Correlation between Olfactory Receptor Cell Type and Function in the Channel Catfish

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The olfactory epithelium of fish contains three intermingled types of olfactory receptor neurons (ORNs): ciliated, microvillous, and crypt. The present experiments were undertaken to test whether the different types of ORNs respond to different classes of odorants via different families of receptor molecules and G-proteins corresponding to the morphology of the ORN. In catfish, ciliated ORNs express OR-type receptors and Goq11. Microvillous ORNs are heterogeneous, with many expressing Goq11, whereas crypt ORNs express Goa. Retrograde tracing experiments show that ciliated ORNs project predominantly to regions of the olfactory bulb (OB) that respond to bile salts (medial) and amino acids (ventral) (Nikonov and Caprio, 2001). In contrast, microvillous ORNs project almost entirely to the dorsal surface of the OB, where responses to nucleotides (posterior OB) and amino acids (anterior OB) predominate. These anatomical findings are consistent with our pharmacological results showing that forskolin (which interferes with Goa/cAMP signaling) blocks responses to bile salts and markedly reduces responses to amino acids. Conversely, U-73122 and U-73343 (which interfere with Goq/phospholipase C signaling) diminish amino acid responses but leave bile salt and nucleotide responses essentially unchanged. In summary, our results indicate that bile salt odorants are detected predominantly by ciliated ORNs relying on the Goq/cAMP transduction cascade. Nucleotides are detected by microvillous ORNs using neither Goa/cAMP nor Goq11/PLC cascades. Finally, amino acid odorants activate both ciliated and microvillous ORNs but via different transduction pathways in the two types of cells.

Key words: olfactory; receptor; G-protein; amino acid; bile salt; transduction

Introduction

The peripheral olfactory system of vertebrates consists of bipolar olfactory receptor neurons (ORNs) that convey information via the first cranial nerve to the olfactory bulb (OB). ORNs may be either ciliated or microvillous (Le Gros Clark, 1956; for review, see Farbman, 2000). In many terrestrial vertebrates, including snakes and rodents, ORNs are segregated into two separate compartments: microvillous ORNs lie in the vomeronasal organ (VNO), whereas ciliated ORNs populate the main olfactory epithelium (OE). Anatomical (Skeen, 1977; Stewart et al., 1979; Strotmann et al., 2000), molecular (Ressler et al., 1994; Vassar et al., 1994; Yoshihara et al., 1997; Mori et al., 1999), and functional studies (Kauer, 1991; Uchida et al., 2000) indicate an odotopic mapping of ORNs onto the OB and the accessory OB, i.e., distinct classes of ORNs respond best to particular molecular features of potential odorants and project to a limited number of glomeruli. Furthermore, in rodents, the type of receptor cell, ciliated or microvillous, correlates with the particular family of receptor molecules and G-proteins used in sensory transduction. The ciliated ORNs of the main OE use OR-type receptors (Buck and Axel, 1991) coupled to the G-protein Goq11 (Jones and Reed, 1989; Shinohara et al., 1992) and project to the main OB. The microvillus receptor cells of the VNO fall into two classes according to the G-protein used (Goa versus Goq) and project to different sites in the accessory OB (Jia and Halpern, 1996; for review, see Mori et al., 2000). A recent study reported that Goq also is expressed in the VNO of rodents (Wekesa et al., 2003).

In vertebrates lacking a VNO, e.g., fish, birds, Old World monkeys, hominoid apes, and humans, microvillous and ciliated ORNs both occupy the same OE [fish (Zeiske et al., 1992); bird, Old World monkeys, apes (Meisami and Bhatnagar, 1998); human (Moran et al., 1982)]. Because these species also lack an accessory OB, both receptor cell types project to the main OB. Additionally, the OE of fish contains, less abundantly, a third type of ORN: the crypt receptor cell (Morita et al., 1996; Hansen and Zeiske, 1998; Hansen and Finger, 2000). Although these three types of ORN are intermingled in one OE, the concept of odotopic representation in the OB may also obtain for fish (Thommesen, 1978; Doving and Selset, 1980; Friedrich and Korsching, 1997; Nikonov and Caprio, 2001). To test this hypothesis in catfish, we used three complementary approaches: (1) retrograde tracing from small, identifiable areas of the OB to label small populations of ORN; (2) in situ hybridization for particular odorant receptors and immunocytochemistry for various G-proteins to test whether the three types of ORN in the OE use

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Asterisks depict the correct band. Other reactive products. Preabsorption with the appropriate peptide resulted in no bands in any lane.

Different transduction cascades and, if so, whether specific glo-meruli of the OB exhibit complimentary immunoreactivity; and (3) electrophysiology of the olfactory epithelium to test whether particular transduction cascades are associated with detection of different classes of odorants.

Preliminary results have been published previously in abstract form (Anderson and Finger, 2000; Anderson et al., 2001; Hansen et al., 2001; Rolen et al., 2003b).

Materials and Methods

Juvenile catfish, Ictalurus punctatus, of both sexes ranging from 11 to 24.5 cm in standard length, were obtained commercially from the Hopper-Stephens Hatcheries (Lenoke, AR) and from the Louisiana State University aquaculture facility. The fish to be processed for histology were kept for at least 1 week in aquaria of the Animal Resources Center, University of Colorado Health Sciences Center. Fish for electrophysiological testing were held in the Louisiana State University Animal Care Facility in a 300 l aquarium filled with charcoal-filtered tap water (CFTW) and maintained on a 12 hr light/dark cycle. All procedures were performed with the approval of the institutional animal care and use committees of the respective institutions.

Immunocytochemistry for G-protein α-subunits

Different types of ORNs in mammals use different second messenger systems, i.e., the microvillous receptor neurons of the VNO express either Go or Gs, whereas the ciliated ORNs of the main olfactory epithelium predominantly express Goα11. We used antisera directed against mammalian forms of G-proteins [Goα OFF, Goα ON, Goα11 (Santa Cruz Biotechnology, Santa Cruz, CA; Calbiochem, La Jolla, CA; Chemicon, Temecula, CA)] (Table 1) to assess the tissue distribution of these forms in catfish.

