Retinal and lenticular ultrastructure in the aestivating salamanderfish, *Lepidogalaxias salamandroides* (Galaxiidae, Teleostei) with special reference to a new type of photoreceptor mosaic

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**Summary.** The salamanderfish, *Lepidogalaxias salamandroides* (Galaxiidae, Teleostei) is endemic to southwestern Australia and inhabits shallow, freshwater pools which evaporate during the hot summer months. Burrowing into the substrate in response to falling water levels allows these fish to aestivate for extended periods of time while encapsulated in a mucous cocoon even when the pools contain no water. Only a few minutes after a major rainfall, these fish emerge into relatively clear water which subsequently becomes laden with tannin, turning the water black and reducing the pH to approximately 4.3. As part of a large study of the visual adaptations of this unique species, the retinal and lenticular morphology of the aestivating salamanderfish is examined at the level of the light and electron microscopes. The inner retina is highly vascularised by a complex system of vitreal blood vessels, while the outer retina receives a blood supply by diffusion from a choriocapillaris. This increased retinal blood supply may be an adaptation for reducing the oxygen tension during critical periods of aestivation. Large numbers of Müller cells traverse the thickness of the retina from the inner to the outer limiting membranes. The ganglion cells are arranged in two ill-defined layers, separated from a thick inner nuclear layer containing two layers of horizontal cells by a soma-free inner plexiform layer. The photoreceptors can be divided into three types typical of many early actinopterygian representatives; equal double cones, small single cones and large rods (2:1:1). These photoreceptors are arranged into a unique regular square mosaic comprising a large rod bordered by four equal double cones with a small single cone located at the corner of each repeating unit. The double cones may optimise perception of mobile prey which it tracks by flexion of its head and “neck” and the large rods may increase sensitivity in the dark tannin-rich waters in which it lives. Each single cone also possesses a dense collection of polysomes and glycogen (a paraboloid) beneath its ellipsoid, the first such finding in teleosts. The retinal pigment epithelium possesses melanosomes, phagocytes and a large number of mitochondria. The anatomy of the retina and the photoreceptor mosaic is discussed in relation to the primitive phylogeny of this species and its unique life history.

**Key words:** Fish, Retina, Vision, Photoreceptors, Mosaic, Retinal pigment epithelium

**Introduction**

The Australian salamanderfish, *Lepidogalaxias salamandroides* is a freshwater fish endemic to the southwest region of Western Australia. After its initial description by Mees in 1961, the phylogeny of this peculiar fish has been debated (see review by Collin and Collin, 1996). However, despite the controversy, it is presently agreed that this species is highly specialised and belongs to an endemic monotypic family, the Lepidogalaxiidae, lying firmly within the galaxioids as a sister group of the osmeroids (Williams, 1987; Begle, 1991). Frankenberg (1968) regards *L. salamandroides* as a “living fossil”, ancestral to the galaxiids making it as primitive as any living teleost. Hence, the salamanderfish and the Australian lungfish, *Neoceratodus forsteri* may comprise an ancient cohort dating back to the “fragmentation of Pangaea” (Rosen, 1973, 1974).

The life history of this species is particularly interesting. Surviving in shallow, freshwater ponds which contain large amounts of tannin (reducing the pH from 7.0 to 4.3) (Christensen, 1982), these small fish burrow into the substrata and aestivate for many months during periods of drought (Pusey, 1981; Berra and Allen, 1989). Covered by mud and leaf litter (to a depth of up to 60 cm) in a pond completely devoid of water, *L.*
salamandroides is thought to utilise cutaneous respiration during these periods given that it possesses an avascular swimbladder incapable of aerial gas exchange (Berra et al., 1989; Martin et al., 1993). A robust wedge-shaped skull (Frankenberg, 1969) and the absence of long ribs (Berra and Allen, 1989) enable the salamanderfish to penetrate the substrate and construct either a pear- or a U-shaped burrow connected to the surface by a thin tube (McDowall, 1981; Pusey, 1981). It also secretes a mucous sheath from goblet cells and mucus-secreting cells located in the epithelium of the skin and cornea which may aid in burrowing and inhibit desiccation during aestivalisation (Collin and Collin, 1996).

