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Sex- and tissue-specific expression of chemosensory receptor genes in a hawkmoth

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For the nocturnal hawkmoth *Manduca sexta*, olfactory and gustatory cues are essential for finding partners, food, and oviposition sites. Three chemosensory receptor families, odorant receptors (ORs), ionotropic receptors (IRs), and gustatory receptors (GRs) are involved in the detection of these stimuli. While many chemosensory receptor genes have been identified, knowledge of their expression profile in potentially chemoreceptive organs is incomplete. Here, we studied the expression patterns of chemosensory receptors in different tissues including the antennae, labial palps, proboscis, legs, wings and ovipositor. We compared the receptors' expression in female and male moths both before and after mating by using the NanoString platform. This tool allowed us to measure expression levels of chemosensory receptor genes in a single reaction using probes designed against 71 OR, 29 IR and 49 GR transcripts. In all tissues investigated, we detected expression of genes from all three receptor families. The highest number of receptors was detected in the antennae (92), followed by the ovipositor (59), while the least number was detected in the hindlegs (21). The highest number of OR genes were expressed in the antennae (63), of which 24 were specific to this main olfactory organ. The highest number of IRs were also expressed in the antennae (16), followed by the ovipositor (15). Likewise, antennae and ovipositor expressed the highest number of GRs (13 and 14). Expression of the OR co-receptor *MsexORCo*, presumably a prerequisite for OR function, was found in the antennae, labial palps, forelegs and ovipositor. IR co-receptors *MsexIR25a* and *MsexIR76b* were expressed across all tested tissues, while expression of the IR co-receptor *MsexIR8a* was restricted to antennae and ovipositor. Comparing the levels of all 149 transcripts across the nine tested tissues allowed us to identify sex-biased gene expression in the antennae and the legs, two appendages that are also morphologically different between the sexes. However, none

of the chemosensory receptors was differentially expressed based on the moths' mating state. The observed gene expression patterns form a strong base for the functional characterization of chemosensory receptors and the understanding of olfaction and gustation at the molecular level in *M. sexta*.

KEYWORDS

hawkmoth, sex-specific, mating status, chemosensory organs, NanoString, chemosensory receptor, *Manduca sexta*

Introduction

Sensory systems help an organism to receive and respond to cues present in its environment and perform essential behaviors such as search for and selection of food, mate and shelter. For insects, the chemosensory system, which detects chemical signals via olfactory and gustatory means, is of great importance (Hansson and Stensmyr, 2011; Depetris-Chauvin et al., 2015; Haverkamp et al., 2018). These chemical signals can be volatile substances (odors) emitted by plants, microorganisms and animals or non-volatile substances (tastants) that remain on the surface of their sources, for example, on leaves. Insects use multiple peripheral organs to detect these signals, such as the antennae and mouthparts on the head, legs and wings attached to the thorax, and male or female genitalia at the tip of the abdomen (Dunipace et al., 2001; Depetris-Chauvin et al., 2015; Xu, 2020; Koutroumpa et al., 2021). These chemosensory organs have on their surfaces several hair-like structures, called sensilla, housing olfactory or gustatory sensory neurons. Different populations of sensory neurons detect different subsets of the chemical space surrounding the organism due to chemosensory receptor proteins expressed in their dendritic membranes (Dahanukar et al., 2005; Joseph and Carlson, 2015; Agnihotri et al., 2016). Three major chemosensory receptor gene families, the odorant receptors (ORs), the ionotropic receptors (IRs), and the gustatory receptors (GRs), encode these receptor proteins. ORs generally detect long-chain pheromone molecules (Zhang and Löfstedt, 2015) and plant-related volatiles of diverse chemical classes (Hallem and Carlson, 2006; de Fouchier et al., 2017). The molecular function of ORs relies on forming heteromeric complexes with the obligate co-receptor ORCo (Sato et al., 2008). IRs can detect both odors as well as tastants (Wicher and Miazzi, 2021). The olfactory function of IRs also requires co-receptors and is usually restricted to detecting acids, aldehydes or amines (Vulpe and Menuz, 2021). The co-receptor IR8a is involved in acid-sensing, while IR25a and IR76b are co-receptors for amine-sensing (Abuin et al., 2011; Vulpe and Menuz, 2021). GRs are mostly known for gustatory functions but are also involved in detecting CO₂ (Jones et al., 2007; Ning et al., 2016) and a few other odorants such as acids (Kumar et al., 2020).

Initially, these chemosensory receptor genes were identified in the genome of *Drosophila melanogaster* (Clyne et al., 1999, 2000; Gao and Chess, 1999; Vosshall et al., 1999; Benton et al., 2009). Then, advancements in genome and transcriptome sequencing methods, and the development of optimized bioinformatic tools to identify homologous genes led to the identification of chemosensory receptor repertoires in many other insect species (Wanner et al., 2007; Grosse-Wilde et al., 2011; Koenig et al., 2015; Walker et al., 2016, 2019; Koutroumpa et al., 2021). Like all gene families, chemosensory receptors evolve by duplication, functional diversification and pseudogenization of their genes. Across insect orders, we find a considerable variation in the total number of receptors, high divergence in amino acid sequences and occurrence of clade-specific receptor subfamilies (Gouin et al., 2017; Pearce et al., 2017; Brand et al., 2018; Xu, 2020; Yin et al., 2021). These findings reveal that chemosensory receptors are under high selection pressure and evolve continuously in a birth and death process. Therefore, functional characterization of these genes in different species is necessary to understand the impact of chemical ecology on insect evolution.