Western blot analysis

To test whether the antibodies against mammalian G-proteins recognized similar molecular species in catfish tissue, we performed Western blots in tissue from catfish OE, OB, and brain, and as a positive control, on tissue from mouse or rat brain. As a negative control the antibodies were preabsorbed with the blocking peptides supplied by Santa Cruz Biotechnology. Briefly, the fish were anesthetized, and dissected olfactory bulbs, olfactory organs, and brains were immersed in protein homogenate buffer (10% SDS, 10 mM EDTA, 100 mM Tris, pH 8.0). The samples were either sonicated or homogenized (Tissue Tearor, Biospec Products). Protein concentrations were determined for each sample in comparison with bovine serum albumin (BSA) standards using the BCA assay (Bio-Rad, Hercules, CA). Western blotting samples were prepared by normalizing the concentration to 2 mg/ml by suspension in 4× sample buffer and dH2O. Samples of mouse and rat brains were prepared in an identical manner for positive controls. Lanes containing 20–40 μg of protein for each sample were separated by SDS-PAGE and transferred to polyvinylidene difluoride membranes (NEN, Boston, MA) and then preincubated in PBS containing 1% goat serum, 3% BSA, 0.1% Tween 20, pH 7.2, for 1 hr. The membranes were probed using antibodies against G-protein subunits overnight at 4°C. After washing, the protein bands were visualized with a peroxidase secondary antibody incubation followed by chemiluminescence using Super Signal West Dura Substrate. Data were captured with a Chemilmager 4400 chemiluminescence detection unit (Alpha Innotech, San Leandro, CA). Alternatively, the bands were visualized with a biotin-labeled secondary antibody followed by ABC (Vector Laboratories, Burlingame, CA) and diaminobenzidine (DAB) (Sigma, St. Louis, MO) reaction in the presence of hydrogen peroxide. The size standards used were biotin-labeled protein standards (Bio-Rad).

The Western blots showed strong double bands in the range of 42–44 kDa for the Goα OFF antibodies (Santa Cruz Biotechnology) in all tissues tested. Western blots performed with the Goα antibodies (Santa Cruz Biotechnology) resulted in a band of 140 μg to 44 kDa in tissue from catfish bulb and brain and a faint band at the same level for OE of catfish. Goα11 antibodies (Santa Cruz Biotechnology, Chemicon, Calbiochem) revealed a band of 41 kDa. After preabsorption of the antiserum supplied by Santa Cruz Biotechnology with the corresponding peptides, no bands appeared for the three antibodies (Fig. 1A–C).

**Light microscopic immunocytochemistry**

Fish were anesthetized, perfused, and fixed as described above. After fixation, the olfactory rosettes and OBs were removed from the tissue block and either processed as whole-mount preparations or cryoprotected with 20% sucrose overnight. Cryostat sections of 10–20 μm were collected onto Superfrost Plus slides (Fisher Scientific, Houston, TX). The tissue was washed three times for 10 min in 0.1 M phosphate buffer, pH 7.2, and, if necessary, incubated in 0.3% H2O2 for 10 min to reduce endogenous peroxidase. Preincubation in blocking solution (0.2% BSA, 1% normal serum, 0.3% Triton X-100 in 0.1 M PBS) for 1–2 hr was followed by incubation in polyclonal primary antibodies overnight for cryostat sections or for one to two nights for whole-mount preparations. Primary antibodies (Santa Cruz Biotechnology) were diluted in blocking solution: Goα, 1:250; Goα11, 1:1000; Gs, 1:500 (Santa Cruz Biotechnology). The antibodies against Goα11 (Calbiochem) were diluted 1:500. Experiments with antibodies directed against Goα did not result in

| Table 1. Antisera and peptides used for G-protein immunocytochemistry |
|----------------|-----------|---------------------------------|
| Supplier       | Article number | Lot number | Sequence          |
| Goα OFF        | Santa Cruz Biotechnology | SC-383 | L-0602, K-189 | Rat 377–394 |
| Goα            | Santa Cruz Biotechnology | SC-387 | K-291 | Rat 105–124 |
| Goα11          | Santa Cruz Biotechnology | SC-392 | H-071 | Mouse common domain 341–359 |
| Goα11          | Chemicon | AB1647 | 2100628 | Rat 283–300 |
| Peptide Goα OFF | Calbiochem | 371751 | B44877 | Rat 115–133 |
| Goα11          | Santa Cruz Biotechnology | SC-383P | B138 | Rat 377–394 |
| Goα11          | Santa Cruz Biotechnology | SC-387P | D171 | Rat 105–124 |
| Peptide Goα11  | Santa Cruz Biotechnology | SC-392P | I041 | Mouse 341–359 |

Figure 1. Western blots of G-protein preparations using tissue of catfish OE, OB, and brain and mouse and rat brain. Bands in A and B were visualized by ABC/DA reaction; C was visualized with chemiluminescence using Super Signal West Dura Substrate. Asterisks depict the correct band. A, Goα OFF, antiseraum reacts with a major band of 42–44 kDa in all catfish tissues as well as in mouse brain. B, Goα11 reveals a consistent band at 40–42 kDa. A smaller band is also evident in some lanes. C, Goα reacts with a band in all lanes at 40–42 kDa. The preparation from catfish OE shows a predominant band at this molecular weight, with few other reactive products. Preabsorption with the appropriate peptide resulted in no bands in any lane.
labeled tissue in catfish. Secondary antibodies were either fluorescence-coupled (dilution 1:400) (Alexa, Molecular Probes, Eugene, OR) or biotinylated (Jackson ImmunoResearch, West Grove, PA) (dilution 1:1000) that were processed with ABC (Vector) and DAB following standard methods. Controls included the omission of primary antibodies, omission of secondary antibodies, and preabsorption for 2 hr at room temperature with the blocking peptide (concentration, 4 μg/ml) supplied by Santa Cruz Biotechnology or Calbiochem. No labeled ORNs were observed under control conditions. Sections and whole-mount preparations were examined in a standard epifluorescence microscope (Zeiss) or a confocal laser scanning microscope (Olympus).

