

Retinal Topography of Ganglion Cells and Putative UV-Sensitive Cones in Two Antarctic Fishes: Pagothenia borchgrevinki and Trematomus bernacchii (Nototheniidae)

Authors: Miyazaki, Taeko, Iwami, Tetsuo, Somiya, Hiroaki, and Meyer-Rochow, V. Benno

Source: Zoological Science, 19(11): 1223-1229

Published By: Zoological Society of Japan

URL: https://doi.org/10.2108/zsj.19.1223

BioOne Complete (complete.BioOne.org) is a full-text database of 200 subscribed and open-access titles in the biological, ecological, and environmental sciences published by nonprofit societies, associations, museums, institutions, and presses.

Your use of this PDF, the BioOne Complete website, and all posted and associated content indicates your acceptance of BioOne's Terms of Use, available at <u>www.bioone.org/terms-of-use</u>.

Usage of BioOne Complete content is strictly limited to personal, educational, and non - commercial use. Commercial inquiries or rights and permissions requests should be directed to the individual publisher as copyright holder.

BioOne sees sustainable scholarly publishing as an inherently collaborative enterprise connecting authors, nonprofit publishers, academic institutions, research libraries, and research funders in the common goal of maximizing access to critical research.

Retinal Topography of Ganglion Cells and Putative UV-Sensitive Cones in Two Antarctic Fishes: *Pagothenia borchgrevinki* and *Trematomus bernacchii* (Nototheniidae)

Taeko Miyazaki^{1†*} Tetsuo Iwami², Hiroaki Somiya^{3‡} and V. Benno Meyer-Rochow⁴

¹Department of Radiation Protection and Safety, National Institute of Radiological Sciences, Ibaraki 311-1202, Japan
²Laboratory of Biology, Tokyo Kasei Gakuin University, Tokyo 194-0292, Japan
³Department of Life Sciences, Faculty of Bioresources, Mie University, Mie 514-8507, Japan
⁴Faculty of Engineering and Science, International University Bremen (IUB), D-28759 Bremen, Germany

ABSTRACT—Accessory corner cones (ACC) have recently been suggested to be UV-sensitive photoreceptor cells. With a view toward explaining prey detection, we examined the topography of retinal ganglion cells and ACCs in two Antarctic nototheniids occupying different ecological niches: the cryopelagic *Pagothenia borchgrevinki* and the benthic *Trematomus bernacchii*. Isodensity maps of retinal ganglion cells showed that the main visual axis, coincident with the feeding vector, was in a forward direction in both species. Visual acuity was determined as 3.64 and 4.77 cycles/degree for the respective species. In *P. borchgrevinki* the highest density of ACCs was associated with the eye's main visual axis. This suggested that this species uses UV-vision during forward-swims and probably in encounters with prey. On the other hand, *T. bernacchii* possessed two horizontal band-shaped high-density areas of ACCs, which stretched from temporal to nasal and ventral to peripheral retinal regions. Therefore, this species appears to use UV-vision to watch prey across the entire circumference of the lateral area and in the water column above its head.

Key words: Antarctic fish, vision, retinal ganglion cell, UV-sensitivity, feeding strategy

INTRODUCTION

The presence of UV-absorbing visual pigment in the single cone at the corner of the retinal square cone mosaic, named accessory corner cone (ACC), has been reported in many freshwater fishes and euryhaline fishes by using microspectrophotometrical and/or molecular analyses (Avery *et al.*, 1983; Hárosi and Hashimoto, 1983; Whitmore and Bowmaker, 1989; Archer and Lythgoe, 1990; Bowmaker *et al.*, 1991; Hawryshyn and Hárosi, 1991; Raymond *et al.*, 1993; Hisatomi *et al.*, 1996, 1997). Furthermore, in juveniles of some salmonids species (*Salmo trutta, Salmo gairdneri* (=*Oncorhynchus mykiss*), *Salmo salar, Oncorhyn*-

* Corresponding author: Tel. +81-59-231-9673; FAX. +81-59-231-9538.

E-mail: taeko@bio.mie-u.ac.jp

[†] Present address: Department of Life Sciences, Faculty of Bioresources, Mie University, Mie 514-8507, Japan. *chus nerka*) and yellow perch *Perca flavescens*, the loss of UV-photosensitivity associated with the almost complete disappearance of ACCs from the retinal cone mosaic has been confirmed (Bowmaker and Kunz, 1987; Hawryshyn *et al.*, 1989; Loew and Wahl, 1991; Kunz *et al.*, 1994; Novales-Flamarique, 2000).

