**Systems/Circuits**

**Generation and Characterization of a Cell Type-Specific, Inducible Cre-Driver Line to Study Olfactory Processing**

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In sensory systems of the brain, mechanisms exist to extract distinct features from stimuli to generate a variety of behavioral repertoires. These often correspond to different cell types at various stages in sensory processing. In the mammalian olfactory system, complex information processing starts in the olfactory bulb, whose output is conveyed by mitral cells (MCs) and tufted cells (TCs). Despite many differences between them, and despite the crucial position they occupy in the information hierarchy, Cre-driver lines that distinguish them do not yet exist. Here, we sought to identify genes that are differentially expressed between MCs and TCs of the mouse, with an ultimate goal to generate a cell type-specific Cre-driver line, starting from a transcriptome analysis using a large and publicly available single-cell RNA-seq dataset (Zeisel et al., 2018). Many genes were differentially expressed, but only a few showed consistent expressions in MCs and at the specificity required. After further validating these putative markers using ISH, two genes (i.e., Pkib and Lbdh2) remained as promising candidates. Using CRISPR/Cas9-mediated gene editing, we generated Cre-driver lines and analyzed the resulting recombination patterns. This indicated that our new inducible Cre-driver line, Lbdh2-CreERT2, can be used to genetically label MCs in a tamoxifen dose-dependent manner, both in male and female mice, as assessed by soma locations, projection patterns, and sensory-evoked responses in vivo. Hence, this is a promising tool for investigating cell type-specific contributions to olfactory processing and demonstrates the power of publicly accessible data in accelerating science.

**Key words:** CRISPR; olfaction; scRNA-seq

**Significance Statement**

In the brain, distinct cell types play unique roles. It is therefore important to have tools for studying unique cell types specifically. For the sense of smell in mammals, information is processed first by circuits of the olfactory bulb, where two types of cells, mitral cells and tufted cells, output different information. We generated a transgenic mouse line that enables mitral cells to be specifically labeled or manipulated. This was achieved by looking for genes that are specific to mitral cells using a large and public gene expression dataset, and creating a transgenic mouse using the gene editing technique, CRISPR/Cas9. This will allow scientists to better investigate parallel information processing underlying the sense of smell.

**Introduction**

The complexity of the brain, in part, originates from the diversity of its components, the rich variety of cells. This diversity is evident in morphology, connectivity, molecular expression profiles, and biophysical properties (Sanes and Masland, 2015; Zeng and Sanes, 2017; Luo et al., 2018), which together give rise to what we refer to as cell types. Because the differences are thought to reflect distinct computational tasks or functions (Masland, 2004; Luo et al., 2018), the ability to selectively identify, and to manipulate, each cell type experimentally is key to understanding how the brain works.

In rodents, complex, synaptic processing of olfactory information in the brain first occurs in the olfactory bulb (OB). The principal cells of the OB, the mitral cells (MCs) and tufted cells (TCs), convey the output of this region and are thought to form parallel information streams. They differ in a variety of anatomic...
and physiological properties (Fukunaga et al., 2012; Igarashi et al., 2012; Phillips et al., 2012; Otazu et al., 2015; Economou et al., 2016; Kapoor et al., 2016; Jordan et al., 2018). MCs, which are the larger of the two, are thought to form distinct circuits with local neurons from those formed by TCs (Mori et al., 1983; Fukunaga et al., 2012; Phillips et al., 2012; Geramita et al., 2016), some of which may explain the differences in how they encode odors. For example, in TCs, the timing of responses adheres strictly to a specific phase of the sniff cycle, whereas MCs modulate the timing widely over the entire sniff cycle (Fukunaga et al., 2012; Igarashi et al., 2012; Ackels et al., 2020). Signal integration over this long temporal window is thought to allow MCs to represent more complex information (Fukunaga et al., 2012). Further, in contrast to TCs whose axons project to a more limited portion of the olfactory lobe, the target areas of MCs range more widely, extending as far as the posterior piriform cortex, the cortical amygdala, and the lateral entorhinal cortex (Haberly and Price, 1977; Igarashi et al., 2012), indicative of a variety of behavioral contexts in which MCs are likely to be important.

Despite the fundamental roles these two cell types play in olfaction, suitable molecular markers and genetic tools are lacking. Molecules commonly used to label the output neurons of the OB include protocadherin-21 (Nagai et al., 2005), T-box transcription factor 21 (Tbx21, also known as Tbet) (Papaioannou and Silver, 1998; Faedo et al., 2002), as well as cholecystokinin (Cck) (Seroogy et al., 1985). In the brain, Tbx21 is expressed from embryonic day 14 (Faedo et al., 2002) and is exclusive to the principal neurons of the OB, labeling both MCs and TCs (Faedo et al., 2002; Mitsui et al., 2011; Haddad et al., 2013). In contrast, Cck is expressed widely in the brain (Larsson and Rehfeld, 1979; Taniguchi et al., 2011). In the OB, expression occurs preferentially in TCs over MCs (Seroogy et al., 1985), which has been used for analyzing the unique physiology of TCs (Economou et al., 2016; Short and Wachowiak, 2019). These overlapping but differential expression patterns between Cck and Tbx21 may be useful in discovering more selective markers to distinguish the two types of principal neurons.

A variety of methods now exist to analyze gene expression patterns in relation to cell types, including In-situ hybridisation (ISH) (Lein et al., 2007), as well as transcriptomic approaches that retain spatial information, with increasing resolution (Stahl et al., 2016). More recently, single-cell RNA sequencing (scRNA-seq) (Sugino et al., 2006; Tang et al., 2009; Pfeffer et al., 2013; Zeisel et al., 2015; Shekhar et al., 2016; Tasic et al., 2016) has seen rapid developments, which have enabled the investigation of cell type-specific gene expression patterns with unprecedented levels of detail and scale (Lein et al., 2017; Zeng and Sanes, 2017). A useful application of this information in turn may be to generate transgenic driver lines that allow a particular cell type to be extensively studied. The availability of Cre-driver lines has been instrumental in revealing unique functions of distinct cell types, across multiple levels of analyses (Gong et al., 2003; Taniguchi et al., 2011; Madsen et al., 2012, 2015; Dhande et al., 2013; Cruz-Martín et al., 2014; Wolff et al., 2014; Sanes and Masland, 2015; Daigle et al., 2018).

Here, we take advantage of a large dataset that has become publicly available (Zeisel et al., 2018), to discover markers that distinguish between MCs from TCs. The results of the analyses allowed us to generate, and characterize, new Cre-driver lines. Such molecular tools will be key to understanding the mechanisms of olfactory perception and behavior.

Materials and Methods

Gene expression data
scRNA-seq data from the mouse brain were obtained from Zeisel et al. (2018) in Loom format. We used the dataset from the level 2 analysis that corresponds to the olfactory neurons. The gene expression table represents the expression levels of 27,998 genes in 10,745 OB cells. The gene expression level, which counts the number expressed, was transformed into log2(count + 1) before analyzing further.

Dimensionality reduction. We screened for genes that have higher variability than expected by calculating a log-transformed Fano factor for each gene, as previously described (Li et al., 2017):

\[
F(x) = \log \left( \frac{\sigma^2(x)}{\mu(x)} \right),
\]

where \(\mu(x)\) and \(\sigma^2(x)\) are the mean and the variance of the expression level across cells, respectively. Then using the mean expression across different cells, we split the genes into 20 subsets and calculate the Z score of the Fano factor within each subset as follows:

\[
Z(x) = \frac{F(x) - \text{mean}(F(x))}{\text{std}(F(x))},
\]

where mean(F(x)) and std(F(x)) are the mean and the SD of F(x) within the subset. The top 500 genes with the highest value of Z(x) were used to cluster the gene expression data. To visualize and cluster the gene expression data corresponding to individual cells in 2D space, we reduce the dimensionality using the principal component analysis (PCA) and t-distributed Stochastic Neighbor Embedding (tSNE) (van der Maaten and Hinton, 2008). We used top 10 principal components to run tSNE with the following parameters: learning rate = 10, perplexity = 33. The result is shown in Figure 1. Since many genes have similar expression patterns across different cells, to increase the power of PCA and tSNE, we extracted overdispersed genes (i.e., the most informative genes).

