

# *In Vivo* Identification of Eugenol-Responsive and Muscone-Responsive Mouse Odorant Receptors

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Our understanding of mammalian olfactory coding has been impeded by the paucity of information about the odorant receptors (ORs) that respond to a given odorant ligand in awake, freely behaving animals. Identifying the ORs that respond *in vivo* to a given odorant ligand from among the ~1100 ORs in mice is intrinsically challenging but critical for our understanding of olfactory coding at the periphery. Here, we report an *in vivo* assay that is based on a novel gene-targeted mouse strain, S100a5–tauGFP, in which a fluorescent reporter selectively marks olfactory sensory neurons that have been activated recently *in vivo*. Because each olfactory sensory neuron expresses a single *OR* gene, multiple ORs responding to a given odorant ligand can be identified simultaneously by capturing the population of activated olfactory sensory neurons and using expression profiling methods to screen the repertoire of mouse *OR* genes. We used this *in vivo* assay to re-identify known eugenol- and muscone-responsive mouse ORs. We identified additional ORs responsive to eugenol or muscone. Heterologous expression assays confirmed nine eugenol-responsive ORs (Olf73, Olf178, Olf432, Olf610, Olf958, Olf960, Olf961, Olf913, and Olf1234) and four muscone-responsive ORs (Olf74, Olf235, Olf816, and Olf1440). We found that the human ortholog of Olf235 and Olf1440 responds to macrocyclic ketone and lactone musk odorants but not to polycyclic musk odorants or a macrocyclic diester musk odorant. This novel assay, called the Kentucky *in vivo* odorant ligand–receptor assay, should facilitate the *in vivo* identification of mouse ORs for a given odorant ligand of interest.

**Key words:** cell sorting; expression profiling; G-protein coupled receptor; odor detection; olfaction; sensory coding

## Introduction

Mammals can detect numerous odorants, and humans may be able to discriminate as many as 1 trillion different odorant mixtures (Bushdid et al., 2014). Remarkably, this enormous capacity for sensory detection and discrimination is not achieved at the expense of specificity. For example, mice have been shown to discriminate every pair of odorant enantiomers tested thus far (Laska and Shepherd, 2007). Underlying this discriminatory power is the expression of a single allele of one odorant receptor

(*OR*) or trace amine-associated receptor (*Taar*) gene in a mature olfactory sensory neuron (OSN; Mombaerts, 2004; Johnson et al., 2012). In addition, this singularity of expression allows receptor-specific axonal inputs to coalesce into glomeruli of the olfactory bulb (Mombaerts et al., 1996), an anatomical organization that contributes to maximizing the capacity for odor discrimination.

The discriminatory power of olfaction rivals that of the visual and auditory systems, but the patterns of receptor activation by odorant ligands remain elusive. Resolution of this problem has been hampered by the sheer number of ORs (~1100 intact *OR* genes in the mouse) and the retention of ORs in the endoplasmic reticulum of heterologous cells as opposed to native mature OSNs (McClintock et al., 1997; Gimelbrant et al., 2001; Lu et al., 2003; Dalton et al., 2013). Nevertheless, odorant ligands for >100 mammalian ORs have been identified. These data indicate that a typical OR is capable of being activated by several structurally related odorants, with some receptors more narrowly and others more broadly tuned (Malnic et al., 1999; Malnic, 2007; Grosmaître et al., 2009; Kato and Touhara, 2009; Saito et al., 2009; Nara et al., 2011). However, translating these data into an understanding of *in vivo* OR activation by odorants is difficult. To advance and refine these ideas further will require *in vivo* assays that go beyond the characterization of individual ORs (Zhao et al., 1998; Malnic et al., 1999; Kajiya et al., 2001; Oka et al., 2006; Shirasu et al.,

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2014) to assay instead simultaneously the entire repertoire of ORs so that sets of ORs activated by odorants may be identified.

Here, we demonstrate the ability of a novel *in vivo* assay in mice, supported by heterologous expression data, to identify ORs that respond to a given odorant ligand. With this approach, we identified or re-identified nine eugenol-responsive ORs, including all four previously identified eugenol-responsive ORs, and four muscone-responsive ORs, including the sole OR known previously to respond to muscone. Eugenol is a major component of the oil of several spice plants, especially the clove plant, *Syzygium aromaticum*. Muscone is a large macrocyclic odorant discovered in scent glands of male deer of the family *Moschidae* and has a long history of use in fragrances (Kraft and Fräter, 2001). The mouse ORs responsive to muscone successfully predicted a human OR that we find to be strongly responsive to macrocyclic ketone and macrocyclic lactone musk odorants.

## Materials and Methods

**Materials.** Sigma-Aldrich was the source of eugenol [4-allyl-2-methoxyphenol (catalog #35995)], mineral oil (catalog #M5904), Tonalid [Chemical Abstracts Service (CAS) 21145-77-7; catalog #W526401], Astrotone (ethylene brassylate; CAS 105-95-3; catalog #W354309), Exaltone (cyclopentadecanone; CAS 502-72-7; catalog #C111201), and isopropyl myristate (IPM; CAS 110-27-0; catalog #172472). Muscone (racemic mixture, 50% in IPM; CAS 10403-00-6) was purchased from Perfumer's Apprentice. Galaxolide (CAS 1222-05-5) was obtained from International Flavors & Fragrances. Mineral oil was the vehicle for eugenol and IPM for muscone.

**S100a5-tauGFP mouse strain.** Using 129/SvJ genomic DNA and the GeneAmp XL PCR kit (Life Technologies), DNA segments flanking the coding exons of *S100A5* were cloned after PCR amplification. The upstream arm was amplified using primers 5'-TATATGCGGCCGCTGC-CATGATGTGCAATGAATTCTTTGAGGG and 5'-TAGGTGGCGCGCCGATATGTACCCTGGACTAGGAGAGAGACAATCAC. The downstream arm was amplified using primers 5'-TATATGGCGCCG-CAGGACACTGGCAGCTCCTGATCCTG and 5'-CGCGCGTTAATTAAGTGGCTGCTCAACAACGTGGTAAGACCAGG. An upstream 4 kb NotI-AscI fragment and a downstream 3.8 kb AscI-PacI fragment were cloned into pBS-SKII that had been modified to accept these sites. An AscI fragment consisting of *IRES-tauGFP* and a *loxP-pgk-neomycin-loxP* sequence was subcloned into the AscI site. A clone of the correct orientation was identified by sequencing. The targeting vector was linearized with PmeI and electroporated into E14 embryonic stem cells that were cultured and selected using G418 as described previously (Mombaerts et al., 1996). Homologous recombination in embryonic stem cell colonies and in mutant mice was determined by PCR using primers 5'-AGCTTTGGCTCCCATCCACGGTG and 5'-TGTTGGACAGCCAGAGGGTCCCC. The *pgk-neomycin* fragment was removed by breeding heterozygotes to *EIIa-Cre* transgenic mice (Lakso et al., 1996), and the *Cre* recombinase transgene was subsequently removed by intercrossing S100a5-tauGFP mice hemizygous for the transgene. S100a5-tauGFP mice were in a mixed 129 × C57BL/6J background. This strain is publicly available from The Jackson Laboratory as stock number 6709, official strain name B6;129P2-S100a5<tm1Mom>/MomJ. Mice homozygous or heterozygous for the *S100a5-tauGFP* mutation (hereafter referred to as S100a5-tauGFP mice) have similar numbers of GFP-positive (GFP<sup>+</sup>) OSNs in the same mosaic pattern across the main olfactory epithelium,  $22 \pm 8/\text{mm}$  along the olfactory epithelium for homozygotes and  $32 \pm 5/\text{mm}$  for heterozygotes in 10- $\mu\text{m}$ -thick tissue sections ( $n = 5$ ;  $p = 0.0741$ ). Both genotypes were used interchangeably in the experiments described herein. As a precaution for hidden effects of genotype, heterozygous and homozygous mice were distributed equally between vehicle control and odorant treatments. All procedures with mice were done according to protocols approved by the Institutional Animal Care and Use Committees of Rockefeller University and the University of Kentucky.

