INTRODUCTION

All olfactory receptor neurons (ORNs) expressing the same receptor protein project their axons to specific glomeruli in the olfactory bulb (OB) where they form excitatory synapses onto a circuit of diverse cell types including mitral/tufted (M/T) cells and several types of juxtaglomerular interneurons (Buck and Axel, 1991; Ressler et al., 1994; Kiyokage et al., 2010). The convergence of receptor input onto OB output neurons forms the basis of an odorant-specific spatial map of M/T cell glomerular activity in which each glomerulus represents the molecular receptive range of the ORN type that innervates it (Mori et al., 2006).

In other sensory systems, learning can often enhance or shift cortical maps of sensory information following learning (Gilbert et al., 2001). Similarly, experiments in the rodents have shown that OB glomerular layer processing of odor information is altered with learning as well. For example, associative conditioning in rat pups leads to increased 2-DG glomerular uptake (Coopersmith et al., 1986; Sullivan and Leon, 1986; Woo et al., 1987; Woo and Leon, 1991) and increased c-fos expression in juxtaglomerular neurons and deeper layer granule cells in regions responding to the trained odorant (Johnson et al., 1995; Funk and Amir, 2000; Salcedo et al., 2005). Together these experiments point toward learning leading to long-term changes in glomerular layer interneuronal responses to learned odors.

While numerous studies have reported changes in M/T cell firing following learning (Wilson et al., 1983; Buonviso and Chaput, 2000; Ravel et al., 2003; Fletcher and Wilson, 2003; Doucette and Restrepo, 2008; Doucette et al., 2011), comparatively little is known about changes in M/T OB spatial representations of odorants following learning. Work from the Freeman lab has also shown that the EEG spatial pattern measured across the bulb for a given odor changes following associative conditioning to that odorant (Freeman and Schneider, 1982; Grajski and Freeman, 1989). More recently, olfactory associative conditioning has also been shown to increase OB dorsal surface intrinsic signal responses to the trained odor (Yuan et al., 2002). While these studies clearly demonstrate long-term changes in both the temporal and spatial patterns of M/T odor responses following learning, it is still not fully clear how M/T cell glomerular odor representations are affected in mammals following associative learning.

Optical imagining in insects however, has recently revealed that olfactory associative conditioning can enhance glomerular level projection neuron representations to the trained odor (Faber et al., 1999; Yu et al., 2004). For example, in bees, pairing an odorant with a sucrose reward lead to increases in the response amplitude of only the glomeruli within the trained odor representation (Faber et al., 1999). While in Drosophila, aversive conditioning has been shown to lead to a recruitment of previously unresponsive glomeruli into the trained odorant glomerular representation (Yu et al., 2004).

In this study, we tested whether aversive associative conditioning alters M/T cell glomerular odor representations in the mammalian OB, and whether the enhanced is driven by new glomeruli...
recruited into the odor representation or altered responses from existing glomeruli. This was accomplished issue using transgenic mice that express the fluorescent Ca$^{2+}$ indicator protein GCaMP2 exclusively in OB M/T cells with no expression in inhibitory interneurons. In these mice, odor-evoked glomerular activity patterns that reflect M/T cell glomerular responses can be visualized on the dorsal surface with individual glomerular resolution (Fletcher et al., 2009). Optical imaging in these transgenic mice allow the unique advantage of comparing glomerular responses from the same glomeruli before and after training in the same animal. Through multi-day optical imaging, glomerular odor responses were compared before and after olfactory associative conditioning with the same odorant. The results suggest that associative conditioning enhances glomerular representation to the trained odor primarily by increasing responses in initially weakly activated glomeruli within the representation.

**MATERIALS AND METHODS**

**ANIMALS AND SURGERY**

Adult transgenic mice expressing the fluorescent calcium indicator protein GCaMP2 under the Kv3.1 potassium channel promoter were used (Diz-Garcia et al., 2005). In these mice, GCaMP2 is expressed primarily in OB output neurons, M/T cells, with no expression in olfactory sensory neuron axons or OB inhibitory interneurons (Fletcher et al., 2009). Odor-evoked changes in glomerular fluorescence in GCaMP2 mice therefore, reflect glomerular level M/T cell odor responses.

