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SPECIALTY SECTION This article was submitted to Chemical Ecology, a section of the journal Frontiers in Ecology and Evolution

RECEIVED 28 June 2022 ACCEPTED 15 August 2022 PUBLISHED 02 September 2022

CITATION

Mahadevan VP, Lavista-Llanos S, Knaden M and Hansson BS (2022) No functional contribution of the gustatory receptor, Gr64b, co-expressed in olfactory sensory neurons of *Drosophila melanogaster*. *Front. Ecol. Evol.* 10:980351. doi: 10.3389/fevo.2022.980351

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No functional contribution of the gustatory receptor, Gr64b, co-expressed in olfactory sensory neurons of *Drosophila melanogaster*

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Chemosensation is essential for the survival of insects. Activities like searching for food, mating, and oviposition in the fruit fly, Drosophila melanogaster are to a great extent governed by chemical cues detected via olfaction and gustation. This chemical information is conveyed to higher brain centers via populations of diverse olfactory sensory neurons (OSNs) and gustatory sensory neurons (GSNs) expressing olfactory receptors (ORs) and gustatory receptors (GRs), respectively. ORs are exclusively expressed in the antenna and in the maxillary palps, while GRs are widely expressed in the labellum, tarsi, genitalia etc. Interestingly, 14 GRs were previously reported to be expressed in the antenna of *D. melanogaster*. However, the spatial expression pattern for all GRs and their functional role are still unclear. Recent data challenge the dogma that single OSNs express a single OR. In the present study, we studied the expression of 12 previously reported GRs among sensory structures on the fly antenna using the Gal4-UAS binary expression system. We observed antennal expression of nine out of the 12 reported. Out of these nine, consistent expression was only apparent for Gr64b, and we reconfirmed its presence in OSNs innervating three glomeruli in the antennal lobe. These glomeruli are known to be innervated by ab5A, ab5B and ab8A OSNs, respectively. Next, we generated double labeling crosses with Gr64b and observed co-expression of Gr64b with Or47a, which is expressed in the ab5B neuron. To elucidate the functional role of Gr64b coexpressed with Or47a, we challenged Or47a-expressing OSNs in wild type and Gr64b^{-/-} mutant flies with odor stimulation using the single sensillum recording technique in two satiation states (fed and starved). Notably, we did not observe any significant odor sensitivity or specificity changes in Gr64b mutants as compared to wild type flies. Taken together, our results reveal coexpression of GRs with ORs in olfactory sensory neurons, while the functional contribution of the GR in this context remains obscure.

KEYWORDS

olfaction, gustation, single sensillum recording (SSR), Gr64b, Or47a, coexpression, olfactory receptors (ORs), gustatory receptors (GRs)

Introduction

Perception of multiple chemical cues is essential for the survival of insects. The fruit fly, Drosophila melanogaster, utilizes such cues to find, e.g., food sources (Stensmyr et al., 2003), suitable mates (Dweck et al., 2015) and oviposition substrates (Dweck et al., 2013) to name a few. The perception of chemical cues is carried out with the help of receptors expressed in peripheral sensory organs of the fly. In D. melanogaster, three gene families code for chemosensory receptors: olfactory receptors (ORs) (Clyne et al., 1999; Vosshall et al., 1999), gustatory receptors (GRs) (Montell, 2009)and ionotropic receptors (IRs) (Benton et al., 2009). Canonically, ORs are expressed in the antenna and maxillary palps, while GRs are expressed in multiple organs such as labellum, tarsi, wings, reproductive organs etc. (Depetris-Chauvin et al., 2015; Montell, 2021). Typically, ORs form a complex with the ubiquitous olfactory receptor co-receptor (Orco) and are responsible for the detection of a unique panel of odorants (Hallem and Carlson, 2006). On the other hand, GRs form heteromeric complexes and their ligand specificity changes depending upon the combination of GRs co-expressed (Jiao et al., 2008). Multiple studies have identified GRs that are required for tasting sweet and bitter compounds and describe diverse functional roles, such as inhibition of feeding and oviposition, inhibition of male to male courtship, etc. (Jiao et al., 2008; Moon et al., 2009; Watanabe et al., 2011; Lee et al., 2012; Freeman and Dahanukar, 2015; Fujii et al., 2015; Dweck and Carlson, 2020; Vernier et al., 2022). The first hint about non-canonical expression of GRs in the antenna of D. melanogaster came in the late 2000s, when two independent studies demonstrated that the perception of CO2 depends on Gr21a and Gr63a co-expressed in the ab1C neuron in the D. melanogaster antenna (Jones et al., 2007; Kwon et al., 2007). Follow-up studies using RNA-seq analysis and binary expression systems demonstrated expression of up to 14 GRs in the antenna, including GRs previously known to be involved in sugar and bitter taste perception (Thorne and Amrein, 2008; Ni et al., 2013; Menuz et al., 2014; Fujii et al., 2015). Based on these results, it was hypothesized that GRs might be co-expressed together with ORs, forming a heteromer and function in detection of novel ligands different from the native tuning profile of the corresponding OR (Fujii et al., 2015).

