Centrifugal Inputs to the Main Olfactory Bulb Revealed Through Whole Brain Circuit-Mapping

Krishnan Padmanabhan1,2,3*, Fumitaka Osakada4,5, Anna Tarabrina6, Erin Kizer6, Edward M. Callaway1,4, Fred H. Gage6 and Terrence J. Sejnowski1,2,7

1 Crick-Jacobs Center for Theoretical and Computational Biology, Salk Institute for Biological Studies, La Jolla, CA, United States, 2 Computational Neurobiology Laboratory, Salk Institute for Biological Studies, La Jolla, CA, United States, 3 Department of Neuroscience, University of Rochester School of Medicine and Dentistry, Rochester, NY, United States, 4 Systems Neurobiology Laboratories, Salk Institute for Biological Studies, La Jolla, CA, United States, 5 Laboratory of Cellular Pharmacology, Graduate School of Pharmaceutical Sciences, Nagoya University, Nagoya, Japan, 6 Laboratory of Genetics, Salk Institute for Biological Studies, La Jolla, CA, United States, 7 Howard Hughes Medical Institute, Salk Institute for Biological Studies, La Jolla, CA, United States

Neuronal activity in sensory regions can be modulated by attention, behavioral state, motor output, learning, and memory. This is often done through direct feedback or centrifugal projections originating from higher processing areas. Though, functionally important, the identity and organization of these feedback connections remain poorly characterized. Using a retrograde monosynaptic g-deleted rabies virus and whole-brain reconstructions, we identified the organization of feedback projecting neurons to the main olfactory bulb of the mouse. In addition to previously described projections from regions such as the Anterior Olfactory Nucleus (AON) and the piriform cortex, we characterized direct projections from pyramidal cells in the ventral CA1 region of hippocampus and the entorhinal cortex to the granule cell layer (GCL) of the main olfactory bulb (MOB). These data suggest that areas involved in stress, anxiety, learning and memory are all tethered to olfactory coding, two synapses away from where chemical compounds are first detected. Consequently, we hypothesize that understanding olfactory perception, even at the earliest stages, may require studying memory and behavior in addition to studying the physiochemical features of odors.

Keywords: olfactory bulb, feedback, olfaction, retrograde tracer, circuits

INTRODUCTION

Understanding the neural basis of olfactory perception remains one of the key challenges of sensory neuroscience. Part of the challenge arises from the inherently high dimensional nature of chemical structure (Secundo et al., 2014). For instance, mapping olfactory percepts onto any single feature of chemical structure such as carbon chain length, remains difficult. A parallel approach has focused on anatomy and physiology. In rodents, the neuronal representations of odors have been investigated by tracking neural activity from the olfactory receptor neurons (ORNs), where components of smell are first detected, to the piriform cortex (STettler and Axel, 2009; Davison and Ehlers, 2011), where they are assembled combinatorially (Babadi and Sompolinsky, 2014) into odor percepts. However, such a feedforward model of olfactory coding, wherein neuronal responses are sequentially marshaled into more complex representations by progressively higher cortical areas glosses over the major feedback or centrifugal projections present in olfaction. As a result,
CMV-eGFP was generated at titers of 1.20 × 10^11 infectious units/ml at the core facility at the Salk Institute. Adeno-associated viruses (AAV) were generated by the GT3 Core facility at the Salk Institute. AAV1-CMV-eGFP and AAV2-CMV-eGFP were generated at titers of 1.20 × 10^11 – 5.13 × 10^12 infectious units/ml. The viruses were stored at −80°C until use.

**METHODS**

Some of the data used in this manuscript has been previously published (Padmanabhan et al., 2016). In this previous work, the focus was exclusively on projections from the AON and the piriform cortex, with the goal being to understand the organization of feedback circuits from these two olfactory regions. Here we provide a quantitative assessment of the organization of the feedback projections from the entire mouse brain. Additionally, this work uses analysis and provides information about the relative distribution of feedback projections that were not previously reported or quantified.

**Rabies Virus**

GFP, mCherry, or BFP was cloned in pSADΔG-F3 as described previously (Wickersham et al., 2007; Osakada et al., 2011; Osakada and Callaway, 2013). SADΔG-GFP, SADΔG-mCherry, and SADΔG-BFP were recovered in B7GG cells with transfection with the corresponding genomic plasmid, pcDNA-SADΔB19N, pcDNA-SADΔB19P, pcDNA-SADΔB19L, and pcDNA-SADΔB19G. Viruses were amplified in B7GG cells in a humidified atmosphere of 3% CO₂ and 97% air at 35°C and concentrated by two rounds of ultracentrifugation. The concentrated rabies viruses were titrated in HEK293t cells. The titers of the rabies viruses used in the present study were 5.0 × 10^8 – 3.0 × 10^9 infectious units/ml.

Adeno-associated viruses (AAV) were generated by the GT3 Core facility at the Salk Institute. AAV1-CMV-eGFP and AAV2-CMV-eGFP was generated at titers of 1.20 × 10^11 – 3.25 × 10^12 infectious units/ml. The viruses were stored at −80°C until use.