For surgery and imaging, mice were anesthetized with sodium pentobarbital (50 mg/kg, i.p.). Mice were secured in a custom stereotaxic apparatus (Narishige) and a chronic imaging window was made on dorsal surface of the skull by thinning the bone overlying the OBs. The imaging window was covered with a thin layer of cyanoacrylate glue (Maxicure, BSI, Inc.) to improve optical transparency and seal the skull. An anchor screw (#000–120, 1/16” length, Small Parts, Inc.) was inserted into the parietal bone. A head post made of 16-gauge syringe needle tubing was placed on top of the skull and secured into place with acrylic dental cement (Stoelting). In some animals, the clarity of optical imaging window significantly degraded over time due to surgical damage or problems with the glue adhering to the surface of the skull. In both cases, these animals were removed from the study. All animal care protocols were approved by the University of Texas and the University of Tennessee Institutional Animal Care and Use Committees.

**ASSOCIATIVE CONDITIONING**

Mice were placed into four groups: a group that received pairings of odor and foot shock (Paired, n = 10), a group that only received foot shocks (Shock only, n = 6), and group that received only odorant presentations (Odor only, n = 6), and a group with no exposure to either MV or foot shock (Naïve, n = 4). The conditioning apparatus consisted of a small plexiglas cylinder placed inside a standard mouse test cage equipped with a floor of parallel steel bars connected to a precision animal shocker (Coulbourn Instruments). Associative conditioning took place in a single session in which the mice received 10 trials of 1% vol/vol methyl valerate diluted in mineral oil (MV) (4 s duration) terminating in 0.25 s duration, 0.4 mA foot shocks. The conditions for the control groups were identical except the mice received either 10 trials of shock only (0.25 s duration, 0.4 mA) or 10 trials MV only (4 s duration). All trials were separated by an inter-trial interval of 2 min. On the day following training, animals were placed in a two-odor choice test. This test involved placing an animal into a standard mouse cage that contained a small cotton swab containing the MV on one side of the cage and a small cotton swab containing a novel odor (peppermint) on the other. Mice were placed along the midline of the cage between the two odor-ant zones and allowed to freely move around the cage for 60 s. The first two sessions (Days 1–4) took place prior to conditioning with all mice naïve to the odor. On Day 7, mice received one of the three conditioning protocols. On the following day (Day 8), mice were imaged again. During each session, MV was presented 8–10 times. Glomerular odor representations were generated from the averages of the individual trials on each day.

Maps of odor-evoked spatial activity were generated by first correcting for photobleaching and then spatially low-pass filtered as described previously (Fletcher et al., 2009). For each trial, the odor-evoked change in fluorescence was calculated by subtracting the average of five frames immediately preceding stimulus onset from the average of five frames centered on the peak of the response from the first respiration for each pixel. Images reflecting the relative change in fluorescence (ΔF/F) were then calculated by dividing the odor-evoked change in fluorescence by the resting fluorescence. The final normalized odor map for each trial was generated by dividing the entire image by the mean response of all activated glomeruli. For each day, all normalized trials were averaged and represented in pseudo color to give the mean normalized response for that day. For pre- versus post-training comparisons, odor maps were overlaid and re-aligned using skull landmarks and blood vessel patterns.

For quantitative analysis, mean normalized individual glomerular responses for each day were calculated from the ΔF/F response measured at the center of each glomerulus (3 x 3 pixel average), normalized to the mean of all glomerular responses, and average across all trials. All individual glomerular responses were calculated from the raw data without any spatial filtering. A glomerulus was considered to be activated if its odor-evoked signal amplitude was five-times greater than the rms noise level calculated from an adjacent non-responsive area (Fletcher et al., 2009). While individual glomeruli could be identified in the same
animal across days, with our limited odor set we were unable to identify the same glomerulus across animals. Due to this, all group analysis was limited to comparing the relative responses of the glomeruli as a population. For comparisons across days, the mean absolute difference (MAD) = (∑(x1 - x2)), where x1 and x2 are the normalized ΔF/F responses measured on the first and second imaging sessions for each glomerulus was calculated (Kent and Mozell, 1992; Verhagen et al., 2007).