However, apart from the deorphanization of Gr28bD expressed in the arista as being involved in avoidance of rapid increases in temperature (Ni et al., 2013), scant information is available regarding the functional significance of GR-OR co-occurrence in the same OSN. In other combinations, it has been reported that detection of sour taste via tarsal gustatory sensory neurons (GSNs) was mediated via co-expressed IRs (Chen and Amrein, 2017). Further, co-expression of broadly tuned IRs such as Ir25a and Ir76b in the labeller GSNs has also been reported. A functional role of such overlap is required for detection of tastants such as amino acids, carboxylic acids etc. via GSNs present in labeller sensilla (Aryal and Lee, 2021; Stanley et al., 2021; Aryal et al., 2022; Shrestha et al., 2022). Two recent studies also demonstrated overlapping expression of multiple gene families, with emphasis on IRs and ORs in the D. melanogaster antenna (Task et al., 2022) and in the yellow fever mosquito, Aedes aegypti (Younger et al., 2022). These studies are particularly important in demonstrating a functional role of IRs and ORs co-expressed in the OSNs. There are at least two possible explanations for such an overlap. First, such co-expression of multiple receptors might lead to heteromeric receptors that could ultimately accommodate novel ligands beyond the usual spectrum of each independent receptor. Secondly, such overlap could help in making the system redundant. If, by any natural incident, one receptor or receptor family fails then the other receptor could still maintain a partial functioning of the neurons. However, both of the studies mentioned above focused on the overlap of ORs and IRs with little focus on the involvement of the GR family.

In the present study, we focused on the co-expression of ORs and GRs in OSNs in the antenna of *D. melanogaster*. To generate a spatial expression map, we attempted to establish expression patterns of 12 GRs previously shown to be expressed in the antenna. Next we focused our attention on one specific sugardetecting GR, Gr64b, due to its general expression pattern in the antenna and due to its co-expression with the well-characterized Or47a (also co-expressing Or33b) in the ab5B OSN. Using single sensillum recording technique, we investigated possible modulatory effects on odor sensitivity and specificity of the ab5B OSN in the absence of Gr64b. However, we did not observe any significant changes neither in odor sensitivity nor in specificity.

Fly line identity	Genotype	Bloomington ID	
Gr21a	Gr21a-gal4	Originally from	
		Vosshall lab	
Gr22e	Gr22e-gal4	BL57608	
Gr28bC	Gr28bC-gal4	BL57618	
Gr28bD	Gr28bD-gal4	BL57620	
Gr43a	Gr43a-gal4	BL57636	
Gr61a	Gr61a-gal4	Originally from Baker lab (ID1190807)	
Gr64a	Gr64a-gal4	BL 57661	
Gr64b	Gr64b.mGFP (direct fusion protein)	BL52630	
Gr64c	Gr64c-gal4	BL57663	
Gr64f	Gr64f-gal4	Originally from Baker lab (ID1191452)	
Gr93a	Gr93a-gal4	BL57679	
mGFP	UAS-mGFP	BL5137	
mRFP	UAS-mRFP	BL27398	
Octuple GR mutant	Δ Gr5a, Gr64a-f, Gr61a null mutant	Originally from Amrein lab	
Or82a	Or82a-gal4	BL 23126	
Or47a	Or47a-gal4	BL 9982	

TABLE 1 A list of all fly lines used during the experiments.

Materials and methods

Fly lines

Flies were reared at 25°C on standard cornmeal agar medium under 12 h light: 12 h dark photoperiod cycle. A mix of 7-15 day old males and females were used for immunohistochemistry and single sensillum recording (SSR) experiments. For starvation experiments, flies were starved for 20 h with access to water *ad libitum* before being used for electrophysiology. Transgenic fly lines were obtained from the Bloomington fly stock center with identities as listed in **Table 1**.