The nocturnal hawkmoth *Manduca sexta* has served for decades as an important Lepidopteran model for research in chemosensory modalities providing deep insight into the physiological, neuroanatomical and behavioral aspects of olfaction and gustation. Recent work has generated a highly curated annotation of OR, IR and GR genes for *M. sexta* (Koenig et al., 2015). This paved the way for the application of CRISPR-Cas9 mediated targeted knock-out experiments revealing the function of the obligate co-receptors MsexORCo for ORs, and MsexIR8a and MsexIR25a for IRs in *M. sexta* (Fandino et al., 2019; Zhang J. et al., 2019). In addition, one sex pheromone receptor and two ORs detecting plant volatiles have been de-orphanized to date (Wicher et al., 2017; Guo et al., 2021; Zhang et al., 2022), whereas the functions of all other chemosensory receptors of *M. sexta* remain elusive to the best of our knowledge. A critical step in understanding any gene's function is knowing when and where it is expressed. There is limited knowledge about the expression of some of these receptors in the antennae (Koenig et al., 2015), the proboscis (Haverkamp et al., 2016), and the female ovipositor (Klinner et al., 2016)

in *M. sexta*. Few studies in other moth species have also investigated the expression of chemosensory genes in different tissues (Jacquin-Joly et al., 2012; Guo et al., 2018; Walker et al., 2019; Koutroumpa et al., 2021).

While input from the antennae can produce odor-directed flight responses in *M. sexta* moths (Schneiderman et al., 1986; Kalberer et al., 2010), a concerted role of information detected at multiple chemosensory tissues is likely for optimal behavioral output, especially regarding feeding and oviposition behaviors. Odor plumes originating from flowers guide moths toward rewarding nectar sources (Riffell et al., 2008). During such foraging, the proboscis which can both smell and taste (Reiter et al., 2015; Haverkamp et al., 2016) is extended, a behavior that is disrupted in knock-out *MsexORCo* mutants (Fandino et al., 2019). In case of oviposition, plant-released volatiles alone can attract gravid *M. sexta* females toward host-plants (Kariyat et al., 2013; Spaethe et al., 2013). When moths approach the host plant, they curl their abdomen even when contact with the plants is prevented (Mechaber et al., 2002). Results from Klinner et al. (2016) suggest that this behavior serves to sample odors with the odor-receptive ovipositor. After landing and before laying eggs, the female moth assesses the chemical profiles of leaves with its forelegs and midlegs, suggesting a gustatory function of these appendages (Yamamoto et al., 1969; Mechaber et al., 2002). Accordingly, putative chemosensory sensilla on the legs of *M. sexta* have been described (Kent and Griffin, 1990). Moreover, CO₂ is an important cue for foraging (Guerenstein et al., 2004b; Thom et al., 2004), where CO₂-detection has been shown to occur in specialized sensilla present on the labial palps (Kent et al., 1986; Guerenstein et al., 2004a). The chemosensory potential of wings has not been reported so far in *M. sexta*; however, it has been observed in other insects like mosquitoes (Yang et al., 2020) and vinegar flies (Raad et al., 2016; He et al., 2019; Yanagawa et al., 2019).

Males and female moths need to extract different information from their environment to perform their sex-specific tasks. A well-known example of sex-specificity in moths is their pheromone communication system. Male moths have specialized antennae to detect female-produced sex pheromones from afar and thereby find mates (Tumlinson et al., 1989). In *M. sexta*, the receptor for the main pheromone component bombykal, *MsexOR1* (Wicher et al., 2017) has a male-specific expression (Grosse-Wilde et al., 2010) and mutant males lacking functional *MsexORCo* are unable to mate (Fandino et al., 2019). Likewise, searching for hostplants for oviposition is a female-specific task, which might require adaptations of the female chemosensory system and expression of female-specific receptors. Apart from behaviors related to reproduction, male and female hawkmoths exhibit different foraging preferences when given a choice between flowers from two plant species (Alarcón et al., 2010).

Moreover, physiological states such as age, nutrition and mating state can modify insect behavior (Gadenne et al., 2016).

Mated female *M. sexta*, for example, shows an increased flight and abdomen curling response compared to virgin females when exposed to hostplant volatiles (Mechaber et al., 2002). Mating can also cause changes in physiological responses toward vegetative and floral plant odors in female *M. sexta* and *Spodoptera littoralis* (Saveer et al., 2012; Bisch-Knaden et al., 2022). For male hawkmoths, feeding after each mating is advantageous for repetitive mating success (Levin et al., 2016), and females benefit from post-mating feeding with increased longevity and more mature eggs than starved females (Sasaki and Riddiford, 1984; von Arx et al., 2013). These mating state-dependent differences might be linked to modulations of chemosensory receptor levels. Mating-dependent up- or downregulation of several ORs was seen in flies, *Drosophila suzukii* (Crava et al., 2019), *Bactrocera dorsalis* (Jin et al., 2017) and the moth *Dendrolimus punctatus* (Zhang et al., 2017).

In the present study, we investigated expression of genes from all three chemosensory receptor families in different chemosensory organs and compared the expression between males and females, pre- and post-mating. We used the NanoString digital platform for gene expression (Geiss et al., 2008; Kulkarni, 2011) to count individual mRNA transcripts using complementary nucleotide probes designed for each gene of interest. The NanoString technology was shown to be highly sensitive and comparable to qPCR and RNA-sequencing techniques (Geiss et al., 2008; Veldman-Jones et al., 2015; Bondar et al., 2020) and has been used to study expression of olfactory receptors in mice (Khan et al., 2011, 2013). Associating gene expression patterns with the known or suspected chemosensory function of the tested tissues enables us to identify potentially important receptors whose function could be revealed in future experiments.

Materials and methods

Insect rearing

Manduca sexta moths were either reared in our lab or pupae were kindly provided by Monika Stengl (University of Kassel). For rearing, we used an artificial diet (156.25 g soy flour, 144.38 g rye flour, 144.38 g wheat flour, 85 g agar-agar, 4.5 ml rapeseed oil, 1.45 g calcium carbonate, 1.45 g sodium chloride, 4.8 g methyl paraben, 4.8 g sorbic acid, 14.5 g ascorbic acid, 38.5 g milk powder, 32 mg nicotinic acid, 16 mg riboflavin, 7.5 mg thiamine, 7.5 mg pyridoxine, 7.5 mg folic acid, 0.64 mg biotin). We kept the larvae in a climate chamber with a 14 h:10 h light:dark cycle, a temperature of 26°C (light cycle) or 24°C (dark cycle), and at 60% relative humidity. Male and female pupae were sorted and moved to separate climate chambers with a 16 h:8 h light:dark cycle, a temperature of 25°C, and at 60% relative humidity (light cycle) or 70% relative humidity (dark cycle). Eclosing adults

were collected daily and held in individual brown paper bags (17 cm × 26 cm) in the pupal climate chambers.