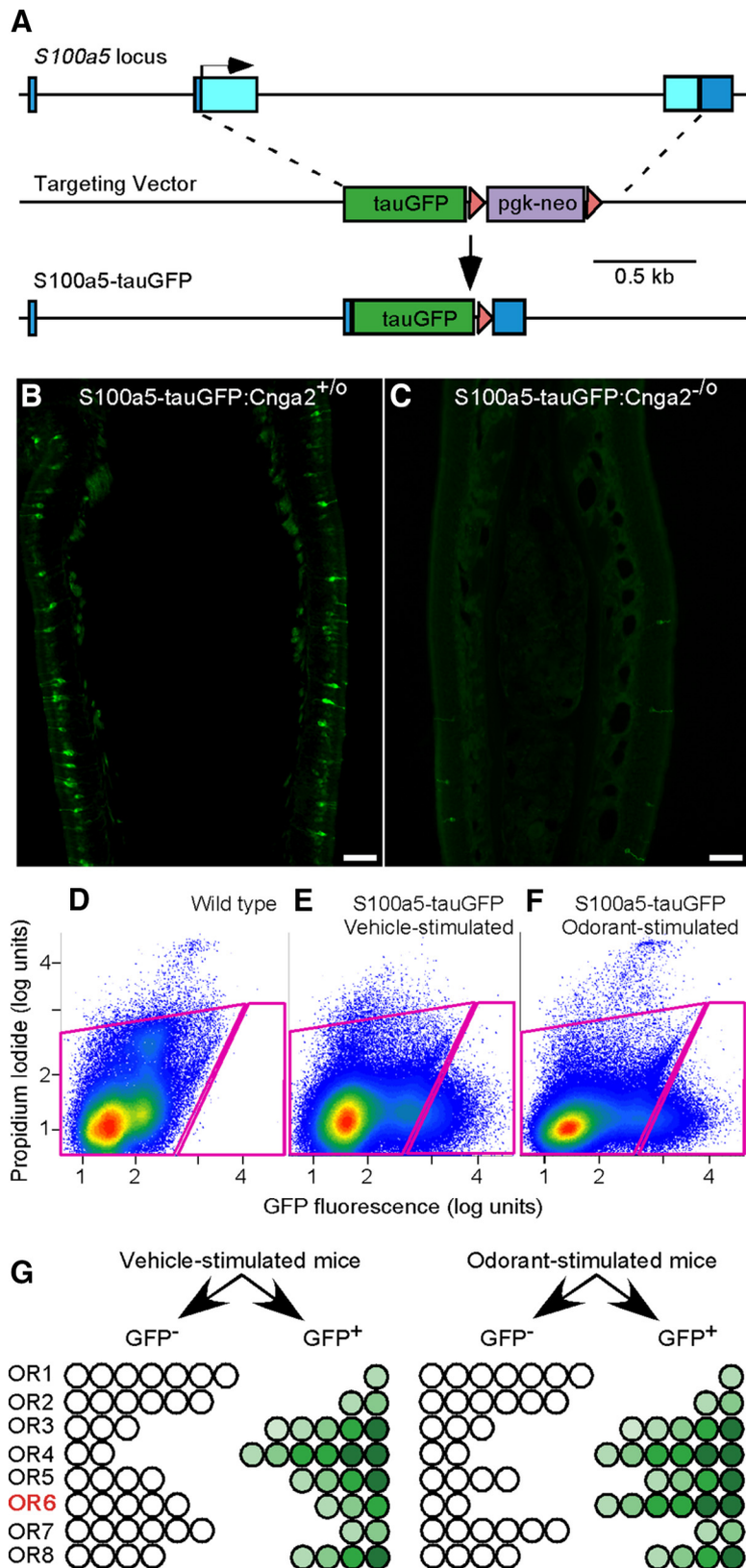
**Odor stimulation in vivo.** Exposing live mice to odor or vehicle was performed with groups of eight S100a5-tauGFP mice of both sexes aged

7–12 weeks. Each mouse was housed individually in chambers under a flow of 1.5 l/min filtered air as described previously (Fischl et al., 2014) for 40 h without food but with *ad libitum* distilled water. This chamber system minimizes ambient odor sufficiently such that the effect of odors on activity-dependent genes in OSNs, which cannot be detected when introduced in a standard cage environment, can be measured (Fischl et al., 2014). This protocol also minimizes the effects of previous odor exposure by allowing for protein turnover to reduce GFP levels. (Naris occlusion data show that GFP fluorescence is significantly reduced by 24 h and reaches a minimum by 48 h.) After a period of 26 h of filtered air in these chambers, intermittent odor exposure was initiated and performed for a period of 14 h. This 14 h period allows odor-stimulated GFP fluorescence to develop (preliminary experiments revealed that increased GFP expression can be detected as early as 6 h) but is too brief for GFP fluorescence to fade (25% reduction in fluorescence 24 h after naris occlusion). Electronically controlled valves were activated for 10 s to divert the flow of air through the headspace of 30 ml tubes containing 200  $\mu\text{l}$  of odorant (a 1:1 mix of odorant and vehicle) or vehicle alone. This switch was activated every 5 min until the completion of the experiment 14 h later. There were thus 168 odor exposures lasting 10 s each, or 28 min total exposure to the odorant ligand over the course of 14 h. Four mice were exposed to the odorant, whereas the other four were simultaneously exposed to vehicle. When mice from two litters were used, litters were distributed equally between the two treatment groups.

**RNA isolation and measurement.** At the completion of odor exposure, olfactory mucosae were dissected and cells dissociated in a procedure involving papain, trypsin, deoxyribonuclease, and low calcium saline as described previously (Yu et al., 2005; Sammeta et al., 2007). Cells from four identically treated mice were pooled, and fluorescence-activated cell sorting (FACS) was performed using an iCyt Synergy cell sorting system (Sony) to collect GFP<sup>+</sup> and GFP-negative (GFP<sup>-</sup>) cell samples in the University of Kentucky Flow Cytometry and Cell Sorting Facility. Total RNA was isolated using the Qiagen RNeasy Micro kit (catalog #74004), and samples were pooled until the GFP<sup>+</sup> sample reached at least 50 ng of RNA, which required 8–16 mice per replicate of each treatment. RNA quantity was measured using Affymetrix Mouse GeneChip 1.0 ST arrays in the University of Kentucky Microarray Facility. This microarray contains 1176 probe sets for mRNAs from ORs and Taars, including all of the 1098 intact OR genes and 14 Taar genes. Data were initially processed using Affymetrix GeneChip Command Console software to generate normalized quantities for each OR and Taar transcript cluster. Additional processing to generate GFP<sup>+</sup>/GFP<sup>-</sup> ratios was done in Microsoft Excel. No OR or Taar transcript clusters were consistently in the bottom 10% of signals or below 2 SDs of the mean signal for all transcript clusters, so all OR and Taar transcript clusters were used in these analyses. The microarray data are available in the Gene Expression Omnibus under the accession number GSE59336.