Chemical stimuli

All chemicals were purchased from with the highest purity (> 97%). Geranyl acetate (CAS: 105-87-3) and pentyl acetate (CAS: 628-63-7) were used as diagnostic odors for ab5A and ab5B neurons, respectively. Both odors were serially diluted in hexane. A panel of 32 odors of ecological significance and representing diverse chemical groups were used to determine any changes in the odor tuning pattern of ab5A and B OSNs. A list of these 32 odors with their suppliers and catalog numbers is available in **Table 2**.

TABLE 2	List of chemicals used for identifying odor-tuning properti	ies
of neuror	s innervating ab5 sensillum class.	

	Common name	CAS	Supplier: Catalogue number
1	Hexane		Stock
2	Ethyl acetate	141-78-6	SA: 270989
3	Ethyl lactate	97-64-3	SA: W244015
4	CO ₂	Aspiration	
5	Methyl salicylate	119-36-8	SA: M6752
6	Methyl acetate	79-20-9	F: 45999
7	Ethyl-3- hydroxybutyrate	5405-41-4	AO: 118540250
8	ethyl hexanoate	123-66-0	SA: 148962
9	2-heptanone	110-43-0	SA: W254401
10	geranyl acetate	105-87-3	Stock (originally SA: 173495)
11	pentyl acetate	628-63-7	SA: W504009
12	1-octen-3-ol	3391-86-4	SA: W280518
13	ethyl benzoate	93-89-0	SA: E12907
14	Ethyl crotonate	623-70-1	SA: 16794-0
15	Acetoin	513-86-0	SA: W200808
16	2 phenyalcohol	60-12-8	SA: 77861
17	benzyl butyrate	103-37-7	Stock (originally A.A: B24241)
18	2-butanone	78-93-3	SA: W217018
19	ethyl butanoate	105-54-4	SA: E15701
20	isopropyl benzoate	939-48-0	Stock (originally ABCR: AB137185)
21	methyl benzoate	93-58-3	SA: 18344
22	6-methyl-5-hepten- 2-one	110-93-0	SA: W270733
23	Hexyl acetate	142-92-7	SA: 10815-4
24	Isopentyl propionate	105-68-0	SA: W208205
25	2-nonanol	628-99-9	SA: N30307
26	2-nonanone	821-55-6	SA: N30307
27	Isopentyl acetate	123-92-2	SA: 30696-7
28	4-methylphenol	106-44-5	SA: 61030
29	Acetophenone	98-86-2	Stock (originally SA: 42163)
30	methyl hexanoate	106-70-7	SA: W270806
31	propyl acetate	109-60-4	SA: 133108
32	nonanal	124-19-6	SA: W278220

Sigma-Aldrich (Steinheim, Germany): SA, Acros Organics B.V.B.A.: AO, Fluka: F, Alfa Aesar: A.A, ABCR GmbH: ABCR and institute stock: Stock.

Electrophysiology: Single sensillum recordings

Single sensillum recordings were performed in order to establish the response profile of OSNs present in individual sensilla on the fly antenna. A 7-15-day old fly was immobilized in a 200 μ l pipette tip and fixed on a glass side with laboratory wax. The funiculus (third antennal segment)

TABLE 3	A list of 14	GRs identified	in the ante	enna based	on previous
studies	(Dunipace et	al., 2001; Men	uz et al., 20)14) .	

No.	Gustatory receptor (GR)	RPKM (average)
1	Gr10a	85.6
2	Gr64f	35.8
3	Gr63a	29.7
4	Gr21a	14.5
5	Gr64c	13.6
6	Gr64b	8.3
7	Gr93a	5.1
8	Gr61a	2.2
9	Gr43a	2.1
10	Gr66a	1.9
11	Gr28b	1.5
12	Gr64a	1.5
13	Gr64d	Not determined
14	Gr22e	Not determined

was fixed in such a position that either the medial or posterior side faced the observer. Extracellular recordings were performed using electrochemically (3M KOH) sharpened tungsten electrodes by inserting a ground electrode in the eye and a recording electrode into the base of a sensillum using micromanipulators (Luigs and Nuemann SM-10). Sensilla were visualized with 1000x magnification using a binocular microscope (Olympus BX51W1). Signals were amplified (Syntech Uni-versal AC/DC Probe)1, sampled (96,000/s) and filtered (3 kHz High-300 Hz low, 50/60 Hz suppression) using a USB-IDAC. Neuronal activity was recorded using AutoSpike software (v3.7) for 3 s pre and 10 s post stimulus. The odor stimulus was delivered for 500 ms using a Pasteur pipette and was added to filtered and rehumidified air being constantly delivered to the fly antenna at 0.6 L/min through an 8 mm id stainless steel tube ending 5 mm from the antenna. Responses were established by calculating the change in spike frequency (spikes/s) 1 s pre and post onset of the stimulus. Stimulus cartridges were prepared by pipetting 10 µl of the desired compound dissolved in hexane onto a filter paper (Rotilabo-round filters, type 601A, Carl Roth GmbH, Germany) with a diameter of 10 mm, which was then placed into a Pasteur pipette. Not more than two sensilla were recorded from each fly and odor stimuli were used for a maximum for two times. An inter-stimulus interval of at least 45 s was maintained and stimulation series always started with the lowest concentration and were then applied in an increasing order of concentrations.

