Journal Club

Editor’s Note: These short, critical reviews of recent papers in the Journal, written exclusively by graduate students or postdoctoral fellows, are intended to summarize the important findings of the paper and provide additional insight and commentary. For more information on the format and purpose of the Journal Club, please see http://www.jneurosci.org/misc/ifa_features.shtml.

Functional Contribution of Adult-Generated Olfactory Bulb Interneurons: Odor Discrimination versus Odor Memory

Katherine G. Akers,1 Masanori Sagakuchi,1 and Maithe Arruda-Carvalho1,2

1Neurosciences and Mental Health, The Hospital for Sick Children, Toronto, Ontario MSG 1X8, Canada, and 2Institute of Medical Science, University of Toronto, Toronto, Ontario M5S 1A8, Canada

Review of Breton-Provencher et al.

Processing olfactory information is critical for an animal’s ability to locate food, detect danger, recognize conspecifics, find mates, and rear offspring. The olfactory bulb (OB) is the initial site of olfactory information processing in the brain. Within the OB, mitral cells receive input from sensory neurons in the olfactory epithelium and send output to higher-order structures such as the piriform cortex. Mitral cell activity is tightly regulated by dendrodendritic synapses with two types of local inhibitory interneurons: granule and periglomerular cells. Interestingly, OB granule and periglomerular cells differ from most neurons in that their production continues well beyond embryogenesis and into adulthood, prompting the question of whether adult-generated OB interneurons make a unique contribution to OB function.

In a recent article in The Journal of Neuroscience, Breton-Provencher et al. (2009) examined the impact of reducing adult neurogenesis on OB function. They decreased neurogenesis by infusing cytosine arabinoside (AraC) via osmotic minipump into the lateral ventricle of mice for 28 d. AraC is an antimitotic drug that incorporates into the DNA of dividing cells and induces apoptotic cell death, thereby disrupting the proliferation of neural progenitor cells in the subventricular zone and their subsequent migration to the OB. After infusion, mice treated with AraC exhibited a 75% reduction in the number of OB cells expressing doublecortin, a marker for immature neurons, compared with control mice treated with saline.

Breton-Provencher et al. (2009) found that although the reduction of adult neurogenesis left the basic structure of the OB intact, it had significant consequences for OB network function. Single-cell recordings revealed that mitral cells from AraC-treated mice exhibited less frequent spontaneous IPSCs and smaller evoked dendrodendritic inhibitory currents compared with control mice, indicating that reduction of neurogenesis decreased mitral cell inhibition. This decreased inhibition was not due to intrinsic changes in the electrophysiological properties of mitral cells, which displayed normal postsynaptic GABA<sub>A</sub> receptor function as well as normal passive membrane properties and spontaneous firing activity. Rather, the decreased inhibition appeared to result from fewer inhibitory synapses made onto the mitral cells, as the lateral dendrites of mitral cells from AraC-treated mice had fewer sites immunopositive for gephyrin, a postsynaptic GABA<sub>A</sub> receptor scaffolding protein. Because the lateral dendrites are where granule cells form inhibitory contacts, this finding suggests that the decrease in mitral cell inhibition resulted from fewer granule-to-mitral cell synapses. As dendrodendritic interactions between granule and mitral cells are largely responsible for driving gamma frequency oscillations (Lagier et al., 2004), the authors predicted that AraC-treated mice would have alterations in this type of oscillatory activity. Indeed, local field potential recordings revealed that AraC-treated mice showed a reduction in the peak frequency of gamma frequency oscillations measured from the mitral cell layer after stimulation of the olfactory nerve (~45 Hz in saline-treated mice vs ~39 Hz in AraC-treated mice).

These findings suggest that adult-generated OB interneurons are critical for shaping the overall pattern of OB network activity and, therefore, may make an important contribution to olfactory-associated behavior. In particular, OB interneurons may play a role in perceptual discrimination between odors. Because each granule cell forms dendrodendritic contacts with multiple mitral cells, the excitation of a granule cell by a mitral cell leads to the inhibition of neighboring mitral cells. This process of lateral inhibition may sharpen the distinction between neural representations of similar odors and thereby facilitate odor discrimination. For example, in the rabbit, individual mitral cells showing excitatory responses to certain odorants showed inhibi-
itory responses to odorants with similar molecular structures, and this odor-specific response pattern was eliminated by blockade of inhibitory neurotransmission (Yokoi et al., 1995). Synchronous oscillatory activity in the OB may also be important for odor discrimination. In the honeybee, pharmacological desynchronization of odor-evoked oscillations impaired perceptual discrimination between structurally similar odorants (Stopfer et al., 1997).