Clustering. Using the two-dimensional data above, hierarchical clustering algorithm HDBSCAN (Campello et al., 2013) was performed with the following parameters: min_clust_size = 5, minPts = 13, which indicates which nearest neighbor to use for calculating the core distances of each point. The cluster with the highest expression level of Tbx21 gene contained 101 cells. Within this group, clustering with HDBSCAN on tSNE space revealed four clusters. We then compared the Cck expression level across the clusters. We found that 18% (2 of 11) of cells from Cluster 4 express Cck above the threshold = 3 (log2(count + 1)), while 64% (58 of 90) from Clusters 1, 2, and 3 express Cck above the threshold; thus, we defined the Cluster 4 to be the putative MC cluster.

Differential expression analysis and identification of molecular markers. We used the Mann–Whitney U test to find differently expressed genes. The test works under the assumption that the samples are independent. p values were adjusted using the Benjamini–Hochberg procedure. We screened for significant genes, with the adjusted p value below the threshold = 0.05, where its median expression level in the MC cluster above the threshold = 3 in >50% of the cells. Of these, genes that are highly expressed in non-MC clusters were eliminated (cutoff of 10% of cells).

Animals
All animal experiments were approved by the animal experiment ethics committee of Okinawa Institute of Science and Technology Graduate University (protocol: 2018-201) and University of Tsukuba. ICR and C56BL/6J mice were purchased from Laboratories International for generation of transgenic mice, and C56BL/6J from Japan CLEA for subsequent breeding. Ai14 (Madsen et al., 2010) were from The Jackson Laboratory, and Ra13-Cre was from the GENSAT project (Gong et al., 2003), via MMRRC (MBP, University of California, Davis). Mice of either sex were used in this study.

Generation of Pkib-IRES-cre and Lbdh2-IRES-CreERT2 mice. Vector construction for knock-in mouse production was as follows: The CRISPR target sequence (5 ’ ATAGAAGCTATGGTATTCTGAGGG-3 ’) was selected for integration of the IRES-Cre sequence just after the stop codon of Pkib and Lbdh2. The pX330 plasmid, carrying both gRNA and
Cas9 expression units, was a gift from Feng Zhang (Addgene plasmid 42230) (Cong et al., 2013). The oligo DNAs (Pkiib-CRISPR F: 5’-caccATAGCAGCTATGTATTCCTG-3’ and Pkiib-CRISPR R: 5’-aacaCAGGAATACATAGCTGCTAT-3’) were annealed and inserted into the entry site of pX330 as described previously (Mizuno et al., 2014). This plasmid was designated as pX330- Pkiib. The donor plasmid pIRES-Cre-Pkiib contained the IRES sequence (Bochkov and Palmenberg, 2006), nuclear translocation signal-Cre, and rabbit globin polyadenylation signal sequence. The 1.6 kb 5’-arm (from 1521 bp upstream to 64 bp downstream of Pkiib stop codon) and the 2.0 kb 3’-arm (from 65 bp downstream to 2038 bp downstream of Pkiib stop codon) were cloned into this vector. DNA vectors (pX330- Pkiib and pIRES-Cre-Pkiib) were isolated with a FastGene Plasmid mini Kit (Nippon Genetics) and filtrated by MILLEX-GV 0.22 µm Filter unit (Merck Millipore) for microinjection. Mice were kept in IVC cages under specific pathogen-free conditions in a room maintained at 23.5 ± 2.5°C and 52.5 ± 12.5% relative humidity under a 14 h light:10 h dark cycle. Mice had free access to commercial chow (MF diet; Oriental Yeast) and filtered water.

Microinjection and genomic DNA analyses. The pregnant mare serum gonadotropin (5 units) and the human chorionic gonadotropin (5 units) were intraperitoneally injected into female C57BL/6J mice with a 48 h interval and mated with male C57BL/6J mice. We collected zygotes from oviducts in mated females, and a mixture of the pX330- Pkiib (circular, 5 ng/µl) and pIRES-Cre-Pkiib (circular, 10 ng/µl) plasmids was microinjected into 148 zygotes. Subsequently, surviving 137 injected

Figure 1. Strategy for identifying mitral-cell specific markers from scRNA-seq data. A, Schematic showing major anatomic differences between the two cell types; MCs (red) are located deeper in the OB layers and project widely in olfactory cortices. TCs (green) are smaller, superficially located principal neurons that project to anterior portions of the olfactory cortex. B, Tbx21 and Cck expression patterns in the main OB; example images showing tdTomato expression patterns in Tbx21-Cre::Ai14 mouse (red) and Cck-IRES-Cre::Ai14 mouse (green). Scale bar, 100 µm. C, Soma positions of tdTomato-expressing cells relative to the EPL boundaries, for the images shown in B. EPL depth was normalized so that it ranged from 0 to 1, with the lower boundary (MCL) corresponding to 0. D, Schematic of workflow; putative mitral cluster from scRNA-seq data is identified by the observation that MCs and TCs both express Tbx21, but Cck is more abundant among TCs. Once putative MC and TC clusters were identified, differential expression analysis was conducted to identify genes that are selectively expressed in MCs. E, OB cells plotted in tSNE coordinates, with Tbx21 and Cck expression levels (left and right, respectively) indicated with color maps shown below. F, Expression levels of common markers for projection neurons of the OB; VGlut1 (Slc17a7) and Cdhr1. G, Tbx21-positive cluster was further analyzed and the subclustered and displayed in new tSNE coordinates. H, Cck expression levels for the subclusters in E. Cluster 2 has the lowest level and is inferred to be the putative MC cluster (red dotted line in E). Statistical significance: *p = 0.05; **p = 0.01. For details, see Experimental design and statistical analysis.
zygotes were transferred into oviducts in pseudopregnant ICR females and 21 pups were obtained.

To confirm the knock-in mutation, the genomic DNA was purified from the tail samples using the PI-200 DNA extraction kit (Kurabo Industries) according to manufacturer’s protocol. Genomic PCR was performed with KOD-Ex (ToyoBo). The primers (Cre forward: 5'-CTCTAGACTACGCTGAAAATGCTTCTGT-3', and Cre reverse: 5'-GTACACCAAGGCTCAAGCACAACCTTACC-3') were used for checking the 5' side correct knock-in, and the primers (Pikb forward: 5'-CTATTTCACAGGTCCAGTTGCTGAAACC-3', and Amp reverse: 5'-ACAGAAAGCTTTTCAGATGATCAGA-3') were used for checking the 3' side correct knock-in. We found that 5 of 21 founders carried the designed knock-in mutation. In addition, we checked random integration of pCX30-Pikb and pLRES-Cre-Pikb by PCR with ampicillin resistance gene detecting primer (Amp detection forward: 5'-TGG CGGGAAGCTAGAGTAA-3', and Amp detection reverse: 5'-TTTGC CTTCCTGTTTTGCT-3') and no founder carried the random integration allele.

**Histology**

ISH. ISH was performed using the RNAscope ISH system (ACDBio) (Wang et al., 2012).

**Brain extraction.** Whole brains were extracted and immediately placed in 4% PFA, dissolved in PB (225.7 mM NaH2PO4; 77.4 mM Na2HPO4; pH 7.4) at 4°C for 24 h. Subsequently, the tissues were sunk in 99.7% ethanol and mixed using a Vortex mixer to allow the tissues to be completely dissolved in 4% PFA. The brain extractions were then fixed in 100% ethanol, and sections were differentiated by quick submersion in 0.2% ammonium hydroxide in distilled water, followed by rinsing in 2 changes of distilled water.

**Probe design.** ISH was conducted using RNAscope (ACDBio), and probes were prepared by ACDBio to be compatible for the procedure. Sequence regions for the Pikb and Ldhld2 probes were selected using the NCBI genetic database. For both probes, regions that were common to all splice variants of each gene were selected. The Pikb probe targeted the region 141-973 bp of the transcript XM_006512605.3. The Ldhld2 probe targeted the region 138-715 bp of the transcript XM_006516048.1. The Tbx21 probe, which targets the region 893-2093 bp of the transcript NM_019507.2, was already commercially available (403331, ACDBio).