To compare changes in the MV odor representation as a whole, each day’s responses from all glomeruli from each animal was represented in a multidimensional odor space (Deisig et al., 2006). The MV representation for each day was expressed as a vector in n-dimensional odor space where each dimension is represented by the responses from each individual glomerulus. To compare changes in the MV representations across days, we calculated their proximity in odor space by comparing their Euclidian distances. The Euclidian distance between the MV representations for each day was calculated as:

\[ d_{ij} = \sqrt{\sum_{k=1}^{p} (X_k - X_j)^2} \]

with i and j indicating days, p the number of glomeruli, and X_k the response of glomerulus k on day i. For comparisons across groups, the mean Euclidian distance between the MV representations on D1 and D4 and D4 and D8 were compared. As no manipulations occurred between D1 and D4 in any group, the Euclidian distances should be similar across groups and represent the day-to-day variability in the representations. We then compared this mean Euclidian distance to the distances between D4 and D8 representations that spanned the training day. For visualization, we then used principle component analysis followed by a varimax rotation to project the representations from each group into a three dimensional odor space. Together these three factors accounted for over 97% of the variance in each group.

During all imaging sessions, animals were freely breathing and respiratory activity was monitored with a piezoelectric device strapped around the chest. Respiratory rates were taken from random trials of each imaging session for each animal. In some sessions, the piezoelectric device did not reliably detect respiration. In these animals, the respiratory rate was calculated from the respiratory oscillation observed in the odor-evoked GCaMP2 odor-evoked signal. Overall, the mean respiratory rate across all animals and days was 1.72 ± 0.06 Hz (range: 1.0–3.1 Hz).

ODORANT PRESENTATION

During imaging, methyl valerate was delivered using a flow-dilution olfactometer previously described (Fletcher et al., 2009). Separate flow controllers for the clean air and the pure odorant vapor were used to mix the flow streams at the end of the odor delivery system to achieve an approximate concentration of 0.25% s.v. at a flow rate of 0.7 L/min. Odor pulse duration was 2 s with an inter-stimulus interval of at least 60 s to prevent any short-term adaptation of the responses. For behavioral training, MV was presented through a small hole at the top of the chamber at a flow rate of 1 L/min using an olfactometer (Vulinitus). For the behavioral training and testing, pure odorsants were dissolved in mineral oil to a 1% vol/vol concentration. This concentration of MV was used for training and testing to account for diffusion of the odorant in the behavioral chamber. The intensity of the odorant in the behavioral chamber was similar to the intensity of the MV delivered in the imaging experiments.

RESULTS

To confirm that the training paradigm used here leads to odor learning, a subset of the imaged mice was placed in a two-odor choice behavioral task on D8 before the final imaging session. Mice in the Paired group spent significantly less time on the MV side compared to naïve non-imaged mice, shock-only, or odor-only mice, suggesting a learned MV aversion (percent time on MV side: Naive: 46.7 ± 2.1%; Shock Only: 51.6 ± 1.3%; Odor Only: 50.0 ± 5.5%; Paired: 30.6 ± 3.4%; ANOVA), Main effect of day: [F(2, 60) = 4.78, p = 0.03] (ANOVA), [F(3, 18) = 7.57, p < 0.05]; post-hoc Fisher’s test, p < 0.05 (Figure 1).

Mice were imaged on D1 and D4 to establish stable baseline responses and imaged again following training (D8). Based on D1 responses from all animals (22 mice, 327 glomeruli), MV activated an average of 14.9 ± 3.6 glomeruli located on the anterior-dorsal region of the OB (Figure 2). The average raw odor response of all glomeruli per animal decreased slightly across days for all three groups, with all groups showing similar decreases (Means: Day 1 = 5.31% ΔF/F; Day 4 = 4.55% ΔF/F; Day 8 = 4.42% ΔF/F) ANOVA, Main effect of day: [F(2, 60) = 4.78, p = 0.03]. Despite this, no significant decrease was observed in any group following training (D4–D8 comparisons; post hoc Fisher’s tests: Shock Only: p = 0.61, ns; Odor Only: p = 0.22, ns; Paired: p = 0.73, ns).

During imaging, normalized responses remained stable across the first two sessions (Day1 and Day4) with the number of glomeruli.
and position of activated glomeruli identical for all groups [Figures 1A, 2A, and 3A (Day1–Day4)]. Similar results were obtained following training, with the number of glomeruli remaining stable [Figures 3A, 4A, and 5A (Day4–Day8)]. However, changes in normalized glomerular responses to MV were observed in the Paired group (Figure 5A). In this case, the relative amplitudes of a few glomeruli within the representation were observed to increase in response to MV.