**Animals**

This study was carried out in accordance with the accordance with the guidelines for care and use of animals by the Institutional Animal Care and Use Committee (IACUC) of the Salk Institute for the Salk Institute for Biological Studies. The protocol was approved by the Institutional Animal Care and Use Committee (IACUC) of the Salk Institute for Biological Studies. Twenty mice (male and female) aged 3–4 months were used each having 1 to 2 injections (N = 23 injections total). The GCL was targeted with stereotaxic coordinates (Bregma +4.25 mm rostral/caudal, +1 mm medial/lateral, −0.6 mm to −1.5 mm dorsal/ventral) using a digital micromanipulator (Kopf, CA, USA). Stereotactic injections to the CA1 region were made relative to bregma (−3.25 mm rostral/caudal, + 3.25 mm medial/lateral, −4 mm dorsal/ventral). Borosilicate micropipettes were pulled on a Sutter P200 and 40–200 nl of virus were delivered to each bulb via pulsed injection from a picospritzer (Parker, OH, USA). Three different fluorescent reporters (GFP, mCherry, and BFP) in the G-deleted rabies viruses were used for the experiments and all 3 labeled neurons extensively, including labeling of dendritic processes. For AAV viral vectors, either eGFP or dsRed were used as fluorescent reporters. Additionally, we used a Cholera-Toxin-β subunit tracer for some anterograde experiments.

**Histology**

Three to 10 days after the viral or fluorescent reporter injection, animals were sacrificed and perfused with 4% paraformaldehyde (PFA) and brains were extracted and transferred to a solution of 4% PFA/30% sucrose (Padmanabhan et al., 2010). Coronal sections, 100 μm in thickness, were made of the mouse brain from the bulb to approximately Bregma −4.25 mm, allowing us to characterize inputs to the bulb. In 3 of the animals, damage while extracting the brain resulted in incomplete reconstructions of some regions such as hippocampus and occipital cortex. In 2 additional animals, tissue warped during perfusion, and though these were qualitatively to determine properties such as injection site size, number of labeled neurons etc., they were not used in calculating the relative distribution of feedback projecting cells from different brain regions.

**Imaging and Data Management**

Images were acquired with an Olympus VS110 slide scanner (Tokyo, Japan). Complete scans of all coronal sections were done at 5 virtual Z-planes for each section. Each coronal section produced a stack of 5 ~30,000 × 50,000 pixel 16-bit images, which were then collapsed into a single maximum intensity projection. Each whole brain scan resulted in ~1 TB of data. For analysis and viewing, each individual section was sorted in a pyramid representation. For each representation

$$G(x, y) = I(x, y) \text{ for level, } l = 0$$

$$G_l(x, y) = \frac{1}{M^N} \sum_{m=-2}^{M} \sum_{n=-2}^{N} G_{l-1}(x + m, y + n)$$

Such that G is the pyramid representation and I the original image where M = 2 and N = 2. Our resolution resulted in representations for l = 0–3.

**Image Dilation and Correlation**

Individual coronal sections were aligned and alignment validated by correlation analysis (using custom functions in MATLAB (MathWorks, MA, USA). First, images were thresholded at different image places using previously described methods (Padmanabhan et al., 2010). This method simply finds an optimal threshold for binary images, or a series of binary image planes corresponding to individual objects of interest. To extent this to identify the boundaries of cells and tissues, we performed additional image processing. First, the Moore-Neighbor tracing algorithm was used to identify the boundaries of cell sections.
as well as boundaries for individual neurons (Gonzalez et al., 2004). Boundaries where then convolved with a 2-dimensional Gaussian to minimize alignment sensitivity to pixel noise, and then a correlation coefficient between all pairs of sections was used to validate the alignment.

**Cell Finding**

For each fluorescent image section, regions of interest were manually identified. Only label within these identified areas was used for analysis, thereby excluding background fluorescence. To identify neurons in an automated way, we defined an image $X$ as where $x_n$ corresponds to the $n$th pixel:

$$x = x_1, x_2, x_3, \ldots x_n$$

We defined another vector $y$ where $y_n$ corresponding to the $n$th pixel:

$$y = y_1, y_2, y_3, \ldots y_n$$

such that

$$y_n \begin{cases} = 1 & \text{if } x_n > i \\ = 0 & \text{else } x_n \leq i \end{cases}$$

where $i$ was a value selected manually for thresholding. Individual sections were thresholded using either a manually selected threshold value, or alternatively, using an optimal image thresholding method (Padmanabhan et al., 2010).