**Hybridization.** On the day of ISH, coronal OB sections (20 μm) were cut on a cryostat (Leica CM3050S, Leica Biosystems) at −20°C, washed in RNase-free PBS (Corning), and immediately mounted on glass slides (Superfrost plus, Thermo Fisher Scientific). Slides were dried for 30 min at 60°C and postfixed for 15 min in 4% PFA at 4°C. Slides then underwent ISH using RNAscope reagents, according to the manufacturer’s protocols. Unless otherwise stated, all reagents were provided in the RNAscope kit (RNAscope Intro Pack 2.5 HD Reagent Kit Brown-Mm,catalog #322365). All slides were dehydrated through an ethanol series (75%, 90%, 100%, 100%, Sigma Millipore) and endogenous peroxidase activity blocked using provided hydrogen peroxide for 10 min at room temperature. Sections then underwent antigen retrieval by submersion in 0.2% ammonium hydroxide in distilled water, followed by quick submersion in 2 changes of 100% ethanol. Subsequently, slides were submerged into 100% ethanol 5 times and air dried. A barrier using an Immend hydrophobic barrier pen was drawn around the sections and left overnight at room temperature to dry. On the following day, slides were treated with Protease Plus and incubated in an oven (HybeEZ II System, ACDBio) for 30 min, followed by a series of incubations in the same oven with provided solutions (AMP1-AMP6) to amplify probes (AMP1 and AMP3: 30 min at 40°C; AMP2 and AMP4: 15 min at 40°C; AMP5: 30 min at room temperature; AMP6: 15 min at room temperature). After amplification, a DAB reaction was conducted (1:1 mixture of DAB-A and DAB-B solutions, Vector Labs) for 10 min at room temperature. Slides were washed by submersion 5 times in 2 changes of distilled water.

**Counterstaining.** OB sections were immersed in Mayer’s hematoxylin solution (MHS16, Sigma Millipore) for 10 min. Excess stain was washed in distilled water, and sections were differentiated by quick submersion in 0.2% ammonium hydroxide in distilled water, followed by washing for 5 min in distilled water. Slides were then dehydrated through a series of ethanol for 5 min each, followed by two 5 min immersions in xylene. Slides were then covered with DPX mountant (06522, Sigma Millipore) for histology and left at room temperature to dry before imaging.

**Virus injection**

Three-week-old, heterozygous Ldhld2-IRES-CreERT2 mice were anesthetized with isoflurane (IsoFlo, Zoetis Japan) and placed on a stereotaxic frame (Kopf). Carprofen (Rimadyil, Zoetis; s.c., 5 mg/kg in saline) was injected subcutaneously for anesthesia. The fur was trimmed, and the skin was disinfected with 10% iodine solution before incision. A craniotomy was made bilaterally over center of the dorsal OB (coordinates relative to bregma: AP: 4.8 mm; ML: ± 0.8 mm); 100 nl of AAV1-pcAG-Flex-EGFP-WPRE (Addgene) was injected from a pulled glass capillary tube (tip diameter ~10 μm) at a depth 0.3 mm relative to the brain surface, at a rate of 2 nl every 5 s, using a Nanoject III injector (Drummond Scientific). Following injection, the glass capillary was left in place for 1 min and then slowly withdrawn. The surgical site was then sutured, and mice allowed to recover in a warmed chamber until fully awake, before being returned to their home cage. It is advisable that AAVs for conditional expression are tested before use, as they can exhibit off-target, “leak” expressions depending on the production protocol (Fischer et al., 2019), especially if not diluted enough.

**Tamoxifen administration**

Tamoxifen solution was dissolved at a concentration of 8 mg/ml in a solvent consisting of 5% ethanol and 95% corn oil (23-0320, Sigma Millipore), for once daily injections of 80 mg/kg (10 ml/kg injection volume). Tamoxifen powder (T5648, Sigma Millipore) was initially suspended in 99.7% ethanol and mixed using a Vortex mixer to allow partial dissolution. Corn oil was subsequently added to make up solution to the final volume, and the solution was heated up to 60°C with agitation on an orbital mixer in an oven, with periodic mixing on the Vortex mixer. When fully dissolved (~30 min), the solution was cooled to room temperature, and mice were injected intraperitoneally using a 26G needle with care taken to avoid bubble formations. Injected mice were housed separately from untreated littermates. The mouse weights were monitored carefully throughout the injection period as well as 3 d after the final injection to ensure recovery. For a proof-of-principle P7 injection, 1 injection of 80 mg/kg was given using a 30G needle. A single-dose protocol was used to minimize disturbance to the pups and the nursing mother. Gloves were rubbed with the cage bedding before handling, and injected pups were returned to the cage with the mother.

**Two-photon functional imaging**

Cal-520 dextran (MW ~11,000, AAT Bioquest) was dissolved to 50 mg/ml in Ringer’s solution comprising the following (in mM): NaCl (135), KCl (5.4), HEPES (5), MgCl2 (1), CaCl2 (1.8). Cal-520 dextran solution was electroporated in the glomerular layer (GL) of the left OB of P24 Ldhld2-CreERT2::Ai14 mice (tamoxifen dose = 3 × 80 mg/kg starting at P21), at a depth ~100 μm below the brain surface, under isoflurane anesthesia. Parameters of electroporation were set according to the low-intensity protocol described by Howis et al. (2010). Immediately after the electroporation, the craniootomy was sealed with an imaging window, and mice were anesthetized with ketamine/xylazine (100 mg/kg−1/20 mg/kg−1, i.p.) and two-photon imaging of dye-loaded TCs and MCs was obtained with a custom two-photon microscope (iNS5) using 980 nm high-power laser ( Insight Deepsee, SpectraPhysics) fitted with a water-immersion 25× objective (CFI75 Apo 25XC W 1300, Nikon) and resonance scanner (30 Hz frame rate; FOV was 256 × 256 μm, 512 × 512 pixels). MCs were those located ~300 μm below the brain surface (labeled represents red fluorescent cells + green fluorescence; unlabeled represents loaded cells without red fluorescence), while TCs were smaller cells located more superficially. Strongly fluorescent cells were excluded from analysis. Five odors were presented in a randomized order using a custom-made, flow-dilution olfactometer (Koldaeva et al., 2019), at 5% of the saturated vapor, while the total flow rate was 2 L/min. Odors used were ethyl butyrate, methyl tiglate, methyl butyrate, acetophenone, and methyl salicylate. Intertrial interval was 30 s during which lines were purged with pressurized air to minimize cross contamination. Because
of bleaching and other time-dependent factors, such as the depth of anesthesia, typically, 3 or 4 presentations were given for each odor. For GCaMP6f imaging, Lbhd2-CreERT2::Ai95D mice were injected with tamoxifen intraperitoneally at P21 (1 × 160 mg/kg). After 2 weeks, the mice were surgically implanted with a cranial window over the left OB, as well as a head plate, and allowed to recover. After 2 further weeks, they were anesthetized with ketamine/xylazine, and head-fixed for imaging. Odors were presented in the manner described above for the electrophoration experiment, but 6 odors were presented with the order randomly permuted. The additional odor was butyl acetate. The body temperatures of the mice were maintained at 36°C using a thermostat.

**Confocal imaging**

Confocal images were acquired on a Zeiss LSM780 confocal microscope with a 10× objective (Carl Zeiss, NA 0.45 Plan-Apochromat) for the whole-brain sagittal sections, and 20× objective (Carl Zeiss, NA 0.8 Plan-Apochromat) for the OB. Using ZEN 2.3 software (Carl Zeiss), images were taken at a resolution of 1024 × 1024 pixels for an FOV of 850.19 μm × 850.19 μm (10×) or 425.1 μm × 425.1 μm (20× objective). To enable comparison and quantification of viral injections, imaging conditions (resolution, gain, laser power, number of averages) were kept consistent. Sequential laser excitation was used to prevent fluorophore bleed-through. Images were taken throughout the whole rostro-caudal extent of viral spread using the 20× objective. For axon projection analysis, images were acquired using a Leica SP8 confocal microscope using a 10× (Leica Biosystems, NA 0.40 Plan-Apochromat) and a 40× (Leica Biosystems, NA 1.3 Plan-Apochromat) objective. Images were taken at a resolution of 1024 × 1024 pixels per FOV (10×: 1163.64 × 1163.64 μm; 40×: 290.91 × 290.91 μm) at sequential excitation to prevent fluorophore bleed-through.

