Immunocytochemical Identification of Primary Olfactory Afferents in Rainbow Trout

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ABSTRACT
We have used a combination of techniques to analyze the primary olfactory projection in trout: anterograde tract tracing with horseradish peroxidase (HRP) and immunocytochemistry with antisera to olfactory marker protein (OMP) and to keyhole limpet hemocyanin (KLH). HRP labeling and the OMP antiserum revealed a subset of ciliated receptor neurons with a wide dendrite that lacked the protruding knob found on other receptor neurons. The organization of the primary olfactory axons was clearly revealed by antiserum to KLH, which reacted with no other neurons. When visualized with anti-KLH, fascicles of olfactory axons penetrated the basal lamina of the olfactory rosette at scattered sites and converged to form the olfactory nerve. Fascicles within the olfactory nerve traveled parallel to the long axis of the nerve until resorted by extensive intermixing as they entered the olfactory bulb. Within the olfactory bulb, most axons terminated in nine discrete terminal fields in the glomerular layer; however, a few olfactory nerve axons projected into the ventral medial telencephalon. Fascicles supplying each terminal field in the glomerular layer followed distinctive trajectories within the olfactory nerve layer. Axons ending in two terminal fields made brush-like terminations rather than the glomerular terminations characteristic of the remaining seven fields. After unilateral olfactory nerve transection, returning olfactory axons reestablished the normal pattern of terminal fields within 14 weeks. It is likely that the organization of afferents in the trout olfactory bulb is similarly well regulated during normal receptor cell replacement.

Key words: axons, regeneration, olfactory marker protein, olfaction, fish

In studies of teleostean olfactory systems, several investigators have described the concentric arrangement of cell types and resulting lamination of the olfactory bulb (Johnston, 1898, '02; Sheldon, '12; Holmgren, '20; Droogleever-Furtuyn, '61; Story, '64; Ichikawa, '76; Bass, '81a; Oka, '83). The laminar organization of teleost olfactory bulbs is less pronounced but generally comparable to that in mammals (reviewed in Allison, '53; Nieuwenhuys, '67; Andres, '70; Alonso et al., '89a). Studies using degeneration, axonal tracing, and electrophysiological techniques have demonstrated that the efferent axons arising from the medial and lateral regions of the olfactory bulb project primarily through the medial and lateral olfactory tracts, respectively, to partially segregated central projections (Scalia and Ebbesson, '71; Ito, '73; Braford and Northcutt, '74; Finger, '75; Satou et al., '79; Dubois-Dauphin et al., '80; Bass, '81b). This division into medial and lateral pathways may reflect the segregation of odor information associated with reproduction and feeding (Doving and Selset, '80; Stacey and Kyle, '83; Sorensen et al., '88; Satou, '90). Given the possibility of such functional segregation of the second order projections in the olfactory system, it is important to understand how the primary olfactory axons are organized and distributed within the glomerular layer of the olfactory bulb.

Recent anterograde and retrograde mapping experiments in trout have demonstrated that each point in the glomerular layer receives information from the entire olfactory rosette (Riddle and Oakley, '91). Here we extend our studies of the organization of the primary olfactory projection in trout, using tract tracing and immunocytochemical labeling to demonstrate major morphological subdivisions within that projection, which may reflect the segregation of axons into functional groups within the olfactory bulb. Olfactory receptor neurons were exposed to HRP applied to the mucosal surface in order to anterogradely label their projections to the olfactory bulb. In other trout, the distri...
bution of olfactory axons was characterized by immunocytochemistry. Neuroanatomical investigations of mammalian and amphibian olfactory systems have benefitted from the use of several antibodies and lectins that react with olfactory receptor neurons (Allen and Akeson, '85a, b; Fujita et al., '85; Hempstead and Morgan, '85a, b; Mori et al., '85; Schwob and Gottlieb, '86; Key and Giorgi, '86a, b; Morgan, '88; Barber, '89; Shinoda et al., '89). The most useful markers permit one to trace the trajectory of an identified group of neurons from source to termination in normal tissue. One of the antisera employed in the present study, anti-olfactory marker protein (anti-OMP), has been shown to bind to olfactory receptor neurons in a variety of vertebrates (Margolis, '80; Chua and Zheng, '87; Baker et al., '89). In fish, OMP-like immunoreactivity (OMP-IR) has been detected previously by radioimmunoassay (Margolis, '80); the present study is the first immunocytochemical demonstration of OMP-IR in fish. In addition to the antiserum against OMP, we used antisera to keyhole limpet hemocyanin (anti-KLH) as a novel marker of olfactory receptor neurons.

