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the map. Although transient ectopic glomeruli may be generated during normal postnatal development, they are pruned in adult animals (30, 31). The pruning process may reflect a refinement process at a later postnatal stage.

Our study puts the olfactory system in line with other sensory systems in that it undergoes a change in circuit plasticity during the critical period of development (32). Although, traditionally, the discussions of critical period have focused on how sensory deprivation affects the development of neural circuits, our study reveals an intrinsic developmental program that unfolds whether or not neural activity is perturbed. Even though the olfactory system is regulated by a critical period, late-generated neurons adopt a different strategy for axon projection. Thus, a developmental critical period may function to restrict the reorganization of the neural circuit and to maintain an established map.

References and Notes


A Critical Period Defined by Axon-Targeting Mechanisms in the Murine Olfactory Bulb

Lulu Tsai* and Gilad Barnea†

The olfactory system remains plastic throughout life because of continuous neurogenesis of sensory neurons in the nose and inhibitory interneurons in the olfactory bulb. Here, we reveal that transgenic expression of an odorant receptor has non–cell autonomous effects on axons expressing this receptor from the endogenous gene. Perinatal expression of transgenic odorant receptor causes rerouting of like axons to new glomeruli, whereas expression after the sensory map is not confined to early development; rather, it is lifelong. In the olfactory system, enhanced plasticity is induced by the continuous recruitment of the inhibitory granule cells and tufted cells, the projection neurons in the olfactory cortex. Thus, an olfactory sensory map is formed in the bulb. In this map, the identity of each odor is encoded by the combination of glomeruli that it activates (3). In contrast to the somatosensory, auditory, and visual maps, neighboring relations between peripheral sensory neurons are not maintained in the olfactory sensory map. Because OSNs continue to integrate into the circuits throughout life, the challenge of axon guidance persists in adulthood (3).

We devised a strategy for ectopic expression of a specific OR, MOR28, in a temporally controlled manner using the tetracycline response element (tetO) to drive its expression. The tetO promoter is activated by the tetracycline-controlled transcription activator tTA, which is inhibited by the antibiotic doxycycline. When doxycycline is removed, expression from the tetO promoter is induced within days (14–16). A similar approach for inducing ectopic expression of ORs was previously used (17–19). Our strategy involved the use of three alleles (fig. S1A). In the first, designated OMP-ires-tTA, the olfactory marker protein (OMP) drives expression of tTA in all OSNs (16). In the second, designated tetO::MOR28-ires-tau-LacZ (TO28), tetO drives the expression of MOR28 and the fusion protein tau–β-galactosidase (β-gal). To distinguish between the OSNs that express MOR28 from its endogenous genomic locus (endogenous MOR28 OSNs) versus OSNs that express MOR28 from the transgene (transgenic MOR28 OSNs), we introduced a third allele, designated MOR28-ires-GFP. OSNs that express MOR28 from this allele also express green fluorescent protein (GFP) (20). Thus, GFP expression marks OSNs expressing MOR28 from its endogenous locus. Because β-gal and GFP are exogenous to mice, staining for each identifies transgenic or endogenous MOR28 OSNs, respectively (fig. S1B).

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This strategy enabled us to induce transgene expression at different developmental points. We generated two founder lines for the TO28 transgene. In the presence of the OMP-IREs-tTA allele, animals from these lines express the transgene only in a small fraction of OSNs, presumably because of position effect variegation. In one of these lines, designated TO28L, the transgene is expressed in ~1% of OSNs, and in the other, designated TO28H, the transgene is expressed in ~5% of OSNs. In both lines, the transgene is expressed throughout the olfactory epithelium, both within and outside the characteristic MOR28 zone of expression (Fig. 1, A and B, and fig. S1B).

Normally, MOR28-expressing OSNs converge on two symmetrical pairs of glomeruli per mouse, one medial and one lateral, all located in the posterior-ventral part of the bulb (13, 16, 20, 21). In both transgenic MOR28 lines, visualization of the projection patterns of transgene-expressing OSNs revealed that they converged on multiple glomeruli that were scattered throughout the bulb. The positions and numbers of these glomeruli varied significantly between animals and also between the two bulbs within the same animal (Fig. 1, C to F). We counted the ectopic glomeruli in sectioned bulbs from both lines and observed 46.8 ± 8.33 β-gal–positive glomeruli per animal (n = 5) in TO28L and 103.25 ± 13.85 glomeruli per animal (n = 4) in TO28H (fig. S1C).