**Statistical analyses.** The stability of OR GFP<sup>+</sup>/GFP<sup>-</sup> ratios makes it possible to screen the entire set of ORs and Taars and identify activated receptors using a relatively small number of replications. We used a Bayesian hierarchical model to obtain normalized measures of odorant effect, accounting for four sources of variation: (1) basal receptor effect, (2) odorant effect, (3) nonspecific effect (change in both odorant and vehicle control), and (4) random measurement error. We find that 86% of the variation in the data are attributable to receptor identity and odorant, and only 14% are attributable to experimental noise. For each odorant effect, the posterior mean divided by the posterior SD provides a measure (Z-statistic) that is approximately normally distributed. Local false discovery rates (FDRs; Efron, 2008, 2010) were used to estimate the probability that each receptor is responsive to the odorant under the conservative assumption that the majority of the receptors are not responsive to the odorant. A 10% FDR was found to be a suitable level of risk for the identification of activated receptors.

**Luciferase assay for OR function in Hana3A cells.** Heterologous expression of ORs was performed as described previously (Saito et al., 2004, 2009; Zhuang and Matsunami, 2007; Dey et al., 2011; Li and Matsunami, 2011; Duan et al., 2012). Briefly, this assay is based on OR-driven increases in cAMP to activate CREs that result in expression of firefly luciferase. Plasmids (pCI; Promega) containing cDNAs encoding



**Figure 1.** The gene-targeted *S100a5*–*tauGFP* mouse strain allows for activity-dependent fluorescent marking of OSNs. **A**, The design of the targeted mutation of the mouse *S100a5* locus, replacing the coding exons with *tauGFP*. Boxes, Exons; light blue, coding sequence; dark blue, noncoding sequence; triangles, loxP sites; arrow, translation start site. **B**, Exposure for 4.5 s detects fluorescent OSNs in a 12  $\mu$ m coronal section at midseptum, showing the mosaic pattern and variable levels of GFP expression in a male mouse, 25 d postnatal, that is wild type for the *Cnga2* locus. **C**, Exposure for 49 s reveals a near absence of fluorescent OSNs in a *Cnga2*-deficient male mouse, 25 d postnatal. Scale bars, 100  $\mu$ m. **D**, FACS of dissociated cells from olfactory mucosae of a wild-type C57BL/6 mouse is used to set sorting gates (circumscribed with magenta lines) for subsequent capture of GFP<sup>+</sup> and

ORs (C57BL/6 sequences) tagged with the N-terminal 20 aa of rhodopsin were transiently transfected along with RTP1S1 (an OR chaperone), *Renilla* luciferase, CRE–firefly luciferase, and pRL–SV40 into Hana3A cells using Lipofectamine2000 (Life Technologies). Cells were plated into 96-well BioCoat plates (BD Biosciences). One day after transfection, the medium in each well was changed to CD293, a chemically defined medium from Life Technologies (catalog #11913  $\times$  019), for 30 min at 37°C. This medium was then exchanged for 25  $\mu$ l of CD293 containing either vehicle or the odorant ligand at concentrations ranging from  $10^{-7}$  to  $10^{-2}$  M and incubated for 4 h at 37°C and 5% CO<sub>2</sub>. Each concentration of odorant ligand was tested in triplicate wells. Luciferase activity was then measured using the Dual-Glo Luciferase Assay System (Promega) according to the protocols of the manufacturer. A Wallac Victor 1420 plate reader (PerkinElmer Life and Analytical Sciences) was used to measure luminescence. To control for transfection efficiency, the firefly luciferase luminescence of each well was divided by the *Renilla* luciferase activity of the well. Normalized luciferase activity was calculated as  $(L_N - L_{min}) / (L_{max} - L_{min})$ , where  $L_N$  is the corrected firefly luminescence,  $L_{max}$  is the maximum firefly luminescence on the plate, and  $L_{min}$  is the minimum firefly luminescence on the plate.

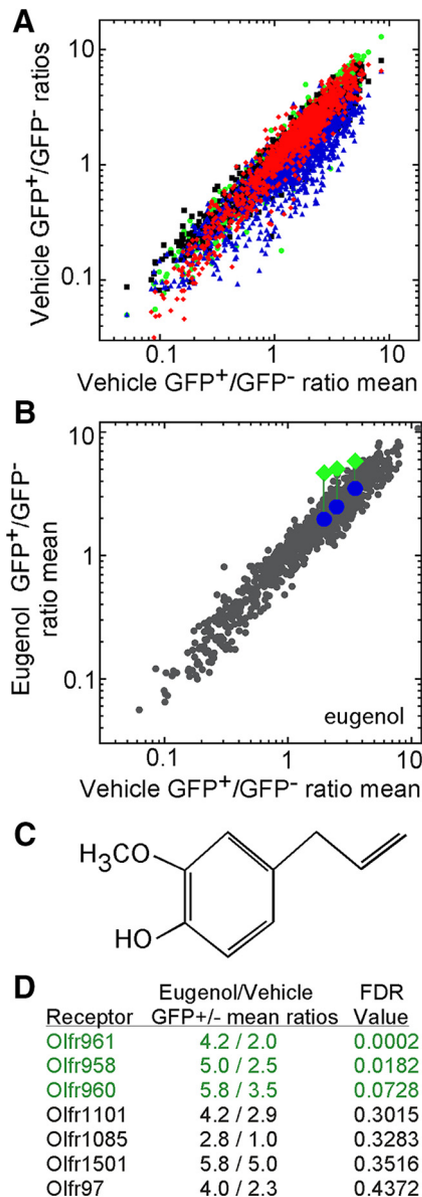
**Results**

**The *S100a5*–*tauGFP* mouse strain**

To screen for activation of ORs in native mouse OSNs *in vivo*, we devised an approach that is based on the activity-dependent expression of GFP from the *S100a5* locus and the expression of one OR gene per OSN (Fig. 1).