To further explore this, we obtained a measure of the mean day-to-day change in glomerular responses prior to any odor training by calculating the mean D1–D4 percent change (Shock Only group: 0.49 ± 10.97%; Odor Only group: −0.12 ± 9.71%; Paired group: 1.66 ± 14.11%), All groups combined: 0.92 ± 12.37%, mean ± 1SD). We found no differences in mean D1–D4 percent change between groups ANOVA, $F(3, 326) = 0.972, p = 0.41$. Because no training took place during the first two imaging sessions, we used the overall mean D1–D4 percent as a range of normal day-to-day variability in glomerular odor responses. Thus, any responses falling out of this range ($−23.82–25.66\%$, Figures 3B, 4B, 5B, gray boxes) could be considered significant changes. For the examples shown here, no post-training responses were found to be outside of this range in either of the mice receiving only foot shocks or presentations of MV alone (Figures 4B and 5B). However, in the mouse receiving pairings of MV and foot shock, two glomeruli showed significant post-training response increases (Figure 5B, glomeruli #1 and #8), and once glomerulus showed a significant post-training response decrease (Figure 5B, glomerulus #19).

In accord with this, we observed more large post-training increases (D4–D8 percent change ≥25%) from glomeruli in the Paired group (9.3%) than either the Shock only (3.2%) or Odor only group (4.1%). The percentage of glomeruli displaying significant post-training response decreases was small for all groups (Shock Only: 1.1%; Odor Only: 1.4%; Paired: 1.9%). The same trend was observed when the cutoff for percent change was lowered to the mean D1–D4 percent change ± 1SD).

To quantify response differences across the glomerular population for each group, the mean day-to-day absolute difference in glomerular response (MAD) for each animal was calculated. This measure gives the mean change in response across all glomeruli. In this analysis, the lower the MAD, the more similar the responses. Since glomerular responses were measured on two separate days prior to training, a comparison of the responses from these days gave a baseline measure of day-to-day variation in responses. The pre-training (D1–D4) MAD was found to be similar across all groups (MAD: Shock only: 0.08 ± 0.006% ΔF/F; Odor only: 0.09 ± 0.005% ΔF/F; Paired: 0.09 ± 0.008% ΔF/F) (Figure 6A, black bars). These values were similar to the mean trial-to-trial MAD calculated from consecutive odor trials taken from each animal on each day (MAD = 0.09 ± 0.01% ΔF/F) (Figure 6B).

To compare changes in glomerular responses following training, the glomerular responses taken from the last pre-training day (D4) and the day following the training (D8) were used to calculate the post-training (D4–D8) MAD for each group (MAD: Shock only: 0.08 ± 0.003% ΔF/F; Odor only: 0.08 ± 0.005% ΔF/F; Paired: 0.14 ± 0.007% ΔF/F) (Figure 6A, red bars). In comparing the pre- and post-training MAD, a significant
Fletcher Learning alters glomerular odor responses

FIGURE 4 | Presentations of odor alone do not significantly alter glomerular G-CaMP2 odor representations in individual mice. (A) Methyl valerate response maps taken from Day 1 and Day 4 prior to any conditioning and are similar across the days. The last map (Day 8) was taken one day after exposing the animal to presentations of odor only. No dramatic changes in response amplitude were seen in glomeruli on the day following the foot shock presentations. The Mean Absolute Day-to-Day Difference (MAD) for each glomerulus remained similar before and after training. Scale bar: 100 μm (B) The left panel indicates the position of the glomeruli. The graph represents the percent change in response for each glomerulus across days. The gray box represents the pre-training percent change (mean ± 2SD) taken from all glomeruli across all three groups. Again, no significant changes in response amplitude were observed across days.

difference between the three groups was found ANOVA: main effect of group: \( F(2, 44) = 4.27, p < 0.05 \), with the Paired group displaying a significant increase between the mean pre- and post-training MAD (post hoc Fisher’s test, \( p < 0.05 \)). This effect was not observed in the other groups.

Within the Paired group, we noticed that the largest post-training changes were often observed in the initially weaker responding glomeruli. To quantify this, we divided all the glomeruli in each group into quartiles based on their initial responses to MV and compared the average post-training percent change in each quartile. In both the Shock only and Odor only groups, average post-training percent changes were small in each quartile (Figure 7). Additionally, no significant differences were observed across quartiles in either group (Shock only: ANOVA: \( F(3, 90) = 0.48, p = 0.7 \); Odor only: ANOVA: \( F(3, 70) = 2.36, p = 0.08 \)). However, in the Paired group, we found that a significant increase in the post-training percent change in glomeruli from the lowest quartile of initial response (Paired: ANOVA: \( F(3, 90) = 7.84, p < 0.05 \); post hoc Fisher’s test, \( p < 0.05 \)). In line with this, we observed more large post-training increases (D4–D8 percent change ≥25%) from first quartile glomeruli in the Paired group (41.7%) than either the Shock only (0%) or Odor only group (9%).