Immunohistochemistry

For whole-mount immunostaining, 7-15 day flies of mixed sexes were anesthetized on ice, and their antennae were dissected into cold Drosophila ringer solution with 0.1% Triton-X. The antennal mix consisted of specimens with only the third segment had been excised and those where both the second and third segments remaining. Dissected antennae (hereafter named "mix") were fixed in 4% paraformaldehyde (with 0.1% Triton-X) for 1 h on ice in a shaker. The mix was washed at least three times with 500 μ l PT (1x PBS + 50 µl Triton) for 15 min each at room temperature. 500 µl Blocking solution (PT + 5% NGS) was added to the mix and incubated for 1 h. Primary antibodies were diluted in 500 µl of fresh blocking solution and added to the mix followed by 48 hour incubation at 4°C. After the incubation, the mix was washed at least 3 times with PT, followed by addition of 500 µl blocking solution with 1hour incubation. Secondary antibodies were diluted in 500 μl of fresh blocking solution and added to the mix followed by 48-h incubation at 4°C. Lastly, the mix was washed at least 3 times with PT and mounted in Vectashield (Vector, Burlingame, CA, United States). Primary antibodies used were: mouse-anti-GFP (1:500), rabbit-anti-RFP (1:100) and rabbitanti-orco (1:500), while secondary antibodies were: goat-antimouse (1:250) and goat-anti-rabbit (1:250). All antibodies used were purchased from Invitrogen (Invitrogen, Carlsbad, CA, United States).

Statistical analysis

SSR traces were analyzed using AutoSpike32 software 3.7 version (Syntech, NL). Changes in action potential (spike count) were calculated by subtracting the number of spikes one sec before (spontaneous activity) from those elicited during one second after the onset of the stimulus. Spike changes for each concentration between wild type and GR mutant fly line were compared using unpaired, non-parametric t-tests with Mann Whitney test. Graphs were generated and statistical tests were performed using GraphPad Prism 9.1.1. Figures were constructed with Adobe Illustrator CS5 and Adobe Photoshop (Adobe system Inc.).

Results

Spatial expression map of gustatory receptors in the antenna

Earlier studies have demonstrated expression of gustatory receptors (GRs) in the antenna of *D. melanogaster* (Table 3). Here, we wanted to establish the spatial expression map

¹ www.syntech.nl

of all previously reported GRs in the antenna. We took advantage of the binary expression system in D. melanogaster, combined 12 GR specific Gal4 drivers independently with membrane expressing GFP (UAS-mGFP) and checked for the spatial expression pattern of each of the GRs in the antenna (Figure 1A). We observed expression of two GRs (Gr93a and Gr28bC) in the second segment (Figures 1 H,J), five (Gr21a, Gr64c, Gr64b, Gr64f, and Gr43a) in the third segment (Figures 1B-G), Gr28bD in the arista (Figure 1I) and Gr64e in both the second and third segments (Figure 1F). Using this data, we were able to generate a spatial expression map of these nine GRs in the antenna (Figure 1K). We proceeded to count the number of cells labeled by each GR driver (Figure 1L). Among these, we observed only a single instance positive for Gr64c out of more than 20 whole mount antenna preparations screened. Although several GRs were expressed in the antenna, only Gr64b displayed a consistent expression pattern throughout multiple replicates with a mean number of 20 Gr64b positive cells (Figure 1L). Therefore, we decided to focus our physiological investigations on Gr64b and proceeded to elucidate its role in olfaction during the next set of experiments.