Recently, we reported the presence of ACCs histologically from the eyes of adults of seven Antarctic nototheniid species as follows: *Trematomus bernacchii, Trematomus newnesi, Trematomus hansoni, Trematomus pennellii, Trematomus scotti, Pagothenia borchgrevinki, Lepidonotothen squamifrons* (=*kempi*) (Miyazaki *et al.*, 2001). Although several specific visual tasks for UV-vision have been proposed (foraging, navigation, and/or communication) and we have very few direct observations on visual feeding in Antarctic fishes, we suggested that Antarctic fishes could increase their feeding efficiency and stock energy by making use of UV-vision during the summer month (Miyazaki *et al.*, 2001).

Topographical analyses of retinal ganglion cells as well as cone photoreceptors can provide us with valuable infor-

[‡] Present address: Department of Biological Resources and Environmental Sciences, School of Agricultural Sciences, Nagoya University, Aichi 464-8601, Japan.

mation on the visual capabilities of fishes. In adult teleosts, the position of the "area centralis (AC)" is related to both habitat and main visual vector of feeding behaviour (Collin and Pettigrew, 1988b, c; Williamson and Keast, 1988; Browman *et al.*, 1990). Furthermore, the distribution of cone photoreceptors determines how the spectral sensitivity of a fish eye changes across its visual field, for different kinds of cones may contain visual pigments with different spectral absorption characteristics (McFarland and Munz, 1975; Beaudet *et al.*, 1997).

In several Antarctic fish species, retinal organizations have been described for specific regions and estimates on visual resolution or photosensitivity were given (Meyer-Rochow and Klyne, 1982; Eastman, 1988; Pankhurst and Montgomery, 1989, 1990; Fanta et al., 1994, Meyer-Rochow et al., 1999). However, the topography of the ganglion cells or cone cells of the whole retina, and in particular the position of the ACs, is still poorly known. Our primary objective was to histologically examine the prev detection system in nototheniids. We also hoped to establish whether the ACCs, as the putative UV-sensitive cones, could assist the fish in prey detection and uptake. The UV-wavelength absorption of ACC's visual pigment has been shown by the previous studies mentioned above, although we have not verified about Antarctic fish yet. In this paper, we therefore examined the retinal topographies of ganglion cells and ACCs for entire retinas of two nototheniids with different habits, namely Pagothenia borchgrevinki and Trematomus bernacchii. We also determined the fish eyes' UV-visual fields. Finally, visual acuity and optical axes were estimated on the basis of retinal ganglion cell distributions, since retinal ganglion cells are providing the link between eyes and behavioural output via the optic nerve.

MATERIALS AND METHODS

Fish

P. borchgrevinki and *T. bernacchii* were caught by line-fishing from a depth of ca. 30 m in Lützow-Holm Bay near Syowa Station (69° S, 39° E). Total lengths of the *P. borchgrevinki* and *T. bernacchii* specimens used were 210 mm and 215 mm, respectively. According to Collin and Pettigrew (1988a), there is no remarkable difference in the topography of cells among specimens of the almost same size of the same species. Based on their observation results, we used a single specimen for each species. Following capture, the fish were anaesthetized with MS222 and then immediately fixed in 10% formalin. Thereafter the eyes were enucleated and cornea, iris, lens, and vitreous humor were removed. The remaining retinae were prepared as follows.

Examination of ganglion cells

The left retinae were radically incised and flatly mounted, and then whole-mount preparations were made following the protocol of Ito and Murakami (1984). The preparations were glued onto slides with the ganglion cell layer facing upward, and stained with crecyl violet. Ganglion cells numbers per 0.13×0.13 mm of each fish retina, about 20 mm in diameter in both fish, were counted at 2mm intervals, under a light microscope, using a calibrated eyepiece. In *P. borchgrevinki* and *T. bernacchii*, 127 and 98 sampling points were used. Values for the ganglion cells were converted into numbers of cells per mm², and then isodensity counter maps of ganglion cells were constructed. The retinal resolving powers were calculated according to the protocol of Collin and Pettigrew (1989), the values thus obtained representing upper limits based on the following equations:

The distance from lens center to retinal focus (posterior nodal distance, PND) was calculated from lens radius (r; mm) and Matthiessen's ratio (2.55):

PND = 2.55r.