Tissue collection and RNA extraction

For virgin samples, males and females remained in separate chambers until tissue collection. For mated samples, males and females were paired 2 days after eclosion during early scotophase and observed every 2 h for successful mating. Tissues from virgin and mated moths (**Supplementary Table 1**) were collected 3 days after eclosion during the scotophase. Moths were first cooled down at 4°C for about 15 min to decrease movements, then tissues were dissected with microscissors, collected in 2 ml microcentrifuge tubes or 15 ml falcon tubes (forewings), immediately immersed in liquid N₂, and stored at −80°C until further processing. The tissues (except forewings) were lysed and disrupted in RLT buffer (Qiagen, Germany¹) using Tissue Lyser LT (Qiagen, Germany) with a steel bead (diameter: 5 mm) at 50 oscillations per second for 15 min. The forewings, due to their large size, were ground to a fine powder in liquid N₂ using a mortar and a pestle. Total RNA was extracted from the lysed tissues using RNeasy Mini Kit (Qiagen, Germany) and RNase-free DNase Set (Qiagen, Germany) was used for the DNase digestion step. The concentration and purity of the extracted total RNA were checked first on a NanoDrop One (Thermo Fisher Scientific²). Three samples for each experimental group, leading to six ovipositor samples (3 virgin female, 3 mated female), and 12 samples for each of the remaining tissues (3 virgin female, 3 mated female, 3 virgin males, 3 mated males) were then run on 2,100 Bioanalyzer (Agilent Technologies, Inc.³) to confirm RNA integrity. Based on initial standardization assays, 200 ng total RNA from each tissue was used for the NanoString gene expression assay.

NanoString gene expression assay

We used the nCounter XT CodeSet gene expression assay (NanoString Technologies, Inc., United States⁴). The custom CodeSet (Zhang et al., 2022) contained 268 probes targeting 71 ORs, 29 IRs, 49 GRs, 47 odorant binding proteins, 5 pickpocket, 3 sensory neuron membrane proteins and 62 candidate reference gene transcripts (**Supplementary Table 2**). The probes were designed using the reference coding sequences of 74 ORs, 21 IRs, and 45 GRs (Koenig et al., 2015) that were submitted to *M. sexta* OGS2.0 (Kanost et al., 2016). Four isoforms of *MsexGR11*, six isoforms of *MsexGR15* and 8 IRs that were identified later

(*MsexIR1.1*, 7d.2, 7d.4, 85a, 100a, 100d, 100f, and 100h) (Liu et al., 2018) were included. Some of the receptor sequences had high homology with sequences of their duplicates, prohibiting the design of unique probes. Therefore, these receptors had to be excluded (*MsexOR23*, *MsexGR15D*, 25, 26, 27, and 36). For all final probes, we performed NCBI BLAST search against *M. sexta* reference mRNA. Except for a few cases with duplicate receptors for which the probes could cross-target, our probes did not have off-targets. The recent *de novo* *M. sexta* genome assembly (Gershman et al., 2021) indicated that *MsexOR84* and *MsexOR89* might be the same gene, and similarly *MsexOR42* and *MsexOR66* might be the same gene. Therefore, we provisionally named these receptors *MsexOR84/89* and *MsexOR42/66*.

We followed the standard protocol from nCounter XT Gene Expression Assay User Manual (MAN-10023-11, page 16⁵). All reagents and buffers were purchased from NanoString Technologies, Inc., United States. An attenuation mix (Eurofins Genomics, Germany⁶) to suppress counts of *MsexABPx*, *MsexOBP1*, *MsexOBP5* and *MsexOBP6* was used as these genes had very high expression levels causing high binding densities for the antennal samples. Briefly, a master-mix of 42 μl Reporter CodeSet, 28 μl Reporter-Plus reagent and 70 μl nCounter SPRINT hybridization buffer was created. For each hybridization reaction we combined 10 μl master-mix, 5 μl total RNA (200 ng), 1 μl attenuation-mix and 3 μl mixture of Capture ProbeSet and Capture-Plus reagent. Hybridization was done at 65°C for 22 h, after which 16 μl Merck water was added to the sample. The entire volume was loaded on nCounter SPRINT Cartridge (NanoString, United States) and processed on nCounter SPRINT Profiler (NanoString, United States).

Data processing

The raw data from the gene expression assay (see Data Availability Statement) was processed using nSolver4.0 (NanoString, United States). Quality control for mRNA data was done on each sample using default parameters for nCounter SPRINT Profiler according to NanoString Gene Expression Data Analysis Guidelines (MAN-C0011-04⁷). The parameters were Imaging QC: 75; Binding Density QC: 0.1 – 1.8; Positive Control Linearity QC: 0.95; Positive Control Limit of Detection QC: 2 standard deviations. Due to high expression levels of chemosensory genes in the antennae, those samples had a slightly higher binding density than the default threshold of 1.8. We visually inspected the raw data from antennae following instructions from the user manual and confirmed that the data

1 <https://www.qiagen.com/de>

2 <https://www.thermofisher.com/de/de/home.html>

3 <https://www.agilent.com/>

4 <https://nanosttring.com/>

5 https://nanosttring.com/wp-content/uploads/MAN-10023-11_nCounter_XT_Assay_User_Manual.pdf