Transmission electron microscopic immunocytochemistry
Olfactory rosettes and bulbs of fixed fish were dissected and embedded in egg yolk. After fixation in 4% paraformaldehyde overnight, vibratome sections of 50 μm were collected in phosphate buffer and processed as floating sections as described for light microscopy, but omitting Triton X-100. After immunocytochemistry, selected sections and whole-mount preparations were postfixed in 2–4% buffered glutaraldehyde followed by 1% osmium tetroxide and embedded in Epon–Araldite following standard protocols as described previously (Hansen and Finger, 2000). Ultrathin sections (silver to gold) were examined in a Philips CM10 electron microscope.

DiI injections
Male and female fishes (n = 44) of the sizes mentioned above were used for 1,1′-dioctadecyl-3,3,3′,3′-tetramethylindocarbocyanine perchlorate (DiI) injections. To test whether specific types of ORN project to discrete regions of the OB, we used DiI as a retrograde postmortem tracer. Animals were anesthetized with MS222 (diluted 1:5000; Sigma) and perfused transcardially with 4% paraformaldehyde in 0.1 M phosphate buffer, pH 7.2. In some cases, the vaginal cavity was gently flushed with a mixture of buffered 0.3% glutaraldehyde and 4% paraformaldehyde after anesthesia and during perfusion to stabilize the epithelial surface of the OE. The top portion of the head containing the olfactory rosettes and the OBs was dissected and left in the fixative for at least one additional night. After fixation, the bone and other covering tissues above the bulbs were removed. Small crystals of DiI were inserted under visual control into different areas of the bulbs by means of a sharpened insect needle or the end of a broken glass micropipette. To prevent the dye from inadvertent spread, the bulbs were covered with liquid agar. The tissue block then was placed into buffered 4% paraformaldehyde at room temperature for 14–48 h. After removal of the dye, the tissue block and the olfactory organs were dissected from the tissue block and embedded in either egg yolk or 15% gelatin (Sigma). The block was fixed in 4% paraformaldehyde overnight. The next day, 50 μm sections were cut on a vibratome. Sections were viewed with epifluorescence in a Zeiss microscope or in a confocal laser scanning microscope (Olympus).

Of 87 injected bulbs, 15 bulbs were rejected because the dye did not travel well or the epithelium was damaged during dissection. The remaining 72 cases showed retrogradely labeled ORNs throughout the olfactory organ, which is consistent with the findings of Morita and Finger (1998). No differences were seen between male and female animals. To test whether specific types of ORN project to discrete regions of the OB, we used DiI as a retrograde postmortem tracer. Animals were anesthetized with MS222 (diluted 1:5000; Sigma) and perfused transcardially with 4% paraformaldehyde in 0.1 M phosphate buffer, pH 7.2. In some cases, the vaginal cavity was gently flushed with a mixture of buffered 0.3% glutaraldehyde and 4% paraformaldehyde after anesthesia and during perfusion to stabilize the epithelial surface of the OE. The top portion of the head containing the olfactory rosettes and the OBs was dissected and left in the fixative for at least one additional night. After fixation, the bone and other covering tissues above the bulbs were removed. Small crystals of DiI were inserted under visual control into different areas of the bulbs by means of a sharpened insect needle or the end of a broken glass micropipette. To prevent the dye from inadvertent spread, the bulbs were covered with liquid agar. The tissue block then was placed into buffered 4% paraformaldehyde at room temperature for 14–48 h. After removal of the dye, the tissue block and the olfactory organs were dissected from the tissue block and embedded in either egg yolk or 15% gelatin (Sigma). The block was fixed in 4% paraformaldehyde overnight. The next day, 50 μm sections were cut on a vibratome. Sections were viewed with epifluorescence in a Zeiss microscope or in a confocal laser scanning microscope (Olympus).

Transmission electron microscopy
To examine the ultrastructure of the retrogradely labeled olfactory receptor cells, selected sections were photoconverted with DAB. Photoconversion was performed according to the protocol of Sandell and Masland (1988). The sections were placed in DAB (1.25 mg/ml) dissolved in Tris buffer, pH 8.5, for 10 min and then, with a fresh supply of DAB solution, exposed to epifluorescence illumination for up to 25 min until a brown precipitate appeared. After photoconversion, the sections were postfixed in buffered 2% glutaraldehyde and 1% osmium tetroxide and embedded in Epon–Araldite for electron microscopy as described by Moran et al. (1992). Ultrathin sections (silver to gold) were examined in a Philips CM10 electron microscope.

In situ hybridization
To describe the expression pattern of receptor molecules in catfish ORNs, we tested four different catfish receptor probes: three different OR probes: CF1, GenBank accession number L09221; CF47, accession number L09221; OR202, accession number L09220 (courtesy of John Ngi, University of California Berkeley); and one V2R-type probe: CF64, accession number AY365191 (courtesy of Michele Rankin and Richard Bruch, Louisiana State University). RNA in situ hybridizations were performed on 12–15 μm cryostat sections of dissected catfish olfactory organs collected onto SuperFrost Plus slides (Fisher Scientific). The sections were pretreated with 3% hydrogen peroxide to quench endogenous peroxidase and digested with Proteinase K (10–20 μg/ml) for 10–20 min to enhance probe penetration. Digoxigenin-labeled cRNA probes were hybridized at 58–60°C overnight followed by high stringency washings in 2× SSC (0.3 M NaCl, 0.03 M Na3HPO4) and 0.2× SSC (0.03 M NaCl, 0.003 M Na2HPO4) at 60°C. To visualize the hybridized probes we used either alkaline phosphatase-labeled sheep anti-digoxigenin antibodies (Roche Biochemicals, Indianapolis, IN) or peroxidase-labeled sheep anti-digoxigenin antibodies (Roche Biochemicals). For alkaline phosphatase-based detection, nitroblue tetrazolium chloride and X-phosphate-5-bromo-4-chloro-3-indolyl-phosphate rendered a light purple chromogenic signal. Alternatively, an enhanced fluorescent signal was produced by the fluorophores tyramide cyanine 3 or tyramide fluorescein. For peroxidase-labeled antibodies the tyramide signal amplification technique (Molecular Probes) was used that enhances the signal of low copy numbers of mRNA up to 1000-fold.