The angle (α), subtending 1mm on the retina was calculated by,

 $\tan \alpha = 1/\text{PND}$ $\alpha = \arctan(1/\text{PND})$ (degree).

When the density of the ganglion cells is D (cells/ mm^2), the linear density is

 \sqrt{D} (cells/mm).

Spatial resolution can be calculated by obtaining the number of cells subtended by one degree of visual arc, i.e.,

cells per degree = \sqrt{D} / α .

Since at least two ganglion cells are needed to distinguish the light and dark boundaries from one cycle of a grating of the highest resolvable frequency, visual acuity is given by

cycles per degree = 1/2 cells per degree

Examination of ACCs

Right retina of the *P. borchgrevinki* and *T. bernacchii* was divided into 24 and 30 topographical locations, respectively, and thus the position of each piece in the original retina was determined accurately. The pieces were dehydrated in alcohol and embedded in paraffin. Tangential sections were cut at 7 μ m thickness and stained with haematoxylin and eosin. Cone photoreceptor layers were observed under the light microscope, and numbers of accessory corner cone cells per 0.1×0.1 mm were counted for respective retinal pieces.

RESULTS

Distribution of ganglion cells

All ganglion cells lying within the retinal ganglion cell layer in each fish species were counted. The averages of cell density, estimated from the counts of ganglion cells and counted number of points, were approx. 1.5×10^3 cells/mm²

Table 1. Summery of ganglion cell counts and calculations of visual acuity in two nototheniids.

	Retinal surface area (mm ²)	Counted points	Av. cell density (cells/mm ²)	Total cell No.	Peak density (cells/mm ²)	Lens diameter (mm)	PND (mm)	α (degree)	cells/ degree	cycles/ degree
P. borchgrevinki	260	127	1.5×10 ³	390×10 ³	3313	5.65	7.20	7.91	7.28	3.64
T. bernacchii	248	98	2.5×10 ³	620×10 ³	6804	5.15	6.57	8.65	9.54	4.77

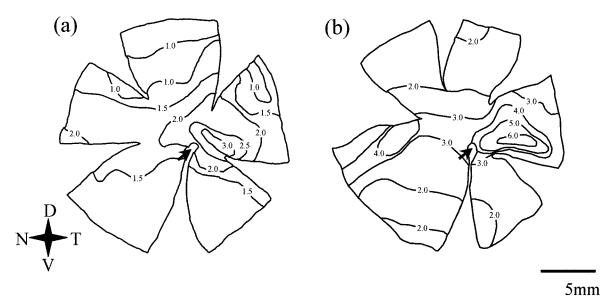


Fig. 1. Isodensity counter maps of the distribution of Nissle-stained neurons (retinal ganglion plus displaced amacrine cells) located within the ganglion cell layer in the left eye of *P. borchgrevinki* (a) and *T. bernacchii* (b). All cell densities $\times 10^3$ per mm². T, temporal; V, ventral; D, dorsal; N, nasal. The arrows indicate the position of optic papilla.

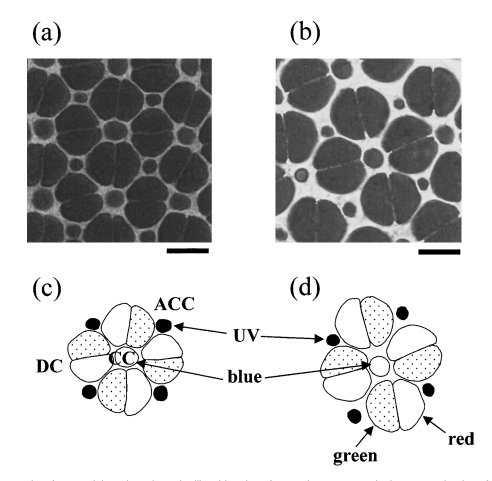


Fig. 2. Photomicrographs of tangential sections through ellipsoid region of cone photoreceptors in the temporal retina of *P. borchgrevinki* (a) and *T. bernacchii* (b). (c) and (d) are tracings of a cone square mosaic on photograph (a) and (b), respectively. ACC, accessory corner cone; CC, central single cone; DC, double cone. The phenotype of the ACC and CC has been identified to be UV- and blue-visual pigment, and that of one of the DC is green- while another of the DC is red-visual pigment (Raymond *et al.*, 1993; Hisatomi *et al.*, 1996, 1997). Scale bar=20 μm.