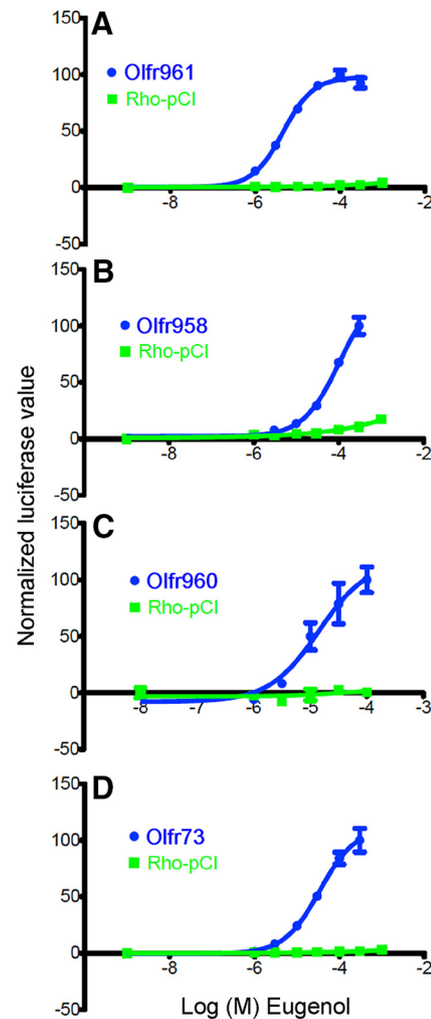
The *S100a5* gene encodes a 93 aa calcium- and zinc-binding protein (Schäfer et al., 2000; Streicher et al., 2010). The abundance of *S100a5* mRNA in OSNs is dependent on odor stimulation (Serizawa et al., 2006; Bennett et al., 2010; Fischl et al., 2014). The *S100a5* protein is located in

GFP<sup>−</sup> cells from *S100a5*–*tauGFP* mice. **E, F**, FACS capture of GFP<sup>+</sup> and GFP<sup>−</sup> cells from a set of four *S100a5*–*tauGFP* mice exposed to vehicle (**E**) and four mice exposed in parallel to muscone (**F**). **G**, In the *in vivo* assay, OSNs that exhibit GFP fluorescence above background are captured in the GFP<sup>+</sup> sample. The distribution of each OSN subtype in GFP<sup>+</sup> and GFP<sup>−</sup> FACS samples from *S100a5*–*tauGFP* mice exposed to an odorant ligand differs from that of *S100a5*–*tauGFP* mice exposed to vehicle only in those few OSN subtypes that express ORs responsive to the odorant ligand, as exemplified by OR6 in this diagram. Odorant-evoked activity in OR6-expressing OSNs results in an increase of the fraction of these OSNs with GFP fluorescence above background, thus causing a redistribution of these OSNs from the GFP<sup>−</sup> FACS sample into the GFP<sup>+</sup> FACS sample. Differing intensities of GFP fluorescence above background exist among OSNs but are not relevant in this assay, which is based on a binary grouping of fluorescence intensities (GFP<sup>−</sup> vs GFP<sup>+</sup>).



**Figure 2.** Eugenol exposure has significant effects on three ORs in *S100a5*- $\tau$ uGFP mice. **A**, The distribution of OR mRNAs between GFP<sup>+</sup> and GFP<sup>-</sup> samples is consistent across experiments. GFP<sup>+</sup>/GFP<sup>-</sup> ratios from mice exposed to vehicle (mineral oil) from four replications, distinguished by color (red, blue, green, and black), plotted against the mean GFP<sup>+</sup>/GFP<sup>-</sup> ratios of these four vehicle treatments. **B**, Mean OR mRNA GFP<sup>+</sup>/GFP<sup>-</sup> ratio values after exposure to eugenol plotted against the mean ratios after exposure to vehicle ( $n = 4$  groups of mice). Three ORs (Olf961, Olf958, and Olf960) show significant elevation (FDR < 0.10) after eugenol exposure. To illustrate the change, their GFP<sup>+</sup>/GFP<sup>-</sup> ratio values in this experiment (green diamonds) are compared with the GFP<sup>+</sup>/GFP<sup>-</sup> ratio coordinates in other experiments that did not use eugenol (blue circles). **C**, Chemical structure of eugenol. **D**, Mean GFP<sup>+</sup>/GFP<sup>-</sup> ratios and corresponding FDR probabilities for the seven ORs with FDR probabilities < 0.5.

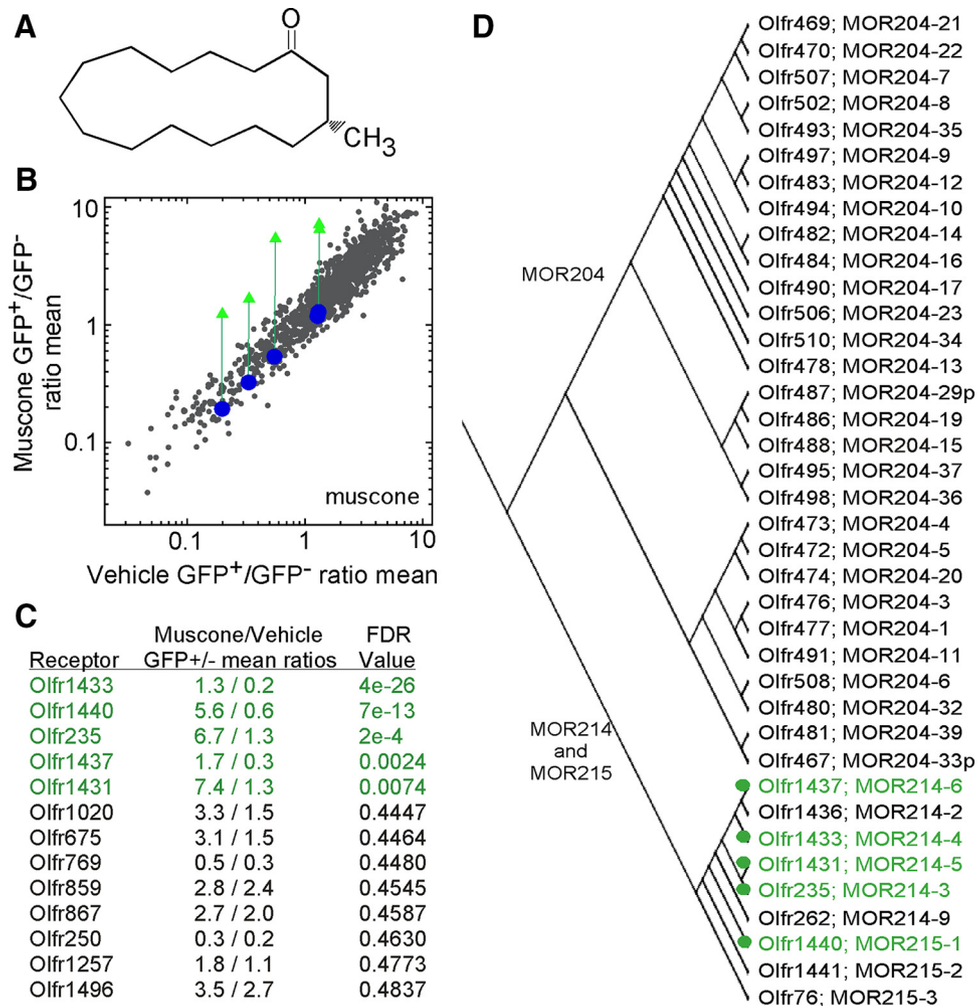
both the dendritic cilia and the axons of OSNs (Schäfer et al., 2000; Kuhlmann et al., 2014). The biological function of *S100a5* is not known. We generated a novel gene-targeted mouse strain with a  $\tau$ uGFP reporter replacing the two coding exons and intervening intron of the *S100a5* gene (Fig. 1A), causing a deletion of the *S100a5* gene. Thus far, we have not identified deficits in mice lacking *S100a5*: homozygous mice are indistinguishable from wild-type littermates. Consistent with the activity dependence of *S100a5* expression, GFP expression from the mutant locus is reflected in a mosaic pattern of fluorescent OSNs in the



**Figure 3.** Heterologous expression assay data for ORs responsive to eugenol. **A–C**, Eugenol concentration–response relationships for Olf961, Olf958, and Olf960, previously identified eugenol-responsive ORs that gave significant differences in the *in vivo* assay. **D**, Eugenol concentration–response relationship for Olf73, the first eugenol-responsive OR to be identified (Kajijiya et al., 2001).

main olfactory epithelium (Fig. 1B). GFP fluorescence is nearly undetectable in a *Cnga2*-deficient genetic background (Zheng et al., 2000), in which nearly all odorant-evoked electrical activity in OSNs is eliminated (Fig. 1C). Unlike the robust activity-dependent expression of *S100a5* in OSNs, neither *S100a5* mRNA in C57BL/6 mice nor GFP fluorescence in *S100a5*- $\tau$ uGFP mice could be detected in vomeronasal sensory neurons (data not shown).