Gr64b is co-expressed with Or47a in ab5B neurons

To establish if Gr64b contributes to the olfactory coding in the cells where it is expressed, we first identified which physiological type of neurons express Gr64b. It is well known that Gr21a along with Gr63a, responsible for detecting CO₂, is expressed in the ab1C neuron and that this neuron does not express the olfactory co-receptor Orco (Jones et al., 2007). We used this information as an internal positive control to verify the function of antibodies used (Supplementary Figure 1). Next, we checked the co-expression pattern of Gr64b along with Orco by using a direct GFP fusion line (Gr64b.mGFP, Table 1) and an anti-Orco antibody. Although we observed co-expression of Gr64b in a few Orco positive cells, a significant population of Gr64b positive cells were Orco negative (Figure 2A). However, the cells positive for both Orco and Gr64b point out toward expression of Gr64b in OSNs. Next, we checked for the innervation pattern of axons of Gr64b-expressing OSNs in the antennal lobe. We could reconfirm that Gr64b positive neurons innervate VA6, DM3 and VM2 glomeruli in the antennal lobe along with canonical innervation in the suboesophageal ganglion as previously reported by Fujii et al., 2015 (Figure 2B). These three glomeruli, VA6, DM3 and VM2 have previously been shown to be innervated by OSNs expressing Or82a, Or47a (co-expressed with Or33b) and Or43b, respectively. Therefore, we hypothesized that the small subpopulation of Gr64b and Orco co-expressing OSNs should also express at least one of the above-mentioned olfactory receptors. To start testing this hypothesis, we generated double cross fly lines with genotypes Or82a-Gal4: UAS-mRFP and Gr64b.mGFP. We did not observe any colocalization of Gr64b and Or82a (**Figure 2C**) but Gr64b positive neurons were located in the immediate vicinity of Or82a positive OSNs with adjacent dendritic projections (**Figure 2D**). The sensillum housing OSNs expressing Or82a and Or47a is named ab5. Next, we checked for the double cross with a genotype Or47a-Gal4: UAS-mRFP and Gr64b.mGFP and could indeed observe colocalization (**Figure 2E**). Notably, only a small subpopulation of Gr64b-expressing cells also express Or47a (**Figure 2E** arrows) as many neuronal somata expressed only Gr64b (**Figure 2E** hand points). Taken together, we show that all Or47a-expressing neurons co-express Gr64b but the converse is not true (**Figure 3E,F**).

Functional implications of Gr64b on olfactory coding in ab5B olfactory sensory neurons

In the previous experiment, we showed that Gr64b often is co-expressed in Or47a expressing neurons. In D. melanogaster, Or47a expression corresponds to the B neuron innervating the ab5 sensillum class. We used single sensillum recording (SSR) technique to investigate if Gr64b contributes functionally to the odor sensitivity threshold and odor specificity of the Or47aexpressing ab5B OSN. We used a GR mutant line [referred to as sugar blind (Yavuz et al., 2014)] that carries a deletion of eight GRs, including Gr64b, all of which are involved in detection of sugars when expressed on the labellum. Even though Gr64b is not co-expressed with Or82a we also checked changes in the ab5A neuron, as a negative control. First, we tested if any changes in the odor sensitivity threshold of ab5B OSNs in the absence of Gr64b. We tested the SSR responses in two physiological states (fed and starved) so as to check for any feeding state dependent olfactory contribution of Gr64b in odor sensitivity. We measured the change in the number of action potentials (spikes) after odor stimulation, and we established the area under the spike frequency curve (Figure 3A). The first analysis gives a measure of quantitative changes, while the latter reveals temporal changes in the firing dynamics of the neurons.

The experiments revealed no significant differences in the odor sensitivity threshold between wild type flies and the octuple GR mutant line (**Figures 3B,C**). Both ab5A and ab5B neurons were equally sensitive to their best ligands (geranyl acetate and pentyl acetate, respectively) at low concentrations (10^{-10} v/v) in both fed and starved states. Nor were any significant differences in spike counts observed at any concentration tested (p > 0.05, unpaired, non-parametric *t*-tests with Mann Whitney test, **Figures 3B,C**).

Next, we studied changes in response specificity in ab5B OSNs lacking Gr64b. Only fed flies were used for these experiments. Once again, we did not observe any significant



changes or shifts in odor tuning to a panel of 30 diverse ligands representing ecologically relevant odors of diverse chemical classes (**Figures 3D-F** and **Table 1**). Taken together, our results suggest that Gr64b plays no significant role in modulating odor sensitivity or specificity in Or47a-expressing OSNs even though they are co-expressed in the very same neurons.

Discussion

Canonical models of odorant receptor (OR) expression in *Drosophila melanogaster* have generally supported a one receptor type - one olfactory sensory neuron (OSN) class model. However, two recent studies challenge this model by showing expression of multiple chemoreceptor families in a single class of OSN in *D. melanogaster* (Task et al., 2022) and also in *Aedes aegypti* (Younger et al., 2022). Both of these studies demonstrated co-expression of the olfactory co-receptor (Orco) and multiple ionotropic co-receptors, Ir8a, Ir76b and Ir25a, in multiple OSN classes in the antenna and maxillary palps. However, the scope of these studies paid little attention to the expression and possible role of gustatory receptors (GRs) in the antenna. Although previous studies have reported expression of multiple GRs in the antenna of *D. melanogaster*, their functional role has not been elucidated (Thorne and Amrein, 2008; Fujii et al., 2015).