As well as characterizing the organization of the primary olfactory projection in trout, this study provides new information regarding two other issues of interest to those studying the chemical senses of fishes—the presence of multiple receptor cell types and of a direct projection of primary olfactory afferents into the telencephalon. A number of investigators have demonstrated that there are at least two distinct types of olfactory receptor neurons in teleosts (reviewed in Yamamoto, '82, see also Rhein et al., '81; Zielinski and Hara, '88) and in mammals (Rowley et al., '89). One cell type bears a distinct olfactory knob with long cilia. The other has a less prominent apical knob bearing numerous microvilli. It has been suggested that in salmonid fishes (Salmo alpinus L., Salvelinus fontinalis, and Oncorhynchus mykiss), the two cell types represent functionally distinct classes (Thommesson, '82, '83), that microvillous cells respond to amino acids and ciliated cells respond to bile salts. Other investigators, however, have been unable to find such a correlation in the catfish (Ictalurus punctatus) between morphology and functional specificity (Erickson and Caprio, '84). Even as the functional significance of these two cells types remains unresolved, other studies have provided evidence for yet a third class of receptor cells. Horseradish peroxidase (HRP) applied to the olfactory nerve or bulb of goldfish (Carassius auratus) and carp (I. punctatus) labeled ciliated cells with a wide dendrite terminating at the surface of the epithelium (Type II ciliar cells, Muller and Marc, '84). We present additional evidence from both HRP and immunocytochemical studies that supports the existence of Type II ciliar receptor cells in trout, suggesting that at least three different olfactory receptor cell types may be present in teleosts.

In addition to finding several morphological types of olfactory receptor neurons, these studies provide further evidence that a small subset of olfactory axons terminates caudal to the olfactory bulb. Each marker of olfactory receptor neurons used in this study also labeled olfactory nerve axons that projected through the olfactory bulb into the ventral-medial telencephalon.

MATERIALS AND METHODS

Rainbow trout (Oncorhynchus mykiss), 18–30 cm long, were obtained from Spring Valley Trout Farm (Dexter, MI) and maintained in 75 gallon aquaria at 14–16°C. Trout were anesthetized for surgery with tricaine methane sulfonate (MS-222, 100 mg/l of aquarium water), wrapped in a wet towel, packed in ice and immobilized in a polystyrene holder. A constant stream of chilled aquarium water containing the anesthetic (MS-222, 80 mg/l) flowed across the gills.

One olfactory mucosa in each of two normal fish was extensively labeled with horseradish peroxidase (HRP, Sigma type VI) so that the distribution of HRP-labeled olfactory axons could be examined. A pledget of gelfoam soaked in 5% HRP was inserted through the anterior naris and placed against the olfactory rosette. The ipsilateral nares were then sealed with Parafilm and cyanoacrylate glue. Three weeks later the fish were re-anesthetized and perfused sequentially with heparinized physiological saline, fixative (1% paraformaldehyde and 1.25% glutaraldehyde in phosphate buffer), and 10% sucrose in phosphate buffer. Cryostat sections of the telencephalon and attached olfactory bulbs, the olfactory nerves, and the olfactory mucosae were thaw-mounted on gelatin-coated slides. To visualize the HRP label, a reaction mixture of tetramethyl benzidine (TMB, Mesulam, '82). Alternate sections of the mucosa were treated with diaminobenzidine (DAB), according to the intensification method of Adams ('81). The somal depth was determined for HRP-filled cells in the material treated with DAB. For a labeled cell that appeared in its entirety in a single section, the distance from the epithelial surface to the center of the nucleus was measured with a calibrated eyepiece reticle.

Seven normal fish were anesthetized with MS-222 and perfused with immunocytochemistry with heparinized physiological saline followed by fixative (6% HgCl2, 1% sodium acetate, 0.1% glutaraldehyde). Six other normal trout were perfused with alternative fixatives (Bouin's fluid, 4% paraformaldehyde, or 70% ethanol/10% acetic acid).

All tissue samples were dehydrated through graded alcohol, cleared in xylene and embedded in paraffin. Sections were cut at 7–10 μm and mounted on gelatin-coated slides. Following rehydration, the HgCl2-fixed tissue was treated with alcoholic iodine followed by 5% sodium thiosulfate to remove residual mercury. All tissue was preincubated in 0.3% H2O2 to block endogenous peroxidase activity, followed by 0.5% normal serum.

Sections of olfactory mucosae and bulbs fixed with Bouin's fluid, ethanol/acetic acid, or HgCl2 were incubated with antisera to olfactory marker protein (anti-OMP, gift of F. Margolis, Roche Institute). The tissue was incubated in normal rabbit serum (NRS) followed by the OMP antiserum diluted 1:200 in NRS. Antibody binding was visualized by means of a biotinylated rabbit anti-goat secondary antibody (Sigma, St. Louis, MO), avidin-biotin-peroxidase complex (ABC; Vector Labs, Burlingame, CA) and DAB (Sigma).

In addition to the OMP antiserum, four commercially available polyclonal antisera were tested for reactivity in the olfactory system of the trout. Three of the antisera were raised against neuropeptides conjugated to KLH: luteinizing hormone-releasing hormone (anti-KLH/LHRH, Incstar, Stillwater, MN), substance P (anti-KLH/SP, Incstar), and vasoactive intestinal peptide (anti-KLH/VIP, Amersham, Arlington Heights, IL). The three antisera were originally obtained to investigate the presence of the neuropeptides in the brain of the trout. After observing that the three antisera produced identical patterns of labeling and that the labeling could not be blocked by preadsorption with the
the KLH to which the peptides had been conjugated. This observed reactivity actually resulted from antibodies against three antisera by preadsorption with KLH and also by using an antiserum against unconjugated KLH (U.S. Biochem, Cleveland, OH). The antiserum against KLH and KLH conjugates were used at final dilutions from 1:1,000 to 1:30,000 in normal goat serum (NGS). Antibody binding was detected with biotinylated goat anti-rabbit IgG (Sigma), ABC, and DAB. Immunocytochemical controls included substitution of the primary antiserum with NGS, substitution of the secondary antibody or ABC with buffer, preadsorption of the antiserum to peptide conjugates with the appropriate neuropeptide (Sigma, 10–100 μg/ml antibody solution), and preadsorption of each antiserum with KLH (U.S. Biochem., 10 μg/ml antibody solution).