$$\text{corr} \left(x, y(i)\right) = \frac{\sum_{j=1}^{m} n_j \ast (i - \bar{x})}{\sqrt{\left(\sum_{j=1}^{m} (j - \bar{x})^2 \ast n_j\right) \ast \left(\frac{N_m - N_i + N_i}{N_m}\right)}}$$

where $x$ is the original image and the thresholded image $y$ at the threshold value $(i-1)$ with $j$ as the bit-depth of the image (for example, between 0 and 255 for an 8-bit image), $n_j$ as the number of pixels with the value $j$ and $\bar{x}$ as the mean pixel value of the image. In addition, $N_m$ was the total number of pixels in the image and $N_i$ was the number of pixels above the threshold value $(i-1)$. We rewrote our correlation where $x$ was the original image and $y(i)$ was the image thresholded at a value of $i$. The optimal correlation:

$$\text{argmax } t(m) = \text{corr} \left(x, y(m)\right)$$

Thresholded images were then run through a series of image processing functions to identify retrogradely labeled neurons. To take thresholded images and segment individual neurons (which constitute multiple pixels), we first, performed a nearest-neighbor pixel grouping to assign contiguous pixels into neurons. Following this step, groups of pixels were sorted based on morphological properties, including the shape (how round to the neurons look, which was biased by the median filtering step) and the number of pixels within each group (which eliminated small pixels that corresponded to either processes or large groups of cells that are clumped together and cannot be easily segmented).

**Data Analysis**

All analysis described in this manuscript, including tissue identification, was done with custom functions written in MATLAB. Unless otherwise noted, error bars are standard deviations.

**RESULTS**

A g-deleted rabies virus expressing a fluorescent reporter was injected into the granule cell layer (GCL) (Haberly and Price, 1978) of the MOB in adult mice (3–6 months of age, $N = 20$, Figure 1A). Retrogradely labeled cells infected with the virus did not express the rabies glycoprotein, thus fluorescence was confined only to neurons at the injection site (Figure 1B) and to those cells whose axonal buttons terminated at the site of the injection (Wickersham et al., 2007; Osakada and Callaway, 2013; Callaway and Luo, 2015). As feedback input to the glomerular layer (Petzold et al., 2009) and the mitral cell layer (Yan et al., 2008; Markopoulos et al., 2012) have been previously studied, we focused our analysis on the GCL. Following 3–10 days of viral infection ($N = 4$ at 3 days, $N = 17$ at 5–10 days), animals were sacrificed and coronal sections of the whole mouse brain were made. To characterize the spatial organization of centrifugal projections to the GCL, we established a whole-brain imaging and analysis method (Figure 1C). First, we collected data from ~160 to 180 µm sections constituting the mouse brain (Figure 1D, we did not section regions of the cerebellum and the most caudal regions of the brainstem).

In order to identify all retrogradely labeled neurons throughout the brain, we needed to perform two segmentation and identification tasks: isolate the tissue to delineate brain areas and identify the location of individual cell bodies (Figure 2A). To do this, we first median filtered the raw images (Figure 2B) to eliminate shot noise in our imaging, as well as to smoothen out the edges of the cell bodies (Figure 2Bi). Although this did reduce the ability to reconstruct individual dendrites (Figure 2Bi), it allowed us to rapidly segment individual neurons and the regions of the brain they inhabited. Next, we used the Maximum Correlation Thresholding (MCT) algorithm (Padmanabhan et al., 2010) which thresholds the image at each single gray-scale value and then calculates the correlation of the thresholded image to the original (Figure 2C, top). The curve represented the correlation of the thresholded image to the original image for each gray-scale value, and had distinct transitions points that were clear when we visualized the derivative of the correlation curve (Figure 2C, bottom). Changes in the correlation corresponding to the edge of the tissue (Figure 2C, red line) and the labeled cells (Figure 2C, purple line) were apparent in both the correlation curve and the derivative curve. While the MCT algorithm provides thresholded images corresponding to various objects (tissue edge, individual neurons, etc.), it does not identify which feature are important, nor does it segment these individual images into objects (such as the various boundaries of tissues, or neurons) that are defined by multiple contiguous pixels in an imaging plane. To address these issues, we first chose thresholding at these two different gray-scale values
Following segmentation of the tissue, a segmentation was performed on the threshold images corresponding to labeled neurons which included shape analysis and cell size identification (See methods). The result of the algorithm appeared to be well-identified neurons. To determine the fidelity of the automated cell finder, in a subset of coronal sections from multiple animals, an expert used identified individual neurons manually and these were compared to the algorithm. Two representative examples of the olfactory bulb (Figure 3C, top) and the piriform cortex (Figure 3C, bottom) show neurons that were identified by the human user manually (red dots), and by the MCT algorithm (blue dots). In both examples, the automated cell finder identified the neurons that an expert human also identified in two distinct regions, with different fluorescent intensities, different neuronal morphologies, and different densities. When we quantified the accuracy of the MCT algorithm to the manual count, we found that across all comparisons, there was an $R^2 = 0.84$, and this was significantly different from chance (Figure 3D, black line is the unit line, gray line is the best-fit for the data). When we examined the correlation between the manual expert and the MCT algorithm further, we found that underestimated occurred when the density of neurons being counted was high (Figure 3E, red area). These corresponded to regions at or near the injection site, where multiple neurons overlapped one-another and the algorithm had difficulty parsing them into individual cells. Despite this limitation, the 0.84 correlation of the algorithm to neurons identified by an expert human, allowed us to study the organization of feedback projecting cells to the bulb.