**Image analysis**

**ISH signal.** Images of DAB- and hematoxylin-stained OB sections were obtained using a wide-field microscope with a 10× objective in RGB, so that the hematoxylin signal could be separated into the blue channel. The same acquisition settings were used for all sections (Tbx21, Lbhd2, and Pkib signals). Dorsal, ventral, medial, and lateral portions of the OB at three anterior-posterior locations were imaged so that all planes were sampled at anterior, dorsal, and ventral locations of the mid-sagittal plane for the tdTomato signal using the Ai14 reporter line, using the DAPI channel to guide sampling, and 10 consecutive planes at a 100 μm interval for the virus injection experiment. Using only the red and green channel for tdTomato labeling and EGFP labeling, respectively, somata were detected manually in ImageJ using the ROI manager and their coordinates exported into MATLAB, without the observer knowing the identity of the mouse. EPL boundaries were demarcated using only the DAPI signals from images, using a custom-written MATLAB routine and the boundary coordinates were stored. The soma depths from above were normalized along the EPL using the boundary coordinates, such that the MCL was defined as 0, and the lower boundary of the GL as 1. One-way ANOVA was used to compare the means, using the anova1 function in MATLAB, and the multcompare function with the crucial value tested with Tukey’s honest significant difference criterion for post hoc multiple comparisons. Cells belonging to MCL were defined as those whose somata are positioned within 30% of the normalized EPL boundary from the MCL. This corresponded, on average, to 43.6 μm, which is equivalent to the lengths of two MC somata (Nagayama et al., 2010). Thus, our measure takes into consideration the displaced MCs.

**Dendrite detection and quantification.** Images used were the same as those used to detect somata above. To emphasize signals originating from dendrites, which are thin processes, background signal was subtracted from the green or red channel using Subtract Background function in ImageJ, with the rolling ball radius set to 5 pixels. Binary masks were created with a single threshold value, and the presence of the signal along the normalized EPL depth at each lateral position was averaged to obtain the density. The dendritic preference index was used to compare the dendritic signal in the upper EPL versus lower EPL, as a proportion of the total dendritic signal detected, calculated as follows:

\[
\text{Signal density}_{\text{upper EPL}} - \text{Signal density}_{\text{lower EPL}} / \left( \text{Signal density}_{\text{upper EPL}} + \text{Signal density}_{\text{lower EPL}} \right)
\]

**Analysis of labeled MCs on a standardized coordinate.** Labeled MCs from coronal sections (1024 × 1024 pixels, 1.2 μm per pixel) were automatically detected in ImageJ by converting the red fluorescence image into binary masks by thresholding and converted into ROIs using the Analyze Particles function (100-600 pixels, circularity 0.1-1). The MCL was delineated using the DAPI channel in MATLAB using the drawpolygon function. The line was interpolated, and labeled MCs were projected on the MCL coordinates. The center of the OB was calculated as the center of the MCL coordinates. To pool data across mice, MCL coordinates were standardized such that it ran from 0 to 2 π radians relative to the center of the OB.

**Whole-brain somata detection.** Positions of somata labeled with tdTomato were automatically detected in the red channel of the stitched confocal images. To automatically detect the labeled somata, background fluorescence was subtracted using ImageJ’s Subtract Background function (100 pixels), then further sharpened to accentuate the somata locally using ImageJ’s Unsharp filter with the radius set to 14 pixels, and mask weight set to 0.6. Then a binary mask was obtained by setting a threshold...
Figure 2. Differential gene expression analysis reveals candidate marker genes for MCs. A, Cluster grouping used for differential gene expression analysis to distinguish TCs from MCs. Tbx21-expressing cells (left) constitute the MC cluster (red) and TC1 cluster (green), which is identical to the combined Clusters 1 and 3 shown in Figure 1G. The OB-wide dataset (right) contains the TC cluster (green), which is equivalent to the Cck-rich clusters shown in Figure 1E without the Tbx21-rich clusters. B, Genes that are significantly enriched in MCs (red) and TC clusters (green). The size of data points indicates the consistency of expression, measured as the fraction of cells in the cluster that express the gene. Mean expression level (log2(count + 1)) is color-coded as shown in the color map above. C, Expression levels of two candidate genes, Pkib and Lbhd2, with the corresponding color maps, superimposed on the three subclusters of the Tbx21-rich cluster (same tSNE coordinate as Fig. 1G), and (D) the whole OB data. Red arrow points to the Tbx21 cluster. E, Example ISH signals revealed by DAB staining for Tbx21 (left), Lbhd2 (middle), and Pkib (right) for the MOB layers indicated. Scale bar, 50 μm. F, ISH signal density relative to the EPL boundary (0-1); hybridization signal was thresholded, and the proportion of pixels above the threshold for each normalized EPL depth was expressed as density. G, Summary of hybridization signals in the superficial locations (depth upper half of EPL). N = 3 mice, with samples from dorsal, ventral, medial, and lateral locations at middle and caudal levels of the antero-caudal axis. H, Quantification of regional variation; average hybridization signal density from the MCL (right) for dorsal, ventral, medial, and lateral samples taken from middle plane (bottom plot) and caudal plane (top plot) of the AP axis. Orange lines indicate data from individual mice. Black lines indicate the average across the 3 mice.
and the Analyze Particles function was used to detect round objects (size = 70–600 pixels, circularity 0.1–1), and detected structures added to the ROI manager, and exported as a list. Using the DAPI signals in the blue channel, boundaries of each nucleus were manually drawn in MATLAB using the drawpolygon function. Finally, for each anatomic region, all detected somas positions within the boundary were counted using the inROI function and normalized by the area to standardize the density of detected cells per mm². Distributions of labeled somata across strains were tested with two-way ANOVA using MATLAB’s anovan function.

**Experimental design and statistical analysis**

The Mann–Whitney U test, t test, Kolmogorov–Smirnov test, and one-way- and two-way-ANOVA were conducted using MATLAB. Unless otherwise stated, t tests were performed unpaired. Paired tests are described as “two-sample t test” in the text. For *post hoc* comparisons following significant ANOVA tests, the p values are given in Table 1 for brevity of figure legends. Only the significant comparisons are listed because of the large number of pairwise comparisons.

Data will be available on request. *Ldbh2-CreERT2* has been donated to The Jackson Laboratory Repository (stock #036054).

**Results**

In search of molecular markers, we sought to compare the gene expression patterns of MCs and TCs. This may reveal candidate markers, which are genes that are selectively enriched in the target cell type of interest, in this case MCs, but not expressed in other cell types. This first requires a method to identify MCs and TCs in a gene expression data, and, second, distinguish their gene expression profiles from each other. Previous studies observed that *Tbx21*, a T-box type transcription factor, labels both MCs and TCs (*Faedo et al., 2002; Mitsui et al., 2011; Haddad et al., 2013*), while the neurotransmitter cholecystokinin (*Cck*) is more abundant in TCs (*Seroogy et al., 1985; Economou et al., 2016*). To verify these distributions in our hands, we crossed *Tbx21-Cre* and *Cck-IRES-Cre* lines (*Taniguchi et al., 2011; Haddad et al., 2013*) with the Rosa-CAG-LSL-tdTomato reporter line, *Ai14* (*Madsen et al., 2012*), for Cre-dependent expression of the red fluorescent protein, tdTomato. We confirm that *Tbx21*-driven expression labels cells in the MCL and the EPL where TCs are located, while *Cck*-driven expression labels a larger number of cells all over the OB (Fig. 1A–C), especially those that extend more superficially in the GL and sporadically in the granule cell layer. Importantly, labeling coupled to *Cck* expression is less consistent in cells that occupy the MCL. These differential expression patterns between *Tbx21* and *Cck* may be used to distinguish MCs from TCs in gene expression data (Fig. 1B,C).