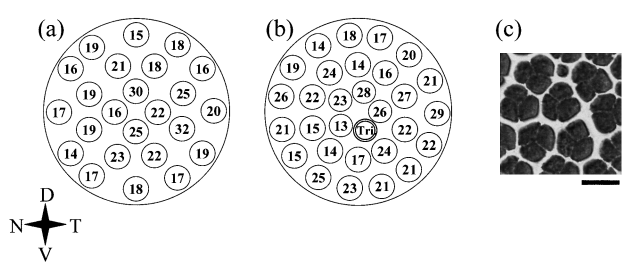


Fig. 3. Maps showing the distribution of ACC densities in the right retina of *P. borchgrevinki* (a) and *T. bernacchii* (b). All cell densities per 0.01mm². Tri. indicates presence of triple cones. (c) Photograph of triple cones in a section of *T. bernacchii* retina. Scale bar=20 μm. T, temporal; V, ventral; D, dorsal; N, nasal.

in *P. borchgrevinki* and 2.5×10^3 cells/mm² in *T. bernacchii* (Table 1). The calculated cell densities were upper estimates, which may include displaced amacrine cells.

Ganglion cells were not uniformly distributed in the retinae of either species. Isodensity maps of *P. borchgrevinki* and *T. bernacchii* resembled each other in that in both species, the region of highest cell density, termed area centralis (AC), was located in the temporal retina (Figs. 1a and 1b). The position of the AC indicates that the visual axis was in a forward direction. In both species, however, a weak horizontal visual streak was also observed a little above the optic papilla (see in Figs. 1a and 1b).

Ganglion cell peak densities were 3313 and 6804 cells/ mm^2 in *P. borchgrevinki* and *T. bernacchii*, respectively, and the corresponding lens diameters were 5.65 mm and 5.15 mm. The PNDs, α s, and the numbers of ganglion cells subtended by one degree of visual arc in the respective species were calculated as shown in Table 1. Consequently, the upper limit of the theoretically derived visual acuity, based on ganglion cell counts, can be given as 3.64 and 4.77 cycles/degree in *P. borchgrevinki* and *T. bernacchii*, respectively.

Distribution of ACCs

Single and double cones were alternately and regularly aligned in the photoreceptor layer, and formed a square mosaic pattern in both species (Figs. 2a and 2b). The ACCs, which were located at the corners of the double cones, could be distinguished from the central single cones by the direction of the axes of the double cones (Figs. 2c and 2d). In both species, the ACCs were present in the entire retina, but the density varied across the retina. Figs.3a and 3b illustrate ACC density distributions in the retinae of *P. borchgrevinki* and *T. bernacchii*, respectively. The highest density of the ACCs was in the temporal retina of both species, which corresponded to the AC. Peak densities of the ACCs were 32 and 29 cells par 0.01mm² in *P. borchgrevinki* and *T. bernacchii*, respectively.

In *P. borchgrevinki*, ACC density was high near the optic papilla of the temporal retina. On the other hand, in *T. bernacchii*, the highest density was seen in the region of the horizontal streak just dorsal of the mid-retina. An additional, high density area was seen in the peripheral part of the ventral retina.

Triple cones were observed near the optic papilla in *T. bernacchii* (Fig. 3c), but not in *P. borchgrevinki*.

DISCUSSION

Both P. borchgrevinki and T. bernacchii belong to the Nototheniidae of the suborder Notothenioidei. However, the former species is cryopelagic, whereas the latter is benthic. Furthermore, the two differ in body shape and position of the eye. The body in *P. borchgrevinki* is laterally compressed and the eyes are positioned laterally. This body shape is commonly associated with fast-swimming fishes and, indeed, this species is often seen as small schools swimming just beneath the sea-ice (Meyer-Rochow, 1982; Meyer-Rochow and Klyne, 1982; Eastman and DeVries, 1985; Foster et al., 1987; Gon and Heemstra, 1990; Pankhurst and Montgomery, 1990). On the other hand, the head of T. bernacchii is broader, and the eye is situated higher on the head and directed anterolaterally. This body shape is typical of benthic fishes, such as gobies and sculpins.

In the present investigation, the main visual axis, thought to provide a clue for the high resolving power of the AC (Shand *et al.*, 2000), was determined to be forward-looking in both species, which is in agreement with the AC's location in the temporal retina. Therefore, the best vision in both species to detect prey items must be considered to lie in front of the fish. Earlier workers have also suggested that

T. bernacchii has a forward-directed feeding vector, since it has a large anterior aphakic space. As for *P. borchgrevinki*, it lacks the anterior aphakic space and may therefore posses an extended visual field (Pankhurst and Montgomery, 1989; Macdonald and Montgomery, 1991; Eastman, 1993).