We first analyzed the precision of our injections to the GCL of the bulb (Figures 4A,B). Boundaries were manually drawn in the tissue (light gray) between the inner plexiform layer (IPL) and the GCL. (Figures 4A,B, dark gray) with the asterisk corresponding to the center of mass of the injection site (taken as the average of the Euclidian position of all the labeled neurons within the bulb). A 3D reconstruction of the bulb from this experiment revealed that although the injection site was confined to the GCL (Figure 4C, asterisk, red points correspond to labeled neurons within the GCL, gray lines correspond to the boundaries of the GCL for each coronal section), a small subset of neurons were also labeled outside the GCL. (Figure 4C, magenta cells), likely due to label being taken up by neurites passing through the injection site and corresponding to mitral/tufted cells (Hovis et al., 2010; Padmanabhan and Urban, 2010). We assessed the ratio of labeled cells inside (Figure 4D, red cells) to cells outside

(Figure 2D), and using existing image segmentation algorithms (Padmanabhan et al., 2016) allowed us to define the tissue shape for each section (Figures 2D–E) as well as identify the individual cell bodies (Figures 2Di–Ei). As different traces have different levels of fluorescence corresponding to different objects (due to experimental variability like injection size, viral titer, cell type, and image parameters) we confirmed that the MCT algorithm produced different correlation curves (Figure 2F, left, red) reflecting differences in fluorescent pixel intensity across the various sections in a single mouse. The approach was robust to differences in pixel intensity across multiple fluorophores including GFP, RFP and, in this example Blue Fluorescent Protein (BFP, Figure 2F, right, blue curves). While the bit-depth for the maximum correlation was different across different experiments (Figure 2G), the shape of the correlation coefficient curve varied smoothly along the rostral caudal axis (Figure 2H), reflecting the overall decrease in mean fluorescence from the injection site in the bulb (where the density of labeled cells was high) to the caudal regions of the mouse brain.

Next, we wished to align the tissue and validate the image segmentation of cell bodies to assess the utility of the method. Using the thresholded images in 2D, we align section edges along the rostro-caudal axis (Figure 3A), which provided a 3D representation of the entire mouse brain. To determine the fidelity of this alignment, we calculated the correlation of each section to all adjacent sections (Figure 3B), with varying degrees of tolerance (as represented by the thickness of the edges of each section = gray lines). Correlation values were high in adjacent sections, but fell of dramatically, confirming that alignment of individual sections was tight (Figure 3B).

We first analyzed the precision of our injections to the GCL of the bulb (Figures 4A,B). Boundaries were manually drawn in the tissue (light gray) between the inner plexiform layer (IPL) and the GCL. (Figures 4A,B, dark gray) with the asterisk corresponding to the center of mass of the injection site (taken as the average of the Euclidian position of all the labeled neurons within the bulb). A 3D reconstruction of the bulb from this experiment revealed that although the injection site was confined to the GCL (Figure 4C, asterisk, red points correspond to labeled neurons within the GCL, gray lines correspond to the boundaries of the GCL for each coronal section), a small subset of neurons were also labeled outside the GCL. (Figure 4C, magenta cells), likely due to label being taken up by neurites passing through the injection site and corresponding to mitral/tufted cells (Hovis et al., 2010; Padmanabhan and Urban, 2010). We assessed the ratio of labeled cells inside (Figure 4D, red cells) to cells outside...
the GCL but still within the bulb (Figure 4D, magenta cells) to determine the precision of our targeting (Figure 4D, black line in each corresponds to the GCL boundary). In this example, 84% of fluorescently labeled neurons were inside the GCL, similar to the percentage of neurons inside the GCL across all experiments, (80 ± 19%, N = 9).

When we examined the distribution of neurons within the injection site, we found that they were confined to the GCL (Figure 4E), with asymmetries in the density of label corresponding to the non-spherical shape of the GCL along the medial-lateral axis (Figure 4E, left). Additionally, our injections were slightly biased to the dorsal and caudal domains of the GCL as evidenced by the skewed distribution of the labeled cells along the dorsal-ventral axis (Figure 4E, middle) and the rostral-caudal axis (Figure 4E, right). While the representative example in Figure 4C shows a dorsal biased injection, additional injections labeled various regions of the GCL (Figure 4E). Although we observed no significant/systematic differences in the label density across position (Figure 4E), this does not rule out the possibility that such differences could exist. Importantly however, these
distributions corresponded to the 3D volume of the GCL in our reconstructions (Figure 4C), further confirming the precision of our targeting. To determine the regions projecting to the GCL, we next examined the distribution of retrogradely labeled neurons throughout the mouse brain. A density map from the injection site in 4C (Figure 4F), and the distribution of label across 10 experiments for which we had reconstructions (Figure 4G, left), revealed that feedback-projecting neurons were almost entirely from ventral regions of the mouse brain (Figure 4G, right), predominantly archicortical in origin (piriform cortex, CA1, etc.) or neuromodulatory (HDB). As we saw label across these regions in an array of experiments (Figure 4G), feedback projections are likely to target both dorsal and ventral bulb.

