Differential axonal projection of mitral and tufted cells in the mouse main olfactory system

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In the past decade, much has been elucidated regarding the functional organization of the axonal connection of olfactory sensory neurons to olfactory bulb (OB) glomeruli. However, the manner in which projection neurons of the OB process odorant input and send this information to higher brain centers remains unclear. Here, we report long-range, large-scale tracing of the axonal projection patterns of OB neurons using two-photon microscopy. Tracer injection into a single glomerulus demonstrated widely distributed mitral/tufted cell axonal projections on the lateroventral surface of the mouse brain, including the anterior/posterior piriform cortex (PC) and olfactory tubercle (OT). We noted two distinct groups of labeled axons: PC-orienting axons and OT-orienting axons. Each group occupied distinct parts of the lateral olfactory tract. PC-orienting axons projected axon collaterals to a wide area of the PC but only a few collaterals to the OT. OT-orienting axons densely projected axon collaterals primarily to the anterolateral OT (aOT). Different colored dye injections into the superficial and deep portions of the OB external plexiform layer revealed that the PC-orienting axon populations originated in presumed mitral cells and the OT-orienting axons in presumed tufted cells. These data suggest that although mitral and tufted cells receive similar odor signals from a shared glomerulus, they process the odor information in different ways and send their output to different higher brain centers via the PC and aOT.

Keywords: mitral cell, tufted cell, olfactory bulb, axon projection, piriform cortex, olfactory tubercle, parallel pathways, two-photon imaging

INTRODUCTION

Olfactory sensation begins with the interaction of odorant ligands and odorant receptor proteins on the cilia of olfactory receptor neurons (ORNs) in the nasal epithelium. In the mouse, one ORN expresses just one type of odorant receptor protein out of a repertoire of approximately 1,000 such receptors (Buck, 1996). ORNs usually send their axons to two glomeruli on the surface of the mouse olfactory bulb (OB), and individual glomeruli receive inputs from ORNs that express a common odorant receptor (Mombaerts, 1999). The stereotypical arrangement of these glomeruli is thought to form odorant receptor maps (Mori et al., 2006). However, it is unclear how this convergent and patterned odor information is represented in the next olfactory processing area through the axonal connection of olfactory sensory neurons to olfactory bulb (OB) glomeruli. However, the manner in which projection neurons of the OB process odorant input and send this information to higher brain centers remains unclear. Here, we report long-range, large-scale tracing of the axonal projection patterns of OB neurons using two-photon microscopy. Tracer injection into a single glomerulus demonstrated widely distributed mitral/tufted cell axonal projections on the lateroventral surface of the mouse brain, including the anterior/posterior piriform cortex (PC) and olfactory tubercle (OT). We noted two distinct groups of labeled axons: PC-orienting axons and OT-orienting axons. Each group occupied distinct parts of the lateral olfactory tract. PC-orienting axons projected axon collaterals to a wide area of the PC but only a few collaterals to the OT. OT-orienting axons densely projected axon collaterals primarily to the anterolateral OT (aOT). Different colored dye injections into the superficial and deep portions of the OB external plexiform layer revealed that the PC-orienting axon populations originated in presumed mitral cells and the OT-orienting axons in presumed tufted cells. These data suggest that although mitral and tufted cells receive similar odor signals from a shared glomerulus, they process the odor information in different ways and send their output to different higher brain centers via the PC and aOT.

In mammals, a few studies have examined the axon projection patterns of mitral and tufted cells, the OB projection neurons. Tracing of single mitral cell axon trajectories has revealed their multiple target projections in the anterior part of the olfactory cortex (OC; Ojima et al., 1984). Dense axonal projection by neighboring mitral cells to the same areas in the anterior OC has also been reported (Buonviso et al., 1991). Retrograde tracing experiments have suggested that mitral cells project to all regions of the OC, while tufted cells project to the anterior part of the OC (Haberly and Price, 1977; Sken and Hall, 1977; Scott et al., 1980). These results have also been supported by electrophysiological studies (Scott, 1981; Schneider and Scott, 1983). However, no report to date has directly compared the axonal projections of mitral and tufted cells.

