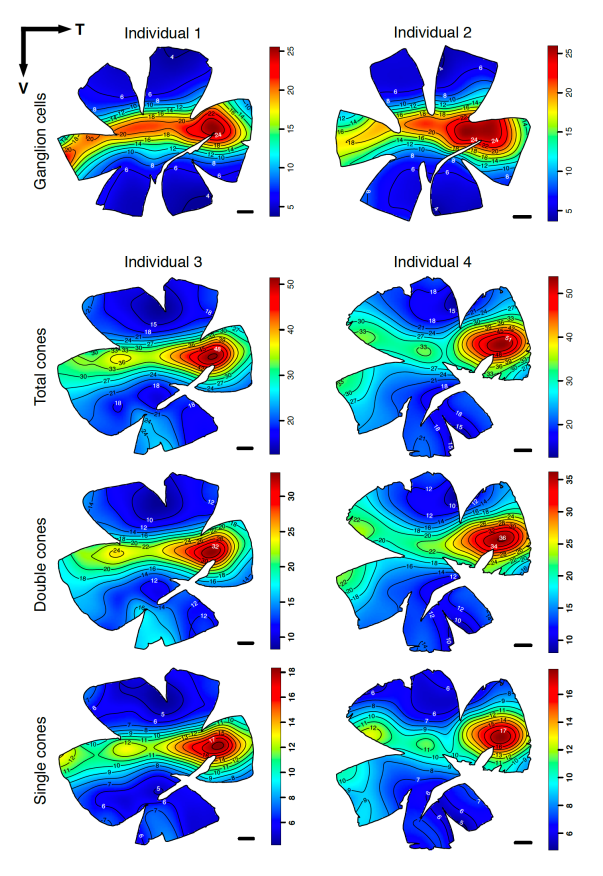


Fig. S1. Intraspecific variability topographic distribution of ganglion cells and cone photoreceptors (single, double and total cones) in *Rhinecanthus aculeatus*. The black lines represent iso-density contours and values are expressed in densities x 10^3 cells/mm². The black arrow indicates the orientation of the retinas. T = temporal, V = ventral. Scale bars: 1 mm.



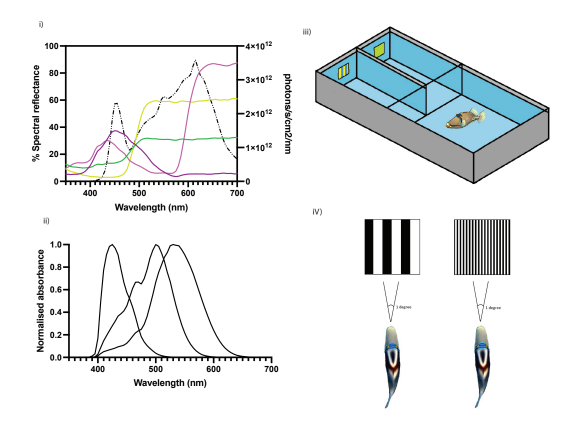


Fig. S2. i) Spectral reflectance of laminated stimuli used in each treatment labelled by color: pink, purple, green and yellow; irradiance of sidewelling lights measured in the testing tanks shown as dashed line; ii) Spectral sensitivities of the three cones in triggerfish, *Rhinecanthus aculeatus*, with a single cone sensitivity of $\lambda_{max} = 413$ nm, and double cone sensitivities of $\lambda_{max} = 480$ nm and $\lambda = 528_{max}$ nm multiplied by yellow cornea (Cheney et al., 2013); iii) Experimental tank setup; iv) An example of the achromatic (black and white) square-wave gratings: a) 1 cycle per degree; b) 5 cycles per degree.

Table S1. Primers used for probe template synthesis of fluorescence in-situ hybridisation. T7 (forward primer) and T3 (reverse primer) RNA polymerase promoter sequences are shown in bold.