In animals that express transgenic MOR28, the endogenous MOR28 axons innervated more than the regular four glomeruli. These glomeruli were always coinnervated by transgenic MOR28 fibers, and they were confined to areas within 650 μm of the typical locations of the wild-type glomeruli (Fig. 1, G and H). In some cases, only a few transgenic MOR28 fibers innervated a glomerulus mainly innervated by endogenous MOR28 fibers, likely the wild-type MOR28 glomerulus (Fig. 1G). In other cases, the transgenic MOR28 fibers and the endogenous MOR28 axons innervated multiple glomeruli. However, the mixing between the two neuronal populations was extensive, and we could not determine which of the glomeruli was the wild-type MOR28 glomerulus (Fig. 1H). Because the endogenous MOR28 axons do not express transgenic MOR28 (fig. S1B), this phenotype is non–cell autonomous and is likely the result of homotypic attraction between the transgenic and endogenous MOR28 fibers (19, 22). We refer to the glomeruli that contain both endogenous and transgenic MOR28 axons as rerouted-MOR28 glomeruli. Roughly the same numbers of rerouted-MOR28 glomeruli are observed in the two TO28 lines: 3.14 ± 0.55 (n = 7) in TO28L and 4.8 ± 0.86 (n = 5) in TO28H (fig. S1D).

We examined whether rerouted-MOR28 glomeruli could be formed throughout the lifetime of the animal, or whether they could only be formed during a specific developmental period. To determine this, we used doxycycline to silence the tetO promoter in mice capable of TO28 expression (mice bearing both the transgene and the OMP-IREs-tTA driver). Pregnant females were fed doxycycline from conception to allow the embryos to form wild-type glomerular maps. We then removed doxycycline from the diet at P0, P7, or P14 to allow transgene expression. Mice were maintained on a doxycycline-free diet to enable continuous transgene expression and were examined after 15 weeks or 32 weeks without the drug, nearly 3 times the half-life of OSNs (fig. S2A). Whereas transgenic MOR28 axons formed ectopic glomeruli and also innervated the wild-type MOR28 glomeruli, endogenous MOR28 axons almost never formed rerouted-MOR28 glomeruli (Fig. 2, A to D, and fig. S3A, C, and F). When doxycycline was removed at P0 or P7, rerouted glomeruli were observed in 1 out of 9 mice, or 1 out of 17, respectively. Rerouting was never observed (0 out of 9 mice) when doxycycline...
was removed at P14 (fig. S3B). We conclude that there is a critical period for the formation of rerouted-MOR28 glomeruli that ends at birth or shortly thereafter.

Among the many differences between the developing and adult olfactory systems is the fraction of OSNs that are immature with outgrowing axons. In embryos, most OSNs are immature, and their axons target the glomeruli within a short period of time. Thus, outgrowing axons do not contact many established axonal tracts. By contrast, at any given time in the adult, only rare OSN axons are actively growing along established tracts made up of mature OSN axons. We therefore tested whether the abundance of immature OSNs with outgrowing axons underlies the critical period for the formation of rerouted-MOR28 glomeruli. Mice capable of TO28 expression were treated with doxycycline beginning at conception to suppress the formation of rerouted-MOR28 glomeruli. As the mice were taken off doxycycline, they were also treated with methimazole, which causes OSN ablation (23). We empirically determined the peak effect of methimazole on MOR28-expressing neurons and found that 5 days after injection only 0.2% of these OSNs remain (fig. S2B). Thus, this paradigm enabled us to examine synchronized regrowth of 99.8% of the axons in the adult. When regrowth occurred in the presence of transgenic MOR28 axons (n = 5 mice), ectopic glomeruli were formed, but we still never observed rerouted-MOR28 glomeruli (Fig. 2, E and F, and fig. S3A). These data indicate that the developmental mechanisms available perinatally, when the olfactory map is established, are not available for regeneration in the adult after olfactory neuron ablation.

We next examined whether continuous transgene expression is required for the maintenance of rerouted-MOR28 glomeruli. Mice expressing the transgene developed until 2 months of age to allow ectopic and rerouted-MOR28 glomeruli to form. These mice were then treated with doxycycline for 21 weeks, or nearly twice the halflife of OSNs, to allow for substantial turnover of MOR28-expressing OSNs (fig. S4). Although no axon fibers expressing transgenic MOR28 were observed at this point, endogenous MOR28 axons continued to innervate the rerouted-MOR28 glomeruli (Fig. 3, A to C). Thus, rerouted-MOR28 glomeruli, once formed, were not dependent on persistent expression of transgenic MOR28.

Finally, we examined whether the rerouted-MOR28 glomeruli would still be targeted by newly growing endogenous MOR28 axons after OSN ablation. Mice expressing TO28L were allowed to form ectopic and rerouted-MOR28 glomeruli until 2 months of age. The OSNs were then ablated by methimazole injection, and the mice were treated with doxycycline for 2 months to allow the glomerular map to be restored without expression of transgenic MOR28 (fig. S4). These mice exhibited as many rerouted-MOR28 glomeruli as mice that were not treated with doxycycline after OSN ablation (Fig. 3, D and E). These results suggest that the previously formed rerouted-MOR28 glomeruli, or the tracts leading to them, are marked as targets for endogenous MOR28 axons.