Electrophysiological experiments
Animal preparation. Each catfish was immobilized with an initial intra-muscular injection of Flaxedil (gallamine triethiodide, 0.03 mg/100 gm body weight). Subsequent injections of Flaxedil were provided as needed during experimentation via a hypodermic needle embedded in the flank musculature. The catfish was wrapped in wet tissue paper and secured with orbital ridge clamps in a custom-made Plexiglas container. The gills were irrigated via a constant flow of CFTW containing the general anesthetic MS-222 (ethyl-m-aminobenzoate methane sulfonic acid; initial concentration, 50 mg/l; Sigma). Minor surgery was performed to remove the skin and connective tissue superficial to the olfactory epithelium to facilitate electrode placement.

Stimulus solutions and delivery. The odorants included t-amino acids (alanine, arginine, glutamate, and methionine), bile salts [sodium salts of taurocholic acid (TCA) and tauroliothiocholic acid (TLCA)], and ATP (Sigma). Test solutions of amino acids, bile salts, and ATP were diluted daily with CFTW to experimental concentrations from stock solutions and pH adjusted to match control CFTW, pH 8.7, bathing the olfactory epithelium. Analysis of the CFTW by the Dionex AAA-Direct Amino Acid Analysis System (sensitivity in the mid femtomole to low picomole range) indicated that free amino acids were not present. Stimuli were delivered via a “gravity-feed” system described previously (Sveinson and Hara, 2000). Briefly, stimulus solutions and the CFTW used to bathe the olfactory epithelium were delivered through separate Teflon tubes (diameter 0.8 mm) at a flow rate of 5–7 ml/min. A foot switch connected to an electronic timer (Model 645, GraLab Instruments Division, Dimco-Gray Corporation, Centerville, OH) triggered a pneumatic actuator valve to introduce the stimulus for 3 sec applications. CFTW continuously irrigated the olfactory epithelium to (1) prevent the epithelium from desiccating, (2) facilitate stimulus delivery, (3) avoid introduction of mechanical artifacts, and (4) rinse the olfactory organ clear of any residual stimuli for a minimum of 2 min between stimulus applications.

Pharmacological agents
Forskolin (an adenylate cyclase activator; Sigma) and 1,9-dideoxyforskolin (inactive analog of forskolin; Calbiochem) were dissolved in dimethyl sulfoxide (DMSO) and added to CFTW. U-73122, a potent inhibitor of agonist-induced phospholipase C (PLC) activation (Yule and Williams, 1992), and U-73343 (Biomol Research Laboratories,
Electrophysiological recording techniques

We recorded underwater electro-olfactograms (EOGs) in vivo by means of calomel electrodes via Ringer’s agar-filled capillary pipettes. The slow DC potential changes in the water above the olfactory epithelium represent the summed generator potentials of the ORNs in response to odorants (Ottoson, 1971). The pipette of the active electrode was positioned near the midline raphe of the olfactory organ; the pipette of the reference electrode was placed against the skin adjacent to the olfactory cavity. The fish was grounded via a hypodermic needle inserted into the flank musculature. The EOG was amplified (Grass P-18; Astro-Med, West Warwick, RI), displayed on an oscilloscope, integrated (0.5 sec) and displayed on a pen recorder, digitized, and stored on a video channel of a high-fidelity VCR.

Pharmacological treatments

Electrophysiological experiments were used to investigate the signaling cascade used by ORNs to transduce odorant information. Each experiment consisted of three stages: odorant responses in the absence of the pharmacological agent (stage 1), odorant responses during pharmacological treatment (stage 2), and responses to the odorants subsequent to removal of the pharmacological agent (stage 3).

During stage 1, CFTW continuously bathed the olfactory epithelium for a minimum of 10 min before stimulus application. For the forskolin experiments, the odorant stimuli were adjusted in concentration to result in an EOG response with approximately the same relative magnitude as the EOG response to forskolin. Some stimulus solutions involved mixtures of odorants, in which case each component of a stimulus mixture was also adjusted in concentration to result in an approximately equal EOG response magnitude when tested individually. The adjusted concentrations ensured that the odorant stimuli and forskolin were equipotent to the CAMP signaling pathway. Neither 1 μM U-73122 nor 1 μM U-73343 elicited appreciable EOG responses; therefore, the concentrations for the odorant stimuli were chosen to reflect those used in the forskolin experiments. CFTW and DMSO served as controls.

During stage 2, the pharmacological agent at the previously adjusted concentration continuously bathed the olfactory epithelium for a minimum of 10 min before stimulus application. All odorants tested during stage 2 were dissolved in the pharmacological agent. Controls were aliquots of the adapting solution, DMSO and CFTW, respectively.

During stage 3, CFTW continuously bathed the olfactory epithelium for 10 min before stimulus application. Stimuli and controls were identical to those described during stage 1.

Electronic figure composition

Light micrographs were either scanned from film images or digitally imaged directly on a SPOT RT (Diagnostic Instruments, Sterling Heights, MI) camera attached to the Olympus or Zeiss microscope. Electron micrographs were scanned either from prints or film negatives on a flatbed scanner (UMAX Powerlook III, Dallas, TX). Images were then processed in Photoshop (Adobe Systems, Mountain View, CA). Adjustments to contrast and brightness were made electronically. Occasional dirt specks were removed electronically from areas of the figures that did not contain images of tissue.