6 <https://eurofinsgenomics.eu/>

7 https://nanosttring.com/wp-content/uploads/Gene_Expression_Data_Analysis_Guidelines.pdf

was robust and usable. However, the raw counts of positive spike-in controls in antennae were lower than those of the other tissues (**Supplementary Figure 1**). To avoid normalization flags, we normalized the antennal samples separate from the samples of the other tissues. For antennae, background subtraction was done first using the raw counts of the 8 negative control probes (mean + 2 standard deviations). After that, two normalization steps were performed, first using the geometric mean counts of the 6 external positive control probes, and second, using the geometric mean counts of three endogenous reference genes that were selected based on their coefficient of variation (CV < 30%). The endogenous reference genes for antennal samples were *melastatin*, *smoothened*, and *dunce*. For the remaining tissue samples, background subtraction and positive control normalization was done similar to the antennae. The endogenous reference genes used for the second normalization were *melastatin*, *smoothened* and two G protein genes (*Gbeta13F* and *concertina*) with a CV < 25%. After these two normalization steps, the minimum normalized count for each sample was defined as “background,” and any chemosensory receptor gene with counts above this background in at least 2 of the 3 samples was considered to be expressed in that experimental group. The normalized data are available online (see section “Data availability statement”).

Data analysis

The double-normalized counts (referred to as “counts” in the Results and Discussion) were \log_2 transformed and used for making heatmaps and for statistical analyses using R (version 4.1.2)⁸ and Rstudio.⁹ The heatmaps were generated using the libraries *heatmaply* and *RColorBrewer* and later edited on Adobe Illustrator CS5. For identifying sex and mating state dependent gene expression in a given tissue (except ovipositor), we performed a nested ANOVA for each gene with sex as the main factor and mating state as the nested factor and a significance level of 0.05. The test was done separately for each tissue only for those genes that were expressed in at least one of the four experimental groups for that tissue. We then calculated the false discovery rate (FDR) using the Benjamini-Yekutieli procedure with the sets of p -values for the two factors for each tissue. Finally, a gene was considered to be significantly differentially expressed between sexes or mating states if $FDR < 0.05$ and \log_2 fold-change ≥ 2 . Ovipositor samples from virgin and mated females were compared with an unpaired t -test. For a p -value < 0.05 and a \log_2 fold-change = 2 between both groups, a gene was considered to be differently expressed.

⁸ <https://www.r-project.org/>

⁹ <https://www.rstudio.com>

Results and discussion

NanoString gene expression assay for *Manduca sexta* chemosensory receptors

We measured the expression of chemosensory receptors in different tissues (antennae, labial palps, proboscis, forelegs, midlegs, hindlegs, forewings, hindwings) of female and male *M. sexta* moths, both before and after mating. In addition, we examined the expression of chemosensory genes in the ovipositor of virgin and mated females. The probe set targeted 71 OR transcripts including the co-receptor *MsexORCo*, 29 IR transcripts including the co-receptors *MsexIR8a*, *MsexIR25a*, *MsexIR76b*, and 49 GR transcripts (see section “Materials and methods” for details). Out of these 149 chemosensory receptors, transcripts for two ORs, seven IRs and 25 GRs were not detected in any of the investigated tissues, that is, counts of these transcripts were in the range of background counts (see section “Materials and methods” for details, **Table 1** and **Supplementary Table 3**). On the other hand, three ORs, three IRs and two GRs were detected in each of the tissues tested.

We identified several genes with tissue-specific expression (**Table 1**). We also found sex-biased expression of several ORs and one GR (FDR corrected $p < 0.05$, nested ANOVA), whereas the expression of IRs seemed to be independent of sex in the investigated tissues (FDR corrected $p > 0.05$, nested ANOVA, **Table 2**). In contrast, the mating status of female or male moths had no statistically significant effect on the expression level of any chemosensory receptor in any tissue (FDR corrected $p > 0.05$, nested ANOVA).

Antennae

In total, in the male and female antennae, we detected transcripts of 63 out of the 71 ORs (89%, **Figure 1A**). Twelve of these antennal ORs had a sex-biased gene expression (**Table 2**). Sixteen out of the 29 IRs (55%) and 13 out of the 49 GR transcripts (27%) were also expressed in the antennae (**Figures 1B,C**).

ORs. The OR co-receptor *MsexORCo* had high counts for the antennal samples (**Figure 1A**). A high expression of *MsexORCo* was expected as its co-expression in all OR expressing neurons is considered necessary for OR-mediated odor detection (Sato et al., 2008) and knocking out *MsexORCo* reduces or completely abolishes antennal, antennal lobe and behavioral responses toward plant odors (Fandino et al., 2019).

We could confirm previous findings that *MsexOR1*, *MsexOR4*, *MsexOR51*, and *MsexOR83* had a male-biased expression in the antennae (Grosse-Wilde et al., 2010; Koenig et al., 2015). *MsexOR1* detects the major female sex-pheromone

TABLE 1 Chemosensory receptor gene transcripts that were tissue-specific, broadly expressed or not detected.

Tissue	ORs	IRs	GRs
Antennae only	<i>OR1, OR4, OR11, OR15, OR18, OR19, OR21, OR22, OR24, OR29, OR31, OR33, OR34, OR36, OR41, OR43, OR49, OR62, OR67, OR69, OR76, OR78, OR85, OR86</i>	<i>IR31a, IR41a, IR75p.1, IR87a</i>	<i>GR8, GR14, GR15B</i>
Labial palps only	–	–	<i>GR1</i>
Proboscis only	–	–	<i>GR11B</i>
Forelegs only	–	–	–
Midlegs only	–	–	<i>GR5</i>
Hindlegs only	–	–	–
Forewings only	–	–	–
Hindwings only	–	–	–
Ovipositor only	–	<i>IR75d</i>	<i>GR15G</i>
Expressed in each of the 9 tested tissues	<i>OR35, OR64, OR84/89*</i>	<i>IR25a, IR76b, IR7d.3</i>	<i>GR2, GR41</i>
Expressed in 8 out of 9 tested tissues	<i>OR8, OR47, OR75, OR77</i>	<i>IR75q.1, IR7d.2</i>	<i>GR3</i>
Not detected in the 9 tested tissues	<i>OR20, OR80</i>	<i>IR3, IR40a, IR64a, IR75p.3, IR85a, IR7d.4, IR100h</i>	<i>GR4, GR6.2, GR9.1, GR9.2, GR10.1, GR10.2, GR11A, GR11C, GR15A, GR16, GR17, GR18, GR19, GR21, GR22, GR24, GR28, GR29, GR30, GR32, GR33, GR37, GR38, GR40, GR43</i>

For information on presence and absence of expression of each receptor in each tissue (see [Supplementary Table 3](#)). *See section “Materials and methods.”