Here, we investigated if GRs contribute functionally to peripheral olfactory coding in *D. melanogaster*. We first tried



to determine the expression patterns of the 12 gustatory receptors that have previously been reported to be expressed in the antenna of *D. melanogaster* (Thorne and Amrein, 2008; Menuz et al., 2014; Fujii et al., 2015) and generated a spatial expression map. Built on these results, we focused our functional study on Gr64b because of its consistent and broad expression pattern and its clear co-expression with Or47a, known to be expressed in ab5B neurons. Using single sensillum recording experiments in Gr64b mutant flies we did not observe any significant changes, neither in odor sensitivity, nor in odor turning properties of Or47a expressing neurons.

non-canonical innervation pattern of the ab5 sensillum class in *D. melanogaster*.

In our initial screening of expression patterns, we selected 12 GRs based on previous reports of expression of GRs in the antenna of *D. melanogaster* (Thorne and Amrein, 2008; Menuz et al., 2014; Fujii et al., 2015). Using the Gal4-UAS binary expression system we could only observe expression of nine out of the 12 GRs tested. Six of the nine GRs expressed can be roughly classified into two groups based on their canonical functions when expressed in the labellum. Gr64b, Gr64c, Gr64e, Gr64f, and Gr43a in different heteromeric combinations have been reported to be involved in detection of sugars, glycerol and fatty acids (Jiao et al., 2008; Wisotsky et al., 2011; Fujii et al., 2015; Kim et al., 2018) while Gr93a is involved in detection of multiple bitter tastants such as caffeine, umbelliferone and coumarin when expressed in combination with other bitter GRs such as Gr33a and Gr66a (Lee et al., 2009; Poudel et al., 2015; Poudel and Lee, 2016). Two GRs, Gr28bD and Gr21a, have been reported to be involved in rapid warmth sensing and CO₂ perception, respectively, while Gr28bC has recently been



magenta line representing the octuple GR mutant fly. n = 4-6/ concentration. Unpaired, non-parametric *t*-tests with Mann Whitney test were performed at each concentration between two genotypes. No significant differences at each concentration were observed across experiments (p > 0.05, unpaired *t*-test). The sensitivity was not changed between two genotypes tested (**D**,**E**). (**F**-**H**) Odor tuning sensitivity of both ab5A and ab5B neurons were checked with a panel of 32 odors representing diverse chemical classes in wild type and octuple GR mutant fly lines in fed state in case of ab5A and ab5B neurons respectively (**F**,**G**) No significant shift in odor tuning curve in case of ab5B neuron was observed in absence of Gr64b (**H**), n = 5.

deorphanized and shown to be involved in detection of saponin (Jones et al., 2007; Kwon et al., 2007; Ni et al., 2013; Sang et al., 2019).

We observed nine out of the twelve tested GRs. Labeling of neurons using binary expression systems greatly depend on the construction of the transgenic line (positional effect). It is thus possible that some essential regulatory elements for the expression could have been replaced during the genetic insertion (Yavuz et al., 2014; Task et al., 2022). This is specifically true in case of GRs from the GR64 cluster due to the complex arrangement of regulatory elements, sometimes also within the locus (Fujii et al., 2015). This limitation of the Gal4-UAS system was resolved by generation of LexA lines, and it could, e.g., be shown that Gr64f has a broad expression in the antenna as well as in the maxillary palps of *D. melanogaster* (Fujii et al., 2015). Furthermore, a broad expression pattern

for Gr64b as well as innervation of Gr64b expressing OSNs in the antennal lobe was reported. However, the authors did not investigate the functional role of any of these non-canonically expressed GRs (Fujii et al., 2015). In our study, using a direct fusion protein (Gr64b.mGFP), we observed a consistent and broad expression pattern for Gr64b. Interestingly, we observed expression of Gr64e only in a small subset of neurons in the funiculus in close vicinity to the base of the arista. This subset of neurons was Orco negative (supplementary Figure 2) however, we cannot rule out the possibility that this subset of neurons could also express one or more IR coreceptor. When expressed in the proboscis, Gr64e has been reported to be involved in the perception of glycerol and fatty acids along with Gr64b (Wisotsky et al., 2011; Kim et al., 2018). We did not check if Gr64b and Gr64e were co-expressed. However, we did not observe a particular cluster of Gr64b positive cells