To provide a more detailed description of the origins of the feedback projections, we aligned all of the brain sections to a mouse brain atlas (Paxinos and Franklin, 2004) and identified the areas corresponding to retrogradely labeled neurons (Figure 5A). We found labeled neurons in the both the ipsilateral (58%) and contralateral (5%) anterior olfactory nucleus (AON), the piriform cortex and the nucleus of the lateral olfactory tract (nLOT; 27.9%), the horizontal limb of the Diagonal Band (HDB; 3.5%), amygdala regions (1.4%), zona inserta (0.001%), the piriform-entorhinal cortex (0.001%) transition and in various nuclei of the hypothalamus (0.08%), consistent with previous reports (Figure 2F, Shipley and Adamek, 1984) as well as projections to the GCL directly from the entorhinal cortex (Figure 5A, light blue cells, 0.3%) and direct centrifugal input from the CA1 region of the hippocampus (Figure 5A, green, 0.75%) as had been hypothesized to exist previously in other species (de Olmos et al., 1978). We quantified the density of these projections from various regions (Figure 5B), and found that while the largest density of retrograde labeled cells came from major olfactory cortical areas (ipsilateral AON, contralateral AON = 63%), 10% of the retrogradely labeled neurons in the brain originated from areas that are not thought of as part of olfactory cortex.

Perhaps most unexpected were the retrogradely labeled neurons we found in the hippocampus and the entorhinal cortex [Figure 5A, light blue cells, and green cells (Leitner et al., 2016)]. Although some recent work has suggested that pyramidal cells in the CA1 region of hippocampus send axons directly to the bulb (Okuyama et al., 2016), it remains unclear if these are projections directly to the MOB or if these target the accessory olfactory bulb (AOB)(de Olmos et al., 1978; Shipley and Adamek, 1984).

To address this, we determined the spatial location of the neurons projecting from CA1. First, we mapped the location of cells onto a 3D model of the CA region of hippocampus (Figures 6A,B) and found retrogradely labeled cells were confined largely to the ventral-lateral region (Figure 6C, N = 6), and could be identified as pyramidal based on their morphology (Figures 6D1, D2). As the rabies label filled dendritic processed, we were able to reconstruct multiple neurons, all of which had soma in the stratum pyramidale, apical dendrites projecting into the stratum radiatum and basal dendritic arbors in the stratum oriens (Figure 6D2). To the best of our knowledge, these are the only direct projections from the CA1 to a primary sensory area (Oh et al., 2014), and they are all the more interesting because they target neurons 2 synapses downstream from olfactory receptors in the sensory epithelium where odors are first detected. To ensure that these vCA1 pyramidal neurons were not retrogradely labeled because our injections accidently spilled over into the AOB, we examined in detail the injection site in the bulb for the experiments (Figure 6E, N = 6) where
vCA1 cells were densely labeled. In these experiments, the MOB (Figure 6F1) was demarcated from the AOB (Figure 6F2). Neurons labeled in the injection site (Figure 6G, MOB cells in black, AOB cells in red) revealed that less than 1% of the fluorescent labeled neurons in the injection site (99.3% = MOB, 0.07% = AOB, Figure 6H1) were in the granule layer of the AOB, further suggesting that projections from vCA1 did directly to the GCL of the MOB. We further confirmed this direct projection by making injections of the anterograde tracer Cholera Toxin-β subunit conjugated to an Alexa 488/555 dye (Figure 6J) into the CA1 region (Figures 6J,K). Clear axonal projections in the GCL of the bulb (Figures 6L1, L2), providing complementary evidence of the link between CA1 and the olfactory bulb.

DISCUSSION

By combining a retrograde virus-labeling strategy with whole-brain 3D reconstruction methods, we describe the structure of centrifugal projections to the GCL of the bulb. First, from a methodological perspective, we provide an approach to segmentation of tissue and neurons that could be used more generally in a number of imaging applications. For instance, in light sheet microscopy application where voxel based approaches have been used (Ahrens et al., 2013), the strategy described here could afford additional insight into the dynamics of single neurons. Additionally, this method may be combined with brain clearing approaches (Liebmann et al., 2016) to reveal the spatial distributions of neurons within and across brain areas. Second,
in addition to known projections from olfactory areas such as the AON and the piriform cortex and neuromodulatory areas such as the HDB, we identified direct CA1 pyramidal cell projections from the ventral/lateral region of the hippocampus and the entorhinal cortex directly to the bulb. Although these tracing methods cannot identify the cell types targeted by centrifugal projections to the GCL, which include granule cells, short axon cells, and the lateral dendrites of mitral/tufted (M/T) cells, they do suggest a diverse group of areas all send centrifugal projections to the bulb and could ultimately influence the neuronal activity of M/T cells in response to incoming stimuli. While this feedback and reciprocal connectivity has been identified previously in other species (Shipley and Adamek, 1984; Van Groen and Wyss, 1990) including the mouse (Mohedano-Moriano et al., 2012), and the findings of this study are in line with these previous descriptions, recent studies have overlooked these projections in detailing hippocampal connectivity in the mouse (Bienkowski et al., 2018). Additionally, our imaging method allowed us to identify the spatial pattern of cells across the brain (Figures 3–5) as well as provide a quantitative description of the relative density of cell projections from all of these feedback regions (Figure 5). Notably, we quantified that nearly 10% of projections come from regions other than olfactory cortex and piriform (Figure 5).