We used FACS to collect separately GFP-fluorescent (GFP<sup>+</sup>) OSNs and nonfluorescent cells (GFP<sup>-</sup>) from olfactory mucosae of *S100a5*- $\tau$ uGFP mice that had been maintained in filtered air chambers for 40 h and exposed intermittently to odorant for the last 14 h of this period (Fig. 1D–F). Because the OSNs that respond to the odorant ligand tested represent only a small fraction of the total number of GFP<sup>+</sup> OSNs, similar numbers of GFP<sup>+</sup> OSNs are captured by FACS from mice exposed to vehicle (Fig. 1E) versus odorant (Fig. 1F). Because of the GFP-dependent capture of more OSNs expressing the few ORs that respond to the odorant ligand, the mRNAs encoding these responsive ORs are the only mRNAs that show an increase in abundance in the GFP<sup>+</sup> FACS sample relative to the GFP<sup>-</sup> FACS sample (Fig. 1G). Odorant-specific effects are confirmed by the absence of effect on



**Figure 4.** Muscone exposure has significant effects on five ORs in S100a5–tauGFP mice. **A**, Chemical structure of muscone. **B**, Mean OR mRNA GFP<sup>+</sup>/GFP<sup>−</sup> ratios after exposure to muscone plotted against the mean ratios after exposure to vehicle (*n* = 4 groups of mice). Five ORs (Olfr1440, Olfr1433, Olfr235, Olfr1431, and Olfr1437) show significant differences (FDR < 0.10) after muscone exposure (green triangles) compared with vehicle exposure. Blue circles reflect the GFP<sup>+</sup>/GFP<sup>−</sup> ratio coordinates of these five ORs in other experiments that did not use muscone. **C**, Mean GFP<sup>+</sup>/GFP<sup>−</sup> ratios and corresponding FDR probabilities for the 13 ORs with FDR probabilities < 0.5. **D**, A portion of the mouse OR phylogenetic tree depicting the relatedness of the members of the MOR214 and MOR215 mouse OR families that gave significant responses to muscone (green).

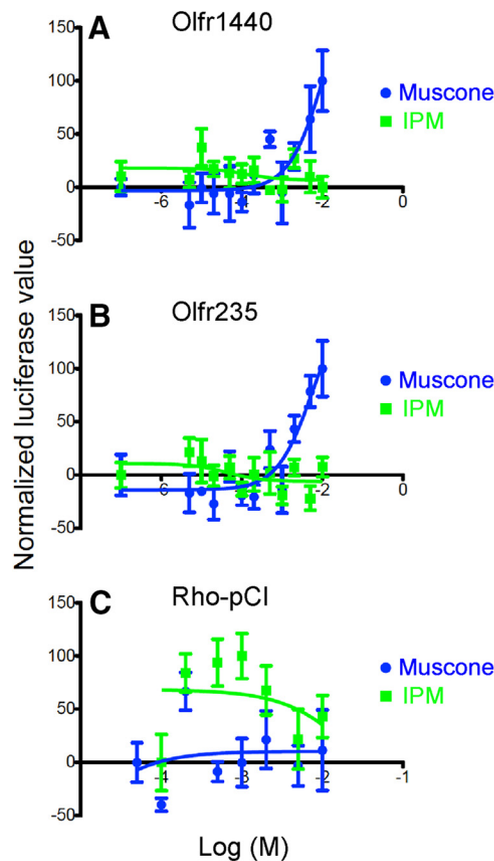
GFP<sup>+</sup>/GFP<sup>−</sup> ratios in samples from mice exposed to vehicle only. To identify responsive ORs in this assay, we measured their mRNAs via GeneChip microarray.

**Eugenol-responsive mouse ORs**

We find that, in the minimal odor conditions in which mice are exposed to vehicle only, each OR mRNA has a characteristic GFP<sup>+</sup>/GFP<sup>−</sup> ratio (Fig. 2A), which may result from the sum of the distinctive basal, constitutive activity of each OR (Imai et al., 2006; Reisert, 2010; Nakashima et al., 2013) and the effects of odors that may arise from the mouse itself or the chamber system. This characteristic GFP<sup>+</sup>/GFP<sup>−</sup> ratio is consistent across experiments: OR identity accounts for 86% of the variation in these data. This consistency is apparent when the mean of GFP<sup>+</sup>/GFP<sup>−</sup> ratios taken from four experiments is plotted against the four individual datasets of GFP<sup>+</sup>/GFP<sup>−</sup> ratios used to generate these mean values (Fig. 2A). When S100a5–tauGFP mice are exposed to an odorant ligand in the *in vivo* assay, only the mRNAs encoding the few responsive ORs will shift up from their characteristic positions near the line of equality, as is shown in Figure 2B for exposure to eugenol. The reason for this upward shift is that an odorant ligand acting as an agonist specifically

elevates GFP fluorescence only among the OSNs expressing an OR that is activated by this odorant, thereby shifting more OSNs expressing this OR from the GFP<sup>−</sup> FACS sample into the GFP<sup>+</sup> FACS sample. Because most mature OSNs express only one OR, statistically significant increases in the GFP<sup>+</sup>/GFP<sup>−</sup> ratio for OR mRNAs thereby allow for the simultaneous identification of the ORs that are activated by the odorant in question.

Eugenol (Fig. 2C) is known to activate at least four mouse ORs (Kajiya et al., 2001; Oka et al., 2006). After exposure to eugenol, the GFP<sup>+</sup>/GFP<sup>−</sup> ratios of three of the four known eugenol-responsive ORs were significantly elevated: Olfr961, Olfr958, and Olfr960 (Fig. 2B, D). When these three ORs were tested in the Hana3A heterologous expression assay (Saito et al., 2004, 2009; Zhuang and Matsunami, 2007; Dey et al., 2011; Li and Matsunami, 2011; Duan et al., 2012), cells separately expressing these three ORs responded robustly to eugenol (Fig. 3A–C). The fourth eugenol-responsive OR, Olfr73 (mouse olfactory receptor MOR174-9, mOR–EG), gave no increase in its GFP<sup>+</sup>/GFP<sup>−</sup> ratio in the *in vivo* assay, but heterologous cells expressing Olfr73 responded to eugenol (Fig. 3D).