TABLE 2 Chemosensory receptor gene transcripts with a sex-biased expression.

Tissue	ORs		IRs		GRs	
	♀	♂	♀	♂	♀	♂
Antennae	<i>OR5, OR6*, OR15*, OR17, OR26, OR33, OR40, OR87</i>	<i>OR1*, OR4, OR51*, OR83*</i>	–	–	–	–
Labial palps	–	–	–	–	–	–
Proboscis	–	–	–	–	–	–
Forelegs	<i>OR6*</i>	–	–	–	–	–
Midlegs	<i>OR6*</i>	–	–	–	<i>GR15C*</i>	–
Hindlegs	<i>OR6*</i>	–	–	–	–	–
Forewings	–	–	–	–	–	–
Hindwings	–	–	–	–	–	–

*Sex-specific, i.e., no expression detected in the opposite sex in the respective tissue.

bombykal (Wicher et al., 2017), which, together with MsexOR4 and MsexOR51, belongs to the expanded pheromone receptor clade (Koenig et al., 2015). Likewise, the fourth male-biased receptor MsexOR83 belongs to a newly discovered lineage of pheromone receptors (Bastin-Héline et al., 2019). Hence, all three orphan, male-biased receptors might be involved in detecting minor components of the female-produced pheromone blend, containing in total 12 components (Kaisling et al., 1989; Kalinova et al., 2001). *MsexOR1*, *51*, and *83* had a male-specific antennal expression, whereas *MsexOR4* transcripts were also detected in female antennae. Such

expression of a pheromone receptor in females is seen in other moth species (Bengtsson et al., 2012; Walker et al., 2016), suggesting that these females can detect their own pheromone (Holdcraft et al., 2016). *M. sexta* female antenna is known to only respond to the minor pheromone component Z11-hexadecanal (Kalinova et al., 2001). However, in male *M. sexta*, *MsexOR4*-expressing olfactory sensory neurons are housed in the most abundant type of sensilla in the male antenna, the long trichoid sensilla (Grosse-Wilde et al., 2010), and neurons in these sensilla do not respond to Z-11-hexadecanal (Kalinova et al., 2001). Therefore, it seems unlikely that MsexOR4 is the

receptor for this minor pheromone component. One OR from the pheromone receptor clade, *MsexOR15*, is female-specific, a rare feature for this subfamily (Bastin-Héline et al., 2019). Its absence of expression in male antennae negates the possibility of *MsexOR15* being the Z-11-hexadecanal receptor, as males can detect this component with neurons housed in short trichoid sensilla (Kalinova et al., 2001). *MsexOR38*, which belongs to the same group as *MsexOR83* (Koenig et al., 2015) has an unbiased expression in males and females, and might therefore be the putative receptor for Z-11-hexadecanal. These ORs of the pheromone receptor clade may also detect pheromones of sympatric hawkmoths, such as *M. quinquemaculata* and *M. rustica*, thus ensuring reproductive isolation (Daimon et al., 2012). However, in the codling moth, *C. pomonella*, an OR belonging to the pheromone clade detects a plant volatile (Bengtsson et al., 2014), opening the possibility that some of the uncharacterized pheromone clade members of *M. sexta* might have a non-pheromonal function.

Apart from *MsexOR15*, we found seven more female-biased ORs. These include those previously identified as female-biased (*MsexOR5*, 6, 15, and 87; Koenig et al., 2015), and four newly identified female-biased ORs (*MsexOR17*, 26, 33, and 40). *MsexOR6*, like *MsexOR15*, was not only female-biased but also female-specific for the antennae. *MsexOR5* and 6 are encoded by two gene duplicates orthologous to the female-specific *Bombyx mori* linalool receptor *BmorOR19* (Anderson et al., 2009; Koenig et al., 2015). Therefore, one of these ORs might be responsible for the enantioselective response to (+)-linalool of the lateral large female glomerulus (latLFG) in the female *M. sexta* antennal lobe (Reisenman et al., 2004) and for the female moth's oviposition preference for (+)-linalool (Reisenman et al., 2010).

It is interesting to note that we have two times more female-biased than male-biased ORs. All male-biased ORs belong to pheromone receptor lineages; however, seven of the eight female-biased ORs are from different clades of potential plant volatile receptors (Guo et al., 2021). These ORs might be important in the context of the female-specific behavior of searching an appropriate oviposition site directed by host-plant volatiles. According to Bastin-Héline et al. (2019), another possible function of these polyphyletic, female-biased receptors might be the detection of male-produced pheromones, such as those presumably emitted from male hairpencils in *M. sexta* (Grant and Eaton, 1973; Birch et al., 1990).

The *MsexORCo*-associated glomeruli on the dorsal surface of the *M. sexta* antennal lobe (Fandino et al., 2019) respond to volatile headspace collections of native, nectar-rich flowers, and host and non-host plant foliage (Bisch-Knaden et al., 2022). This suggests that ORs might be responsible for the antennal detection of a large variety of floral and vegetative odors in *M. sexta* (Fraser et al., 2003; Spaethe et al., 2013). Ligands can be predicted based on homology to other de-orphanized Lepidopteran ORs for highly conserved OR clades. Such an

approach has led to identification of the floral odor phenyl acetaldehyde as a ligand for *MsexOR67* (Guo et al., 2021). The second identified plant volatile-detecting OR, *MsexOR35*, detects α -copaene, a volatile abundant in the headspace of the beetle-infested host-plant *Datura wrightii* that attracts ovipositing *M. sexta* (Zhang et al., 2022).