Despite these efforts, it still remains unclear in mammals how the odorant information that converges in the glomerulus is represented in the next olfactory processing area through the axonal
projections of mitral and tufted cells. To address this question, we traced the trajectories of OB neurons associated with a single glomerulus on the ventrolateral surface of the whole brain using two-photon microscopy. Overall projection patterns were found to comprise an overlay of two distinct axon branching patterns, those predominantly to the piriform cortex (PC) and those to the olfactory tubercle (OT). Segregated dual-color labeling was then used to further demonstrate that these two axon groups originate from presumed mitral and tufted cell populations, respectively.

**MATERIALS AND METHODS**

All experimental protocols were approved by the University of Texas Medical School at Houston and Yale University Institutional Animal Care and Use Committee in accordance with the National Institutes of Health guidelines. A total of 19 mice (11 OMP-Synaptophysin knock-in mice, Jackson Lab, and 8 C57BL/6, Charles River) were anesthetized with nembutal (i.p. 50 mg/kg body weight). The animals were kept on a heating pad with circulating water set at 40°C. The bone covering the dorsal OB was carefully thinned, and a small opening was made (200–500 μm diameter) for glass pipette penetration. Dextran-conjugated Alexa 488 or Alexa 594 (Invitrogen) was diluted in Ca++-free Ringer solution (140 mM NaCl, 5 mM KCl, 3 mM MgCl, 0.01 mM EDTA, 10 mM HEPES, and 10 mM glucose, pH 7.5) and used as 10% dye solution. Glomeruli were visualized by two-photon fluorescence imaging of OMP-SpH knock-in mice (Bozza et al., 2004). A dye-filled glass pipette (tip I.D., 2.5 μm) was placed into the target glomerulus under visualization with a two-photon microscope, or into the superficial or deep external plexiform layer (EPL) according to micromanipulator penetration depth (100–150 or 200–250 μm depth, respectively). Small square-wave pulses were delivered via the pipette for dye electroporation (3–5 μA, 23-ms pulse width, 2 Hz, 1200–3600 pulses). The electroporation effective area using this technique has been previously reported to be about 30 μm in diameter (Nagayama et al., 2007), and was effective to label a single glomerulus or superficial/deep EPL in the OB.

Three to 5 days after the dye injection, animals were deeply anesthetized by nembutal injection (i.p. 150 mg/kg) and perfused transcardially with 0.9% NaCl followed by 4% paraformaldehyde in 0.1 M phosphate buffer. Brains were carefully withdrawn from the skull, and the ventrolateral aspect of the whole brain was imaged using a custom-built two-photon microscope system. The imaging system consisted of a mode-locked laser, operating at a 100-fs pulse width, 80-MHz pulse frequency, and 810-nm wavelength (Tsunami Millennia Xs or Mai Tai DeepSee, Spectra Physics), and an Olympus Fluoview scan box mounted on an upright BX51WI microscope. Images were captured with a 10× Olympus water immersion-type objective lens (NA = 0.3). Images were collected at different focus planes with a 10-μm interval. The images were stacked and tiled for enlarged views using Image-J and Photoshop software.

Some brains were post-fixed overnight and sectioned with a vibratome to observe the dye injection site. The 50-μm sections were mounted on glass slides using mounting medium that included DAPI (Vectorshied, VECTOR), and images were captured using an Olympus microscope with a CCD camera (Sensicam). In total, single-glomerulus dye injections were carried out in six brain hemispheres, and six superficial/deep EPL dye injection samples were used in this study for data analysis.

**RESULTS**

**AXON PROJECTION PATTERN FROM A SINGLE GLOMERULUS**

To determine how neurons projecting from the same glomerulus transmit odor information to higher brain centers, those neurons were labeled with a local dye electroporation method (Nagayama et al., 2007). The relationship between the dye-filled pipette and the target glomerulus, which expressed Synaptophysin–pHluorin protein, was monitored using two-photon microscopy (Figure 1A, inset).

In some cases, we could clearly observe the labeled projection neurons within 1 h after dye injection (Movie S1 in Supplementary Material). Three to five days after dye injection, the ventrolateral surface of the fixed whole brain was imaged by the two-photon system (Figure 1A). The labeled single-glomerulus axons could not be traced within the OB because they extended deep into the structure. They appeared on the brain surface near the border between the OB and anterior OC (AOC; this area includes the lateral, dorsal and ventroposterior divisions of the anterior olfactory nucleus; Haberly, 2001), then coursed along the lateral olfactory tract (LOT), and finally branched into the AOC, anterior/posterior PC (a/pPC), and OT. Some labeled axons extended to more caudal or ventral brain areas, most likely toward the lateral entorhinal cortex and amygdaloid cortex.