The binocular visual field, based on eye position and body morphology in *T. bernacchii*, is broader than that of *P. borchgrevinki* but less far-reaching. On the contrary, *P. borchgrevinki* appears to be able to detect prey further away from the body than *T. bernacchii* (Fig. 4). Consequently, it is likely that *P. borchgrevinki* can focus earlier on prey during cruising, while *T. bernacchii* waits and watches prey over a wide range before it initiates an attack. A similar difference in visual fields has been noted between two other nototheniids, namely *Gobionotothen gibberifrons* and *Trematomus newnesi*. The latter two species share the same environmental depth of 40–80 m bottom, but they differ in their feeding-behavioural strategies (Fanta *et al.*, 1994). The expansion of fish visual field should also be discussed based on the eye movement. Regarding eye movement of

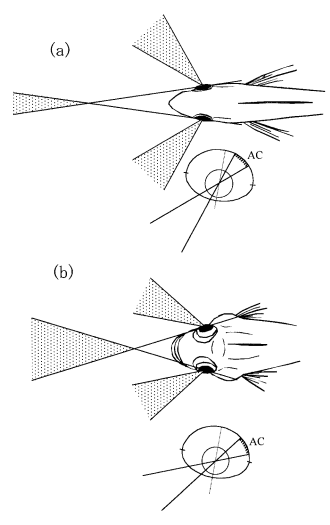


Fig. 4. Schematic diagrams of the binocular visual fields based on eye position, body orientation and position of AC in the retina of *P. borchgrevinki* (a) and *T. bernacchii* (b).

nototheniids, the maximum eye rotation has been reported as about 15 and 20 degree for *P. borchgrevinki* and *Dissoticus mawsonii*, respectively (Montgomery and Macdonald, 1985). Nevertheless, although we don't know how nototheniids use monocular and binocular vision for their visual interest, the eye movement might allow fish to observe its surroundings without moving head or body (Fritsches and Marshall, 2002). Both of *P. borchgrevinki* and *T. bernacchii* may be able to aim prey with the expansion of visual covering range by eye movement.

Regarding visual acuity, there was no remarkable difference in the present two species. The correlation between the visual acuity level of fishes and their swimming types has not been clearly shown in Tamura (1957). Retinal resolving power in cycles per degree can be converted to that of minutes of arc (Murayama and Somiya, 1998), e.g., according to the equation

min of arc = $1/\sqrt{D \times \alpha \times 60}$.

If that is done, values of 8.2' and 6.3' result for *P. borchgrevinki* and *T. bernacchii*, respectively. These values are almost the same as those in other marine teleosts, which were derived from maximum cone densities (*Pagrus major*. 6.4' for a fish of 200 mm in body length; *Lateolabrax japonicus*: 8.5' for an 180 mm long fish; *Sebastiscus marmoratus*: 6.7' also for an 180 mm long fish, Tamura, 1957).

Meyer-Rochow and Klyne (1982) have counted cone and ganglion cells in both species (P. borchgrevinki: approx. 8300 cells/mm² for a fish of approx. 200 mm total body length; T. bernacchii: approx. 4200 cells/mm² for a fish of approx. 240 mm total body length: converted from original data). Furthermore, there have been studies on the theoretical visual acuity calculated from retinal cone densities for both species (P. borchgrevinki: 25-50' for fish of 63-220 mm total body length: Pankhurst and Montgomery, 1990; T. bernacchii: 10' in the temporal retina for a fish of 192 mm total body length; Miyazaki, unpublished data). Some differences between earlier values and our present values might have been caused by differences in fish size or retinal region analyzed for each examination. Nevertheless, a close agreement between values based on cone cell and ganglion cell estimates, as well as behavioural tests clearly exists (Collin and Pettigrew, 1989; Arrese et al., 2000).

The maximum distance (L; cm), at which a fish could recognize a prey item (ϕ ; cm), can be estimated from the fish's visual acuity according to

D=($\phi \times 1$ /min of arc $\times 180 \times 60$)/ π ,

provided optical properties in the sea (turbidity and light intensity) are neglected (Miyazaki and Nakamura, 1990; Miyazaki *et al.*, 2000). On that basis, for example, the recognizable distance for a prey of 3 cm diameter would be about 12.6 m in *P. borchgrevinki* and 16.4 m in *T. bernacchii*.