A perplexing finding is that although AraC-treated mice showed no evidence of memory for a previously encountered odor after 60 min, they were able to remember an association between an odor and a food reward for at least 7 d. If mice cannot exhibit poor discrimination between odors (100% almond vs 100% coconut) based on their chemical composition, what is the basis for their ability to associate odors with food reward? One possibility is that discrimination is the developmental requirement for even the most difficult odor discriminations. A second factor that might influence whether reduced OB neurogenesis impairs odor discrimination may be the difficulty of the discrimination because the odors were too similar, leaving open the possibility that using more similar odors could reveal impairments. A recent study (Lazarini et al., 2009), however, demonstrated that irradiated mice with reduced OB neurogenesis showed normal odor discrimination despite being extensively tested in tasks with highly similar odors, providing evidence that adult OB neurogenesis is not necessary for even the most difficult odor discriminations.

This study is the latest in a growing collection of studies investigating the impact of reduced adult OB neurogenesis on olfactory-associated behavior in mice (Table 1). These studies, however, appear to provide conflicting results. Whereas some report that reduced neurogenesis impairs odor discrimination (Gheusi et al., 2000; Enwere et al., 2004; Bath et al., 2008), others report that reduced neurogenesis leaves odor discrimination intact but impairs odor memory (Breton-Provencher et al., 2009; Lazarini et al., 2009), and yet another reports that reduced neurogenesis impairs neither odor discrimination nor odor memory (Imayoshi et al., 2008).

One factor that may influence whether reduced OB neurogenesis impairs odor discrimination may be the developmental timing of the experimental manipulation. As proposed by Lazarini et al. (2009), impairments in odor discrimination may be observed when OB neurogenesis is disrupted during embryogenesis but not during adulthood. That is, the ability to discriminate between odors may depend on the initial formation of OB neural circuitry brought about by the early wave of neurogenesis during the prenatal and early postnatal periods, but may not depend on the continued addition of new neurons to that circuitry during adulthood. Consistent with this idea, odor discrimination was impaired by genetic mutations that likely affected OB neurogenesis throughout development (Gheusi et al., 2000; Enwere et al., 2004; Bath et al., 2008) but was not impaired by manipulations that targeted neurogenesis specifically during adulthood (Imayoshi et al., 2008; Breton-Provencher et al., 2009). A recent study (Lazarini et al., 2009) did not find deficits in odor discrimination because the odors were too distinct, leaving open the possibility that using more similar odors could reveal impairments. A recent study (Lazarini et al., 2009), however, demonstrated that irradiated mice with reduced OB neurogenesis showed normal odor discrimination despite being extensively tested in tasks with highly similar odors, providing evidence that adult OB neurogenesis is not necessary for even the most difficult odor discriminations.

A perplexing finding is that although AraC-treated mice showed no evidence of memory for a previously encountered odor after 60 min, they were able to remember an association between an odor and a food reward for at least 7 d. If mice cannot recognize an odor after a short span of time, it is difficult to understand how they could form long-term odor associations. One possible reason for this discrepancy is that short- and long-term odor memories were assessed by two different behavioral tasks, the former involving odor habituation and the latter involving odor–food associations, which may have differential requirements for adult-generated OB interneurons. For instance, odor–food associative memory might be unaffected by reductions in adult OB neurogenesis because associations between odors and nonolfactory stimuli rely on piriform and orbitofrontal cortices (Wilson et al., 2006) more than the OB.

This study is the latest in a growing collection of studies investigating the impact of reducing adult OB neurogenesis on olfactory-associated behavior in mice (Table 1). These studies, however, appear to provide conflicting results. Whereas some report that reduced neurogenesis impairs odor discrimination (Gheusi et al., 2000; Enwere et al., 2004; Bath et al., 2008), others report that reduced neurogenesis leaves odor discrimination intact but impairs odor memory (Breton-Provencher et al., 2009; Lazarini et al., 2009), and yet another reports that reduced neurogenesis impairs neither odor discrimination nor odor memory (Imayoshi et al., 2008).

One factor that may influence whether reduced OB neurogenesis impairs odor discrimination may be the difficulty of the discrimination. As noted by Breton-Provencher et al. (2009), adult-generated OB interneurons may be exclusively involved in discrimination between highly similar odors (e.g., odors with similar molecular structures or similarly comprised mixtures of odors). For instance, aged or mutant mice with reduced OB neurogenesis discriminated between two distinct odors (100% almond vs 100% coconut) but not two similarly comprised mixtures of the odors (58% almond/42% coconut vs 42% almond/58% coconut) (Enwere et al., 2004). Perhaps some studies (Imayoshi et al., 2008; Breton-Provencher et al., 2009) did not find deficits in odor discrimination because the odors were too distinct, leaving open the possibility that using more similar odors could reveal impairments. A recent study (Lazarini et al., 2009), however, demonstrated that irradiated mice with reduced OB neurogenesis showed normal odor discrimination despite being extensively tested in tasks with highly similar odors, providing evidence that adult OB neurogenesis is not necessary for even the most difficult odor discriminations.