Non-neuronal cells also took up the fluorescent tracers and contributed to spotty background fluorescent noise. Despite this noise, the labeled individual axons could be clearly separated and traced. Surprisingly, each axon from a single glomerulus showed a distinct branching pattern and projected to multiple brain areas in a different way (Figures 1B–G). Overall, the axons from a common glomerulus covered an extensive area of the OC (Figure 1H). These results suggest that the projection neurons innervating the same glomerulus do not necessarily display the same axonal trajectory and are also consistent with recent experiments in zebrafish (Miyasaka et al., 2009).

According to their length and branching patterns, these axons could be largely categorized in two groups. One group of axons occupied the dorsolateral part of the LOT and split many axonal branches into the dorsolateral OC, predominantly to a/pPC (Figures 1B–C). These axons emitted only a few axon collaterals toward the OT. The other group occupied the ventrolateral part of the LOT and mainly projected to the ventrolateral OC, predominantly to the OT (Figures 1D–G). Such OT-orienting axons tended to project more densely to the anterolateral OT (a/OT). A total of 19 axonal trajectories from six brain hemispheres are summarized in Figure 1I, focusing on their secondary axon branching patterns. Each axon was categorized into one of five subgroups based on its axon projection orientation index: (OT branch number)/(OT + PC total branch number). This quantification clearly demonstrates an uneven branching pattern of projection neurons, with axons preferentially targeting either OT or PC. These data suggest that these two types of neurons sharing a glomerulus send information to different higher brain areas, such as a/OT or PC.

**DIFFERENTIAL AXON PROJECTION PATTERNS AFTER SUPERFICIAL OR DEEP EPL DYE INJECTION**

Projection neurons in the OB consist of mitral and tufted cells. Retrograde tracing and electrophysiological experiments suggest that mitral cells project to the entire OC while tufted cells project...
FIGURE 1 | Axonal trajectories of neurons from a single glomerulus. (A)
Two-photon images of the ventrolateral aspect of the whole brain. A schematic
Diagram of the dye injection and observation areas from the same ventrolateral
View of the whole brain is shown in the upper left. Inset: One-shot image of the
dye injection pipette and the target glomerulus before dye injection. Main figure:
Two-photon images of the olfactory cortex in the same animal. Labeled axons
appeared on the brain surface near the border between OB and AOC (upper left
part of the image) and tracked along the LOT and branch into a/pPC and OT. An
area of dense axonal projection appears in the anterior OT (yellow dotted circle).
Non-neuronal cells on the surface of the brain also uptake dye and show spotty
noise in the image. AOC, anterior olfactory cortex; aPC, anterior piriform cortex;
pPC, posterior piriform cortex; OT, olfactory tubercle. (B–G) Highlights of the
Individual axon trajectories in (A). (H) Superimposition of the images (A–G).
PC-oriented axons mainly run along the dorsal part of the LOT, while OT-oriented
axons mainly run along the ventral part of the LOT. The population of the
Single-glmomenuis axons covers a wide area of the OC. Presumed cortical borders
are drawn in yellow. (i) Axon projection orientation index. Nineteen axons were
categorized into five subgroups on the basis of their axon projection orientation
index calculated as (OT projection branches)/(OT + PC branches). PC, OT, and
neutral projection groups are shown as green, red, and gray columns, respectively.
to the anterior part of the OC (Haberly and Price, 1977; Sken and Hall, 1977; Scott et al., 1980; Scott, 1981; Schneider and Scott, 1983). However, no report has directly observed and compared the axonal trajectories of mitral and tufted cells in the same brain. One hypothesis is that the two observed types of axonal trajectories originate from these two types of OB projection neurons. To address this possibility, these neurons were labeled with different colored dyes. Mitral cells are known to have large cell bodies in the mitral cell layer and extend their secondary dendrites within the deep part of the EPL. In contrast, tufted cells have relatively small cell bodies in the EPL and the bottom of the glomerular layer (GL), and extend secondary dendrites within the superficial EPL (Mori et al., 1983; Orona et al., 1984; Shepherd et al., 2004). Therefore, differential dye injection into the superficial or deep EPL may increase the likelihood of labeling the two types of projection neurons distinctively.