Overall, our results for the antennal expression of ORs largely agree with a previously performed RNA sequencing analysis, especially for the sex-biased ORs (Koenig et al., 2015), providing support for using the NanoString assay as a valid method for the study of insect chemosensory gene expression (Supplementary Figure 2).

IRs. Several IRs of the conserved antennal IR group (Croset et al., 2010; Yin et al., 2021) were expressed in the *M. sexta* antennae. These included the IR co-receptor genes *MsexIR8a*, *25a*, *76b* which had the highest counts among all IRs expressed in the antennae (Figure 1B). *MsexIR8a* and *MsexIR25a* are required for native responses of antennal coeloconic sensilla to carboxylic acids and amines, respectively (Zhang J. et al., 2019). For other antennal IRs, functions can be predicted based on *D. melanogaster* orthologs. *MsexIR76b* is ortholog to *DmelIR76b*, which is involved in both smell and taste of polyamines (Hussain et al., 2016) and also mediates gustatory detection of low salt concentrations and sour substances (Zhang et al., 2013; Chen and Amrein, 2017). Similarly, *MsexIR31a*, together with the co-receptor *IR8a*, is likely involved in acid detection, while *MsexIR41a*, when co-expressed with the co-receptor *MsexIR25a* or *MsexIR76b*, might be involved in detecting amines (Silbering et al., 2011; Hussain et al., 2016). IRs can also have non-chemosensory roles. Based on homology to *Drosophila* IRs, *MsexIR21a*, *MsexIR93a* and *MsexIR25a* could have cool-temperature-sensing functions (Knecht et al., 2016; Ni et al., 2016). Alternatively, *MsexIR93a* and *MsexIR25a* might be involved in hygrosensation (Knecht et al., 2016) by neurons present in the styliform sensilla of the antenna (Dahake et al., 2022).

We also found expression of nine Lepidoptera-specific IRs in the antennae (Koenig et al., 2015; Yin et al., 2021). Seven of these, *MsexIR1.1*, *MsexIR1.2*, *MsexIR4*, *MsexIR75p.1*, *MsexIR75p.2*, *MsexIR75q.1*, and *MsexIR87a* putatively encode IRs nested within the antennal IRs and are considered to have an olfactory function (Yin et al., 2021). All except *MsexIR87a* belong to a monophyletic group of putative acid-detectors (Hou et al., 2022). This group includes two acid-sensing IRs of *Agrotis segetum*, *AsegIR75q.1*, which is tuned specifically to octanoic acid and *AsegIR75p.1*, which has a broader tuning and detects acids, aldehydes and alcohols. Two Lepidoptera-specific IRs, *MsexIR7d.3* and *MsexIR100d*, lie among the divergent IRs (Yin et al., 2021; Hou et al., 2022), which are defined as IRs that are not expressed in the antennae but in other tissues (Croset et al., 2010; Yin et al., 2021). However, the expression pattern of *MsexIR7d.3* resembles that of its *A. segetum* ortholog *AsegIR7d.3*, which is highly expressed in