In addition to our studies of normal trout, we lesioned the olfactory system in 18 trout in order to verify the identity of the neuronal elements labeled by anti-KLH and to compare the organization of the normal and regenerated systems. The fish were anesthetized as described above and a small hole was drilled in the midline of the skull at the level of the posterior margin of the orbit. The hole was enlarged with fine rongeurs to expose the olfactory bulbs and posterior portion of the olfactory nerves. In 9 of 18 trout, one olfactory nerve was transected approximately 1 mm anterior to the olfactory bulb and the cut ends of the nerve reapposed. In seven other trout, the olfactory bulb was resected and removed along with the posterior portion of the olfactory nerve. Gelfoam controlled occasional bleeding and was used to fill the hole in the skull. The incision was closed with 5-0 nylon sutures (Ethicon). Finally, in each of the two remaining trout, one olfactory rosette was exposed and extirpated with microscissors. The flow of fresh aquarium water across the gills revived the fish and restored vigorous opercular movement. After 2 to 14 weeks of recovery, the fish were re-anesthetized with MS-222 and perfused with the HgCl₂ fixative as described above. Sections from both the lesioned and contralateral sides of the olfactory system of experimental fish were processed simultaneously so that immunoreactivity could be compared in lesioned and control tissue processed under identical conditions.

Following immunocytochemical labeling, we made camera lucida drawings of the glomerular layer from every fourth section of one olfactory bulb from each of four normal trout. Using a digitizing tablet and a microcomputer, we determined the total volume of the glomerular layer, the volume of each of the nine terminal fields, and the percentage of the total glomerular layer volume in each terminal field (See Table 1 footnote). In addition, for two fish in which the olfactory nerve had been transected 9–14 weeks prior, two-dimensional representations of the size and position of the normal and regenerated terminal fields were prepared and analyzed with a digitizing tablet and computer (See Fig. 10 legend).

RESULTS

The gross anatomy of the primary olfactory system in trout is similar to that in other salmonoids (Pfeiffer, '63; Yamamoto, '82). The olfactory epithelium is located on the broad surfaces of 12–16 lamellae that radiate from the base of each of the bilaterally paired olfactory rosettes. The edges and tip of each lamella are covered with nonsensory epithelium, as are secondary folds that separate strips of olfactory epithelium on each face of the lamellae. The olfactory nerve, approximately 1 cm long in a 25 cm trout, emerges from the ventral posterior region of each rosette and projects to the ipsilateral olfactory bulb. The olfactory bulbs are sessile, i.e., directly attached to the telencephalon.

The results of the HRP and anti-OMP studies will be presented first, followed by the investigations of the normal and regenerated olfactory system using the novel marker of olfactory receptor neurons, anti-KLH.

HRP labeling

HRP applied to the olfactory epithelium labeled two distinct morphological classes of cells. Each cell had a fine basally directed process (arrows in Fig. 1A,B). The apical process of some cells ended in a 1–2 μm diameter knob that extended slightly above the surface of the epithelium (small arrowhead in Fig. 1B). The remaining HRP-filled cells had an apex approximately 4 μm wide that ended flush with the epithelial surface (large arrowheads in Fig. 1A,B). All HRP-labeled cells fit into these two classes; there were no labeled cells with wide protruding apices or with narrow flat apices. The somata of labeled cells with wide apical processes were usually deeper in the olfactory epithelium than those of the cells bearing olfactory knobs. In two animals, the average somal depth of 280 HRP-filled receptor cells with dendritic knobs was 26 ± 6 μm (mean ± S.E.M.), while the mean somal depth of 400 cells with wide, flat apices was 38 ± 8 μm. HRP-filled axons were distributed throughout the olfactory nerve layer and glomerular layer of the ipsilateral olfactory bulb but were absent from the contralateral olfactory bulb (Riddle and Oakley, '91). The distribution of labeled axons within the glomerular layer was not continuous; rather, the labeled fibers appeared to terminate in several discrete fields (e.g., Fig. 1C). The pattern of terminations appeared similar in both of the labeled fish. Fascicles of labeled axons projected through the olfactory bulb and into the ventral medial telencephalon.

Olfactory marker protein

Cells immunoreactive for the OMP antiserum were unevenly distributed in the olfactory epithelium. In some regions, immunoreactive cells were present every 3–5 μm, while in other areas, they were absent from stretches of olfactory epithelium more than 100 μm long. All immunoreactive cells were intensely and similarly labeled. Each OMP-like immunoreactive cell had a stained axon, unstained nucleus, and a darkly stained dendrite that extended to the surface of the epithelium. The dendrites of a few immunoreactive cells ended in apical knobs, but most immunoreactive dendrites had a wide apex approximately 4 μm in diameter, which terminated flush with the epithelial surface (Fig. 2A). OMP-like immunoreactivity was also evident in axonal bundles beneath the olfactory epithelium, in the olfactory nerve, and in the olfactory nerve layer and glomerular layer of the olfactory bulb (Fig. 2B). Some immunoreactive axons projected from the olfactory nerve through the olfactory bulb and into the ventral medial telencephalon, following the same path as the forebrain projection labeled by HRP. No other neurons in the forebrain were immunoreactive.