at the base of the arista where Gr64e is expected. It is thus likely that these cells represent an independent set of neurons positive for Gr64e. The arista of D. melanogaster houses six cells out of which three are responsible for cold and rapid heat sensing. These cells express Ir25a (along with other IRs) and Gr28bD, respectively (Ni et al., 2013; Budelli et al., 2019). We observed efferents of Gr64e expressing neurons leading to the arista. A possibility open for future investigation is that Gr64e would be involved in modulating the function of Gr28bD. The second segment of the antenna (pedicle) houses the auditory center, Johnston's organ, and has been demonstrated to be innervated by Gr28bC and Gr68a-expressing neurons (Ejima and Griffith, 2008; Thorne and Amrein, 2008). Silencing of Gr68a-expressing neurons resulted in hampering the courtship behavior in D. melanogaster and thereby demonstrated another non-canonical expression and functional involvement of GRs when expressed in the auditory center of the antenna. However, it would be interesting to question if this phenotype was a result of all Gr68a expressing neurons or only because of silenced neurons in the second segment. We observed expression of Gr93a, Gr28bC and a few cells labeled with Gr64e on the second segment. Future investigations should establish if Gr28bC, Gr64e and Gr93a are generally expressed in the same set of neurons and if they contribute to the auditory and specifically mating behaviors of the fly.

Gr64b-expressing OSNs innervate three (VA6, DM3, and VM2) and four (VA6, DM3, VM2, and DA1) glomeruli in the antennal lobes of females and males, respectively (Fujii et al., 2015). In our experiments we observed innervation of only three glomeruli. Our preparations contained brains from flies of both sexes. It is thus possible that brains showing positive staining came exclusively from the females and therefore, DA1 innervation was not observed. Next, we expected that due to the innervation of both the VA6 and the DM3 glomeruli, Gr64b should be co-expressed in both Or82a- and Or47a-expressing OSNs, known to innervate these two glomeruli (Couto et al., 2005). However, we observed co-expression of Gr64b only with Or47a (Figure 2E), while cells positive for Gr64b and Or82a were separate but in close vicinity, suggesting innervation of the same sensillum (Figure 2D). This mismatch raises a question regarding labeling of the VA6 glomerulus (innervated by Or82a) by Gr64b-expressing OSNs even though there is no co-expression at the OSN level. Such mismatch of receptor expression and glomerular innervation has been studied in detail in the recent two papers showing co-expression of Orco and IR25a (Task et al., 2022; Younger et al., 2022). In case of D. melanogaster, innervation of multiple glomeruli innervated by Ir25a positive neurons was observed (Task et al., 2022). It was earlier shown that the V glomerulus is innervated only by neurons expressing Gr21a and Gr63a, which are involved in detection of CO₂ (Jones et al., 2007; Kwon et al., 2007). However, surprisingly, it was in the recent study by Task et al. (2022) observed that a few Orco and Ir25a positive neurons also innervate the V glomerulus sparsely, pointing to a wider possibility of overlap between these three gene families, but which could be revealed only after using more effective expression systems (Task et al., 2022). Similarly, the VM6 glomerulus, which was earlier considered a singular entity, was shown to be innervated by subpopulations of different neuron classes. A subdomain was innervated by a new type of OSNs expressing an Rh50 ammonium transporter protein (Vulpe et al., 2021) and also by Ir25a positive OSNs (Task et al., 2022). Further, it was shown that the VA6 glomerulus (corresponding to Or82a) was innervated by OSNs expressing multiple co-receptors, including Orco, Ir25a, Ir8a, and Ir76b, while, interestingly, the DM3 (corresponding to Or47a and Or33b) was innervated only by Orco positive neurons. It is also interesting to note that Or47a-expressing neurons do not co-express any of these widely expressed IR co-receptors, yet express Gr64b. We did not check if there is co-expression of IR coreceptors and Gr64b. However, it would be interesting and informative for the future to check if these two gene families overlap in the antenna of D. melanogaster. It must also be noted that in our experiments we used the Gr64b.mGFP transgenic line as a proxy for expression of the Gr64b receptor protein. In the future it would be interesting to investigate the expression pattern of Gr64b also by in situ hybridization or by generating antibodies for Gr64b protein.