Identification of molecular markers by differential expression analyses requires a robust and large dataset, especially when distinguishing similar cell types, such as in the case of MCs and TCs. We turned to a public, large-scale scRNA-seq dataset of the mouse brain (*Zeisel et al., 2018*). This contains data from ~0.5 million cells, 10,745 cells of which are from the OB. We clustered the data based on the similarity of gene expression patterns. To achieve this efficiently, we identified the top 500 overdispersed genes of 27,998 genes in the dataset (see Materials and Methods; Fig. 1D). Such genes are highly informative for determining genetic differences among the cells. Using this reduced dataset, we performed PCA, followed by tSNE on the first 10 principal components to further reduce the dimensionality of the gene expression space to two. The combination of the two algorithms preserves both the global and local structures of the data (*Kobak and Berens, 2019*). To obtain clusters, hierarchical density-based special clustering algorithm (HDBSCAN) (*Campello et al., 2013*)

| Gene name | Mean expression (log2(count + 1)) | Adjusted p |
|-----------|----------------------------------|------------|
| Myh8      | 2.14                             | 1.57E-09   |
| Pckl      | 3.40                             | 7.02E-06   |
| Fayd7     | 3.66                             | 1.41E-05   |
| A23006516Rik (Ldbh2) | 3.77     | 1.55E-05   |
| Ebf1      | 2.55                             | 1.55E-05   |
| Cab2      | 4.60                             | 1.55E-05   |
| Snca      | 5.10                             | 1.55E-05   |
| Ctg1      | 2.53                             | 1.55E-05   |
| Cdx1 1    | 0.42                             | 1.55E-05   |
| Sostdc1   | 0.36                             | 2.07E-05   |
| Tmsb10    | 6.12                             | 3.05E-05   |
| Spp1      | 2.20                             | 3.45E-05   |
| Ntng1     | 4.76                             | 4.25E-05   |
| RP23-407N2,  | 1.81                             | 4.70E-05   |
| 1110008F14Rik | 2.45     | 0.000163822 |
| Tspan17   | 1.99                             | 0.000211554 |
| Uchl1     | 3.24                             | 0.000253744 |
| Shisa3    | 1.98                             | 0.000253744 |
| Gap43     | 3.69                             | 0.000439148 |
| Ppm1j     | 1.85                             | 0.000440468 |
| Rph3a     | 0.74                             | 0.000440468 |
| Crta1c    | 1.97                             | 0.000457781 |
| Tmnc1     | 0.27                             | 0.000585813 |
| Mmp17     | 0.45                             | 0.000707168 |
| Nov       | 0.18                             | 0.001406527 |
| Gm13      | 1.03                             | 0.001599665 |
| Rab15     | 1.62                             | 0.001978521 |
| Gm27199   | 1.11                             | 0.002521565 |
| Vsnt2     | 0.83                             | 0.003524909 |
| Npr1      | 1.30                             | 0.003848843 |
| Doc2g     | 4.52                             | 0.004924184 |
| Nptx4     | 0.78                             | 0.005760704 |
| Kcnq3     | 0.94                             | 0.005928465 |
| Slc1a2    | 1.29                             | 0.007685702 |
| Sncb      | 4.97                             | 0.00781742 |
| Eph4      | 1.37                             | 0.009298688 |
| Aph8      | 1.91                             | 0.009830705 |
| Nrsn1     | 2.65                             | 0.010055383 |
| Diras2    | 1.50                             | 0.01049975 |
| Rgs4      | 0.76                             | 0.010986111 |
| A1413582  | 2.09                             | 0.0124281 |
| Abizbp    | 0.09                             | 0.01393499 |
| Ling01    | 1.67                             | 0.01393499 |
| Tshz2     | 3.22                             | 0.01393499 |
| Mal2      | 0.18                             | 0.01393499 |
| FlkB1b    | 0.00                             | 0.01393499 |
| Prkcb     | 1.23                             | 0.01504482 |
| Cdh4      | 1.02                             | 0.01504482 |
| Meg3      | 7.01                             | 0.01660475 |
| Chma3     | 0.61                             | 0.01660475 |
| Ilftm10   | 0.74                             | 0.01905793 |
| Adk       | 0.34                             | 0.02020242 |
| Stx1a     | 0.89                             | 0.02020242 |
| Igf1bp5   | 2.75                             | 0.02381658 |
| Fam19a1   | 0.09                             | 0.0261028 |
| Grin1os   | 0.09                             | 0.0261028 |
| Cdc20     | 0.09                             | 0.0261028 |
| Gm26803   | 0.09                             | 0.0261028 |
| Resp18    | 1.14                             | 0.0261028 |
| Pantr1    | 1.12                             | 0.02678842 |
| Lmo4      | 0.91                             | 0.02690157 |
| Pwrl1     | 1.07                             | 0.03585761 |
| Kitl      | 0.00                             | 0.0397583 |
Figure 3. OB-wide tSNE data and Allen Brain Atlas ISH data used to screen candidate MC markers that were not analyzed further. Differential expression analysis indicates that *Fxyd7*, *Ebf1*, *Snca*, *Calb2*, and *Myh8* are significantly enriched in MCs relative to TCs. **A**, To screen candidates, the expression pattern (color map) in the whole OB data was analyzed. The same tSNE coordinates as in Figure 1E, F are used, with *Lbhd2* expression pattern shown for comparison. **B**, Further, the ISH database of the Allen Brain Atlas was used to assess the spatial expression patterns. *Fxyd7* seems to be expressed by neurons deep in the granule cell layer as well as superficial cells. *Snca* is expressed by some superficially located neurons as well as some neurons of the anterior olfactory nucleus. *Ebf1* is hardly detectable in the OB, although it is present in the Purkinje cell layer of the cerebellum. Dense *Calb2* hybridization signal is visible in the GL, EPL, MCL, as well as the granule cell layer. *Myh8* signal was not described in the ISH database, but it is a marker for somatostatin-positive cells in the subventricular zone (Lim et al., 2018), which is the source of SST-positive interneurons of olfactory cortices. The expression data in **A** show low levels of *Myh8* expression in many neurons outside of the Vglut1- and 2-positive clusters. Image credit: Allen Institute.
was applied on the two-dimensional tSNE space to cluster the data (see Materials and Methods). Within the OB dataset, we found that 1682 cells belong to \textit{Cck}-positive clusters, while \textit{Tbx21}-expressing cluster comprised 101 cells. Generally, expression patterns of \textit{Cck} and \textit{Tbx21} together mirror those of \textit{Slc17a7} (VGluT1) and \textit{Slc17a6} (VGluT2), indicating that they are mainly glutamatergic populations (Fig. 1E,F), with the largest portion of glutamatergic, \textit{Cck}-positive clusters residing outside of the \textit{Tbx21}-positive cluster. Further, a small set of \textit{Cck}-expressing neurons did not overlap with the \textit{Slc17a7}-positive cluster (Fig. 1E,F). To identify a putative MC cluster from the scRNA-seq data, we took advantage of the observation that MCs and TCs both express \textit{Tbx21}, but \textit{Cck} is more abundant among TCs. In the \textit{Tbx21}-positive cluster, the second largest cluster (Cluster 2; Fig. 1G) showed the lowest \textit{Cck} expression level (Fig. 1H). We thus refer to this as the putative MC cluster, and refer to the remaining as TC Cluster 1 (TC1).

An ideal molecular marker should be expressed abundantly and consistently in the cell type of interest, while having minimal expression levels in other cell types. To search for candidates with these properties, gene expression patterns of putative MCs were compared against the rest of \textit{Tbx21}-expressing neurons (TC1; Fig. 2A), as well as glutamatergic, \textit{Cck}-positive clusters outside of the \textit{Tbx21}-cluster (TC2; Fig. 2A). First, the Mann-
Whitney U test was used to screen genes that are differentially expressed, with \( p \) values adjusted using the Benjamini–Hochberg procedure. This procedure identified several differentially expressed genes (Table 2), at the adjusted \( p = 0.05 \) level. Among these were \textit{Calb2} (calbindin 2), \textit{Ntng1} (netrin G1), \textit{Ppm1j} (protein phosphatase 1J), \textit{Rph3a} (rabphilin 3A), \textit{Kcnq3} (voltage-gated potassium channel subfamily Q member 3), and \textit{Chrna2} (cholinergic receptor nicotinic \( \alpha_2 \)). Of the differentially expressed genes, we focused on those that are present in the majority (>50%) of cells in the putative MC cluster, but in <10% of the cells outside of this cluster (Fig. 2B–H; Table 2). Only a small number of the differentially expressed genes fulfilled these criteria, and even fewer showed minimal expression levels outside of MCs, as judged by the OB-wide expression patterns (Fig. 3A), as well as by the ISH data in the Allen Brain Atlas (Lein et al., 2007). Candidate genes that showed clear hybridization signals outside of the MCL were therefore not pursued further (Fig. 3B).