During aestivalisation, L. salamandroides survives on stores of fat and by reducing its metabolic rate. However, within minutes of a major rainfall, it actively emerges from its burrow (Berra and Allen, 1988; Berra et al., 1989) to go in search of food. Initially, prey capture is aided by the clarity of the water but this is impeded by the tannin-rich substrate which stains the water black, placing enormous constraints on the visual system. Thus far, little is known of the visual capabilities of this species and how the eye has adapted to changes in the physical constraints of temperature, pH, light levels and desiccation.

A detailed ultrastructural description of the cornea has shown that, in addition to mucus-secreting cells, sutureal fibres, which link stromal collagen lamellae, may provide a physical constraint to combat the extreme changes in pH during the aestivalisation phase and/or the ionic changes in the water following emergence from the burrow (Collin and Collin, 1996). However, a number of other findings suggest that the eyes are not only functional but optimised for vision in bright light. These include two types of pigment granules in the dermal stroma of the cornea which suggests that coloured filters may be effective in eliminating light scatter, the reduction of chromatic aberration and possibly the improvement of visual acuity (Kondrashov et al., 1986) especially when the tannin concentrations are lowest after heavy rain (Collin and Collin, 1996). A corneal iridescence layer may also play a role as a selective coloured filter (Collin and Collin, 1996) to take advantage of the transmitted light not absorbed by the dark tannin-rich water it inhabits as it sits at the entrance of its burrow, perched on its pelvic fins in search of its prey of chironomid larvae (90% of the total stomach contents) and crustaceans (Collin S.P. and Gill H., unpublished observations). Not an active predator, L. salamandroides chooses to lie-in-wait for its benthic mobile prey. With only limited eye mobility (Collin and Collin, 1996), prey are tracked with the aid of a flexible spine or “neck”, made possible by a number of spaces between the anterior vertebrae (Frankenberg, 1969; Berra and Allen, 1989), which allows both side-to-side and up-and-down movements of the head.

As part of a continuing study of the visual capabilities of L. salamandroides, the retina is examined at the levels of the light and electron microscopes. Our findings show that the retina is thin, optimised for low oxygen tensions and possesses three types of photoreceptor adapted for optimising vision in low light levels. A unique regular photoreceptor mosaic may also increase the perception of movement without the advantage of an area centralis or fovea. Retinal features in common with other primitive gnathostomes also indicate that the current phylogenetic placement of this species can be confirmed.

Materials and methods

Twelve individuals (30-58 mm in length) of the Australian salamanderfish, Lepidogalaxias salamandroides (Galaxiidae, Teleostei) were collected from three small freshwater pools, 10 km south of Northcliffe and approximately 300 km south of Perth, Western Australia. Specimens were collected by either a fine-mesh handnet or a seine net (0.5 cm mesh size) but only in winter and spring since most of these pools are dry during the summer months. Fish were transferred to aerated tanks and transported to the University of Western Australia, where they were maintained in a constant 12:12 hr. day/night cycle in aquaria housed in the Department of Zoology.

All specimens were sacrificed in the light with an overdose of tricaine methane sulphonate (MS222, 1:2,000) under the ethical guidelines of the National Health and Medical Research Council of Australia. Three heads, 12 whole eyes and 6 eyecups (with their corneae and lenses removed) were immersion-fixed in a mixtare of fresh 2.5% glutaraldehyde and 2% paraformaldehyde in 0.1M sodium cacodylate buffer (pH 7.4) for between 12 and 24 hours, and later stored in 0.1M sodium cacodylate buffer (pH 7.4). The whole heads and 6 whole eyes were dehydrated and embedded in Historesin (Reichert-Jung) and cut in transverse section (1-2 µm) on an American Optical rotary microtome using a glass knife. The eyecups were embedded in araldite and cut in both the transverse and tangential planes (1-2 µm) using an LKB Nova motorised ultramicrotome. Semi-thin sections were stained with either Toluidine blue or Richardson’s stain and viewed using an Olympus compound light microscope (BH-2).