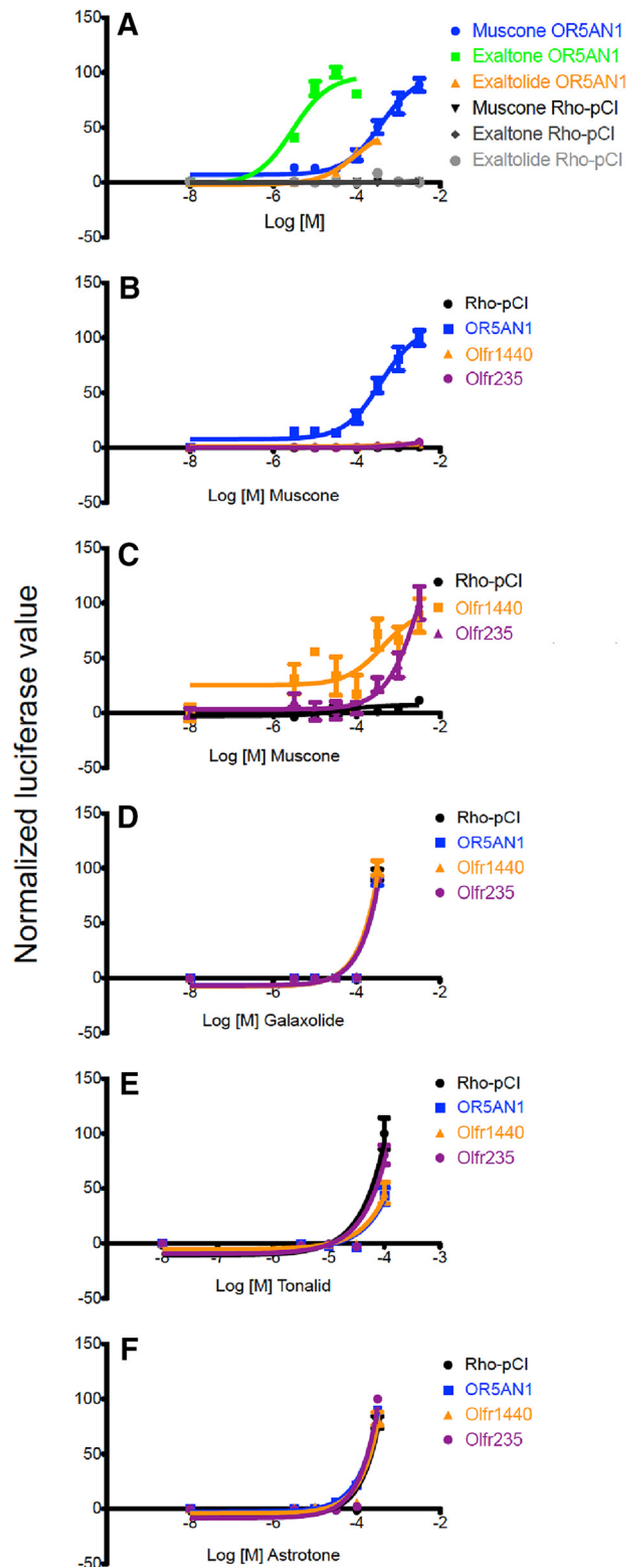


**Figure 5.** Confirmation of muscone responses from two mouse ORs by heterologous expression. *A*, Olfr1440. *B*, Olfr235. *C*, Rho-pCI, empty plasmid vector control. IPM used as vehicle.

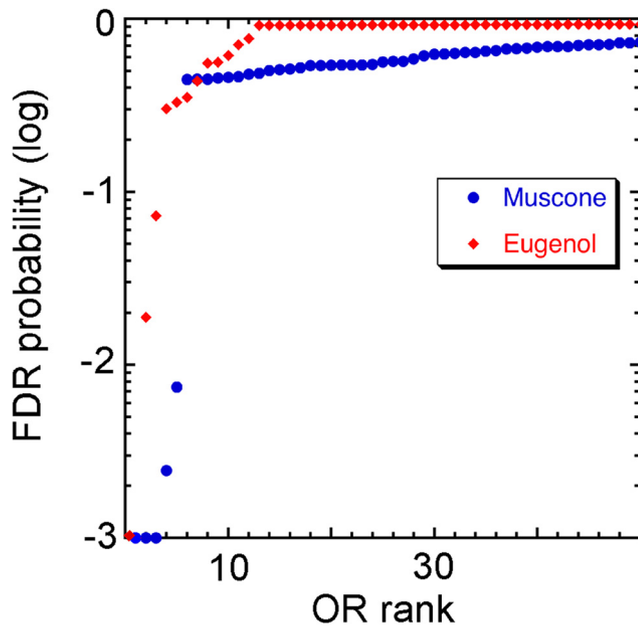
### Muscone-responsive mouse and human ORs

We next exposed S100a5–tauGFP mice to muscone, a macrocyclic ketone musk odorant (Fig. 4*A*). A single mouse OR, Olfr1440, has been shown recently to respond to muscone (Shirasu et al., 2014). We find that *in vivo* exposure to muscone has significant effects on five ORs, all from the MOR214 and MOR215 families (Fig. 4*B–D*). *In vitro* testing using the heterologous expression assay confirmed the ability of muscone to activate Olfr1440 and Olfr235 (Fig. 5). Limitations in the odorant concentrations that can be applied in this assay prevent these dose–response relationships from reaching saturation. Cells transfected with plasmids encoding Olfr1431, Olfr1437, and Olfr1433 did not respond to muscone (data not shown).

Based on the sequence similarity between the muscone-responsive mouse ORs identified by our *in vivo* assay, we determined that the most similar human OR is OR5AN1: its percentage identity to the five mouse OR proteins ranges from 68% (Olfr1440) to 81% (Olfr1433). Predicting that OR5AN1 is responsive to musks, we tested both macrocyclic and polycyclic musk odorants against OR5AN1 *in vitro*. Hana3A cells expressing OR5AN1 responded to three musks: muscone and Exaltone, which are macrocyclic ketone musks, and Exaltolide, a macrocyclic lactone musk (Fig. 6*A*). OR5AN1 gave robust responses to these macrocyclic musks but did not give responses different from pCI plasmid vector controls when exposed to two polycyclic musks, Galaxolide and Tonalid, or to Astrotone, a macrocyclic diester musk (Fig. 6*B–F*), confirming and extending previous observations (Shirasu et al., 2014).



**Figure 6.** Activation of muscone-responsive mouse and human ORs by certain macrocyclic musk odorants in heterologous expression assays. *A*, OR5AN1, a human OR, responded to the macrocyclic ketone musks muscone and Exaltone and to the macrocyclic lactone musk Exaltolide. *B*, The OR5AN1 response to muscone is stronger than responses of mouse ORs. *C*, Olfr1440 and Olfr235 responses to muscone. *D, E*, Polycyclic musk odors fail to evoke responses different from the plasmid vector negative control, Rho-pCI, in cells transfected with OR5AN1 or the mouse ORs responsive to muscone. Data for some ORs cannot be seen because of overlap of the curves. *F*, Astrotone, a macrocyclic diester musk odorant, also fails to evoke responses different from the plasmid vector negative control, Rho-pCI, in cells transfected with OR5AN1 or the mouse ORs responsive to muscone. Data for some ORs cannot be seen because of overlap of the curves.



**Figure 7.** Distribution of the 50 lowest FDR probabilities from the *in vivo* eugenol and muscone experiments illustrate the ability of the assay to separate odorant-responsive receptors from nonresponders. Receptors with FDR probabilities <0.001 were truncated at 0.001.