Omission of the OMP antiserum, the secondary antibody, or the ABC eliminated labeling. No attempt was made to...
Fig. 1. Horseradish peroxidase (HRP) labeling in the primary olfactory system. A, B: Several HRP-filled cells are visible in these sections through the olfactory epithelium. Two distinct morphological classes of labeled cells are evident, one characterized by a narrow apex topped by an olfactory knob (small arrow), the other characterized by a wider apex terminating flush with the epithelial surface (arrowheads). Bar = 10 μm. C: Horizontal section through one olfactory bulb three weeks after the ipsilateral olfactory rosette was labeled with HRP (darkfield). The olfactory nerve enters from the left. At this level along the dorsal-ventral axis of the olfactory bulb, HRP-labeled axons project into three discrete terminal fields, anterior medial (small arrowhead), lateral (large arrowhead) and posterior lateral (small arrow). No HRP-labeled axons extend into the deeper laminae of the olfactory bulb. Bar = 250 μm.
block the binding of the OMP antisera by preadsorption, thus it remains to be determined whether the molecule in trout recognized by the antisera is the same as mammalian OMP.

**KLH-like immunoreactivity**

*Specificity of labeling.* Antisera against KLH and against KLH conjugated to neuropeptides reacted intensely with trout olfactory receptor axons (see Fig. 3A). The four antisera produced almost identical labeling; anti-KLH/VIP produced slightly lower background reactivity than other equidiluted antisera. Reactivity was minimally affected by variation in fixation, although paraformaldehyde reduced the background slightly. Omission of the primary antibody, or ABC, eliminated labeling. Preadsorption of each antisera to a KLH/peptide conjugate with the corresponding peptide (e.g., preadsorption of anti-KLH/VIP with VIP) did not affect reactivity, even at an antigen concentration of 100 μg/ml of antiserum (Fig. 3A). We were unable to confirm the findings of Alonso et al. ('89b, '90), who, using anti-VIP from another source, reported specific VIP-like immunoreactivity in rainbow trout olfactory nerve and bulb. Preadsorption of each of the four antisera with KLH (10 μg/ml antiserum) completely eliminated labeling (Fig. 3B). Since the four antisera produced identical patterns of reactivity, and since all immunoreactivity was eliminated by preadsorption with KLH while preadsorption with the neuropeptides had no effect, we concluded that all labeling was due to antibodies against KLH, and hence, we refer to all reactivity as KLH-like immunoreactivity (KLH-IR). Characterizing the antigen(s)
recognized by the KLH antisera lay outside of the scope of the present study.

**KLH-IR in the normal olfactory system.** Although KLH-IR was evident in both sensory and nonsensory epithelia in the olfactory mucosa, the pseudostratified columnar olfactory epithelium was easily distinguished from the less organized nonsensory epithelium covering the lamellar ridges. KLH-IR in the olfactory epithelium was prominent in the membranes of spindle-shaped cellular processes oriented perpendicularly to the surface and extending from the middle of the epithelium to the apical surface (Fig. 4A). The intense immunoreactivity at the surface of the olfactory epithelium made it difficult to resolve the apical endings of individual cells. Nor could we easily distinguish olfactory receptor cell somata from those of supporting cells. Scattered large mucus cells in the olfactory mucosa were also immunoreactive (arrowhead in Fig. 4A). KLH-IR was intense in a few 6–8 µm diameter spheroidal cells scattered throughout the core of each lamella and in the base of the olfactory rosette. We could not resolve individual immunoreactive axons within the olfactory epithelium, but fascicles of immunoreactive axons, 2–8 µm in diameter, were located at intervals of twenty to several hundred micrometers directly beneath the basal lamina (small arrows in Fig. 4A). Each immunoreactive fascicle projected from the basement membrane into the central core of the lamella and joined larger bundles of axons running parallel to the long axis of the lamella (large arrow in Fig. 4). The axonal bundles from all lamellae merged in the base of the rosette to form the olfactory nerve. KLH-IR was intense throughout the length of the olfactory nerve. Cross-sections of the nerve contained thousands of dispersed immunoreactive elements 2–5 µm in diameter, approximately the same size as the glially limited axonal bundles apparent in transmission electron micrographs of the olfactory nerve of pike (Kreutzberg and Gross, '77) and trout (Riddle and Oakley, unpublished observations).

Within the trout brain, KLH-IR was largely limited to the primary olfactory afferents in the olfactory bulb and a small bundle of fibers that extended into the telencephalon. Immunoreactivity was intense in the olfactory nerve layer and glomerular layer (see Fig. 3A). No labeled neurons were apparent in the deeper layers of the olfactory bulb, but blood vessels were lightly labeled throughout the brain. A few small immunoreactive cells that lacked processes were scattered in the meningeal and periventricular regions of normal brains. Such cells were more common in lesioned fish. Their greater concentration around the lesion site suggests reactive gliosis.