While these figures do not seem to vary too greatly between the two species, the distribution of the ACCs in the

retina, on the other hand, was quite dissimilar. In P. borchgrevinki, the region in which the density of the ACCs was highest coincided with the position of the AC. Therefore, UVvision in P. borcharevinki must be assumed to be most effective in a forward direction, and consequently P. borchgrevinki ought to principally recognize prey in the swimming direction from a larger distance than T. bernacchii (Cronin et al., 1994; Losey et al., 1999). As for T. bernacchii, the highdensity area was the horizontal streak, which stretched from the temporal to the nasal retina. An additional area of high density was developed in the peripheral parts of the ventral retina. UV-vision in T. bernacchii, therefore, appears to be effective not only along the main visual axis of the forwardmovement but also across a vast lateral area and into the upper water column itself. This extensive field could improve prey discovery, allowing T. bernacchii to remain relatively motionless in anticipation of prey. All of these conclusions are supported by the eye's morphology. We believe P. borchgrevinki detects pelagic prev, e.g. copepods (Hoshiai and Tanimura, 1981; Hoshiai et al., 1989) as silhouettes against downwelling light, whereas T. bernacchii relies mainly on laterally incident light to feed, since T. bernacchii in contrast to P. borchgrevinki displays an obvious corneal iridescence (Macdonald and Montgomery, 1991; Eastman, 1993), which screens out bright downwelling light.

The benefit for fish using UV-vision for feeding, lies in enhancing the contrast of prey against a UV-background and it is thought that UV-sensitivity is widespread in zooplanktivorous fishes (e.g., juvenile rainbow trout Oncorhynchus mykiss, pumpkinseed sunfish Lepomis gibbosus: Browman et al., 1994; juvenile yellow perch Perca flavescens: Loew and Wahl, 1991; Loew et al., 1993; pomacentrid fish Dascyllus trimaculatus, Pomacentrus coelestis, Chromis punctipinnis: McFarland and Loew, 1994). In Antarctic nototheniids, UV-vision may represent one strategy to increase foraging efficiency during the Antarctic summer, when light conditions are appropriate for visual feeding and a visual threshold of 2×10^9 photons cm⁻² s⁻¹ is reached (Morita et al., 1997). However, other and additional means of detecting predators and prev are entirely possible, especially if it is remembered that individuals with aberrant eyes can, apparently, reach full adulthood (Meyer-Rochow, 1990). To (a) determine the extent that nototheniids can make use of UV-light and (b) understand the role of triple cones, present in the retina of T. bernacchii (but not P. borchgrevinki), are thus our next goals.

ACKNOWLEDGEMENTS

We are thankful to members of 34th and 35th Japanese Antarctic Research Expedition for their kind support during respective research in Antarctica. This study was partly supported by the research fund of NIRS Brain Research Project.