Results

Even at the light microscopic level, we could distinguish three types of ORN retrogradely labeled by DiI injections into the OB (Fig. 2C–F). Ciliated ORNs possess long, undulating apical processes usually radiating from an olfactory knob at the distal ends
of tall ORNs (Fig. 2C,D). Crypt receptor cells were obvious because of their typical ovoid shape and location in the top half of the olfactory epithelium (Fig. 2C,E). Most of the short (but not ovoid) and intermediate height ORNs bore small apical endings on olfactory knobs that were less pronounced than in ciliated ORNs. Ultrastructural analysis demonstrated that these small apical endings belonged to microvillous ORNs (Figs. 2C,F, 3). Despite this general morphological pattern, some cells of short and intermediate height bore cilia, whereas a few tall cells had small microvillous-like apical endings. To investigate the ultrastructure of the labeled ORNs, selected vibratome sections were photoconverted after injections into dorsal OB, midventral OB, and the two specific ventral sites for crypt ORNs. The photoconverted ORNs examined at the electron microscopic level matched our presumed morphological categorizations on the basis of the light microscopic findings. Transmission electron microscopy sections proved that ciliated ORNs were retrogradely labeled by injections into the ventral areas (Fig. 3A), and microvillous ORNs were labeled by injections into the dorsal areas (Fig. 3C). Crypt ORNs (Fig. 3B) occurred only when Dil crystals were placed into the two specific ventral sites.

To determine the pattern of projections of different ORN types into the OB, we divided the OB into different injection regions. These regions were defined not only by topographical location, e.g., anterior, posterior, and lateral, but also by morphological features such as obvious fiber bundles or the irregular shape of the bulb. This was important because in some animals the bulb was slightly rotated either to the lateral or medial side. Dil injections into different sites of the OB resulted in a distinct pattern of cell projections. We divided the types of ORN into three categories on the basis of their relative location in the height of the olfactory epithelium and, in the case of the crypt ORNs, on their unmistakable shape. Tall cells correspond to different sizes of ciliated ORNs with cell bodies in the bottom one-third of the epithelium; medium and short cells are considered microvillous ORNs in the top two-thirds of the epithelium; crypt ORNs occupy the top half of the epithelium.

The main difference in projections of ciliated and microvillous ORNs related to the dorsal versus the ventral regions of the bulb (Fig. 4). Tall ciliated ORNs were labeled by injections into the areas on the ventral side of the OB. In contrast, injections into the dorsal-most areas labeled medium and short presumed microvillous ORNs in the epithelium. Injections into the medial or lateral areas of the OB showed a less segregated pattern of label. Only injections into the medial middle area of the bulb resulted in predominantly tall ciliated ORNs in the olfactory epithelium, as summarized in Figure 4. The areas anteromedial and posteromedial and the three lateral injection sites anterolateral, middle lateral, and posterolateral revealed a mixed population of short, intermediate, and tall ORNs in the olfactory epithelium, bearing either cilia or microvilli.

Labeled crypt ORNs occurred in the olfactory epithelium only after Dil injection into two discrete areas in the ventral bulb, one more anteriorly and one more posteriorly. Injections into the medial and lateral areas that led to mixed populations of labeled cells did not label crypt ORNs, nor were crypt ORNs detected in any injection into dorsal sites.

Counting types of ORN retrogradely labeled from particular dorsal or ventral sites in the OB proved that significantly different populations of ORN were labeled depending on the location of the Dil placement into the OB (Table 2). Injection sites on the ventral and medial OB resulted mostly in labeled ciliated ORNs, whereas injections in the dorsal anterior and posterior OB labeled mostly microvillous ORNs. The total number of crypt cells labeled from the two specific ventral sites of the OB was not as high as the number of microvillous and ciliated ORNs labeled by injections in dorsal and ventral OB (Table 2), indicating that the crypt ORNs are a relatively rare cell type projecting only to small regions in the OB. Consequently, ciliated ORNs projecting to neighboring glomeruli were also labeled by injections targeted to the crypt cell recipient zones.

In summary, microvillous ORNs project mostly to the dorsal and lateral portions of the olfactory bulb, whereas ciliated ORN projections predominate ventrally and along the medial side of the OB. Crypt cell ORNs project to two small areas of the ventral OB (Fig. 4).

**In situ hybridization with receptor probes**

The procedure of in situ hybridization has a strong impact on tissue preservation. Fragile structures such as membranes and especially microvilli and cilia are easily and often damaged, e.g., by the incubation in proteinase K necessary to permeabilize the tissue. On the basis of light and electron microscopic experience (see above), the tall ORNs are ciliated ORNs, whereas ORNs with cell bodies higher in the epithelium tend to be microvillous ORNs. Most of the intermediate ORNs bear microvilli, although some intermediate ORNs may bear cilia instead. Crypt ORNs that would be easily distinguished even in in situ preparations because of their ovoid cell body and their location in the uppermost portion of the OE were not reactive with the in situ probes used in this study. As also seen in G-protein experiments (see below), some cells located near the base of the epithelium reacted with the in situ probes. These cells did not reach the top portion of the epithelium and were considered to be basal cells or ORNs in the process of differentiation.
The absolute number of labeled ORNs varied slightly between only a small fraction of the numerous intermediate and tall cells. Therefore, are interpreted to be ciliated ORNs. Each probe labeled bodies located predominantly in the bottom half of the OE, and intermediate and tall ORNs. These cells were slender, with cell bodies located predominantly in the bottom half of the OE, and therefore interpreted to be ciliated ORNs. Each probe labeled only a small fraction of the numerous intermediate and tall cells. The absolute number of labeled ORNs varied slightly between individuals.

OR-labeled cells occurred in all areas of the lamellas from the midline raphe to the peripheral and basal areas close to the nonsensory epithelium (Fig. 5A, C). Within the lamellas, the OR-positive ORNs were spaced more or less evenly.