The six remaining eyes were prepared for electron microscopy by post-fixing in 2% osmium tetroxide with 1.5% potassium ferricyanide in 0.1M sodium cacodylate buffer (the reduced osmium method of Collin and Allansmith, 1977). Tissue was then dehydrated in acetone and embedded in resin (Polybed/812, Polysciences Inc). Thin sections were stained with lead citrate and uranyl acetate and examined on a Siemens Elmiskop 1A or an Hitachi H500 transmission electron microscope.

All measurements were made on enlargements of electron micrographs using a magnifier and graticule. Photographs were taken on either 35 mm Kodak Technical Pan film (rated at 50 ASA, light microscopy) or Kodak 4489 electron microscope film.
Results

The eyes of the salamanderfish, *Lepidogalaxias salamandroides* occupy a lateral position in the head with a binocular overlap of approximately 45° (Collin and Collin, 1996). The horizontally-elongated eyes comprise approximately 23% of the total head length (McDowall and Pusey, 1983, Fig. 1A). The lens, which fails to fill a rostrally-tapered pupil, is spherical. Examination of the lens using transmission electron microscopy shows that its structure is typical of other teleosts and comprises a granular lens capsule (0.95 μm in thickness), an anterior layer of epithelial cells (2.3 μm in thickness) and a concentric arrangement of cells surrounding an ill-defined lens nucleus (Fig. 1C). The lens is suspended dorsally by a suspensory ligament and ventrally by a ligament attached to the retractor lenticulus muscle which contains smooth muscle fibres and mitochondria.

The retina of *L. salamandroides* is approximately 135 μm thick and is covered by a pattern of vitreal blood vessels closely apposed to the inner limiting membrane (ILM, Fig. 1B, D). The inner surface of the ILM is almost entirely covered by the endfeet of Müller cell processes, which divide the ganglion cell axons into bundles, traversing the retina to the level of the outer limiting membrane (OLM; Fig. 1D). The optic nerve head is situated in the centro-temporal region of the retina, at the dorsal end of a falciform process which extends toward the ventro-nasal retinal margin. A large proportion of the optic axons entering the retina at the optic nerve head are myelinated. However, the myelinated axons are rare in other regions of the retinal nerve fibre layer.

The ganglion cell somata (up to 5 μm in diameter) lie in two ill-defined layers and possess rough endoplasmic reticulum organised in Nissl-bodies, mitochondria, Golgi apparatus and lysosomes. The inner plexiform layer is approximately 35 μm in thickness and is identified by its complex arrangement of dendritic processes and synaptic connections (Fig. 1B). Amacrine and bipolar cells and two layers of horizontal cells (7-9 μm in diameter), together with scattered Müller cell nuclei, comprise the inner nuclear layer (INL, 25 μm in thickness).

The outer plexiform layer is 5.5 μm thick and has a complex arrangement of synaptic connections between horizontal and bipolar cell processes and rod and cone terminals. The terminals of the rod (spherules) and cone (pedicles) photoreceptors are tightly-packed and make many contacts with other cell processes by way of surface contacts and synaptic ribbons located at the end of spherul-directed invaginations. The synaptic ribbons are coated with an amorphous substance and are surrounded by vesicles which are always aligned along the length of the ribbon. Rod spherules are consistently found to have a single synaptic ribbon (up to 3 μm in length and 60 nm in width) adjacent to an arciform density, in contrast to cone pedicles, which have multiple (two to four) synaptic ribbons of similar dimensions (Fig. 1E). Due to the complex arrangement of processes, it is difficult to ascertain how many invaginating processes penetrate either the rod or cone terminal, but in rods and in most cones, the synaptic ribbon projects from the juncture of two lateral processes and a single central process to form a triad. In the centre of the ribbon there is an electron lucent material bordered on each side by a dark band. The rod synaptic vesicles are more concentrated throughout the rod terminal than the vesicles in the cone pedicle, although the size of the vesicles in both types appears similar (approximately 65 nm in diameter). Desmosomal and gap-like junctions appear to join the apical processes of the Müller cells and the myoid region of the photoreceptors at the OLM.

Numerous processes or microvilli of the Müller cells project sclerad from the OLM and surround the inner segments of the photoreceptors.