REFERENCES

Archer SN, Lythgoe JN (1990) The visual pigment basis for cone

polymorphism in the guppy, *Poecilia reticulata*. Vision Res 30: 225–233

- Arrese C, Archer M, Runham P, Dunlop SA, Beazley LD (2000) Visual system in a diurnal marsupial, the numbat (*Myrmecobius fasciatus*): retinal organization, visual acuity and visual fields. Brain Behav Evol 55: 163–175
- Avery JA, Bowmaker JK, Djamgoz MBA, Downing JEG (1983) Ultra-violet sensitive receptors in a freshwater fish. J Physiol 334: 23–24
- Beaudet L, Novales-Flamarique I, Hawryshyn CW (1997) Cone photoreceptor topography in the retina of sexually mature Pacific salmonid fishes. J Comp Neurol 383: 49–59
- Bowmaker JK, Kunz YW (1987) Ultraviolet receptors, tetrachromatic colour vision and retinal mosaics in the brown trout (*Salmo trutta*): age-dependent changes. Vision Res 27: 2101–2108
- Bowmaker JK, Thorpe A, Douglas RH (1991) Ultraviolet-sensitive cones in the goldfish. Vision Res 31: 349–352
- Browman HI, Gordon WC, Evans BI, O'Brien WJ (1990) Correlation between histological and behavioral measures of visual acuity in a zooplanktivorous fish, the white crappie (*Pomoxis annularis*). Brain Behav Evol 35: 85–97
- Browman HI, Novales-Flamarique I, Hawryshyn CW (1994) Ultraviolet photoreception contributes to prey search behaviour in two species of zooplanktivorous fishes. J Exp Biol 186: 187–198
- Collin SP, Pettigrew JD (1988a) 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
- Collin SP, Pettigrew JD (1988b) Retinal topography in reef teleosts. I. Some species with well-developed areae but poorly-developed streaks. Brain Behav Evol 31: 269–282
- Collin SP, Pettigrew JD (1988c) Retinal topography in reef teleosts. II. Some species with prominent horizontal streaks and highdensity areae. Brain Behav Evol 31: 283–295
- Collin SP, Pettigrew JD (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
- Cronin TW, Marshall NJ, Quinn CA, King CA (1994) Ultraviolet photoreception in mantis shrimp. Vision Res 34: 1443–1452
- Eastman JT, DeVries AL (1985) Adaptations for cryopelagic life in the Antarctic notothenioid fish *Pagothenia borchgrevinki*. Polar Biol 4: 45–52
- Eastman JT (1988) Ocular morphology in Antarctic notothenioid fishes. J Morphol 196: 283–306
- Eastman JT (1993) Antarctic Fish Biology, Evolution in a unique environment. Academic Press Inc, San Diego
- Fanta E, Meyer AA, Grötzner SR, Luvizotto MF (1994) Comparative study on feeding strategy and activity patterns of two Antarctic fish: *Trematomus newnesi* Boulenger, 1902 and *Gobionotothen gibberifrons* (Lönnberg, 1905) (Pisces, Nototheniidae) under different light conditions. Nankyoku Shiryô (Antarctic Record) 38: 13–29
- Foster BA, Cargill JM, Montgomery JC (1987) Planktivory in *Pag-othenia borchgrevinki* (Pisces: Nototheniidae) in McMurdo Sound, Antarctica. Polar Biol 8: 49–54
- Fritsches KA, Marshall NJ (2002) Independent and conjugate eye movements during optokinesis in teleost fish. J Exp Biol 205: 1241–1252
- Gon O, Heemstra PC (1990) Fishes of the southern ocean. JLB Smith Institute of Ichthyology, Grahamstown
- Hárosi FI, Hashimoto Y (1983) Ultraviolet visual pigment in a vertebrate: a tetrachromatic cone system in the dace. Science 222: 1021–1023
- Hawryshyn CW, Arnold MG, Chaisson DJ, Martin PC (1989) The ontogeny of ultraviolet photosensitivity in rainbow trout (*Salmo gairdneri*). Visual Neurosci 2: 247–254
- Hawryshyn CW, Hárosi FI (1991) Ultraviolet photoreception in carp:

microspectrophotometry and behaviorally determined action spectra. Vision Res 31: 567–576

- Hisatomi O, Satoh T, Barthel LK, Stenkamp DL, Raymond PA, Tokunaga F (1996) Molecular cloning and characterization of the putative ultraviolet-sensitive visual pigment of goldfish. Vision Res 36: 933–939
- Hisatomi O, Satoh T, Tokunaga F (1997) The primary structure and distribution of killifish visual pigments. Vision Res 37: 3089–3096
- Hoshiai T, Tanimura A (1981) Copepods in the stomach of a nototheniid fish, *Trematomus borchgrevinki* fry at Syowa Station, Antarctica. Mem Natl Inst Polar Res 34: 44–48
- Hoshiai T, Tanimura A, Fukuchi M, Watanabe K (1989) Feeding by the nototheniid fish, *Pagothenia borchgrevinki* on the ice-associated copepod, *Paralabidocera antarctica*. Proc NIPR Symp Polar Biol 2: 61–64
- Ito H, Murakami T (1984) Retinal ganglion cells in two teleost species, Sebastiscus marmoratus and Navodon modestus. J Comp Neurol 229: 80–96
- Kunz YW, Wildenburg G, Goodrich L, Callaghan E (1994) The fate of ultraviolet receptors in the retina of the Atlantic salmon (*Salmo salar*). Vision Res 34: 1375–1383
- Loew ER, Wahl CM (1991) A short-wavelength sensitive cone mechanism in juvenile yellow perch, *Perca flavescens*. Vision Res 31: 353–360
- Loew ER, McFarland WN, Mills EL, Hunter D (1993) A chromatic action spectrum for planktonic predation by juvenile yellow perch, *Perca flavescens*. Can J Zool 71: 384–386
- Losey GS, Cronin TW, Goldsmith TH, Hyde D, Marshall NJ, McFarland WN (1999) The UV visual world of fishes: a review. J Fish Biol 54: 921–943
- Macdonald JA, Montgomery JC (1991) The sensory biology of notothenioid fish. In "Biology of Antarctic fish" Ed by G Prisco, B Maresca, B Tota, Springer-Verlag, Heidelberg, pp 145–162
- McFarland WN, Munz FW (1975) Part III: The evolution of photopic visual pigments in fishes. Vision Res 15: 1071–1080
- McFarland WN, Loew ER (1994) Ultraviolet visual pigments in marine fishes of the family Pomacentridae. Vision Res 34: 1393–1396
- Meyer-Rochow VB (1982) Lifeforms under Antarctic ice. Kagaku Asahi 42: 16–19 [in Japanese]
- Meyer-Rochow VB (1990) A case of abnormal eye enlargement in the Antarctic fish *Pagothenia borchgrevinki* (Pisces, teleostei, notothenioidei). NZ Antarctic Rec 10: 28–31
- Meyer-Rochow VB, Klyne MA (1982) Retinal organization of the eyes of three nototheniid fishes from Ross Sea (Antarctica). Gegenbaurs Morphol Jahrb 128: 762–777
- Meyer-Rochow VB, Morita Y, Tamotsu S (1999) Immunocytochemical observations of the pineal organ and retina of the Antarctic teleosts *Pagothenia borchgrevinki* and *Trematomus bernacchii*. J Neurocytol 28: 125–130