Target	Primer	Sequence
gana		
gene		
RH2A	RH2A_F1	5'-TAATACGACTCACTATAGGGATGTACAAGCTCCTGGCTTG-3'
	RH2A_R1	5'-AATTAACCCTCACTAAAGGGTCTTTGCAAAGAAGGCGCAC-3'
RH2C-	RH2C_F1	5'-TAATACGACTCACTATAGGGCCATCAACTTCCTGACGCTA-3'
1 &	RH2C_R1	5'-AATTAACCCTCACTAAAGGGAGCCAAACACCATCAGGACA-3'
RH2C-		
2		
SWS2B	SWS2B_F1	5'-TAATACGACTCACTATAGGGAGAGGGCCCCAGATGACTTCT-3'
	SWS2B_R1	5'-AATTAACCCTCACTAAAGGGACGGACTGACTCGATGAGGA-3'

			Mapping						Proportional opsin expression % (normalized to coding sequence length)						
			Rods	Single cones (SC)		Double cones (DC)			Rod vs Cone SC		DC				
Origin	ID	# filtered transcriptome reads	<i>RH1</i> # reads	<i>SWS2B</i> # reads	<i>RH2A</i> # reads	<i>RH2C-1</i> # reads	<i>RH2C-1</i> # reads	<i>LWS</i> # reads	R	С	SWS2B	RH2A	RH2C-1	RH2C-2	LWS
Field	FZ9	10736408	14826	560	1488	1037	1901	44	74.62	25.38	100	33.29	15.65	50.08	0.97
	FZ10	9975040	23638	1390	1876	1894	2376	0	75.77	24.23	100	30.52	26.33	43.15	0.00
	FZ11	6864530	15196	798	654	458	1170	14	83.04	16.96	100	28.49	22.01	48.90	0.60
	1LIT	22709442	136187	18897	26723	3429	22709	3014	79.07	20.93	100	47.86	5.38	41.44	5.32
	Mean	12571355	47696.50	5411.25	7685.25	1704.50	8751.67	768.00	78.13	21.87	100	35.04	17.34	45.89	1.72
	Se	3481577.01	29879.20	4498.64	6351.03	646.09	5229.22	748.72	1.89	1.89	-	4.39	4.55	2.12	1.22
	LY	21584148	273003	40177	82429	18303	35983	52	60.61	39.39	100	60.27	11.87	27.83	0.04
	SQ	21581834	272743	34687	79679	28871	34329	212	60.47	39.53	100	55.69	6.19	37.97	0.15
Aquarium	PE	24905208	305271	40809	81017	20063	26989	124	64.30	35.70	100	63.20	14.43	22.28	0.10
	NE	20168834	387049	66207	168363	22443	79481	1170	53.34	46.66	100	62.03	16.11	21.44	0.43
	Mean	22060006	309516.50	45470.00	102872	22420	44195.50	389.50	59.68	40.32	100	60.30	17.34	45.89	0.18
	<i>S.e</i> .	1005269.25	26948.78	7047.68	21837.55	2311.58	11923.10	262.22	2.29	2.29	-	1.65	2.17	3.81	0.09

Table S3. Predicted *R. aculeatus* visual pigment peak spectral sensitivities (λ_{max}) compared to reference visual pigments (*O. latipes* RH1; *Oreochromis niloticus*, SWS2B, RH2B, RH2Aalpha, LWS), *R. aculeatus* λ_{max} determined via MSP, and tuning sites and effects considered for predictions. ¹ (Matsumoto et al., 2006), ²(Spady et al., 2006), ³(Dungan et al., 2016), ⁴(Fasick and Robinson, 1998), ⁵(Yokoyama et al., 2007), ⁶(Yokoyama and Tada, 2003), ⁷(Chinen et al., 2005), ⁸(Yokoyama and Jia, 2020), ⁹(Cheney et al., 2013)