The photoreceptors can be differentiated into three types; equal, double (twin) cones, small single cones and large rods (with a ratio of 2:1:1 Figs. 2A, B, 3A, B, 6A). These photoreceptors are arranged in a regular square mosaic. Within each repeating unit, a large rod is bordered by four equal double cones with the single cone located at each corner (Figs. 4, 5). Units overlap where each double cone forms part of two units. The common surface between the two components of each double cone is oriented perpendicular to its neighbours, whereas double cones from the inner surface of the OLM (Figs. 2B-D, 5B-C).

Each double cone is comprised of two cones of equal size (up to 30 μm in length and 3.0 μm in width measured at the base of the outer segment). One component of the double cone lies slightly sclerad of the other. The double cones are closely apposed along their inner segment surfaces (Fig. 2A, B), immediately adjacent to which lies a complex series of membranes. These subsurface cisternae consist of membrane-bound structures (approximately 20 nm in diameter) which run longitudinally for the length of each inner segment. The cisternae of the mitochondria located adjacent to the inner segment borders of each apposing cone in the doublet are often oriented parallel to the subsurface cisternae (Fig. 2C, D). The tapered outer segments of the equal double cones are up to 12 μm in length and contain a series of discs comprising two membranes, both approximately 5 nm in thickness, with an intradisc space of 5 nm. Each disc is between 14 and 28 nm from its nearest neighbouring disc. A thin triplae membrane surrounds the outer segments of the cones. Each outer segment is surrounded by 12 to 14 calycal processes which do not appear to indent the disc membrane. The double cones possess a short myoid and their lightly-staining nuclei, which lie in the scleral region of the outer nuclear layer, are often invaginated.

The single cones are smaller (up to 27 μm in length and 1.5 μm in width measured at the base of the outer segment) than the double cones and possess a tapering outer segment surrounded by 10 to 12 calycal processes (Figs. 3A, B, 6A). The mitochondria within the ellipsoid
Fig. 1. **A**, Lateral view of the head of *Lepidogalaxias salamandroides* showing the large, lateral eyes and ventral mouth. Bar: 1.5 mm. **B**, Transverse section of the light-adapted retina showing the thick inner nuclear layer and the large row of rods (r) scleral to the equal double cones (dc). gc, ganglion cell layer; iNL, inner nuclear layer; ipi, inner plexiform layer; oNL, outer nuclear layer; rpe, retinal pigment epithelium. Bar: 15 μm. **C**, Electron micrograph of the anterior lens showing the granular lens capsule (gc) surrounding a thicker epithelial cell layer of interdigitating cells (ec) and a concentric arrangement of inner lenticular cells. Bar: 1.5 μm. **D**, Electron micrograph of a vitreous blood vessel (vb) shown in transverse section indented and opposed to the inner limiting membrane (iml, arrowheads). Note that the end feet of the Müller cells (m) line the iml, dividing the ganglion cell axons (a) into fascicles. Bar: 2.5 μm. **E**, Electron micrograph of the outer plexiform layer showing the synaptic ribbons (arrowheads) within the rod spherules (rs) and cone pedicles (cp). Bar: 1 μm.
Fig. 2. A. Transverse section of three equal double cones (dc) lying in the light-adapted position. Bar: 2.5 μm. B. Higher magnification of an equal double cone showing the disc membranes and mitochondria within the outer (os) and inner (is) segments, respectively. Bar: 1 μm. C. Tangential section of an equal double cone at the level of the ellipsoid. m: mitochondria. Bar: 1 μm. D. Higher magnification of the complex of subsurface cisternae lying internal to the opposing inner segment membranes (small arrowheads) of each component of a double cone. The membranes of the inner and outer cisternae are depicted by large arrowheads and arrows, respectively. m: mitochondria. Bar: 0.2 μm.
Retinal morphology of the salamanderfish

appear tightly-packed, are more lightly-staining than those of the double cones and show a gradient in size with the smaller mitochondria located sclerad. Unique to the single cones, is a dense collection of what appear to be polysomes (38 nm in diameter) scattered in a region of the myoid beneath the mitochondria. Within this region lie numerous membrane-bound vesicles containing clusters of granules (50 nm in diameter). These granules, thought to be aggregations of glycogen, together with the surrounding ribosome-rich tissue may constitute a paraboloid (Fig. 6B,C).