Table 1. Impairment in olfactory-associated behavior after reduction of adult OB neurogenesis

<table>
<thead>
<tr>
<th>Study</th>
<th>Method of reducing OB neurogenesis</th>
<th>Impact on immature adult-generated neurons</th>
<th>Impact on mature adult-generated neurons</th>
<th>Odor discrimination</th>
<th>Odor memory</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gheusi et al., 2000</td>
<td>NCAM+/−</td>
<td>Migration in RMS: impaired</td>
<td>GCL width: 35% reduction</td>
<td>Impaired</td>
<td>Impaired short-term (80–100 min)</td>
</tr>
<tr>
<td>Enwere et al., 2004</td>
<td>Aging, Lir+/−, TGFα+/−/−/−/−</td>
<td>BrdU+/−/−/−/−+cells (4 wk postinjection); 59% reduction in GL in aged mice Proliferation: no change</td>
<td>BrdU+/−/−/−/−+ cells (4 wk postinjection); 41% reduction in GL, 55% reduction in GCL in aged mice</td>
<td>Impaired</td>
<td>Impaired</td>
</tr>
<tr>
<td>Bath et al., 2008</td>
<td>BDNF+/−/−/−, TrkB+/−/−/−, DNM2+/−/−/−</td>
<td>DCM+ cells in RMS: reduced DCX+ cells in GL and GCL: 70% reduction</td>
<td>Neun+ cells in GCL: 10% reduction No deficit</td>
<td>No deficit</td>
<td>No deficit</td>
</tr>
<tr>
<td>Imayoshi et al., 2008, Lazarini et al., 2009</td>
<td>Nestin-CRE-ER−/−×NSE-DTA Focal SVZ irradiation</td>
<td>DCX+ cells in RMS: reduced DCX+ cells in GL and GCL: 70% reduction</td>
<td>Neun+ cells in GCL: no change No deficit</td>
<td>No deficit</td>
<td>No deficit</td>
</tr>
<tr>
<td>Breton-Provencher et al., 2009</td>
<td>LV AraC infusion</td>
<td>DCX+ cells in GCL: 75% reduction</td>
<td>Neun+ cells in GCL: no change No deficit</td>
<td>No deficit</td>
<td>Impaired short-term (60–120 min)</td>
</tr>
</tbody>
</table>

BrdU, Bromodeoxyuridine; CRE, cAMP response element; GCL, granule cell layer; GL, glomerular layer; LV, lateral ventricle; RMS, neural migratory stream; SVZ, subventricular zone.
A third factor that may influence the type of behavioral impairment observed after reduction of OB neurogenesis is the age of the affected neurons. After their generation in the subventricular zone, newborn neurons destined for the OB undergo several migrational and maturational stages before fully integrating into OB circuitry. In the study by Breton-Provencher et al. (2009), AraC-treated mice possessed the same number of mature OB granule cells as control mice yet exhibited alterations in OB network function and odor memory, suggesting that immature granule cells play a special role in OB function. In support of this possibility, immature adult-generated granule cells display greater immediate early gene expression in response to novel odors compared with mature cells (Magavi et al., 2005), and only immature adult-generated granule cells exhibit long-term potentiation of synaptic strength (Nissant et al., 2009). To determine how different ages of adult-generated OB interneurons might differentially contribute to olfactory-associated behavior, it will be critical to target the cells at specific maturational stages.

The study by Breton-Provencher et al. (2009) is valuable in that it uses a multi-level approach from cell to behavior to demonstrate the importance of adult neurogenesis in olfactory information processing. Their finding of impaired odor memory after reduction of adult neurogenesis invites future studies to focus on precisely how adult-generated OB interneurons are involved in the retention of odor memories. Furthermore, as a complement to studies showing that reduction of OB neurogenesis impairs olfactory function, future studies could examine whether enhancement of OB neurogenesis improves olfactory function. Such studies would increase our understanding of how new neurons can be successfully integrated into existing neural circuits, which could propel efforts to improve functional recovery from brain injury or disease. A comprehensive knowledge of how brain pathology and/or the recovery of brain function might involve adult neurogenesis in the OB, hippocampus, and neocortex will be fundamental for the translation of basic research findings to clinically relevant situations.

References