By careful monitoring of the depth of the pipette tip, dextran-conjugated Alexa 488 (green) and Alexa 594 (red) were injected into the deep and superficial EPL, respectively (Figure 2A, insets). The dyes were localized to a small area of the EPL and labeled neurons that extended their dendrites to the injected area, as reported previously (Nagayama et al., 2007). Mitral and tufted

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**FIGURE 2 | Superficial and deep EPL cell axon projection pattern. (A)** Inset: Coronal image of the tracer injection site in the OB. Dextran-conjugated Alexa 488 (green) and Alexa 594 (red) were injected into the deep and superficial parts of the EPL, respectively, in the same area of the dorsal OB. The majority of the red-labeled neurons were tufted cells, and most green-labeled cells were mitral cells. DAPI nuclear labeling is blue. Main figure: Cortical observations in the same animal. The majority of green-labeled presumed mitral cell axons run along the dorsal part of the LOT and mainly branch into the a/pPC, although faintly green-labeled axons can also be observed in the OT. The majority of red-labeled presumed tufted cell axons run along the ventral part of the LOT and mainly project into the OT, although a small number of red-labeled axons can also be observed in the a/pPC. (B) Distribution of the labeled neurons in the OB. Neurons labeled by the superficial EPL (S-EPL; n = 37) and deep EPL (D-EPL; n = 82) dye injections were categorized in two groups, mitral cell layer cells (MCL cells; Green) and external plexiform layer/glomerular layer cells (EPL/GL cells; Red). (C) Axon projection orientation index. The majority of superficial EPL neurons (red column; n = 30) were grouped in the higher index number categories (right side of the graph), while deep EPL neurons (green columns; n = 34) tended to have lower index numbers (left side of the graph).
cells were successfully labeled selectively with different colored dyes. The labeled neurons were identified and grouped as mitral cell layer cells (MCL cells; presumably mitral cells) and EPL/GL cells (presumably tufted cells), by their distribution patterns near the dye injection area (six sections). Small number of neurons (21 cells) was labeled by both colored dyes. Those neurons were not used for the analysis. The segregation of the labeled cells into superficial and deep EPL injections is summarized in Figure 2B. Of the total number of cells labeled by deep EPL injections, 79% were MCL cells. Of the total number of cells labeled by superficial EPL injections, 86% were EPL/GL cells. As the background near the EPL dye injection sites was too bright in some sections to identify the labeled EPL neurons, the EPL/GL cells numbers could be underestimated. The labeled axons of presumed mitral cells and tufted cells were also clearly segregated within the LOT and in the OC. The majority of superficial EPL neuronal axons (red) mainly occupied the ventral part of the LOT and projected into the a/OT. The majority of deep EPL neuronal axons (green) occupied the dorsal part of the LOT and projected into the a/pPC (Figure 2A).

The axon pathways appeared segregated even in the anterior portion of the LOT; in the area of the AOC, and maintained this segregation to the anterior portion of the OT and a/PC. Some faintly green-labeled deep EPL axons could also be observed in the posterior part of the OT, and a few red-labeled superficial EPL axons could be observed in the PC. These results clearly suggest that, even in the same area of the OB, the majority of mitral and tufted cells show differential axonal trajectories and mainly project to different target areas, PC or aOT, respectively. Figure 2C shows the axon projection orientation index of the 30 superficial- and 34 deep EPL neurons in the six brain hemispheres, as well as the distinctive axonal projection patterns of deep and superficial EPL neurons, presumably mitral and tufted cells, respectively.

**DISCUSSION**

**LIMITATIONS OF THE TRACING METHOD**

To interpret the data accurately, the physical drawbacks of this tracing method must first be discussed. This method has the following difficulties and limitations: (1) tracing of obscured brain surface areas, such as the rhinal sulcus and septal regions of the OC, lateral entorhinal cortex, and amygdaloid cortex; (2) detection of weakly labeled and small branches; (3) detection of the axons that extend into deeper brain areas such as the ventroralstral anterior PC (APC_{vr}) region beneath the LOT (Ekstrand et al., 2001); and (4) identification of individual axons in axon-congested areas such as the anterior LOT in the area of the AOC. Comparatively, although classical single-neuron approaches have a higher quality of individual axon tracing in regard to the limitations above, our method allows the tracing and comparison of multiple axons simultaneously within the same sample. This advantage is critical for the exploration of large neuroanatomical circuits and allowed us to compare the individual axon trajectories arising from a common glomerulus as well as from differentially labeled presumed mitral and tufted cell populations. Further use of this method could reveal OB–OC connections more clearly and completely in future investigations.