	RH1	SWS2B	RH2C-1	RH2C-2	RH2A	LWS
Similarity to reference	94.1	86.1	85.5	85.5	91.5	89.9
amino acid sequence	-					
(%)						
Total variable amino	21	49	51	51	30	36
acid						
Variable amino acid	15	32	30	29	18	19
in transmembrane						
regions						
Variable amino acids	1	5	12	11	8	0
at known tuning sites						
Reference pigment	502 ¹	425 ²	47	2^{2}	528 ²	560^{2}
peak absorbance (nm						
λ_{max})						
Known tuning sites	S299	F46V (+8) ⁵	M88C (+3) ⁷	M88C (+3) ⁷	C88A (-3) ⁷	-
and applied tuning	A (-	A109G $(-2)^5$	I112V (+1) ⁷	$I112V (+1)^7$	$I112V (+1)^7$	
effects (nm)	$2)^{3,4}$	G164A	$T266V (-2)^7$			
		$(+1)^{6}$				
		W265T (-				
		29) ⁵				
Candidate tuning sites	S166	C163F	C98A	C98A	A151T	-
– no effects	А	S166F	V185C	V185C		
documented		S168A				
Predicted peak	500	403	474	476	526	560
absorbance (nm λ_{max})						
MSP peak absorbance	498 ⁹	413 ⁹	48	80 ⁹	528 ⁹	-
$(nm \lambda_{max})$						

Table S4. Summary of stimuli presented to each fish and in which order, whether the fish was trained to receive a food reward from the Distractor (11 cpd) or the test gratings (0.5-6 cpd), the total number of trials conducted by each fish for each colour (total number of trials = 2438) and calculated discrimination thresholds at 62% correct choice. NA indicates not tested due to time taken to complete treatment 1.

Fish ID	Size (SL, cm)	Treatment 1	S +ve	Threshold (cpd)	Treatment 2	S +ve	Threshold (cpd)
Billy	16	Green- yellow (179)	Control (11 cpd)	2.19	NA	NA	NA
Bitey	10	Pink-purple (120)	Grating	2.45	Green- yellow (167)	Grating	2.17
Diego	16	Achromatic (203)	Grating	4.89	Pink-purple (75)	Grating	2.73
Ernie	17	Achromatic (205)	Control (11 cpd)	5.30	NA	NA	NA
Gilbert	16.5	Green- yellow (184)	Grating	2.71	Achromatic (235)	Grating	5.03
Lyra	10	Pink-purple (120)	Grating	2.95	Green- yellow (167)	Grating	3.04
Mike	15	Green- yellow (188)	Grating	2.58	Pink-purple (167)	Grating	3.27
Sophie	15	Pink-purple (193)	Control (11 cpd)	5.44	Achromatic (235)	Control (11 cpd)	5.44

Further references

Cheney, K. L., Newport, C., McClure, E. C. and Marshall, N. J. (2013). Colour vision and response bias in a coral reef fish. *J Exp Biol* **216**, 2967-2973.

Chinen, A., Matsumoto, Y. and Kawamura, S. (2005). Reconstitution of ancestral green visual pigments of zebrafish and molecular mechanism of their spectral differentiation. *Mol Biol Evol* **22**, 1001-10.

Dungan, S. Z., Kosyakov, A. and Chang, B. S. W. (2016). Spectral tuning of killer whale (orcinus orca) rhodopsin: Evidence for positive selection and functional adaptation in a cetacean visual pigment. *Mol Biol Evol* **33**, 323-336.

Fasick, J. I. and Robinson, P. R. (1998). Mechanism of spectral tuning in the dolphin visual pigments. *Biochemistry* **37**, 433-438.

Matsumoto, Y., Fukamachi, S., Mitani, H. and Kawamura, S. (2006). Functional characterization of visual opsin repertoire in medaka (oryzias latipes). *Gene* **371**, 268-278.

Spady, T. C., Parry, J. W., Robinson, P. R., Hunt, D. M., Bowmaker, J. K. and Carleton, K. L. (2006). Evolution of the cichlid visual palette through ontogenetic subfunctionalization of the opsin gene arrays. *Mol Biol Evol* **23**, 1538-47.

Yokoyama, S. and Jia, H. (2020). Origin and adaptation of green-sensitive (rh2) pigments in vertebrates. *FEBS Open Bio* **10**, 873-882.

Yokoyama, S. and Tada, T. (2003). The spectral tuning in the short wavelengthsensitive type 2 pigments. *Gene* **306**.

Yokoyama, S., Takenaka, N. and Blow, N. (2007). A novel spectral tuning in the short wavelength-sensitive (sws1 and sws2) pigments of bluefin killifish (lucania goodei). *Gene* **396**, 196-202.