- Miyazaki T, Nakamura Y (1990) Single line acuity of 0-year-old Japanese parrotfish determined by the conditioned reflex method. Nippon Suisan Gakkaishi 56: 887–892 [in Japanese]
- Miyazaki T, Shiozawa S, Kogane T, Masuda R, Maruyama K, Tsukamoto K (2000) Developmental changes of the light intensity threshold for school formation in the striped jack *Pseudocaranx dentex.* Mar Ecol Prog Ser 192: 267–275
- Miyazaki T, Iwami T, Yamauchi M, Somiya H (2001) "Accessory corner cones" as putative UV-sensitive photoreceptors in the retinas of seven adult nototheniid fishes. Polar Biol 24: 628– 632
- Montgomery JC, Macdonald JA (1985) Oculomotor function at low temperature: Antarctic *versus* temperate fish. J Exp Biol 117: 181–191
- Morita Y, Meyer-Rochow VB, Uchida K (1997) Absolute and spectral sensitivities in dark- and light-adapted *Pagothenia borchgrevinki*, an Antarctic nototheniid fish. Physiol Behav 61: 159– 163
- Murayama T, Somiya H (1998) Distribution of ganglion cells and object localizing ability in the retina of three cetaceans. Fisheries Sci 64: 27–30
- Novales-Flamarique I (2000) The ontogeny of ultraviolet sensitivity, cone disappearance and regeneration in the sockeye salmon *Oncorhynchus nerka*. J Exp Biol 203: 1161–1172
- Pankhurst NW, Montgomery JC (1989) Visual function in four Antarctic Nototheniid fishes. J Exp Biol 142: 311–324
- Pankhurst NW, Montgomery JC (1990) Ontogeny of vision in the Antarctic fish *Pagothenia borchgrevinki* (Nototheniidae). Polar Biol 10: 419–422
- Raymond PA, Barthel LK, Rounsifer ME, Sullivan SA, Knight JK (1993) Expression of rod and cone visual pigments in goldfish and zebrafish: A rhodopsin-like gene is expressed in cones. Neuron 10: 1161–1174
- Shand J, Chin SM, Harman AM, Moore S, Collin SP (2000) Variability in the location of the retinal ganglion cell area centralis is correlated with ontogenetic changes in feeding behavior in the black bream, *Acanthopagrus butcheri* (Sparidae, teleostei). Brain Behav Evol 55: 176–190
- Tamura T (1957) A study of visual perception in fish, especially on resolving power and accommodation. Bull Jap Soc Sci Fish 22: 536–557
- Whitmore AV, Bowmaker JK (1989) Seasonal variation in cone sensitivity and short-wave absorbing visual pigments in the rudd *Scardinius erythrophthalmus.* J Comp Physiol A 166: 103–115
- Williamson M, Keast A (1988) Retinal structure relative to feeding in the rock bass (*Ambloplites rupestris*) and bluegill (*Lepomis macrochirus*). Can J Zool 66: 2840–2846

(Received April 30, 2002 / Accepted August 27, 2002)