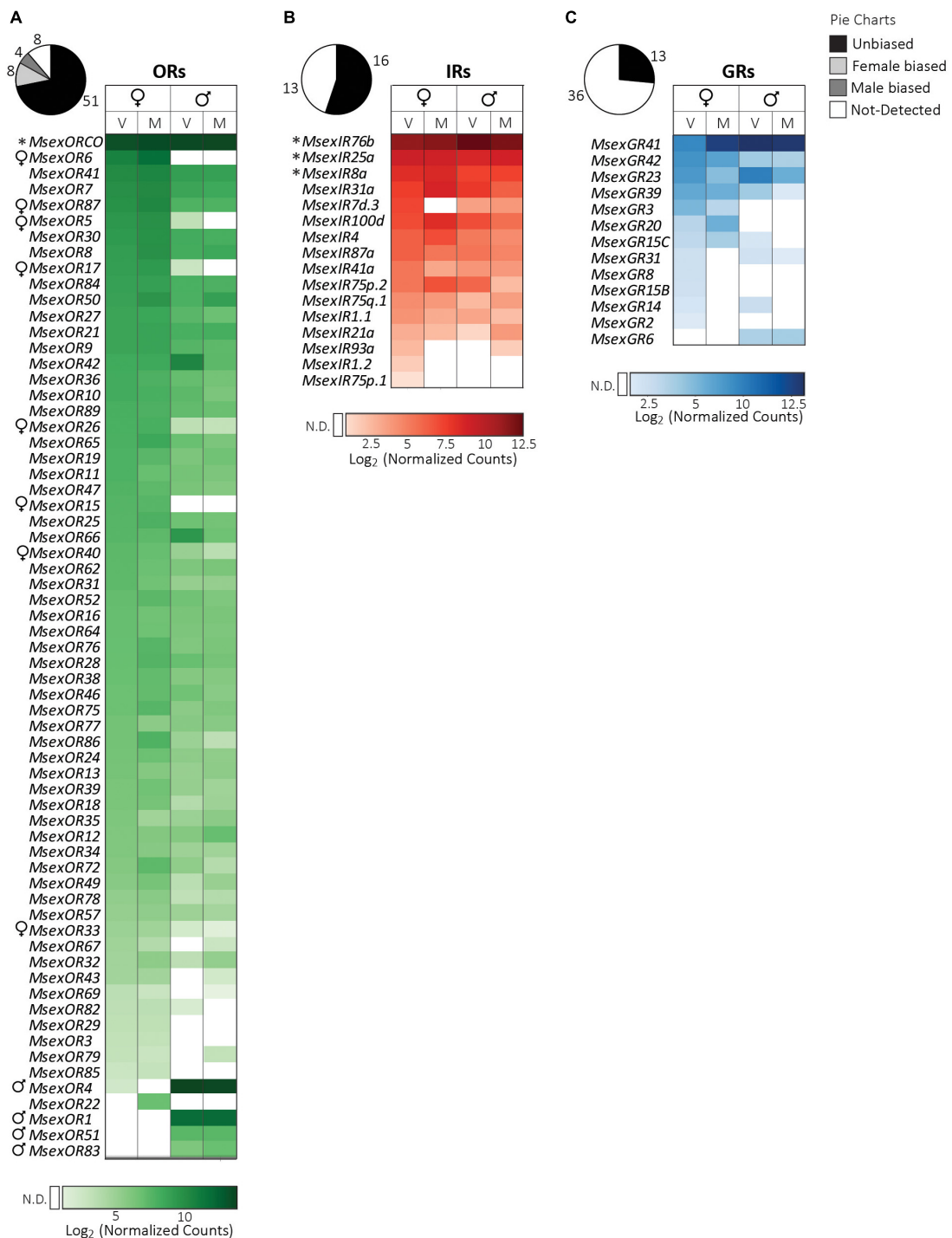


FIGURE 1

Expression of chemosensory receptor genes in the antennae of adult *M. sexta*. (A) Odorant receptors, (B) ionotropic receptors, (C) gustatory receptors. Pie charts depict the percentage of expressed genes (absolute numbers next to the charts) of each receptor family (see legend on top right). Heatmaps were created separately for each receptor family using the \log_2 of the geometric mean ($n = 3$ biological replicates) of normalized counts obtained from the NanoString assay (see section "Materials and methods"). Light to dark shades indicate low to high counts (see color bar at bottom of each heatmap); white cells indicate no detection (N.D.) of transcripts (for a definition see section "Materials and methods"). Columns represent female and male moths of different mating states (V: virgin, M: mated). Rows represent receptors that were expressed in the antennae of at least one of the four groups, and are sorted first according to values in virgin females, followed by mated females, virgin males and mated males. Symbols next to receptor gene names: *, OR or IR co-receptors; ♀, female-biased receptors; ♂, male-biased receptors (FDR corrected $p < 0.05$, nested ANOVA). Mating states of males and females did not have a significant effect on receptor gene expression (FDR corrected $p > 0.05$, nested ANOVA).

the antennae (Hou et al., 2022). The function of receptors in the *IR7d* subfamily is unknown; however, the phylogenetically related *DmelIR7a* is expressed in neurons of the gustatory organs of larval and adult *D. melanogaster* (Croset et al., 2010), suggesting a gustatory function for these receptors.

Sex-dependent *IR* expression has been observed in other moth species. For instance, both female- and male-biased *IRs* have been identified in *B. mori* (Yin et al., 2021). In contrast, we did not find any significant effect of sex on *IR* expression in the antennae of *M. sexta*, although *MsexIR8a*-mediated detection of acids from larval feces is crucial for oviposition deterrence behavior, which prevents larval crowding (Zhang J. et al., 2019). This female-specific behavior seems to be independent of differential expression of the *IRs* investigated in our study.

GRs. Consistent with a previous report, we detected antennal expression of genes of two putative CO₂-receptors *MsexGR2* and *MsexGR3*, sugar receptor *MsexGR6* and bitter receptors *MsexGR41* and *MsexGR42* (Koenig et al., 2015). However, we could not confirm the antennal expression of the fructose receptor genes *MsexGR9.1*, *9.2* and *10.2*. In addition, we found transcripts of seven more bitter receptors and another putative sugar receptor *MsexGR8*, whose ortholog *BmorGR8* detects the sugar alcohol inositol, which is ubiquitously found in plants (Zhang et al., 2011).

MsexGR41 had the highest counts among all *GRs* expressed in the antennae (Figure 1C) and encodes a highly conserved and unduplicated putative bitter *GR* in the Lepidopteran order. The ortholog of *MsexGR41* in the silkworm (*BmorGR63*) is broadly expressed in different chemosensory tissues of adults and larvae (Guo et al., 2017). Similarly, *MsexGR41* transcripts were detected not only in the adult and larval antennae but also in non-chemosensory tissues such as gut, fat bodies, brain and gonads (Koenig et al., 2015), suggesting additional roles for this receptor. Functional studies of bitter receptors in Lepidopteran species are sparse (reviewed in Xu, 2020). Moreover, predictions based on sequence similarity might be difficult since bitter receptors, like *ORs*, are highly divergent and often go through lineage-specific expansions within Lepidoptera (Koenig et al., 2015).

Although putative gustatory sensilla chaetica are present on the antennae of *M. sexta* (Lee and Strausfeld, 1990), the functional roles of antennal gustatory neurons are unknown. In the moths *Heliothis virescens* and *Agrotis ipsilon*, sugar and bitter detecting gustatory neurons in the antennae influence the proboscis extension behavior (Jørgensen et al., 2007; Hostachy et al., 2019). Similarly, antennal *GRs* might also influence proboscis extension in *M. sexta*, a behavior which is dependent on both olfactory and visual cues (Raguso and Willis, 2002) and *MsexORCo* function (Fandino et al., 2019).