V2R-type receptor probe

The only V2R-like probe (CF64) available labeled short or intermediate ORNs. Their nuclei were predominantly located in the top half of the OE. Because of their position and morphology, these cells were considered microvillous ORNs (Fig. 5B, D). Within the lamellas, the pattern of distribution of V2R-positive ORNs varied between individuals.

Table 2. Distribution of labeled ORN cell types according to site of injection in OB

<table>
<thead>
<tr>
<th>Injection site</th>
<th>Ciliated ORNs</th>
<th>Microvillous ORNs</th>
<th>Crypt cell ORNs</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dorsal OB</td>
<td>3</td>
<td>13</td>
<td>0</td>
<td>16</td>
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<tr>
<td></td>
<td>4</td>
<td>17</td>
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<td>21</td>
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<td>13</td>
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<tr>
<td></td>
<td>5</td>
<td>13</td>
<td>0</td>
<td>18</td>
</tr>
<tr>
<td>Total</td>
<td>16 (20%)</td>
<td>64 (80%)</td>
<td>0 (0%)</td>
<td>80</td>
</tr>
<tr>
<td>Ventral/medial</td>
<td>24</td>
<td>2</td>
<td>0</td>
<td>26</td>
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<td></td>
<td>21</td>
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<td>12</td>
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<tr>
<td>Total</td>
<td>80 (86%)</td>
<td>13 (14%)</td>
<td>0 (0%)</td>
<td>93</td>
</tr>
<tr>
<td>Ventral/crypt</td>
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<td>2</td>
<td>8</td>
<td>11</td>
</tr>
<tr>
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<td>7</td>
</tr>
<tr>
<td></td>
<td>8</td>
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<td>5</td>
<td>18</td>
</tr>
<tr>
<td></td>
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<td>4</td>
<td>8</td>
<td>21</td>
</tr>
<tr>
<td></td>
<td>14</td>
<td>5</td>
<td>4</td>
<td>23</td>
</tr>
<tr>
<td>Total</td>
<td>32 (40%)</td>
<td>17 (21%)</td>
<td>31 (39%)</td>
<td>80</td>
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</table>

Counts of the type of ORN retrogradely labeled by different injections of Dil into the olfactory bulb are shown. Each row is a different case. The total number of labeled cells is given for a representative sample of 300 μm of epithelium from each selected section. This distribution is significantly different from chance (χ² test; p < 0.001). Because the crypt cell projecting areas are small (on the basis of the size of Goα,s-immunoreactive glomeruli) relative to the size of the Dil injections, other cell types are frequently labeled by injections into the crypt cell target zone of the ventral OB, but crypt cells are never labeled by injections remote from their specific small targets in the ventral OB.

GOα,s immunoreactivity

In the sensory areas of the olfactory epithelium, the GOα,s-immunoreactive fibers were present in the olfactory nerve layer in ventral and medial areas of the OB as well as whole-mount preparations of single olfactory lamellas and OBs revealed distinct labeling for Goα,s, Goq, and Goαq (Fig. 6).

GOα,s-immunoreactive fibers were present in the olfactory nerve layer in ventral and medial areas of the OB (Fig. 6G). In addition, smaller numbers of GOq-immunoreactive fibers were present in the olfactory nerve layer of the rostral lateral portion of the OB. Some small bundles of labeled fibers coursed dorsoventrally in the caudal portion of the bulb.

Gq, immunoreactivity

Antisera directed against different G-protein α-subunits were used to examine whether the expression of these transduction-related proteins correlated with the morphological type of ORN and, if so, whether the distribution of these G-proteins in the glomeruli of the olfactory bulb corresponded to our findings on the basis of Dil injections. Of the four antisera used, only three (Goα,s, Goq, and Goαq) resulted in unambiguous label in catfish tissue. Tissue sections of the olfactory epithelium and the OB as well as whole-mount preparations of single olfactory lamellas and OBs revealed distinct labeling for Goα,s, Goq, and Goαq (Fig. 6).
against GORNs. Many, but by no means all, microvillous dorsal aspects of the bulb (Fig. 6). Examination of ultrathin sections showed Gq/11-labeled microvilli in a subset of microvillous ORNs (Fig. 7C), presumably corresponding to the punctate apical label seen at light microscopic levels. Somatic membranes of these cells were seldom labeled and then only faintly. The height of these cells within the epithelium was intermediate, consonant with these cells being microvillous ORNs. Gq/11 antisera labeled many, but by no means all, microvillous ORNs.

In the olfactory bulb, the antibodies against Gq/11 mostly labeled glomeruli in dorsal aspects of the bulb (Fig. 6H). Large glomeruli located anteriorly and posteriorly were Gq/11 positive. Some smaller glomeruli appeared in lateral and medial locations.

In summary, ciliated ORNs express Buck and Axel (1991) OR-type receptors with Gq/11, and project mostly to ventrally situated glomeruli. The V2R-type receptor probe hybridizes to intermediate height receptor cells, which are apparently microvillous ORNs. Most of the microvillous ORNs express Gq/11 and project to dorsal OB. Crypt ORNs express Gq, and project to two unique glomerular territories along the ventral midline of the OB; the receptor molecules used by this cell type are unknown.

Electrophysiology and pharmacological agents

Effects of forskolin on odor-evoked responses
We used the EOG to assess ORN population responses to various odorants and drugs. Simultaneous recordings of the EOG and multunit responses in the epithelium show that the magnitude of the negative-going component of the EOG correlates well with the level of multiunit activity in the OE (Fig. 8A). Accordingly, we used the magnitude of the negative-going component of the EOG as a basis for comparison across treatments.