**ODOR SIGNAL PROCESSING IN THE Olfactory CORTEX**

From the nasal epithelium to OB, olfactory information is transferred via genetically and anatomically sophisticated pathways to converge onto individual glomerular modules. From the OB to OC, however, our data indicate that individual projection neurons innervating a single glomerulus show quite divergent axonal projection patterns, and in total cover an extremely wide area of the OC. Therefore, the axonal output of each glomerulus should overlap with the projections of other glomeruli. These results strongly suggest that OB to OC projections may be organized according to different rules than those governing nasal epithelium to OB projections. Olfactory cortical neurons have been shown to receive multiglomerular inputs (Illig and Haberly, 2003; Murakami et al., 2005; Zou et al., 2005; Lei et al., 2006; Zou and Buck, 2006; Rennaker et al., 2007; Poo and Isaacson, 2009; Stettler and Axel, 2009), and individual OC neurons may reasonably collect such multiglomerular inputs in a different manner in different OC areas (Kadohisa and Wilson, 2006; Yoshida and Mori, 2007; Kikuta et al., 2008, 2010; Wesson and Wilson, 2010). As a result of this, a vast reorganization of the encoding and processing of olfactory information may occur in the OC (Miyasaka et al., 2009).

**PARALLEL PROCESSING IN THE MAIN olfactory SYSTEM**

Previous anatomical experiments have shown that PC neurons project to brain surface cortical areas, such as the orbitofrontal, insular, entorhinal, and perirhinal cortex, in addition to the amygdala (Luskin and Price, 1983; Johnson et al., 2000; Illig, 2005). In contrast, the OT is categorized as part of the ventral striatum and densely projects to other striatal areas including the ventral pallidum, caudate putamen, nucleus accumbens, ventral tegmental area, and lateral hypothalamic area (Scott and Chaffin, 1975; Millhouse and Heimer, 1984; Ikemoto, 2007). Moreover, topographic connections between OT subareas and the nucleus accumbens and ventral pallidum have been reported (Ikemoto, 2007). The aOT could potentially establish a specific stream among the pathways between OT and striatum. Taken together with the present findings, these studies suggest that the segregation of distinct olfactory processing pathways originating in mitral and tufted cells is conserved not only from OB to OC but also in higher brain centers.

One attractive hypothesis is that mitral and tufted cells initiate parallel processing streams in the olfactory system. An important component of parallel processing in other sensory systems is that each pathway in the parallel streams has different functions, handles different aspects of the same sensory input, and processes information in different brain areas. Compared with mitral cells, tufted cells have a low spike threshold, weak and narrow tuning range of lateral inhibition, high firing frequency, and strong respiratory locking activity (Schneider and Scott, 1983; Ezeh et al., 1993; Nagayama et al., 2004; Griff et al., 2008). These data suggest that mitral cells may be involved in comparing their own and surrounding glomerular activities, while tufted cells may in turn be involved in rapid odor detection over the respiratory rhythm, even at low concentrations. Moreover, tufted cell circuits have been reported as highly sensitive and plastic to sensory deprivation (Imamura et al., 2006). OT neurons receive substantial dopaminergic inputs from the ventral tegmental area, which are related to reward signals (Ikemoto, 2007). These reports suggest that the tufted cell pathway may have a critical
function in monitoring the odor environment and changing animal behavioral states in response to external environmental changes. In contrast, a mitral cortical pathway may be adequate to establish an accurate odor image. For a clearer characterization of these two olfactory pathways, a direct functional comparison of mitral and tufted cells common to the same glomerulus, as well as of PC and OT, should be conducted in the future.

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SUPPLEMENTARY MATERIAL

MOVIE S1 | Representative example of the labeled neurons associated with a single glomerulus. Dorsal part of the OB was imaged 1 h after dye injection by two-photon microscopy. Images were aligned from the superficial to the deep part of the OB. Green fluorescence indicates olfactory receptor neuronal axons that express Synapto-pHluorin protein. Red colored neurons and their dendrites are labeled by dextran-conjugated Alexa 594. All of the labeled neurons (periglomerular, tufted and mitral cells) are associated with a single glomerulus.

MOVIE 1

In vivo two-photon imaging of the dye-labeled neurons associated with a single glomerulus.