Based on the initial screening, \textit{Pkib} (protein kinase inhibitor \( \beta \)) and \textit{Lbhd2} (LBH domain containing 2; Fig. 2C,D) genes fit the criteria for an MC marker. To confirm that these genes indeed are selectively expressed in MCs, we conducted ISH for \textit{Pkib} and \textit{Lbhd2} (Fig. 2B–D) on OB sections. Indeed, probes for \textit{Pkib} and \textit{Lbhd2} gave rise to monolayer-like signals at the lower boundary of the EPL, corresponding to the location of the MCL. For quantification, \textit{Pkib} and \textit{Lbhd2} signals were expressed as density (see Materials and Methods) and plotted relative to the boundaries of the EPL. This revealed that \textit{Pkib} and \textit{Lbhd2} both label cells in the MCL, with significant reduction in the superficial signals corresponding to TCs, especially compared with \textit{Tbx21} (Fig. 2D; mean signal densities in the upper EPL: \textit{Tbx21} = 0.39 ± 0.007; \textit{Lbhd2} = 0.005 ± 0.0002; \textit{Pkib} = 0.01 ± 0.001, \( p = 0.0007 \), one-way ANOVA, \( F = 17.9 \), degrees of freedom = 2). Hybridization signal in the MCL was relatively uniform throughout the OB (Fig. 2H), while residual expression patterns of \textit{Pkib} and \textit{Lbhd2} in non-MC cells differed somewhat, with faint signals.
Figure 6. *Pkb-IRES-Cre* labels a wide variety of non-MC neurons, with late onset labeling in MCs. **A**, Strategy for CRISPR/Cas9-mediated generation of the *Pkb-IRES-Cre* transgenic mouse. The CRISPR-target sequence (inset) was just after the stop codon of the *Pkb* gene. Construct included sequences for IRES, Cre-recombinase with a nuclear localization signal, and rGpA. **B**, Cre-mediated recombination pattern in a 30-d-old *Pkb-IRES-Cre::Ai14* mouse. Structures visible in this sagittal plane are revealed by DAPI (top), and the corresponding pattern of recombination revealed by tdTomato signal. Scale bar, 0.5 mm. **C**, tdTomato expression pattern relative to the OB layers from the same animal. ONL, Olfactory nerve layer. Note the dense labeling of the ONL. **D**, A higher magnification of image in **C**. Signals in the MCL are faint or lacking at this developmental stage. Scale bar, 0.1 mm. **E**, A confocal image showing the EGFP expression pattern 2 weeks after an injection of AAV1-flex-EGFP into the MOB in a 42-d-old *Pkb-IRES-Cre* mouse. **F**, Summary of soma location (left) and dendritic signal density (right) relative to the EPL boundaries. Note the heavy presence of somata outside of the MCL and dendrites in the upper portion of the EPL.
Figure 7. *Lbhd2-CreERT2* line achieves MC-specific labeling in the OB even in adulthood. **A**, Constructs for CRISPR/Cas9-mediated knock-in line. IRES-CreERT2 cassette was targeted to a region immediately following the stop codon of the *Lbhd2* gene. **B**, Schematic of the tamoxifen injection protocol: tamoxifen was injected intraperitoneally at P21, and the recombination pattern in the OB was examined 3 weeks later. Cohorts of mice received injection for either 1 or 3 d at 80 mg/kg per day. **C**, Example recombination patterns. From left: *Tbx21-Cre::Ai14*, *Ra13-Cre::Ai14*, and *Lbhd2-CreERT2::Ai14* (one injection) and *Lbhd2-CreERT2::Ai14* (three injections). Scale bar, 100 μm. **D**, Summary of labeled structures at P42 for the corresponding mouse lines, showing the proportion of cells in the MCL relative to all labeled cells (left), comparison of labeled dendrites in the superficial versus deep portions of the EPL (middle), and density of labeled somata in the MCL (right). **E**, Comparison of labeling patterns between *Tbx21-Cre::Ai14*, *Ra13-Cre::Ai14*, and *Lbhd2-CreERT2::Ai14* lines. Left, Labeled cells in the MCL as a percentage of total number of labeled cells. Middle, tdTomato signal density in the upper half of EPL subtracted by the signal density in the lower half of EPL. Right, Number of labeled cells detected per mm of MCL. *N = 3* mice per transgenic line for all plots. Data are mean ± SEM. Statistical significance: *p = 0.05; **p = 0.01. For details, see Experimental design and statistical analysis.
Cre-independent, “leak expression” depending on the production protocol (Fischer et al., 2019) or if not diluted sufficiently. Since the two independent Cre-driver lines (Ra31-Cre vs Ra13-Cre) show similar recombination patterns, we chose to analyze the line Ra13-Cre. As above, we crossed Ra13-Cre mice with Ai14 reporter mice to analyze the pattern of Cre-mediated recombination in the brain (Fig. 5). At postnatal day 7 (P7), red fluorescence was highly selective, showing dense and restricted expression in the cells of the MCL of the OB (Fig. 5A–C; mean number of fluorescent cells as a proportion of fluorescent cells in the MCL = 0.09 ± 0.04 for P7; p = 0.18, t test for mean = 0, t-statistic = 2; n = 3 mice). Correspondingly, labeled dendrites were observed preferentially in the lower portion of the EPL (fluorescence signal density = 0.20 ± 0.03 for lower EPL vs 0.10 ± 0.01 for upper EPL; p = 0.03, two-sample t test for equal means, n = 3 mice each), consistent with MCs having dendrites that ramify in the deeper portion of the EPL. At this developmental stage, red fluorescence was observed only sparsely in the rest of the brain, except for the lateral septum and the dorsomedial nucleus of the hypothalamus. However, in older mice, the residual recombination becomes widespread and is observed throughout the brain. In the OB at this stage, while the labeling is still restricted to the projection neurons, a substantial number of TCs also become labeled (mean number of fluorescent cells in the upper EPL as a proportion of fluorescent cells in the MCL = 1.05 ± 0.08 for P21 and 1.33 ± 0.12 for P42). A Cre-driver line that we generated for the second marker candidate, Pkib, was deemed unsuitable for MC-specific labeling because of late-onset expression in MCs, as well as a widespread recombination in neurons other than MCs (Fig. 6).

The developmental accumulation described above makes Ra13-Cre unsuitable for investigating MCs in adult mice. However, the ISH signal for Lbhd2 mRNA indicates a clear preference for MCs in adulthood. Therefore, it is possible that, when the recombination efficiency is calibrated appropriately, a more selective labeling of MCs may be feasible. To this end, we generated a new knock-in line (Fig. 7A) using CRISPR/Cas9, where the inducible Cre-recombinase, Cre-ERT2 (Feil et al., 1997), is inserted into the 3′-UTR of the Lbhd2 gene (the target sequence: 5′-ACCAAGAGGACCTTCAT-3′; Fig. 7A).

To test whether selective labeling is maintained beyond P7 in the new, inducible Cre-driver line, we injected tamoxifen intraperitoneally at P21 in Lbhd2-CreERT2::Ai14 mice, and analyzed the GL and EPL for Pkib and Lbhd2, respectively. Thus, Pkib and Lbhd2 are promising candidates for selectively labeling MCs. On the other hand, the same analysis failed to reveal clear molecular markers for subclasses of TCs (Fig. 4).