**Table 1.** ORs tested in the Hana3A cell heterologous expression assay

Eugenol			Muscone		
Olfr	MOR	FDR	Olfr	MOR	FDR
Olfr961*	MOR224-5	0.0002	Olfr1433	MOR214-4	0.0000
Olfr958*	MOR224-9	0.0188	Olfr1440*	MOR215-1	0.0000
Olfr960*	MOR188-5	0.0728	Olfr235*	MOR214-3	0.0000
Olfr1101	MOR179-3	0.3015	Olfr1437	MOR214-6	0.0024
Olfr1085	MOR191-1	0.3283	Olfr1431	MOR214-5	0.0074
Olfr1501	MOR212-3	0.3516	Olfr1020	MOR201-2	0.4447
Olfr97	MOR156-2	0.4372	Olfr769	MOR114-4	0.4480
Olfr1160	MOR173-1	0.9179	Olfr867	MOR143-2	0.4587
Olfr610*	MOR9-2	0.9197	Olfr250	MOR170-14	0.4630
Olfr1052	MOR172-1	0.9245	Olfr1257	MOR232-1	0.4773
Olfr1234*	MOR231-2	0.9832	Olfr1496	MOR127-1	0.4837
Olfr73*	MOR174-9	1.0000	Olfr74*	MOR174-4	0.5204
Olfr151	MOR171-2	1.0000	Olfr488	MOR204-15	0.5354
Olfr1031	MOR200-1	1.0000	Olfr816*	MOR113-1	0.6397
Olfr178*	MOR184-6	1.0000	Olfr394	MOR135-8	0.8547
Olfr922	MOR161-3	1.0000	Olfr1213	MOR233-7	1.0000
Olfr1274	MOR228-4	1.0000			
Olfr821	MOR109-1	1.0000			
Olfr811	MOR110-6	1.0000			
Olfr1090	MOR188-4	1.0000			
Olfr1325	MOR102-1	1.0000			
Olfr1496	MOR127-1	1.0000			
Olfr904	MOR167-3	1.0000			
Olfr591	MOR24-1	1.0000			
Olfr215	MOR119-2	1.0000			
Olfr432*	MOR123-2	1.0000			
Olfr1054	MOR188-2	1.0000			
Olfr1014	MOR213-5	1.0000			
Olfr360	MOR159-1	1.0000			
Olfr913*	MOR165-10	1.0000			

\*Positive responses.

**Weakly eugenol- or muscone-responsive mouse ORs**

In both the eugenol and muscone experiments, a few dozen ORs had elevated GFP<sup>+</sup>/GFP<sup>-</sup> ratios, but only eight exceeded our criterion for statistical significance. The *in vivo* assay thus produces strong distinctions between responsive ORs and unrespon-

sive ORs, with relatively few ORs showing intermediate responses (Fig. 7). ORs that failed to reach significance in the *in vivo* assay likely fall into one of three categories: (1) ORs that are insensitive or nonresponsive to the odorant (i.e., true negatives), (2) ORs that are capable of responding to the odorant but responded too weakly *in vivo* to achieve statistical significance, and (3) ORs that responded in some replications *in vivo* but too inconsistently to achieve significance. The reasons why ORs fall into the latter two categories are likely attributable to effects related to odorant concentrations. These ORs may require higher concentrations than they can experience in native OSNs during the *in vivo* assay, and the actual concentrations reached may differ across mice as a result of breathing patterns or other biological variables.

Using heterologous expression, we tested 36 ORs likely to fall into these latter two categories (Table 1). One, Olfr73, is known to respond to eugenol, and we confirmed this (Fig. 3D). Among the other 35 ORs, we identified only seven responsive ORs: five responsive to eugenol (Fig. 8A–E) and two responsive to muscone (Fig. 8F, G). All seven of these were previously orphan receptors. These findings argue that the majority of ORs lacking a significant response in the *in vivo* assay are not responsive to the odorant ligand and that these nonsignificant *in vivo* responses are true negative responses, as one would expect from the FDR calculations. In contrast, heterologous expression confirmed the responses of five of the eight significant ORs from the *in vivo* assay (Figs. 3A–C, 5A, B). These results demonstrate the ability of the *in vivo* assay to discriminate response from nonresponse.

**Discussion**

To identify chemosensory receptors that are activated by a given odorant ligand, we developed a novel *in vivo* assay in live, unanesthetized, freely breathing and freely behaving mice. We demonstrate that this assay re-identified ORs known to be responsive to eugenol and muscone. We identified ORs for muscone and eugenol that respond *in vivo* and *in vitro* to these odorants. The ORs that are responsive to eugenol *in vivo* in S100a5–tauGFP mice encompass three of the four previously identified eugenol-responsive ORs, all of which are related to each other and all of which have been confirmed to respond to eugenol in heterologous expression systems (Kajiya et al., 2001; Katada et al., 2003, 2005; Oka et al., 2006). We do not understand the lack of an *in vivo* response from the other eugenol-responsive OR, Olfr73, in our assay. We have not yet excluded any of the possible explanations, which range from the presence of an antagonist in the experiments to a polymorphism or differences in expression of the *Olfr73* gene in the S100a5–tauGFP strain.

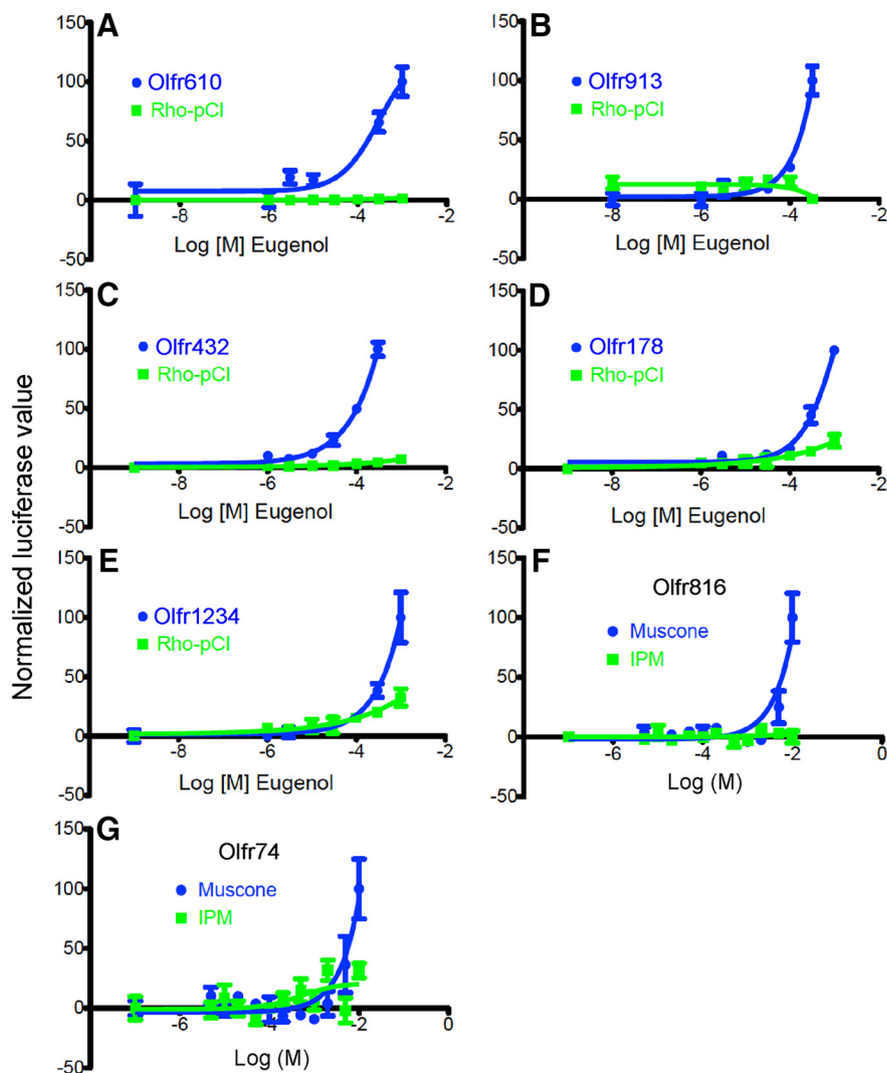
The ORs that are responsive to muscone *in vivo* encompass five related ORs, and two of these ORs gave muscone responses in heterologous expression assays: Olfr235 and Olfr1440. Our data confirm and extend previous evidence that Olfr1440 is a muscone-responsive receptor and also help explain the activation of a small number of olfactory bulb glomeruli by muscone (Shirasu et al., 2014). In the absence of positive responses in a heterologous expression assay, the other three ORs, Olfr1431, Olfr1433, and Olfr1437, must be considered only as candidate muscone-responsive ORs. The absence of responses does not prove they are unable to respond to muscone, because confounding factors may have prevented them from responding in heterologous cells. Confounding factors, especially involving membrane trafficking, affect a fraction of ORs expressed in heterologous expression systems, even when OR chaperones and sequence modifications are used to enhance delivery of ORs to the plasma membrane (Peterlin et al., 2014). Only if proof of function of these ORs in the