The immunoreactive fibers in the glomerular layer ended in nine separate terminal fields (Figs. 8-10). The positions of the nine fields were similar in all of the fish that we examined, and the relative size of each field was constant across the four individuals in which the size of the fields was measured. The largest terminal field comprised 35% of the total volume of the glomerular layer and the smallest about 1% (Table 1). Axonal fascicles, 2–6 µm in diameter, often ended cohesively in spheroidal glomerular terminations up to 50 µm in diameter (Fig. 8A, B), resembling the terminations described in other species (reviewed in Allison, '53; Nieuwenhuys, '67; Andres, '70). In contrast, olfactory axons in the two posterior lateral terminal fields ended in a diffuse brush pattern (e.g., Fig. 5C). To reach a given field, the fibers followed distinctive trajectories through the olfactory nerve layer. For example, the axons terminating in the posterior lateral field (PL) always coursed posteriorly around the lateral face of the olfactory bulb. In contrast, the axons that supplied the ventral posterior lateral field (VPL), which lies immediately adjacent to PL, coursed along the ventral aspect of the bulb.

A discrete bundle of KLH-like immunoreactive fibers projected through the ventral medial olfactory bulb into the telencephalon (arrows in Fig. 6A), along the same path as axons labeled by HRP (Fig. 6B) and by anti-OMP (Fig. 6C). In some individuals, a few fibers from the ventral lateral olfactory nerve layer joined the bundle as it entered the
MARKER FOR Olfactory Receptor Axons in Trout

TABLE 1. Relative Size of the Nine Terminal Fields in the Trout Olfactory Bulb

<table>
<thead>
<tr>
<th>Terminal field</th>
<th>Volume in percent (mean ± S.D.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dorsal lateral (DL)</td>
<td>17 ± 2.0</td>
</tr>
<tr>
<td>Anterior medial (AM)</td>
<td>30 ± 2.0</td>
</tr>
<tr>
<td>Dorsal posterior medial (DPM)</td>
<td>1 ± 0.2</td>
</tr>
<tr>
<td>Dorsal posterior lateral (DPL)</td>
<td>2 ± 0.6</td>
</tr>
<tr>
<td>Lateral (L)</td>
<td>35 ± 0.8</td>
</tr>
<tr>
<td>Posterior lateral (PL)</td>
<td>2 ± 0.6</td>
</tr>
<tr>
<td>Ventral posterior lateral (VPL)</td>
<td>1 ± 0.2</td>
</tr>
<tr>
<td>Ventral posterior (VP)</td>
<td>1 ± 0.3</td>
</tr>
<tr>
<td>Ventral medial (VM)</td>
<td>10 ± 0.6</td>
</tr>
</tbody>
</table>

*The volume of each field was determined by measuring its area in every fourth horizontal section, multiplying by the section thickness (10 μm) and section interval (4), and summing the sections for each field. For each of four animals, the volume of each field was divided by the total volume of the glomerular layer, i.e., the sum of all nine fields. The mean percentage volume for each field was then calculated for the four animals.

The bundle of KLH-like immunoreactive fibers coursed into the ventral medial telencephalon, where some axons appeared to terminate diffusely, a few crossed the anterior commissure and others proceeded posteriorly.

**KLH-IR after transection of the olfactory nerve, olfactory bulbectomy, and extirpation of the olfactory rosette**

The changes in KLH-IR detected after unilateral lesions of the olfactory system were limited to ipsilateral structures.

**Olfactory mucosa and nerve.** Olfactory nerve transection produced a moderate decrease in KLH-IR in the olfactory epithelium and a marked, but transient, decrease in the axons in the lamellar core and in the olfactory nerve (Fig. 7). The maintenance of some immunoreactivity after nerve transection was likely due to the labeling of supporting cells (see Discussion). The loss of immunoreactivity in olfactory axons was most apparent four weeks after transection of the nerve (n = 2). Axonal bundles within the olfactory rosette regained their normal KLH-IR by 9 to 14 weeks after transection (n = 5), while KLH-IR within the olfactory nerve was still reduced. There were similar changes in KLH-IR in the olfactory mucosa and nerve after olfactory bulbectomy; KLH-IR decreased markedly in the axonal bundles within the rosettes and in the olfactory nerves.

**Olfactory bulb.** Olfactory nerve transection eliminated KLH-IR in the olfactory bulb within two to four weeks (Fig. 8A), except for a few KLH-IR fibers in the dorsal olfactory bulb and in a discrete group of glomeruli in the ventral medial terminal field (VM). Partial reinnervation of the olfactory bulb was evident six weeks after nerve transection, with the immunoreactive fibers most evident in the ventral and dorsal lateral regions of the olfactory bulb (n = 2). With longer survival periods, the axonal density increased and scattered fibers appeared laterally and posteriorly (9–10 weeks, n = 3, Fig. 8B). By nine weeks, some olfactory axons were present in all terminal fields. Even fields remote from the entry point of the olfactory nerve, such as the posterior lateral field, received some

See Fig. 5. KLH-IR in the olfactory bulb. A: Horizontal section through the olfactory bulb, approximately in the middle along the dorsal-ventral axis. Axons terminating in the anterior (enlarged in B) and lateral terminal fields form glomerular endings (arrowheads in B). Those that terminate in the posterior lateral field form non-glomerular brush-like terminations (enlarged in C). Bars = 250 μm (A), 100 μm (B) and 50 μm (C).
MARKER FOR Olfactory RECEPTOR AXONS IN TROUT

583

axons relatively early in the process of reinnervation. Thus, reinnervation did not proceed as a smooth wave that filled the anterior sites before advancing to the more posterior fields. Reinnervating axons appeared to return more or less contemporaneously throughout the glomerular fields in rough proportion to the normal density (Fig. 8B).