The rods are large (up to 60 μm in length and 6.0 μm in width measured at the base of the outer segment) with a short ellipsoid which contains lightly-staining, elongated (in the vitread-sclerad axis) mitochondria arranged into a closely-packed hexagonal array (Figs. 5D, 6A). The outer segments comprise two-thirds of the receptor length and do not taper sclerad. Each cylindrically-shaped rod outer segment possesses up to 8 incisures which appear as spaces or gaps in the discs which run longitudinally, often for a large proportion of the length of the outer segment. The rod disc spacing appears less than that measured in either of the other two types of cone photoreceptors. Up to 42 calycal processes lie closely apposed to the membrane enveloping the discs of the large rod outer segments. The rod nuclei are smaller than the cone nuclei and lie in the vitread region of the outer nuclear layer. All three photoreceptor types possess accessory outer segments.

The retinal pigment epithelium (RPE) possesses pigment granules which appear circular in tangential section (approximately 0.5 μm in diameter) but elongate

Fig. 3. A. Single (sc) and double (dc) cones showing the differential staining of the mitochondria in the ellipsoid, the difference in myoid thickness and the relative positions of the nuclei. Note that the nuclei (n) of some double cones are bi-lobed. Arrowheads depict the level of the outer limiting membrane. Bar: 2 μm. B. Small single cone (sc) in between two equal double cones (dc). Arrowhead depicts a calycal process. m: mitochondria. Bar: 1.5 μm.
Fig. 4. A. A tangential section across the regular photoreceptor mosaic at the levels of the inner (is) and outer (os) segments. Bar: 10 μm. B. Higher magnification of the mosaic at the level of the inner segments showing the square mosaic of four double cones (d) surrounding a central rod (r). s: single cone. Bar: 10 μm. C. The mosaic at the level of the double cone outer segments. Bar: 10 μm. D. The mosaic at the level of the outer segments of the large rods (r). Bar: 15 μm.
Fig. 5. Series of electron micrographs showing the changes in the appearance of tangential sections of the photoreceptor mosaic in progressive levels from the myoid to the outer segments. **A.** The myoid level. Bar: 1.5 μm. **B.** The vitread border of the double cone ellipsoid. Bar: 5 μm. **C.** The scleeral border of the double cone ellipsoid. Bar: 10 μm. **D.** The level of the double cone outer segment ultras. Note the dense collection of mitochondria within the rod ellipsoid. Arrowheads depict calycal processes surrounding the outer segment of a double cone outer segment. d: double cone; r: rod; s: single cone. Bar: 2.5 μm.
Fig. 6. A. Electron micrograph showing the three types of photoreceptors: equal double cone (d), single cone (s) and large rod (r) in the scleral region of the light-adapted retina. m: mitochondria of the rod ellipsoid. Bar: 3 μm. B. Collections of polysomes (arrows) surrounded by aggregations of glycogen in the myoid region of the single cones thought to comprise a paraboloid. Bar: 1.5 μm. C. Higher magnification of the polysomes in the paraboloid. Bar: 0.5 μm. D. Transverse section of the retinal pigment epithelium showing numerous inclusions including myeloid bodies (mb), mitochondria (m) and melanosomes (g). bm: Bruch's membrane; p: phagosome or lamellated body; pe: pericyte. Bar: 2 μm.
in transverse section (up to 3.0 μm in length) and surround the outer segments of the photoreceptors in the light-adapted state (Fig. 6A, D). Each mononucleate RPE cell is joined to its neighbour by various types of junctions including zona adhaerens and is rich in mitochondria, Golgi apparatus and phagosomes. Bruch's membrane is 0.33 μm in thickness and is pentalaminate. Its vitread and scleral borders comprise the basement membranes of the retinal pigment epithelium and the choriocapillaris, respectively, with a layer of collagen fibrils separating a broken layer of elastic tissue.

The choriocapillaris possesses two to three layers of melanocytes containing oval and circular-shaped melanosomes (up to 1.3 μm in diameter) interspersed between large and small thin-walled capillaries (Fig. 6D). The endothelial cells of the choriocapillaris are moderately fenestrated.