heterologous expression assay were obtained, for example by demonstrating responses to a different odorant, would we have evidence sufficient to conclude that these ORs are unresponsive to muscone.

The subsets of ORs we identified as responsive to either muscone or eugenol *in vivo* each consist of related ORs. Some odorants, including eugenol and muscone, tend to activate clusters of neighboring glomeruli, which are often innervated by OSNs expressing related ORs (Malnic et al., 1999; Oka et al., 2006; Johnson and Leon, 2007; Matsumoto et al., 2010; Nara et al., 2011; Falasconi et al., 2012; Shirasu et al., 2014). However, because these data are based on just two odorants, they are insufficient to conclude that odorants typically activate sets of related ORs. Future experiments will reveal if some odorants instead activate unrelated ORs.

The sequence similarity of the muscone-responsive ORs identified by the *in vivo* assay predicted an orthologous human OR that is highly sensitive to muscone and some other macrocyclic musk odorants, confirming recent results obtained independently (Shirasu et al., 2014). Musks are a particularly interesting class of odorants because deficits in detecting these odorants are among the most common anosmias and hyposmias in humans (Whissell-Buechy and Amore, 1973; Gilbert and Kemp, 1996). These sensory deficits are typically not generalized to all musks but rather to subclasses of musks (Triller et al., 2008), consistent with the activation of OR5AN1 by macrocyclic ketone and macrocyclic lactone musks but not polycyclic musks. We speculate that the sets of mouse ORs responsive to other classes of musk odorants, such as polycyclic musk odorants, intersects poorly with those responding to muscone.

Odorant concentration is often positively correlated with the number of OSNs or glomeruli activated (Malnic et al., 1999; Ma and Shepherd, 2000). This correlation predicts that a single moderate concentration of an odorant should show strong activation of a few ORs and weaker activation of other, less sensitive, ORs. Our data are consistent with this prediction. Whether our approach identifies all ORs capable of being activated by any given odorant is difficult to determine, but this completeness can be estimated by comparing the number of responsive ORs to the number of responsive glomeruli per olfactory bulb. We identified four muscone-responsive ORs, which is very similar to the one to three dorsomedial glomeruli and the occasional few ventral glomeruli per olfactory bulb reported to respond to muscone (Shirasu et al., 2014). The nine eugenol-responsive ORs we detected are significantly fewer than the 37–56 eugenol-responsive glomeruli per olfactory bulb reported recently (Shirasu et al., 2014), but the concentration of eugenol was twofold lower in our experiments. The S100a5–tauGFP assay is able to identify many of the ORs capable of responding to an odorant, but it may not identify some ORs, especially weakly responding ORs that are



**Figure 8.** Heterologous expression assay data for additional ORs responsive to eugenol or muscone. **A–E**, Eugenol concentration–response relationships for five eugenol-responsive ORs that were previously orphan receptors. **F, G**, Muscone concentration–response relationships for two muscone-responsive ORs that were previously orphan receptors. Rho–pCl, Expression plasmid used as a negative control. IPM used as vehicle.

capable of evoking detectable responses in glomeruli because of the amplification effect of axonal convergence, or in heterologous expression assays in which higher concentrations can be applied. When odorants are applied as volatiles in air to live mice, the odorant concentration reaching dendritic cilia of OSNs is not known and is potentially variable because of factors differentially affecting the access of odorants to ORs in the dendritic cilia. In other respects, however, the *in vivo* assay is resistant to factors that could limit the breadth of the set of responsive ORs detected. The assay gives stable and consistent outcomes in the absence of odor exposure, no OR mRNA is found solely in the GFP<sup>+</sup> sample, and even ORs with high GFP<sup>+</sup>/GFP<sup>−</sup> ratios after exposure to vehicle are capable of further elevation in their ratio during exposure to an odorant. In fact, ORs with significant responses to the few odorants tested thus far already span control GFP<sup>+</sup>/GFP<sup>−</sup> ratios of 0.2–3.9, a range that includes 89% of ORs (T.S.M. and W.B.T., unpublished data). In general, the S100a5–tauGFP assay successfully identifies sets of ORs responsive to an odorant ligand and is resistant to false positives.

In summary, we have developed a novel *in vivo* assay to identify ORs responding to a given odorant ligand in awake, unanesthetized mice.

thetized, freely breathing and freely behaving mice. Mimicking a mouse encountering an ambient odor in its environment, this assay begins with an odorant ligand and returns the identities of ORs that responded. Our assay covers the entire OR repertoire simultaneously in a multiplex manner. Therefore, it has a higher capacity than *ex vivo* approaches that also rely on the expression of one OR gene per OSN but are based on individual odorant-responsive OSNs (Malnic et al., 1999; Touhara et al., 1999; Kajiya et al., 2001; Nara et al., 2011). Another recently developed *in vivo* approach differs in that it targets single OSNs and relies on anesthetized mice (Shirasu et al., 2014). Our assay should in principle detect responses from any OR. It is stable and robustly distinguishes a response from the lack of a response. To reduce variability and further increase sensitivity, we are backcrossing the S100a5–tauGFP mice to C57BL/6 mice to place the reporter gene into the inbred genetic background that is the basis for most expression profiling methods. This backcrossing will also facilitate testing of whether other mRNA quantification methods, such as NanoString (Khan et al., 2011) or RNA sequencing, may improve performance. Given the place of origin of our new assay, we have named it the Kentucky *in vivo* odorant ligand–receptor assay.

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