To investigate whether the changes in individual glomerular responses following training are enough to alter the overall representation of the odor, we compared the Euclidian distances in putative olfactory space between the D1 and D4.
representations to that between the D4 and D8 representations in each animal. As no manipulations occurred between D1 and D4 in any group, the Euclidian distances were similar across groups (Shock only = 0.40; Odor only = 0.32; Paired = 0.45). Following the training, only the Paired group showed a shift in the representation as measured by the Euclidian distance (Shock only = 0.44; Odor only = 0.34; Paired = 0.66). Paired t-tests revealed a significant difference only in the Paired group (Shock only: $t_{(5)} = -0.98, p = 0.37$; Odor only: $t_{(5)} = -0.61, p = 0.56$; Paired: $t_{(5)} = -2.40, \ p < 0.05$) (Figure 8A).

To visualize this shift in the representation, we performed PCA analysis on the all the glomerular responses from each group across days. MV representations from each group were projected onto a 3-dimensional odor space based on the first three principle components (Figures 8B–D). As can be seen in the two control groups, the post-training D8 odor representations are clustered close to the D1 and D4 responses. However, in the Paired group only, the D8 response shifted away from the D1 and D4 responses.

**DISCUSSION**

Using *in vivo* imaging, this study investigated the impact of olfactory aversive learning on OB M/T cell glomerular odor responses in adult mice. The results demonstrate that olfactory learning significantly alters OB M/T cell glomerular odor representations to the trained odor by altering the response amplitudes of glomeruli within the representation. Overall, we find that associative conditioning alters M/T cell responses following training primarily by increasing the responses in the initially weakest responding glomeruli. Overall, this study provides further evidence that the neuronal representations of odors that predict aversive or emotional events can be enhanced at the first synapses of central olfactory processing. These changes are potentially large enough to alter the representation of the odorant at the glomerular level and could serve as the basis for learning-induced changes in higher olfactory centers.

These findings expand upon previous studies reporting enhanced glomerular responses following learning in rodents (Coopersmith et al., 1986; Johnson et al., 1995; Yuan et al., 2002; Salcedo et al., 2005). In these studies, subsequent exposure to the trained odorant lead to increased c-fos activation in populations of juxtaglomerular neurons (Johnson et al., 1995; Salcedo et al., 2005; Funk and Amir, 2000), as well as increased 2-deoxyglucose uptake in glomerular layer neurons in regions known to be activated by the learned odorant (Woo et al., 1987; Woo and Leon, 1991). Since the calcium indicator used is this study is expressed only in M/T cells (Fletcher et al., 2009), the changes that we observed in glomerular responses reflect changes in M/T cell glomerular tuft responses to OSN input. Our results suggest that aversive conditioning enhances M/T cell glomerular responses in some glomeruli and that this enhancement is driven largely by the weakest responding glomeruli of the representation. As calcium increases measured within a glomerulus reflect a population M/T cells, the increased responses following learning seen in some glomeruli could reflect increases in individual M/T cell activity, recruitment of additional M/T cells associated with the glomerulus, or both. Similar results have been reported in insects, with associative conditioning leading to enhanced principle neuron activity into the antennae lobe glomerular representation of the trained odorant (Faber et al., 1999; Daly et al., 2004; Yu et al., 2004). However, unlike similar studies in *Drosophila* where previously unresponsive glomeruli were recruited into the odor representation following associative conditioning (Yu et al., 2004), we found no instances of recruitment of new glomeruli following conditioning.
Fletcher Learning alters glomerular odor responses

FIGURE 8 | Olfactory associative conditioning significantly alters the representation of the trained odor in odor space. (A) Euclidian distances in putative olfactory space between the D1 and D4 representations and between the D4 and D8 representations were compared for each group. Following the training, the only the Paired group showed a shift in the representation as measured by the Euclidian distance. (B–D) MV representations in odor space before and after conditioning in all three groups (Day 1: blue; Day 4: green; Day 8: red). In both the Shock only and Odor only groups, the post-training D8 odor representations are clustered close to the D1 and D4 responses. However, in the Paired group, the D8 response has shifted away from the D1 and D4 responses. *Denotes a significant difference between the groups (p < 0.05).