We have shown that OSNs expressing a particular OR are affected non-cell autonomously by other OSNs that express the same OR. This supports the notion that homotypic attraction exists between OSNs expressing the same OR (19, 22). It is unclear whether these interactions are directly mediated by the axonal OR itself (13), or by any of the cell adhesion molecules implicated in OSN guidance (3). A role for OR-mediated, non-cell autonomous interactions in glomerular formation adds a layer of complexity to the prevailing model that focuses on cell-autonomous mechanisms by which ORs affect guidance (3, 24). Homotypic attraction between OSNs leads to convergence of like axons on the same glomerulus. This is in contrast to the homotypic repulsion between neurons expressing the same splice form of the adhesion molecule DSCAM1 in Drosophila and self-avoidance between neurites expressing the same type of protocadherin in mammals (25, 26). Should similar non-cell autonomous mechanisms participate in the organization of other circuits, it would pose a challenge for the development of neuronal regeneration therapies.

The confinement of rerouted-MOR28 glomeruli to areas within 650 m of the typical locations of the wild-type glomeruli and the cap in the number of rerouted-MOR28 glomeruli in the two founder lines suggest that axon rerouting can occur only within a “neighborhood” of glomeruli. This “neighborhood” likely corresponds to the area where the final sorting of axons expressing the same OR into discrete glomeruli occurs (3).

What is the nature of the guidance signal? It is formally possible that the axons of the ~30 endogenous MOR28 OSNs per mouse remaining after methimazole ablation serve as the structural substrate of the memory for the tract to the rerouted-MOR28 glomeruli. However, a model in which the few residual endogenous MOR28 OSNs act as “pioneers” is unlikely in view of the sparse distribution of these neurons. Alternatively, fragments of the axons of the dead neurons could mark the tracts to the glomeruli. In this case, the axons of regenerating OSNs would follow this trail to the correct glomerulus by interacting with these fragments. As another possibility, axons may leave molecular traces in the surrounding extracellular matrix, or may induce expression of specific marks in postsynaptic neurons. Regardless of the precise mechanism, our results show that these marks are put in place only during a developmental critical period. This may ensure the stability of the olfactory sensory map in spite of the continuous regeneration of OSNs throughout life.

References and Notes
Acquisition of Germ Plasm Accelerates Vertebrate Evolution

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Primordial germ cell (PGC) specification occurs either by induction from pluripotent cells (epigenesis) or by a cell-autonomous mechanism mediated by germ plasm (preformation). Among vertebrates, epigenesis is basal, whereas germ plasm has evolved convergently across lineages (epigenesis) or by a cell-autonomous mechanism mediated by germ plasm (preformation). Among Primordial germ cell (PGC) specification occurs either by induction from pluripotent cells by extracellular signals, a process referred to as epigenesis (6, 7), which also employ epigenesis (8). In contrast, in frogs the master regulators of pluripotency as employed in mammals have been deleted (6, 9, 10), and the GRN for mesoderm underwent expansions of key regulatory molecules (7, 11). Similar genetic innovations evolved in the GRNs for zebrafish development (12), which also uses preformation (13). The correlation of germ plasm with genetic change has been proposed to result from the relaxation of constraints on somatic development imposed by maintaining the PGC induction pathway (1, 3, 4). To investigate this possibility, we compiled available expressed sequence tag, mRNA and cDNA sequences from vertebrates (fig. S1A and table S1) identifying ortholog pairs shared between sister taxa with different modes of PGC specification and an appropriate mammal and outgroup sequence (14) (fig. S1B). To increase sequence numbers from organisms using epigenesis, we generated transcriptions from the axolotl and an Acipenseriforme, Acipenser ruthenus (the sterlet) (14), identifying 82,954 sequence clusters across all vertebrates. All analyses were performed with protein coding DNA sequence, excluding the saturated third position (14) (figs. S2 and S3).

Of the 56 published gene trees involving an anuran and a urodele, 29 do not recapitulate the known species phylogeny (table S2). The majority of the incongruent gene trees group urodele

Fig. 1. Amphibian four-taxon tree topologies. (A) Number of significant trees by bootstrapping (>70%) and SH test (P < 0.05) for each topology rooted with a Teleostei sequence. (B and C) The proportions of species phylogeny (black), mammal-urodele (gray), and mammal-anuran (white) topologies per species. (D) The likelihood of each species grouping with mammals when the tree is incongruent; species using preformation are shown in red, those using epigenesis in blue. Dashed lines indicate equal probability of species grouping with mammal or outgroup. (B) to (E) Only species with >20 significant trees are shown. The results excluding the transcriptome are shown in fig. S4.