**Discussion**

The presence of vitreal blood vessels, a choriocapillaris and an extensive system of Müller cell processes indicates that the retina of *Lepidogalaxias salamandroides* is either metabolically-active and/or may be under extreme oxygen tension. Vitreal vessels normally provide an effective transport mechanism for metabolic exchange in retinas containing high densities of retinal neurons often concentrated into a specialised zone of acute vision e.g. in some marine (Hanyu, 1962; Collin, 1989) and freshwater (Kohbara et al., 1987; Collin et al., 1996a,b) teleosts. The retina in *L. salamandroides* is thin (135 μm) when compared to other teleosts, for example 265 μm in the cutlips minnow, *Exoglossum maxillatum* (Collin et al., 1996a), 330 μm in the sand lance, *Limnichthys fasciatus* (Collin and Collin, 1988) and 570 μm in the snake mackerel, *Gempylus serpens* (Munk, 1985), and both the photoreceptor and ganglion cell layers possess relatively low densities of neurons (Collin S.P. and Gill H., unpublished data). Therefore, the increased blood supply to the retina of *L. salamandroides* may be an adaptation for reducing the oxygen tension during critical periods of aestivation when metabolic rate, and therefore oxygen exchange, may be low.

The presence of at least two morphological types of photoreceptor terminals with single (rod) and multiple (cone) synaptic ribbons provides evidence for the differentiation of photoreceptor types in *L. salamandroides*, although double and single cone terminals could not be differentiated. As in other species of teleosts, the cone terminals in the salamanderfish are larger and contain more synaptic sites (ribbon synapses and surface contacts) than the rod terminals (Cohen, 1972; Braekevelt, 1982) with the lateral and central invaginations corresponding to the horizontal and bipolar cell processes, respectively (Stell, 1965; Dowling, 1968). Similarly, the scleral and vitread location of the cone and rod nuclei within the outer nuclear layer has been noted as a differential criterion in the teleosts (Brackevelt, 1985; Collin et al., 1996a) and mammals (Cohen, 1972; Braekevelt, 1982).

The confirmation of three photoreceptor types in *L. salamandroides* has been found in a number of early actinopterygian representatives such as the redfish, *Calamoichthys calabaricus* (Munk, 1969), the bowfin, *Amia calva* (Munk, 1968) and the garfish, *Lepisosteus sp.* (Munk, 1968; Collin and Collin, 1993) which all possess double cones, single cones and large rods. Similarly, although not closely related to the teleostean fishes, the Australian (*Neoceratodus forsteri*) and African (*Protoperus aethiopicus*) lungfishes both possess double cones, single cones and large rods. Therefore, in support of the phylogenetic inferences proposed by Fraenkel (1968) and Rosen (1973, 1974) on osteological criteria, who both agree that *L. salamandroides* is as primitive as any living teleost and together with the Australian lungfish are “living fossils”, morphological features of the photoreceptors of the salamanderfish may also provide a basis for phylogenetic comparisons.

Double cones are common components of the photoreceptor mosaic in a number of teleost species. The advantage of the cone doublet over single cones is unknown but sensitivity may be raised due to its increased retinal coverage and therefore enhanced quantum capture in low light environments (Boehlert, 1979; Lythgoe, 1979). The arrangement of the double cones into a regular mosaic may also increase the detection of moving objects without the advantage of an area centralis or fovea (Collin S.P. and Gill H., unpublished data) and provide a means of chromatically sampling the same part of the visual field with two receptors of different spectral sensitivity. Increasing sensitivity would be of particular importance for *L. salamandroides* since it survives in darkly-stained water and needs to optimise its perception of small mobile prey which it tracks by flexion of its head and “neck” (Berra and Allen, 1989).