Furthermore, that feedback from regions such as vCA1 and EC are pronounced across multiple experiments (Figures 5, 6) suggests that these circuits could play important roles in shaping odor representations. Although these regions constitute ~1% of the feedback projections we identified, they number of cells corresponds to the ratios observed in amygdala (1.3%) and ~1/3 of the cell density of feedback from HDB (3%). Other previous work has shown the role that cholinergic neurons from the basal forebrain play in shaping mitral cells responses (Rothermel et al., 2014), suggesting that even small number of connections, such as those from vCA1 and entorhinal cortex may have a functional role. Furthermore, as vCA1 also projects to piriform cortex, both monosynaptic and disynaptic feedback projections may pay direct and indirect roles across multiple timescales to influence the firing of mitral and tufted cells. Finally, while our method examines the feedback to the GCL, distinctions in between dorsal and ventral bulb (Kobayakawa et al., 2007) may emerge functionally based on asymmetries in the feedback. The functional role of these projections shall be an important focus of future experiments.

While M/T cells, the principal neurons of the bulb are responsive to odor stimulation, their activity can be modulated by other sources (Kay and Laurent, 1999), including centrifugal input, both from olfactory regions (Wesson et al., 2008; Boyd et al., 2012; Markopoulos et al., 2012) and from neuromodulatory centers (Rothermel et al., 2014). In addition to these known projections, we characterized additional feedback from the ventral region of CA1 hippocampus, whose spatial location may provide a hint of their functional role (Moser and Moser, 1998; van Strien et al., 2009; Fanselow and Dong, 2010). Coherence in the electrical activity of Local Field Potentials (LFPs) between the MOB and the hippocampus has been previously described (Martin et al., 2007), including oscillatory coupling in the beta (15–35 Hz) frequencies (Gourévitch et al., 2010), suggesting that these feedback circuits could play a role in LFP coherence between the two regions. Furthermore, ventral/lateral regions of CA1 are important for encoding stress responses (Henke, 1990) and, accordingly, feedback could carry this behaviorally relevant information to gate odor processing at the level of the bulb.
The limitations of adducing function from structure alone notwithstanding, our description of centrifugal projections suggests unique features of olfactory system. First, unlike other sensory systems, olfactory circuits are highly compact. They target the amygdala and cortex without a thalamic way point and receive direct feedback at the earliest processing stages from a variety of brain regions. Second, circuits tasked with encoding chemical sensation are tethered to evolutionarily old circuits such as those of the hippocampus. Finally, centrifugal feedback to the bulb is diverse.

Unlike other sensory modalities, like vision or audition, where stimulus features have clear equivalents in perceptual space, the link between chemical structure and odor perception remains an open question (Secundo et al., 2014). We propose that one reason for this difference is that the olfactory system has privileged access to the information about prior olfactory representations, the animal’s behavioral state, and input from learning and memory systems at the level of the mitral and tufted cells in the bulb via direct centrifugal innervation (Restrepo et al., 2009; Gire et al., 2013). Consequently, coding at the bulb may be far less of a
purely sensory operation and more about the interplay between chemical sensation and memory (Wilson and Stevenson, 2003).

AUTHOR CONTRIBUTIONS

KP conceived the project, designed, carried out the experiments and performed the analysis, and wrote the manuscript. FO provided reagents and assisted with experiments. AT and EK performed histology and imaging. FO, AT, and EK provided feedback on the manuscript. EC, FG, and TS provided assistance with project development and experiment design and contributed feedback to the manuscript writing and editing.

FUNDING

KP is supported by a NIMH K99 MH1101634, R01 MH113924, a NSF CAREER Award, a NARSAD Young Investigator Award, the Lieber Foundation, and a Crick-Jacobs Junior Fellowship. FO was supported by the Japan Society for the Promotion of Science, the Japan Science and Technology Agency, the Japan Agency for Medical Research and Development, the Kanae Foundation for the Promotion of Medical Science, the Uehara Memorial Foundation, the Naito Foundation and the Pioneer Fund. EC was supported by the National Institutes of Health and the Kavli Institute for Brain and Mind at University of California San Diego. Support for FG was provided by The Mathers Foundation, The JPB Foundation, and the Helmsley Foundation. TS was supported by Howard Hughes Medical Institute, ONR N000141310672, and ONR N000141210299.