Reinnervation of all regions of the olfactory bulb increased with longer survival, but the apparent density of immunoreactive fibers in all terminal fields was still clearly below normal after 14 weeks (n = 2, Fig. 9). The denervated olfactory bulb was noticeably smaller than the contralateral bulb due to marked thinning of the olfactory nerve layer (Fig. 9A). From 9–14 weeks, the pattern of terminal fields in the reinnervated olfactory bulbs matched that on the control side (Figs. 8–10).

Two fish euthanized six weeks after removal of one olfactory rosette had no KLH-IR fibers in the ipsilateral olfactory bulb, except in a small region in the ventral medial olfactory bulb where a few glomeruli remained. It is possible that the surviving fibers were from the contralateral nerve; contralateral projections by olfactory fibers have been reported in Xenopus and Rana (Ebbesson et al., '86; Duncan et al., '90). There was, however, no loss of immunoreactive fibers in the corresponding region of the olfactory bulb contralateral to the lesioned nerve. Further, we observed no contralateral projections in the HRP preparations, although it is technically possible that the ventral medial glomeruli originated from receptor cells that failed to take up HRP and were therefore not visualized by that method.

**Telencephalic fibers.** The telencephalic fascicle of KLH-immunoreactive fibers was absent ipsilaterally from fish euthanized four weeks after nerve transection. The telencephalic fascicle reappeared by six weeks after surgery, but remained small even after 14 weeks (see Fig. 8C). The ipsilateral telencephalic fascicle was absent in all five bullectomized fish, and was also eliminated by ablation of the olfactory rosette.

**DISCUSSION**

**Morphological types of olfactory receptor neurons**

Previous investigations of the teleost olfactory system have demonstrated two receptor cell types, microvillous and ciliated. Both cell types have distinct apical knobs of roughly similar diameter, although the knob of microvillous cells is somewhat less prominent (Evans and Hara, '78; Rhein et al., '81; Muller and Marc, '84; Zielinski and Hara, '88). While some evidence from studies of salmonids has suggested that the two morphological types are also functionally distinct (Thommessen, '82, '83), studies using the channel catfish have not supported this functional distinction (Erickson and Caprio, '84). Thus, the functional significance of multiple cell types remains unclear. In the present study, HRP applied to the olfactory mucosa labeled some cells with olfactory knobs, and another class of cells with wider dendrites that lacked apical knobs. Rhein et al. ('81) identified similar “Type II ciliated cells” in the olfactory mucosa of trout without commenting on their function. Muller and Marc ('84) concluded that similar cells in the olfactory mucosa of goldfish and catfish were receptor neurons since they were retrogradely labeled by HRP injected into the olfactory bulb or applied to the olfactory

![Image](https://example.com/image.png)
nerve. Since HRP was applied to the surface of the mucosa in this study, it is possible that some labeled cells with flat heads were supporting cells. Most of the cells, however, appeared to have axons. Further, we noted in an earlier study that cells with similarly wide, flat apices were labeled by fluorescent beads injected into the olfactory bulb (Riddle and Oakley, '91). Moreover, in the present study, cells with identical morphology were intensely stained by antisera to OMP. OMP-IR has been recognized as a relatively specific marker for mature olfactory receptor neurons in other vertebrates (Margolis, '80; Baker et al., '89). It remains to be determined, of course, whether the molecule recognized by the OMP antisera is the piscine homologue of mammalian OMP. Regardless, the fact that axons throughout the olfactory nerve layer and glomerular layer were OMP-positive while there were virtually no OMP-positive cells with the classical morphology of receptor neurons (i.e., with knobs), suggests that the Type II ciliar cells in the OMP preparations were mature receptor neurons. This argument assumes only that the OMP antiserum recognizes the same population of receptor cells in the epithelium and in the olfactory bulb. Taken together, the observations with OMP antiserum, anterograde HRP labeling and retrograde labeling with fluorescent beads make it likely that the flat-headed cells are olfactory receptor cells, with some sufficiently mature to have extended axons into the olfactory bulb.

We considered the possibility that the olfactory receptor cells with dendritic knobs and those with wide, flat apices represented the extremes of a morphological continuum associated with receptor cell maturation. It is often noted that the dendrites of olfactory receptor cells in mammals become stouter as the cells mature and their somata migrate toward the epithelial surface. This cannot explain the wide dendrites of the Type II cells since their somata were located more deeply in the epithelium than the somata of the cells with distinct dendritic knobs. Thus, if the different morphologies represent different maturational states, in fish the Type II cells with wider dendrites would be the younger cells.