To determine whether transduction of amino acid, bile salt, or nucleotide odorant information in catfish involves the Gi/o/cAMP second messenger pathway, forskolin (an adenylyl cyclase activator) was applied continuously to the olfactory epithelium on the assumption that the treatment would either decrease the number of adenylyl cyclase molecules available for G-protein-coupled receptor activation or desensitize certain components, resulting in an attenuation of responses to odorants that activate this pathway. Forskolin, when applied to the olfactory epithelium, elicited a robust EOG response (Fig. 8B). By contrast, 5 μM 1,9-dideoxyforskolin (an inactive analog of forskolin), which is equivalent to the highest concentration of forskolin tested, did not evoke an appreciable EOG response above that of control (n = 3 fish) (Fig. 8D). Therefore, forskolin was adjusted in concentration to result in an EOG response approximately equivalent to the EOG response to the odorant stimuli (Fig. 8B1). This process equalizes the potency of the test stimuli to the cAMP signaling pathway. During forskolin (3–5 μM) treatment, the magnitude of the EOG responses to a mixture of L-amino acids (alanine, arginine, glutamate, and methionine) and bile salts (TCA and TLCA) was reduced by 70 and 74%, respectively, of the responses obtained in the absence of forskolin, whereas the magnitude of the EOG responses to ATP remained relatively unaffected (Fig. 8B, C). Importantly, adaptation to the mixture of bile salts did not attenuate the response to forskolin (n = 2; data not shown), which indicates that forskolin, the structure of which grossly resembles that of bile salts, did not compete for the bile salt receptors.

Effects of U-73122 and U-73343 on odor-evoked responses
To determine whether transduction of amino acid, bile salt, and nucleotide odorant information in catfish involves the IP3 second messenger pathway, U-73122 (a membrane-permeant potent inhibitor of agonist-induced PLC activation) was applied continuously to the olfactory epithelium. The assumption was that U-73122 treatment would inhibit Gi, coupling with PLC resulting in an attenuation of the response to odorants using this path-
the EOG response to amino acids than U-73122, whereas the magnitude of the EOG responses to bile salts and ATP remained unaffected (Fig. 9B, D).

Taken together, these physiological studies are consistent with the interpretation that ORN responses to bile salts are mediated predominantly by ORNs using the Go\textsubscript{olf/s} transduction pathway. In contrast, responses to nucleotides are unaffected by blockade of this pathway. Because blockers of both Go\textsubscript{olf/s} and PLC pathways diminish responses to amino acids, both transduction pathways are implicated in the detection of amino acids by different populations of ORNs.

In summary, antisera to the different G-protein α subunits labeled different types of ORNs and different glomerular territories in the OB. Taken together with the findings of Nikonov and Caprio (2001), the immunocytochemical and retrograde Dil tracings indicate that Go\textsubscript{olf/s}-reactive, ciliated ORNs project to glomeruli in the medial and ventral portions of the OB responding to either amino acids or bile salts. Go\textsubscript{q} antisera strongly label crypt cell ORNs and two glomerular regions on the ventral surface of the OB; however, nothing is known about the function of these ORNs. In addition, some microvillous ORNs exhibit fainter reactivity for Go\textsubscript{q}, Finally, Go\textsubscript{q11} immunoreactivity is present in a subpopulation of short microvillous ORNs and in the glomerular layer of the anterior dorsal and dorsolateral areas of the OB, areas that are responsive to amino acids (Fig. 10). Responses to nucleotides are mediated by microvillous ORNs but via a mechanism that does not involve PLC activation.

Discussion

The present study demonstrates that in catfish, as in rodents, the ORNs of the extended olfactory system are divisible into classes defined both by morphology and by transduction cascade. Furthermore, each of these classes of receptor cells projects to a restricted set of glomeruli within the OB. We identify three classes of ORNs in catfish: (1) tall ciliated cells using Go\textsubscript{olf/s} and projecting predominantly to medially and ventrally situated glomeruli; (2) microvillous ORNs projecting to dorsal glomeruli, some of which use Go\textsubscript{q11}; and (3) crypt-cell ORNs using Go\textsubscript{q} and projecting to two restricted glomerular territories at the anterior and posterior extremes of the ventralmost OB (Fig. 10).

Odorant specificity and ORN type

Fish respond to various classes of olfactory stimuli: amino acids, nucleotides, bile salts, and pheromones (Thommesen, 1983b; Carr, 1988; Hara and Zhang, 1996; Hara and Zhang, 1998; Sorensen and Caprio, 1998). The former two are generally feeding
stimulants, whereas the latter two are implicated in social interactions (Sorensen and Caprio, 1998). Whether a particular type of ORN is invariably activated by a particular class of stimulus may vary according to the different species of fish. Comparison of the results of our DiI and immunocytochemistry experiments with a recent electrophysiological study in catfish (Nikonov and Caprio, 2001) leads to three conclusions regarding the correlation between receptor cell type and function. First, a subset of microvillous ORNs projecting to the dorsal OB mediates the responses to nucleotides. Second, ciliated ORNs projecting to the medial face of the OB mediate the response to bile acids. Finally, other ciliated ORNs mediate much, but not all, of the responsiveness to amino acids. Some responses to amino acids appear to rely on a PLC-based transduction pathway in microvillous ORNs projecting to the anterior part of the dorsolateral segment of the OB.

**Nucleotides**

Our studies suggest that transduction of nucleotide odorants by microvillous ORNs does not rely on a Go<sub>q/11</sub>/PLC transduction cascade. The physiological studies of Nikonov and Caprio (2001) demonstrate that response to nucleotides occurs primarily within the dorsal and caudolateral OB. Placement of DiI into this same area of the OB predominantly labels microvillous ORNs; however, only a few Go<sub>q/11</sub>-positive glomeruli are present in this region, suggesting that nucleotide-sensitive microvillous ORNs may not use Go<sub>q/11</sub>. Similarly, treatment with U-73122 and U-73343 (inhibitors of PLC activation) does not attenuate responses to nucleotides, indicating that nucleotide-sensitive ORNs use a transduction mechanism that does not act via a PLC mechanism. Polysine odorants also appear to be detected via a transduction cascade not involving either cAMP/Go<sub>olf</sub> or PLC/IP<sub>3</sub> systems (Rolen et al., 2002, 2003a). Additional studies are necessary to determine the transduction cascade underlying olfactory detection of nucleotides in catfish.