REFERENCES

Ahrens, M. B., Orger, M. B., Robson, D. N., Li, J. M., and Keller, P. J. (2013). Whole-brain functional imaging at cellular resolution using light-sheet microscopy. Nat. Methods 10, 413–420. doi: 10.1038/nmeth.2434

Babadi, B., and Sompolinsky, H. (2014). Sparseness and expansion in sensory representations. Neuron 83, 1213–1226. doi: 10.1016/j.neuron.2014.07.035

Bienkowski, M. S., Bowman, I., Song, M. Y., Gou, L., Ard, T., Cotter, K., et al. (2018). Integration of gene expression and brain-wide connectivest reveals the multiscale organization of mouse hippocampal networks. Nat. Neurosci. 21, 1628–1643. doi: 10.1038/s41593-018-0241-y

Boyd, A. M., Sturgill, J. F., Poo, C., and Isaacson, J. S. (2012). Cortical feedback control of olfactory bulb circuits. Neuron 76, 1161–1174. doi: 10.1016/j.neuron.2012.10.020

Callaway, E. M., and Luo, L. (2015). Monosynaptic circuit tracing with glycoprotein-deleted viruses rabies. J. Neurosci. 35, 8979–8985. doi: 10.1523/JNEUROSCI.0409-15.2015

Davison, I. G., and Ehlers, M. D. (2011). Neural circuit mechanisms for pattern detection and feature combination in olfactory cortex. Neuron 70, 82–94. doi: 10.1016/j.neuron.2011.02.047

de Olmos, J., Hardy, H., and Heimer, L. (1978). The afferent connections of the main and the accessory olfactory bulb formations in the rat: an experimental HRP-study. J. Comp. Neurol. 181, 213–244. doi: 10.1002/cne.901810202

Fanselow, M. S., and Dong, H.-W. (2010). Are the dorsal and ventral hippocampus functionally distinct structures? Neuron 65, 7–19. doi: 10.1016/j.neuron.2009.11.031

Gire, D. H., Restrepo, D., Sejnowski, T. J., Greer, C., De Carlos, J. A., and Lopez-Mascaraque, I. (2013). Temporal processing in the olfactory system: can we see a smell? Neuron 78, 416–452. doi: 10.1016/j.neuron.2013.04.033

Gonzalez, R. C., Woods, R. E., and Eddins, S. L. (2004). Digital Image Processing Using MATLAB. New Jersey, NF: Pearson Prentice Hall.

Gourévitch, B., Kay, L. M., and Martin, C. (2010). Directional coupling from the hippocampus to the olfactory bulb during a go/no-go odor discrimination task. J. Neurosci. 30, 2633–2641. doi: 10.1152/jn.01075.2009

Haberly, L. B., and Price, J. L. (1978). Association and commissural fiber systems of the olfactory cortex of the rat. J. Comp. Neurol. 178, 711–740. doi: 10.1002/cne.901780408

Henke, P. G. (1990). Hippocampal pathway to the amygdala and stress ulcer development. Brain Res. Bull. 25, 691–695.

Hovis, K. R., Padmanabhan, K., and Urban, N. N. (2010). A simple method of in vivo electroporation allows visualization, recording, and calcium imaging of local neuronal circuits. J. Neurosci. Methods 191, 1–10. doi: 10.1016/j.jneumeth.2010.05.017

Kay, L. M., and Laurent, G. (1999). Odor- and context-dependent modulation of mitral cell activity in behaving rats. Nat. Neurosci. 2, 1083–1089. doi: 10.1038/14801

Kobayakawa, K., Kobayakawa, R., Matsumoto, H., Oka, Y., Imai, T., Ikawa, M. et al. (2007). Innate versus learned odor processing in the mouse olfactory bulb. Nature 450, 503–508. doi: 10.1038/nature06281

Leitner, F. C., Melzer, S., Lütcke, H., Pinna, R., Seeburg, P. H., Helmchen, F., et al. (2016). Spatially segregated feedforward and feedback neurons support differential odor processing in the lateral entorhinal cortex. Nat. Neurosci. 19, 935–944. doi: 10.1038/nn.4303

Liebmann, T., Renier, N., Betayeb, K., Greengard, P., Tessler-Lavigne, M., and Flajolet, M. (2016). Three-dimensional study of Alzheimer’s disease hallmark using the isdclo clearing method. Cell Rep. 16, 1138–1152. doi: 10.1016/j.celrep.2016.06.060

Markopoulos, F., Rokni, D., Gire, D. H., and Murthy, V. N. (2012). Functional properties of cortical feedback projections to the olfactory bulb. Neuron 76, 1175–1188. doi: 10.1016/j.neuron.2012.10.028

Martin, C., Beshel, J., and Kay, L. M. (2007). An olfacto-hippocampal network is dynamically involved in odor-discrimination learning. J. Neurophysiol. 98, 2196–2205. doi: 10.1152/jn.00524.2007

Mohamed-Moriano, A., de la Rosa-Prieto, C., Saiz-Sanchez, D., Ubeda-Bañon, I., Pro-Sistiaga, P., de Moya-Pinilla, M., et al. (2012). Centrifugal telencephalic afferent connections to the main and accessory olfactory bulbs. Front. Neuroanat. 6:19. doi: 10.3389/fnana.2012.00019

Moser, M. B., and Moser, E. I. (1998). Functional differentiation in the hippocampus. Hippocampus 8, 688–619.