Having identified candidate markers for MCs, we sought to test whether Cre-recombinase expression from these loci would allow MC-specific labeling. Screening several public depositories, we found a Cre-driver line for Lbhd2 under a synonymous gene symbol (A230065H16Rik) on GENSAT, a large repository of BAC-mediated transgenic mouse lines (Heintz, 2004; Gong et al., 2007). Since the two independent Cre-driver lines (Ra31-Cre vs Ra13-Cre) show similar recombination patterns, we chose to analyze the line Ra13-Cre. As above, we crossed Ra13-Cre mice with Ai14 reporter mice to analyze the pattern of Cre-mediated recombination in the brain (Fig. 5). At postnatal day 7 (P7), red fluorescence was highly selective, showing dense and restricted expression in the cells of the MCL of the OB (Fig. 5A–C; mean number of fluorescent cells as a proportion of fluorescent cells in the MCL = 0.09 ± 0.04 for P7; p = 0.18, t test for mean = 0, t-statistic = 2; n = 3 mice). Correspondingly, labeled dendrites were observed preferentially in the lower portion of the EPL (fluorescence signal density = 0.20 ± 0.03 for lower EPL vs 0.10 ± 0.01 for upper EPL; p = 0.03, two-sample t test for equal means, n = 3 mice each), consistent with MCs having dendrites that ramify in the deeper portion of the EPL. At this developmental stage, red fluorescence was observed only sparsely in the rest of the brain, except for the lateral septum and the dorsomedial nucleus of the hypothalamus. However, in older mice, the residual recombination becomes widespread and is observed throughout the brain. In the OB at this stage, while the labeling is still restricted to the projection neurons, a substantial number of TCs also become labeled (mean number of fluorescent cells in the upper EPL as a proportion of fluorescent cells in the MCL = 1.05 ± 0.08 for P21 and 1.33 ± 0.12 for P42). A Cre-driver line that we generated for the second marker candidate, Pkib, was deemed unsuitable for MC-specific labeling because of late-onset expression in MCs, as well as a widespread recombination in neurons other than MCs (Fig. 6).

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the distribution of red fluorescence 3 weeks after injection to assess the recombination pattern (Figs. 7B–E, 8A). At the lowest dose tested (one injection of 80 mg/kg), the labeling was sparse (average density of labeled MCs = 5.7 ± 0.9 cells per mm), but 83.7 ± 3.7% of labeled somata were located in the MCL (Fig. 7C, D). Other labeled cells were mostly TCs, save for sporadic labeling in granule cells, which constituted ~1% of the labeled cells. When the dose was increased to three intraperitoneal injections of tamoxifen (at 80 mg/kg per day, over 3 d), denser labeling was achieved (mean density = 10 ± 4 cells per mm; 78.0 ± 11.0% of labeled somata were in MCL) while maintaining specificity, indicating that the tamoxifen dose can be calibrated to titrate the specificity and density of labeling. Compared with the patterns of recombination observed with existing lines, namely, Tbx21-Cre and Ra13-Cre mice, overall, the new line achieves a labeling that is substantially more selective for MCs, as measured by the positions of somata (p = 0.016, F = 6.01, one-way ANOVA; n = 3 mice for Tbx21-Cre and Ra13-Cre, 4 mice for Lbhd2-CreERT2::Ai14) and dendrites (p = 2.99 × 10⁻⁶, F = 59.31, one-way ANOVA; n = 3 mice for Tbx21-Cre and Ra13-Cre, 4 mice for Lbhd2-CreERT2::Ai14). Consistent with the recombination pattern observed with the Ra13-Cre line, tamoxifen injection at P7 also resulted in an MC-specific labeling (Fig. 8B). To test whether AAV-mediated conditional labeling is possible, AAV1-pCAG-Flex-EGFP-WPRE (100 nl) was stereotactically injected into the dorsal OB at a depth of 300 μm below the brain surface in P21 Lbhd2-CreERT2::Ai14 mice. Tamoxifen injections (3 × 80 mg/kg, i.p.) overlapped such that the first of the three injections occurred immediately after the AAV injection. Three weeks later, labeling pattern was analyzed, which showed predominantly MC-selective labeling similar to the pattern obtained with the Ai14 reporter line (Fig. 8C).

Having achieved MC-selective labeling with the Lbhd2-CreERT2 line, we wished to further characterize the properties of the labeled cells in the OB, as well as the distribution of labeled fibers in the olfactory cortices (Fig. 9). Specifically, we wished to assess whether the labeled MCs are present uniformly in all domains of the OB. To this end, confocal images from coronal sections from anterior, middle, and caudal levels of the OB from Lbhd2-CreERT2::Ai14 mice were analyzed. This revealed consistent MCL labeling in all regions of the OB, except for the most anterior level, which tended to show a sparser labeling on the medial side (Fig. 9B–D), although this was not statistically significant (p = 0.56, two-way ANOVA; n = 3 mice). In terms of the projection patterns of labeled fibers in the olfactory cortices, we detected red fluorescent fibers as fascicles throughout the antero-caudal extent of the lateral olfactory tract (Fig. 9A,B), as well as...
thin fibers with bouton-like structures in the molecular layers of olfactory cortices, including in the anterior olfactory nucleus, olfactory tubercle, and the anterior and posterior piriform cortices (Fig. 9E–G).

In addition to the anatomic traits, to assess the labeled cells functionally, we loaded a synthetic calcium indicator, Cal-520 dextran, by electroporation (Madisen et al., 2015) and the dye-loaded cells showed detectable responses to any of the 6 odors presented (Fig. 10F). This time, a fraction of cell-odor pairs (22.2%; n = 270 cell-odor pairs, 45 cells; 4 mice) exhibited robust fluorescence increases locked to odor presentations (Fig. 10F), although the majority of cells did not show detectable responses to any of the 6 odors presented (Fig. 10G). Overall, the results here indicate that the new inducible Cre-driver line, Lbhd2-CreERT2, achieves a highly specific labeling of functional MCs in the OB.

Finally, we examined the recombination pattern in the brain at large, beyond the OB. To assess this, we analyzed the distribution of labeled somata in the anterior olfactory nucleus, olfactory tubercle, anterior and posterior piriform cortex, and tenia tecta, as well as other, commonly studied regions, including the thalamus, cerebellum, hippocampus, and cerebral cortex. In the Lbhd2-CreERT2 mice, for both doses of tamoxifen tested, consistent labeling was observed unexpectedly in a small number of nuclei, including the ventromedial nucleus of the hypothalamus and the lateral septum (Figs. 11, 12). Compared with the Ra13-Cre driver line, the olfactory cortices were devoid of fluorescent cells (Fig. 11B,C); no labeled cells were detected for aPCx, pPCx, AON, and OT in Lbhd2-CreERT2::Ai14 mice; in Ra13::Ai14 mice, mean density of labeled cells = 91.2, 38.5, 490.2, and 818.0 cells below the brain surface, red fluorescence is pseudo-colored magenta (labeled represents red fluorescent cells + green fluorescence; unlabeled represents loaded cells without red fluorescence), whereas TCs were smaller cells located more superficially. Strongly fluorescent cells were excluded from analysis. Scale bar, 50 µm. C. Normalized fluorescence (ΔF/F) from TCs (left) and labeled and unlabeled MCs (right, top and bottom, respectively), shown as color map (n = 2 mice). Excitatory responses are more prevalent in TCs. Cal-520 has a lower affinity to Ca$^{2+}$ than GCaMP6 variants, which may make hyperpolarizing responses to odors less detectable. D. Cumulative histogram of response amplitude, for TCs (black), labeled MCs (magenta), and unlabeled MCs (gray). Overall distributions are not significantly different (two-sample Kolmogorov–Smirnov test; p = 0.56). E. Top, Experimental configuration. Bottom, Example FOV at a depth ~280 µm below the brain surface. Scale bar, 50 µm. F, Top, Example transients from two ROIs indicated in E. Calibration: vertical, 0.5 ΔF/F; horizontal, 1 s. Dotted line indicates onset of the first inhalation after final valve opening. Bottom, Summary of fluorescence change in response to odor presentations shown with a color map (n = 228 cell-odor pairs, 38 cells, 3 mice). G, Summary statistics of evoked responses. Left, Histogram of mean fluorescence change during odor (1 s from inhalation onset) expressed as Z score for the data in E. Right, Number of odors that each cell responds with a fluorescence increase (Z score >2).