**Bile salts**

Responses to bile salts occur predominantly in the medial part of the olfactory bulb (Nikonov and Caprio, 2001), including dorsal and ventral aspects. Our anatomical results demonstrate that this medial face of the OB receives input predominantly from ciliated ORNs expressing Go<sub>olf</sub>. Accordingly, the ciliated Go<sub>olf</sub>-expressing ORNs projecting to the medial OB likely mediate detection of bile salts. In studies on salmonid, Thommesen (1983a) likewise attributed sensitivity to bile salts to ciliated ORNs. This conclusion also is supported by our pharmacological results: only forskolin, which disrupts the Go<sub>olf</sub>cAMP transduction system, attenuated the EOG response to bile salts, whereas inhibitors of the PLC pathway had no effect.

**Amino acids**

Responses to amino acids are prevalent in the rostral, ventral, and dorsolateral OB (Nikonov and Caprio, 2001), where ciliated cell projections predominate (present study). The conclusion that some ciliated ORNs mediate detection of amino acids is corroborated by studies in trout, goldfish, and zebrafish. Zielinski and Hara (1988) reported responses to amino acids in trout larvae at a time when only ciliated ORNs are present in the OE. Likewise, during regeneration after axotomy and bulbectomy in goldfish, responses to amino acids were correlated with the presence of ciliated ORNs (Zippel et al., 1996, 1997). That forskolin significantly reduces responses to amino acid odorants further supports this conclusion.

Other studies, however, attribute amino acid responses to microvillous ORNs. In salmonids, Thommesen (1983a) found responses to amino acids in the epithelium correlated with the presence of microvillous ORNs. Speca et al. (1999) postulated...
that microvillous ORNs mediate amino acid signals in goldfish. Hamandi et al. (2001a,b) report that in a carp, microvillous ORNs project to the posterolateral OB, which is crucial for feeding-related behaviors, thereby implicating microvillous ORNs in responses to food odors. In the present study, a subset of microvillous ORNs, immunoreactive for Gαq/11, project to the anterodorsal areas of the OB that are responsive to amino acids. These findings are consistent with several other studies that support the concept that subsets of both microvillous and ciliated ORNs respond to amino acids. In salmonids (Hara and Zhang, 1996, 1998), amino acids elicit responses in lateroposterior sites of the OB, sites in which ciliated ORNs project to the anterodorsal areas of the OB that are responsive to amino acids. In salmonids, biochemical studies (Lo et al., 1993) show that amino acids activate an IP3 pathway. Microvillous ORNs. In salmonids, biochemical studies (Lo et al., 1993; Lo, 1994) show that amino acids activate an IP3 pathway. Studies on other species of fish also support the hypothesis that amino acids stimulate some ciliated ORNs as well as some microvillous ORNs. In salmonids, biochemical studies (Lo et al., 1993; Lo, 1994) show that amino acids activate an IP3 pathway. The results of our study on goldfish are consistent with our electrophysiological results. Treatment with either forskolin (which disrupts the Gαq/cAMP pathway) or U-73122 and U-73343 (which disrupts the PLC transduction cascade) led to partial attenuation of EOG responses to amino acids. Nikonov and Caprio (2001) recorded amino acid responses from anterodorsal areas of the OB where we find projections of microvillous ORNs expressing Gαq/11 and from ventral areas of the OB where we find Gαo—a positive ciliated ORNs. Two different second messengers involved in the transduction of amino acids have also been reported for salamanders (Delay and Dionne, 2002) and Xenopus (Manzini et al., 2002).

Studies on other species of fish also support the hypothesis that amino acids stimulate some ciliated ORNs as well as some microvillous ORNs. In salmonids, biochemical studies (Lo et al., 1993; Lo, 1994) show that amino acids activate an IP3 pathway. The activation of the IP3 pathway by amino acids is likewise supported by a study on zebrafish reporting the elimination or attenuation of amino acid-evoked responses after application of an IP3-gated channel blocker or a PLC inhibitor (Ma and Michel, 1998). Finally, Sato and Suzuki (2001) used single-cell patch methods to determine that both ciliated and microvillous ORNs can detect amino acids. Our results suggest that a similar situation obtains in catfish.

G-proteins and cell type
The different types of ORNs in the VNO and main OE of rodents use different transduction mechanisms and attendant G-proteins (Schandar et al., 1998; Wekesa and Anholt, 1999). Reports on fish including catfish also indicate the expression of different G-protein subunits (Abogadie et al., 1995; DellaCorte et al., 1996) using different transduction mechanisms (Miyamoto et al., 1992; Restrepo et al., 1993). Likewise in catfish, we find that different types of ORNs use various G-proteins: Gαq/11 in ciliated ORNs, Gαq in crypt...
ORNs. A previous study of G-proteins in catfish (DellaCorte et al., 1996) suggests that ciliated ORNs use Goq. We cannot account for this report, which is at odds with our findings in the same species. Both our light and electron microscopic analyses show that Goq immunoreactivity occurs in cells situated in the top half of the epithelium, which possess microvilli. Perhaps the antisem was used by DellaCorte et al. (1996) cross-reacted with proteins other than Goq. Our attempts to replicate their findings using an antisem generated against the same peptide sequence yielded no specific staining in the OE. The ciliated ORNs in our preparations react only with the specific Goq antisem and not with those directed against either Goq or Goq.

In summary, the present study demonstrates that morphologically distinct classes of ORNs use different transduction cascades and maintain distinct patterns of projection to the OB. Ciliated ORNs using Gq and maintain distinct patterns of projection to the OB. Ciliated ORNs use different transduction cascades and not with those directed against either Gq or Goq. Perhaps the antisem used by DellaCorte et al. (1996) cross-reacted with proteins other than Goq. In our preparations react only with the specific Goq antisem and not with those directed against either Goq or Goq.

References
Manzini I, Røssler W, Schild D (2002) cAMP-independent responses of ol-
factory neurons in *Xenopus laevis* tadpoles and their projection onto olfactory bulb neurons. J Physiol (Lond) 545:475–484.