Oh, S. W., Harris, J. A., Ng, L., Winslow, B., Cain, N., Mihalas, S., et al. (2014). A mesoscale connectome of the mouse brain. Nature 508, 207–214. doi: 10.1038/nature13186

Okuyama, T., Kitamura, T., Roy, D. S., Itohara, S., and Tonegawa, S. (2016). Ventral CA1 neurons store social memory. Science. 353, 1536–1541. doi: 10.1126/science.aaf7003

Osakada, F., and Callaway, E. M. (2013). Design and generation of recombinant rabies virus vectors. Nat. Protoc. 8, 1583–1601. doi: 10.1038/nprot.2013.094

Osakada, F., Mori, T., Cetin, A. H., Marshel, J. H., Virgen, B., and Callaway, E. M. (2011). New rabies virus variants for monitoring and manipulating activity and gene expression in defined neural circuits. Neuron 71, 617–631. doi: 10.1016/j.neuron.2011.07.005

Padmanabhan, K., Eddy, W. F., and Crowley, J. C. (2010). A novel algorithm for optimal image thresholding of biological data. J. Neurosci. Methods 193, 380–384. doi: 10.1016/j.jneumeth.2010.08.031

Padmanabhan, K., Osakada, F., Tarabrina, A., Kizer, E., Callaway, E. M., Gage, F. H., et al. (2016). Diverse representations of olfactory bulb feedback control of olfactory bulb circuits. Neuron 76, 1161–1174. doi: 10.1016/j.neuron.2012.10.020

Acknowledgments

We wish to thank J. Fitzpatrick and J. Kasuboski for assistance with imaging. We thank members of the EC, FG, and TS lab for providing helpful comments on this manuscript, and M. L. Gage and S. Skinner for feedback on the manuscript.
information in centrifugal feedback projections. *J. Neurosci.* 36, 7535–7545. doi: 10.1523/JNEUROSCI.3358-15.2016

Padmanabhan, K., and Urban, N. N. (2010). Intrinsic biophysical diversity decorrelates neuronal firing while increasing information content. *Nat. Neurosci.* 13, 1276–1282. doi: 10.1038/nn.2630

Paxinos, G., and Franklin, K. (2004). *The Mouse Brain in Stereotaxic Coordinates, 2nd ed.* San Diego, CA: Elsevier Academic Press.

Petzold, G. C., Hagiwara, A., and Murthy, V. N. (2009). Serotonergic modulation of odor input to the mammalian olfactory bulb. *Nat. Neurosci.* 12, 784–791. doi: 10.1038/nn.2335

Price, J. L., and Powell, T. P. (1970). An experimental study of the origin and the course of the centrifugal fibres to the olfactory bulb in the rat. *J. Anat.* 107, 215–237.

Restrepo, D., Doucette, W., Whitesell, J. D., McTavish, T. S., and Salcedo, E. (2009). From the top down: flexible reading of a fragmented odor map. *Trends Neurosci.* 32, 525–531. doi: 10.1016/j.tins.2009.06.001

Rothermel, M., Carey, R. M., Puche, A., Shipley, M. T., and Wachowiak, M. (2014). Cholinergic inputs from Basal forebrain add an excitatory bias to odor coding in the olfactory bulb. *J. Neurosci.* 34, 4654–4664. doi: 10.1523/JNEUROSCI.5026-13.2014

Secundo, L., Snitz, K., and Sobel, N. (2014). The perceptual logic of smell. *Curr. Opin. Neurobiol.* 25, 107–115. doi: 10.1016/j.conb.2013.12.010

Shipley, M. T., and Adamek, G. D. (1984). The connections of the mouse olfactory bulb: a study using orthograde and retrograde transport of wheat germ agglutinin conjugated to horseradish peroxidase. *Brain Res. Bull.* 12, 669–688.

Stettler, D. D., and Axel, R. (2009). Representations of odor in the piriform cortex. *Neuron* 63, 854–864. doi: 10.1016/j.neuron.2009.09.005

Van Groen, T., and Wyss, J. M. (1990). Extrinsic projections from area CA1 of the rat hippocampus: olfactory, cortical, subcortical, and bilateral hippocampal formation projections. *J. Comp. Neurol.* 302, 515–528. doi: 10.1002/cne.903020308

van Strien, N. M., Cappaert, N. L., and Witter, M. P. (2009). The anatomy of memory: an interactive overview of the parahippocampal-hippocampal network. *Nat. Rev. Neurosci.* 10, 272–282. doi: 10.1038/nrn2614

Wesson, D. W., Carey, R. M., Verhagen, J. V., and Wachowiak, M. (2008). Rapid encoding and perception of novel odors in the rat. *PLoS Biol.* 6:e82. doi: 10.1371/journal.pbio.0060082

Wickersham, I. R., Finke, S., Conzelmann, K.-K., and Callaway, E. M. (2007). Retrograde neuronal tracing with a deletion-mutant rabies virus. *Nat. Methods* 4, 47–49. doi: 10.1038/nmeth999

Wilson, D. A., and Stevenson, R. J. (2003). The fundamental role of memory in olfactory perception. *Trends Neurosci.* 26, 243–247. doi: 10.1016/S0166-2236(03)00076-6

Yan, Z., Tan, J., Qin, C., Lu, Y., Ding, C., and Luo, M. (2008). Precise circuitry links bilaterally symmetric olfactory maps. *Neuron* 58, 613–624. doi: 10.1016/j.neuron.2008.03.012

**Conflict of Interest Statement:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.