One question that is crucial to our understanding of the fly olfactory system is thus whether non-canonically co-expressed GRs contribute functionally to peripheral olfactory coding. Till date, a functional role has been assigned only to three antennal GRs (Gr21a, Gr63a and Gr28bD). Amongst those, Gr21a and Gr63a are co-expressed in the ab1C neuron in the funiculus and are involved in detection of CO_2 (Jones et al., 2007; Kwon et al., 2007) while Gr28bD is expressed in the arista and has been shown to be involved in rapid heat sensing (Ni et al., 2013). Co-expression of Gr10a with Or10a in the ab1D neuron in the funiculus has also been reported without any further study regarding its role in odor perception (Jones et al., 2007). In the study by Task et al. (2022, see above) the authors checked for functional changes in OSNs co-expressing Ir25a using SSR technique. They found small but significant changes in a few sensillum classes in Ir25a mutant genotypes. However, functional changes were different for individual OSN types, as, e.g., depression of spike numbers in case of ab3A OSN type when stimulated with 1-octen-3-ol, while increase in spike numbers in the pb1A and pb3A types when stimulated with the same compound. This demonstrates that removing Ir25a affects functionality in different OSN types independently. However, if this leads to any behavioral changes is yet to be explored. What ecological benefit could such overlap confer to an insect? A possible answer to this question lies in the formation of redundancy in the olfactory system and was demonstrated in another recent study in Aedes aegypti (Younger et al., 2022). There have been multiple attempts to generate genetically modified strains of anthropophilic mosquitoes by masking human odors. This was attempted either by deleting receptors involved in CO2 perception or by those involved in human odor sensing (Degennaro et al., 2013; McMeniman et al., 2014; Raji et al., 2019). However, deletion of either of these receptors did not completely hamper the host seeking abilities of mosquitoes under semi field conditions. A possible explanation to these observations was recently presented, revealing an extensive overlap between multiple gene families in OSNs of the Aedes aegypti antenna, maxillary palps and, to some extent, proboscis (Younger et al., 2022). The authors showed co-expression of Orco and Ir25a in multiple OSNs. The striking feature of these recent findings is that the olfactory system contains subsets of neurons positive for either only Orco or only Ir25a, while some neurons are positive for both Orco and IRs. This holds true also in the case of CO₂ sensing Gr1-3-expressing neurons. Such redundancy is probably one of the key factors explaining the highly successful host seeking seen in mosquitoes, even in the absence of Orco. In this context, we were curious if there is any functional significance for the overlap between GR64b and Or47a expression and investigated if Gr64b has a role in odor sensitivity as well as in the tuning profile of Or47aexpressing OSNs on the D. melanogaster antenna. The odor tuning curve of an OSN is primarily determined by the ORs (Hallem and Carlson, 2006) but another study demonstrated amplification of Or67d neuronal responses in the combination of another receptor family protein, ppk25 (Ng et al., 2019). We did not observe any functional involvement of Gr64b in modulating odor sensitivity of Or47a OSNs in case of change in sensitivity threshold or changes in odor tuning specificity, suggesting no functional involvement at least in the odor detection characteristics of Or47a. We cannot yet rule out the possibility of other non-olfactory functional roles of Gr64b. Other non-canonical roles for Gr64b have also been suggested as one of the downstream factors for fatty acid sensing via Gr64e (Kim et al., 2018). Recently, a role of Gr64 cluster genes was demonstrated in proteostasis and is crucial for the survival of ribosomal protein mutant epithelial cells in D. melanogaster (Baumgartner et al., 2021). Lastly, a pleiotropic role of GRs has been well documented in case of expression in the fly brain, gastrointestinal tract and reproductive sites (Park and Kwon, 2011; Miyamoto and Amrein, 2013; Vernier et al., 2022). Therefore, it can be hypothesized that Gr64b may not perform a chemosensory role but rather act as a GPCR involved in cellular functions when expressed in ab5B neurons.

Data availability statement

The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation.

Ethics statement

This study on the fruit fly *Drosophila melanogaster* was performed in Germany where the research on invertebrates does not require a permit from a committee that approves animal research.

Author contributions

SL-L, MK, and BH conceived the project. VPM and SL-L designed the experiments. VPM conducted the experiments, made the figures, and data analysis. All authors discussed the results and wrote the article.

Funding

This study was supported by the Max Planck Society (SL-L, MK, and BH) and the International Max Planck Research School (IMPRS) at the Max Planck Institute of Chemical Ecology (VPM).

Acknowledgments

We thank Regina Stieber, Silke Trautheim, Sin Prelic, Carolin Hoyer, and Dr. Veit Grabe for comments, discussions, and technical support.

Conflict of interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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Supplementary material

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/ fevo.2022.980351/full#supplementary-material

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