While we did observe some glomeruli in the Paired animals that significantly increased, we did not observe any large-scale changes in response amplitude across all glomeruli following learning. This is in contrast to earlier studies in rat pups that have reported an overall decrease in odor-evoked M/T cell responses following associative conditioning (Wilson et al., 1985; Wilson and Leon, 1988). Unlike these studies, we found very few instances of decreases in M/T glomerular odor responses following learning. There are several potential reasons for this difference. First, increased fluorescence in calcium indicators observed in M/T dendritic glomerular tufts reflects a combination of calcium influx from OSN-driven activation of NMDA receptors, dendritic-initiated spiking, and back-propagating somatic-initiated action potentials (Chen et al., 2002; Yuan et al., 2004; Zou et al., 2006; Fletcher et al., 2009). It is likely that the learning-induced increases in M/T cell odor-evoked suppression reported in the earlier electrophysiological studies are driven by changes in deeper granule cell mediated inhibition (Brennan and Keverne, 1997). Given the strong calcium signals observed in M/T dendritic tufts in response to OSN input (Chen et al., 2002; Zou et al., 2006), it is possible that this deeper level inhibition in M/T cell output is masked when only observing activity in the tuft. Another potential explanation lies in the fact that the studies mentioned above (Wilson et al., 1985; Wilson and Leon, 1988) were performed in rat pups and our study was performed in adult mice. As it is clear that infants rodents possess a unique and immature olfactory system (Greer et al., 1982; Raineki et al., 2010), it is possible that the olfactory learning in infants and adults utilizes different mechanisms. Finally, it is important to note that our technique only allows us to observe glomerular odor representations located across the dorsal surface. 2-DG mapping studies in mice have shown that methyl valerate activates a cluster of glomeruli in the posterior-lateral region of the OB as well (Johnson et al., 2009). With our technique, we can only observe the glomerular representation on the dorsal surface and we may be missing the effects of training on glomeruli located in other regions.

POTENTIAL MECHANISMS OF GLOMERULAR PLASTICITY

As M/T cell odor responses reflect a combination of input from sensory neurons, interneuronal inhibition, and neuromodulatory input from centrifugal cortical and neuromodulatory regions (Kosaka and Kosaka, 2005; Fletcher and Chen, 2010; Kiyokage et al., 2010), one possible mechanism behind this effect could
involve Hebbian plasticity at the level of ORN input into the bulb (Ennis et al., 1998). In this case, it is possible that the simultaneous activation of OB neurons by ORN input and centrifugal activation via foot shock creates the conditions necessary to induce long-term changes in the sensitivity of the ORN-M/T cell synapse.

Additionally, it is possible that the enhanced glomerular responses following associative conditioning could be caused by learning-dependent structural changes within the glomerular layer. For example, associative conditioning has also been shown to increase the size of glomeruli responsive to the trained odorant (Woo et al., 1987; Jones et al., 2008). In one case, this size increase appears to be due to increases in the number of ORN axons innervating the glomeruli (Jones et al., 2008). While interesting, it is unlikely that similar anatomical reorganization is driving the increased responses seen in glomeruli in this study, as the changes observed in these studies required several days of associative conditioning and our study look at responses the day immediately following training.

**POTENTIAL IMPACT ON ODOR DISCRIMINATION AND PERCEPTION**

The post-training enhancement of glomerular odor response seen here could have several consequences on M/T cell odor coding. First, the increased responses following training could serve to increase sensitivity in the weaker glomeruli, possibly decreasing behavioral detection thresholds. Second, enhanced glomerular responses may lead to a shift in glomerular odor receptive fields (ORFs). In the auditory system, neuronal RFs can be shifted away from their preferred frequency and toward the paired frequency following tone-foot shock pairings (Lennartz and Weinberger, 1992; Edeline et al., 1993). Similarly, odorant experience has been shown to shift M/T cell ORFs toward the experienced odor (Fletcher and Wilson, 2003). Together these changes could lead to a decorrelation of the representation of the trained odor and other similar odors, possibly making the trained odor more discriminable. In line with this, behavioral discrimination experiments in both rats and humans studies have demonstrated increased olfaction acuity following associative conditioning (Rabin, 1988; Fletcher and Wilson, 2002; Linster et al., 2002; Li et al., 2008).

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Learning alters glomerular odor responses

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