Further investigations will be required to characterize the different morphological types of olfactory receptor neurons, and to evaluate the significance of those putative olfactory receptor cells that displayed OMP-like immunoreactivity. It remains to be determined whether these morphologically distinct cells are equally distinctive in their response properties. Future investigations may be facilitated by the use of markers like those applied in this study, since our observations indicate that the different cell types differ somewhat in their antigenic properties.

Axons of the olfactory nerve make direct projections into the telencephalon

A number of studies using immunocytochemical labeling and tract tracing have demonstrated neural connections between the olfactory mucosa and various telencephalic regions posterior to the olfactory bulb. Demski and Northcutt reported that HRP applied to the olfactory mucosa of

Fig. 8. KLH immunoreactivity in horizontal sections of the olfactory bulb 2–10 weeks after unilateral olfactory nerve transection. A: The ipsilateral olfactory bulb (ob) had no KLH-IR 14 days after transection of the right olfactory nerve. B: Significant reinnervation is evident 9 weeks after transection of the right olfactory nerve. Some axon terminations are present even in the lateral field (arrowheads) and in the posterior lateral field (arrow), which corresponds exactly with the position of the posterior lateral field on the normal side (arrow). C: Ten weeks after transection of the left olfactory nerve, some KLH-immunoreactive axons are present (small arrows) that resemble the more robust telencephalic projections on the normal side (large arrows). Bar in C = 40 μm for A–C.
Fig. 9. Four sequential (dorsal to ventral) horizontal sections through the olfactory bulbs of a trout 14 weeks after unilateral (left) olfactory nerve transection. Nine terminal fields can be identified in these sections in the right (control) olfactory bulb: A: The most ventral regions of the dorsal posterior medial (DPM) and dorsal posterior lateral (DPL) terminal fields are visible along with the anterior medial (AM) and dorsal lateral (DL) fields. B: Field AM continues ventrally where the lateral (L) and posterior lateral (PL) fields come into view. C: the goldfish labeled axons that projected into the ventral medial telencephalon. They concluded that the fibers belonged to the terminal nerve (Demski and Northcutt, '83; Demski, '84). Bazer et al. ('87) reported that cobaltous lysine applied to the olfactory mucosa of salmon fry labeled axons that followed a similar trajectory, but concluded that the fibers may have been labeled by transynaptic transport of the label. In the present study, telencephalic fibers were labeled by HRP applied to the olfactory mucosa of mature trout. It is unlikely that the labeled fibers were the result of transynaptic labeling since unconjugated HRP is not thought to cross synapses. Thus, our findings raise the possibility that the telencephalic fibers observed in Bazer's study may have been labeled directly, and that axons linking the olfactory mucosa and the ventral medial telencephalon may be present in both young and mature salmonids as well as in a variety of other fish (see von Bartheld et al., '88). The pathway followed by the fibers into and within the telencephalon of the trout is generally consistent with the location of terminal nerve axons in other fishes (Munz et al., '82; Munz and Claas, '87; Capron de Caprona et al., '86; von Bartheld et al., '88). If the fibers belong to the terminal nerve, however, it is somewhat surprising that we could not demonstrate gonadotropin releasing hormone (GnRH) reactivity, since immunoreactivity for GnRH is characteristic of teleostean, amphibian and mammalian terminal nerve axons (Jennes and Stumpf, '80; Schwanzel-Fukuda and Silverman, '80; Munz et al., '82; Stell et al., '84; Wirsig and Getchell, '86; Munz and Claas, '87; Schwanzel-Fukuda et al., '87; Stell and Walker, '87; von Bartheld et al., '88). Perhaps the axons projecting into the trout telencephalon would react to antibodies against teleostean GnRH (Kah et al., '86). However, two other observations suggest that the labeled telencephalic fibers were not part of the terminal nerve. Each marker of olfactory receptor neurons also labeled the telencephalic fascicle, consistent with a direct projection of primary olfactory axons into the ventral medial telencephalon. Of course it is possible that the antibodies recognized different antigens in receptor neurons and in the telencephalic fibers or that olfactory and terminal nerve fibers share the same antigen(s). The loss of the telencephalic fibers after ablation of the olfactory mucosa, however, implies that the somata associated with the telencephalic fibers were not located in the olfactory
Figure 10
nerve or in the telencephalon (in contrast to the ganglion cells of the known terminalis system). Rather, it seems more likely that the fibers were primary olfactory afferents. Such direct projections of primary olfactory afferents past the olfactory bulb and into the telencephalon were recently described in several gymnotid fishes (Gymnotus carapo, Hypopomus sp., Apteronotus leptorhynchus, Szabo et al., '91) and in the three-spined stickleback (Electrophorus electricus, Honkanen and Ekstrom, '90). Thus, our observations represent the third family of fishes in which such a direct projection of primary olfactory axons into the ventral medial telencephalon has been revealed.