The presence of a parabolid situated below the ellipsoid in the single cones of *L. salamandroides* emphasises the primitive origin of the photoreceptors in this species. The existence of a parabolid appears to be a pleiomorphic (primitive) character in the phylogeny of the Actinopterygii due to its presence in all the living representative outgroups i.e. the Polypteroidei (bichirs, *Munk, 1969*), Acipenseridae (sturgeons, although lost in the rods, *Munk, 1969*), Polyodontidae (paddlefishes, *Munk, 1969*), the Ginglymodi (garfishes, Collin and Collin, 1993) and the Halecomorphi (bowfins, *Munk, 1969*). Until now, there have been no known representatives of the Teleostei to have retained a parabolid in their retinal photoreceptors. Alternatively, the presence of a parabolid may be an apomorphic (derived) trait for gnathostomes, since it has not been found in the photoreceptors of agnathan and chondrichthyan and persists in the Dipnii (lungfishes, Pfeiffer, 1968; Munk, 1969; Locket, 1970), Amphibia, Reptilia and Aves (Wals, 1942; Detwiler, 1943). The two types of granules found in the parabolid of *L.*
salamandroids putatively contain glycogen. Although the site of this glycogen synthesis and its metabolic pathway are not well understood, it may serve as an energy store for the myoid (Rodieck, 1973).

The high proportion of large rods and their unique position within the square mosaic of the retina of L. salamandroides is based on the assumption that the morphological characteristics traditionally used to classify rods and cones are appropriate. The features that separate the large rods from the other two cone types in the salamanderfish include the close position to the retinal pigment epithelium, a long, non-tapering outer segment, longitudinal incisures, a long myoid region, a wider spacing between the disc membranes (Cohen, 1969), the enclosure of the outer segment discs by an external membrane and a mitochondrial size gradient in the inner segment (vitreo-scleral in the single cones and centro-peripheral in the double cones but absent in the rods). Although, some features described here for rods may be common to cones in other species of teleosts, the full complement of criteria suggest our classification is correct. However, physiological and microspectrophotometric analyses are planned to further characterise these photoreceptor types.

The presence of a large number of rods is usually associated with a nocturnal habitat or life in deep water. However, for species living in turbid (labrids, Finner and Nicol, 1974) or darkly-stained water (lepidogalaxids, this study) where the light levels may be constantly changing and often low, a rod-rich retina would be a distinct advantage. The rods of L. salamandroides are also very large in comparison to other vertebrates. In the light-adapted state, these rods are 0 µm in length and 6.0 µm in width. This compares to 30 µm in length and 2 µm in width in the marine European eel Anguilla anguilla (Braeckevelt, 1988), 40 µm in length and 4 to 5 µm in width in both the short-tailed sargray, Dasylus breviscudata (Braeckevelt, 1994) and the northern pike, Esox lucius (Braeckevelt, 1975) but falls short of the long rods of the goldeye, Hiodon alosoides (135 µm, Braeckevelt, 1982) and some deep-sea teleosts (340 µm in Sculpelarchus anolis, Collin et al., 1998). The environmental influences which have induced the evolution of a large rod in L. salamandroides is not understood but due to the increase in optical density of the outer segment, these large rods would effectively lower the threshold of visual sensitivity (Bassi and Powers, 1990) and may act as a macrorceptor.

The function of the rod incisures in L. salamandroides is unknown, but there may be some advantage to increasing the length of the disk perimeter and decreasing its surface area, thereby aiding in the diffusion of substances to and from the disk membrane (Rodieck, 1973). In some amphibians, the incisures are particularly numerous and deep (Nilsson, 1965; Cohen, 1972), while in some reptiles (Dunn, 1966), birds (Cohen, 1963) and mammals (Kroll and Machemer, 1968), the disks show a single deep incisure or a series of scallops.

The calycal processes that emanate from the edge of the inner segment and sit in the grooves of the rod incisures in L. salamandroides are also little understood. These processes are widely distributed in all vertebrate classes and may have a supportive role in preventing the outer segment from rotating about the eccentrically situated connecting cilium (Rodieck, 1973).

This is the first detailed report of a rod that lies at the centre of a square mosaic. This position is usually reserved for a single cone which adopts the central position of the repeating unit where the orientation of the contiguous membrane of each double cone radiates from the centre like a four-spoked wheel. The closest description is one by Karsten (1923) who describes the photoreceptor mosaic in another amphibious species, the mudskipper, Periophthalmus koelreuteri, as a "quadangular" mosaic with four double cones bordering a rod. Further research is necessary to establish whether this photoreceptor pattern is common to an amphibious lifestyle, as may be predicted for the Australian lungfish, or simply a developmental phenomenon designed to maximise chromatic sampling using this complement of spectrally-sensitive receptors.

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Retinal morphology of the salamanderfish


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