Figure 10. Odor response properties of labeled MCs compared with TCs. A, Schematic showing low-intensity electroporation of Cal-520 dextran solution in the GL. B, MCs were those located ~300 µm below the brain surface, red fluorescence is pseudo-colored magenta (labeled represents red fluorescent cells + green fluorescence; unlabeled represents loaded cells without red fluorescence), whereas TCs were smaller cells located more superficially. Strongly fluorescent cells were excluded from analysis. Scale bar, 50 µm. C, Normalized fluorescence (ΔF/F) from TCs (left) and labeled and unlabeled MCs (right, top and bottom, respectively), shown as color map (n = 2 mice). Excitatory responses are more prevalent in TCs. Cal-520 has a lower affinity to Ca$^{2+}$ than GCaMP6 variants, which may make hyperpolarizing responses to odors less detectable. D, Cumulative histogram of response amplitude, for TCs (black), labeled MCs (magenta), and unlabeled MCs (gray). Overall distributions are not significantly different (two-sample Kolmogorov–Smirnov test; p = 0.56). E, Top, Experimental configuration. Bottom, Example FOV at a depth ~280 µm below the brain surface. Scale bar, 50 µm. F, Top, Example transients from two ROIs indicated in E. Calibration: vertical, 0.5 ΔF/F; horizontal, 1 s. Dotted line indicates onset of the first inhalation after final valve opening. Bottom, Summary of fluorescence change in response to odor presentations shown with a color map (n = 228 cell-odor pairs, 38 cells, 3 mice). G, Summary statistics of evoked responses. Left, Histogram of mean fluorescence change during odor (1 s from inhalation onset) expressed as Z score for the data in E. Right, Number of odors that each cell responds with a fluorescence increase (Z score >2).
from Ra13-Cre; \( n = 3 \) mice per group). Thus, the results indicate that the labeling is, overall, relatively specific to MCs in the whole brain, suggesting that the new inducible Cre-driver line may be suitable for a variety of studies to investigate olfactory processing.

**Discussion**

A wide variety of neuron types that exist in the sensory systems are thought to reflect diverse components for information processing in the brain (Masland, 2004; Luo et al., 2018). Availability of Cre-driver lines have led to a multitude of fundamental insights into unique, cell type-specific contributions to sensory processing and perception (Münch et al., 2009; Dhande et al., 2013; Cruz-Martín et al., 2014; Takahashi et al., 2020). Recent progress in the acquisition, analyses, and applications of large-scale gene expression data have allowed efficient analysis of the differences between cell types of interest (Birnbaum, 2018; Luo et al., 2018). In this study, we used a publicly available gene expression dataset to discover candidate molecular markers for the key second-order cells of the olfactory system, MCs, which we validated with histology, and finally with new Cre-driver lines generated by CRISPR/Cas9-mediated gene editing. We report that one driver line in particular provides a substantial improvement in the ability to selectively label MCs.

Among the several candidates identified from our differential expression analysis, we found Lbhd2 to be the most promising. Specifically, at postnatal day 7, the recombination pattern for the noninducible, Ra13-Cre::Ai14 is restricted mainly to MCs in the OB. This pattern is consistent with the description on the GENSAT expression database (Gong et al., 2007). Since TCs are already present and located superficially in the EPL at this stage of development (Mizuguchi et al., 2012), this pattern likely reflects genuine MC specificity in neonatal mice. Despite an increase in the sporadic Lbhd2-driven labeling of TCs and other regions of the brain at later stages, the preferential expression in MCs over TCs that persists in adulthood can be used to our advantage. Thus, with our new inducible Cre-driver line, the expression can be targeted selectively to MCs even in adulthood. It is notable that, despite the fact that many genes were differentially expressed across the two cell types, markers suitable as genetic tools are harder to identify, especially when selective expression is required across developmental stages. This difficulty may partly be because of the similarity between MCs and TCs, and that cell types are often defined by a combination of genes, rather than single genes (Luo et al., 2018).

In the OB, labeled MCs were present in most domains of the OB, except for a small patch on the medial, anterior OB that showed a curious lack of labeling. Whether or not these correspond to subclasses of MCs, for example, those that differ in the glomerular association (Li et al., 2017) or cortical projection patterns (Zeppilli et al., 2020), will be intriguing for future investigation. Outside of the OB, labeled somata were sparse, if not absent, especially in the areas that MCs target, including in the anterior olfactory nucleus, olfactory tubercle, and the anterior and posterior piriform cortices. This makes the Lbhd2-CreERT2...
line suitable for investigating the downstream, decoding mechanisms of mitral-specific activity in all these areas, and also when imaging from boutons of MCs (Pashkovski et al., 2020). Beyond these areas, however, we observed a small number of specific regions that showed the presence of labeled somata. The areas include the lateral septum, ventromedial nucleus of the hypothalamus, and the medial amygdala, even at the lowest dose of tamoxifen used. Thus, future studies using this line need to take this into account when interpreting data, in particular for investigating innate, social behavior, which involve these areas (Stowers and Liberles, 2016). However, the fact that only a subset of the nuclei in the pathways are labeled may make this line unexpectedly useful for investigating mechanisms of social behavior.

This study was aided by publicly accessible data, speeding up discovery. One limitation, if any, in using this dataset for this study may have been the data size, where only a small fraction of OB cells expressed Tbx21, and even fewer belonged to the putative MC cluster. The relatively small MC cluster size may partly be biological. Of the neurons present in the OB, ~80% are GABAergic. Further, glutamatergic neurons comprise heterogeneous groups, including those that lack lateral dendrites (Hayar et al., 2004; Antal et al., 2006). Thus, MCs comprise only a small proportion (~1%) of OB neurons (Burton, 2017; Schwarz et al., 2018). Our histology indicates that superficially located, Tbx21-expressing cells are located below the GL, unlike the Cck-expressing population that includes a dense population located more superficially. It should also be noted that a large proportion of glutamatergic and Cck-expressing cells were found outside of the Tbx21-positive cluster. Some of this latter group may correspond to external TCs, which are glutamatergic but lack lateral dendrites (Macrides and Schneider, 1982). It is possible that protocols used to obtain the scRNA-seq data may have been inadvertently biased against large cells with prominent dendrites, such as the filtering step involving a pore size of 30 μm (Zeisel et al., 2018).

Despite the need for tamoxifen, this new method for labeling MCs has several advantages over the existing methods. Currently, MC labeling and manipulations are achieved predominantly by depth, birthdate, or retrograde viral expression using the differential projection targets of MCs versus TCs (Haberly and Price, 1977; Imamura et al., 2013; Rothermel et al., 2013; Economo et al., 2016). While this can indeed bias expression patterns, the overlap in somatic and dendritic locations (Schwarz et al., 2018), as well as projection targets (Haberly and Price, 1977; Igarashi et al., 2012), means that it is not trivial to achieve a highly selective labeling. In contrast, our transgenic mouse line described here allows for reproducible and selective labeling of MCs over TCs, with the added advantage that labeled MCs are located throughout the OB. Even in imaging applications that can distinguish the cell type based on the soma depth, with the new driver line, it will be possible to investigate the physiology of subcellular compartments, such as the long lateral dendrites, without the need to painstakingly trace back the structures to soma for cell type identification. Similarly, investigations of downstream decoding mechanisms, such as one involving precise optogenetic activations of OB projections using patterned light stimuli (Chong et al., 2020), may now be done in a cell type-specific manner. Thus, our new tool may bring us closer to

**Figure 12.** Recombination pattern outside of the OB in Lbhd2-CreERT2 mice. A, Confocal image at a sagittal plane ~0.36 mm from the midline in a Lbhd2-CreERT2:Ai14 mouse. Scale bar, 1 mm. B, Coronal view at ~1.46 mm posterior to the bregma, showing (B1) labeled cells in the basolateral amygdaloid nucleus, fiber endings in the molecular layer medial amygdaloid nucleus and posterior piriform cortex; (Bii) densely labeled somata in the ventromedial nucleus of the hypothalamus; and (Biii) labeled fibers in the medial habenular nucleus. C, Summary of labeled structures with respect to distinct pathways. Top, MCs of the main OB are labeled, but not their cortical targets. Middle, Principal neurons of the accessory OB are labeled. Labeled fibers, but not somata, are visible in the medial amygdaloid nucleus. The target of the medial amygdaloid nucleus, namely, the ventromedial nucleus of the hypothalamus, has densely labeled cells. Bottom, Lateral septum densely contains labeled cells; the output fiber tracts are strongly labeled (stria medullaris), and labeled fibers are clearly visible in the target structure, namely, the medial habenular nucleus.
understanding how parallel olfactory processing contributes to mechanisms of sensory perception and, ultimately, behavior.

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