**Organization of the normal and regenerated primary olfactory projections labeled with anti-KLH**

The antisera to KLH greatly facilitated the evaluation of the organization of the normal and regenerated primary olfactory projection in trout. The evidence of KLH-IR in both supporting cells and olfactory receptor cells may reflect their common derivation from the olfactory placode. Alternatively, it is possible that a molecule recognized by the antisera is secreted by one cell type and taken up by the other, or that different molecules in the two cell types share a common epitope. In any case, the maintenance of KLH-IR in the olfactory epithelium following olfactory nerve transection was probably due in part to the immunoreactivity of supporting cells, but may also reflect on-going and experimentally induced olfactory receptor neuron replacement. While KLH-IR was nonspecific in the mucosa, its high specificity in the brain for olfactory axons made it a valuable probe for characterizing the organization of these afferents. We infer from the pattern of anti-KLH labeling that olfactory axons undergo little fasciculation until they collect near exit points scattered along the basal lamina, since immunoreactive axons were not evident in the superficial olfactory epithelium. (Individual axons would probably be too small to detect—see below.) The trajectory of fascicles from the basal lamina to the olfactory bulb was easily followed in the anti-KLH preparations. With the possible exception of a few axons terminating in the VM terminal field, the KLH-like immunoreactive axons from each olfactory rosette reached the olfactory bulb through the ipsilateral olfactory nerve layer. Thus, immunoreactive fibers in the terminal field is slightly reduced in size in the reinnervated bulb; however, the relative sizes and the pattern of the terminal fields are quite similar. Bar = 0.5 mm for A–C.

**Fig. 10.** Schematic maps of the nine terminal fields shown in one normal olfactory bulb (A) and in two olfactory bulb 9 weeks (B) and 14 weeks (C) after transection of the olfactory nerve. Abbreviations are the same as in Figure 9. The maps in A and C were prepared from the olfactory bulbs shown in Figure 9, the positions of the horizontal sections shown in that figure (A through D) are indicated by dotted lines. These flattened two-dimensional representations of glomerular terminal fields were prepared from anti-KLH-stained horizontal sections. Camera lucida drawings were made of the boundaries of the terminal fields evident in every fourth horizontal section. The linear extent of each terminal field and the distance between fields was measured using a digitizing tablet. This produced for each horizontal section a straight line as long as the circumference of the glomerular layer with the terminal fields and inter-terminal regions indicated by tick marks. These lines were arranged from dorsal to ventral, using the center of the anterior face of the olfactory bulb as a fiducial mark. Terminal field boundaries were drawn by interconnecting the tick marks in this stack of lines. The left end of the dashed lines in each figure is the posterior medial extent of the glomerular layer, while the right end is the posterior lateral margin. Compared to the normal side each anterior fields was still subnormal while the posterior fields were being reformed. Despite the slow rate of olfactory nerve regeneration (Cancalon, '85), by three to four months after nerve transection the reinnervating axons terminated in the normal pattern of terminal fields, albeit with fewer axons (Figs. 8–10). While it is unknown how axon-target interactions contribute to terminal field formation, it seems likely that terminal field reconstitution following transection utilizes mechanisms similar to those employed during normal receptor cell turnover.

It is interesting to consider whether the parcellation into glomerular terminal fields in trout corresponds to functional segregation comparable to the macroglomerulus in the insect olfactory lobe (Boeckh and Boeckh, '79; Matsu-
moto and Hildebrand, '81; Hansson et al., '92) and the modified glomerular complex in the rodent olfactory bulb (Teicher et al., '80; Greer et al., '82). The macroglomerulus and the modified glomerular complex are believed to process information about specific odors of behavioral importance. Electrophysiological studies have demonstrated functional segregation in the olfactory bulb of aca-, char and rainbow trout (Salmonidae) that might correspond to the dorsal lateral (DL) and anterior medial (AM) terminal fields. Recording from the dorsal surface of the olfactory bulb, Thommesen and Doving found that amino acids elicited greater responses laterally while the re-

sponses to bile salts were greater medially (Thommesen, '78; Doving et al., '80). It should be possible to combine electrophysiological mapping with anti-KLH immunocytochemistry in order to characterize the responses within different terminal fields. It would be particularly interesting to evaluate the response characteristics of fibers in the posterior lateral glomerular layer. With their unusual brush-like terminations, these axons may have a special functional role. It should be noted that morphological differences in olfactory terminals are not limited to the trout; two regionally segregated classes of terminals have been reported in the olfactory bulb of the goldfish (Finger, '88).

Summary and conclusions

Antergrade tract tracing with HRP and immunocytochemistry with antisera against olfactory marker protein and keyhole limpet hemocyanin revealed the pathways of olfactory axons and the pattern of their terminations in the olfactory bulb of rainbow trout. These methods distinctly labeled olfactory receptor neurons, including a class of apparent receptor cells with a wide dendrite that lacked an olfactory knob and a subset of olfactory nerve axons that projected directly into the ventral medial telencephalon. The bulk of the labeled axons terminated in the glomerular layer of the trout olfactory bulb and were organized into nine distinctly positioned terminal fields, ranging in size from 1 to 35% of the glomerular layer. In some terminal fields, the olfactory axons made glomerular-like terminations, in others brush-like terminations. The nine terminal fields, which may reflect functional parcellation of olfactory information, were substantially reconstituted by 14 weeks after unilateral transection of the olfactory nerve. These observations on olfactory receptor cell types and the organization of their projections provide a foundation for continuing investigations relating the anatomical organization of the olfactory system to its function. In addition, the molecular markers described here will be useful for studying the development of the terminal fields and the means by which this organization is maintained during the continual replacement of olfactory receptor cells.

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