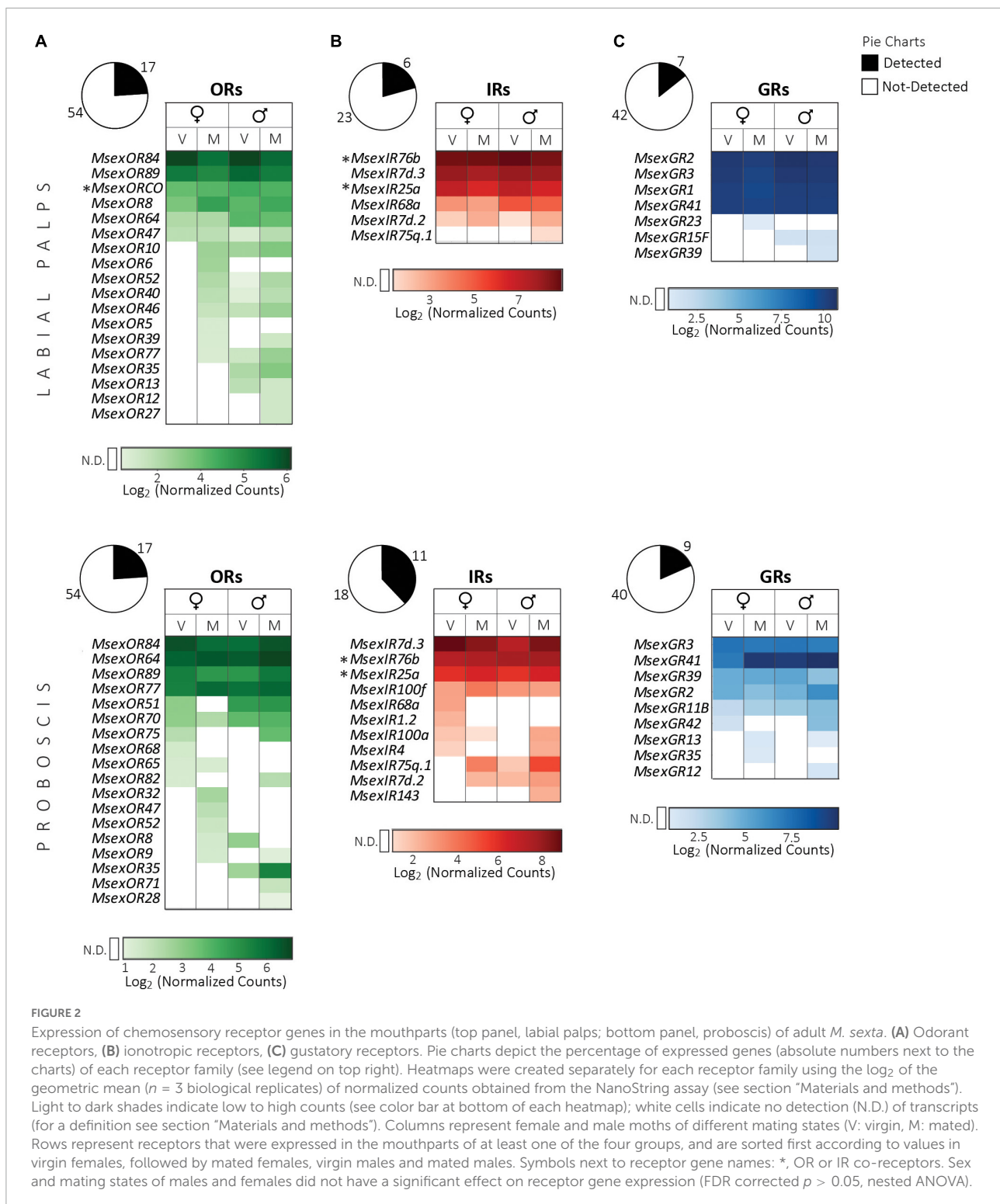
In *Drosophila*, co-expression of the two CO₂ receptors, *DmelGR21a* and *DmelGR63a*, in the fly antennae is required for CO₂ detection (Jones et al., 2007). Similarly, expression of two putative CO₂ receptor genes, *MsexGR2* and *MsexGR3*,

suggests that the *M. sexta* antennae could also detect CO₂; a function that is not reported so far. In moths, however, CO₂ detection is generally assigned to labial palps. In fact, the CO₂ sensitive glomerulus in the antennal lobe in *M. sexta* is targeted by sensory neurons in the labial pit organ of the labial palp (Guerenstein et al., 2004a). In contrast to *Drosophila*, Lepidopteran species have three *GRs* (*GR1*, *GR2* and *GR3*) in the CO₂ receptor subfamily (Robertson and Kent, 2009). Functional analysis in *Helicoverpa armigera* revealed that co-expression of *HarmGR1* and *HarmGR3* is necessary for CO₂ reception (Ning et al., 2016). Assuming that the functions of these *GRs* are conserved, the absence of expression of *MsexGR1*, the ortholog of *HarmGR1*, might actually indicate the inability of *M. sexta* antennae to detect CO₂. *MsexGR2* and *MsexGR3* might be involved in olfaction as *GRs* of the CO₂ receptor family were shown to be activated or inhibited by various odorants in *D. melanogaster* and *Aedes aegypti* (MacWilliam et al., 2018; Kumar et al., 2020).

Mouthparts

In contrast to the antennae, transcripts of only 17 *ORs* each were detected in labial palps and proboscis (Figure 2A). The labial palps in addition expressed six *IRs* and seven *GRs*, whereas in the proboscis 11 *IRs* and nine *GRs* were expressed (Figures 2B,C).

ORs. Of the 17 *ORs* expressed in each mouthpart, only seven *ORs* were common between both tissues, indicating that labial palps and proboscis have different chemosensory functions (Figure 2A). *MsexORCo* expression was consistently detected in male and female labial palp samples at high levels, whereas in the proboscis, it seemed absent. In *M. sexta*, like in several other moth species, the labial palps are so far known only to detect CO₂ (Guerenstein et al., 2004a; Ning et al., 2016). The expression of several *ORs*, like the potential linalool-receptors *MsexOR5* and *MsexOR6* (Grosse-Wilde et al., 2010), and more importantly, *MsexORCo*, strongly indicates that the labial palps of *M. sexta* might have an olfactory function beyond the detection of CO₂. We found expression of 13 *ORs* in labial palps of mated females compared to only five *ORs* in virgins. Although this difference was not significant according to the nested ANOVA we applied, it might reflect a slightly increased olfactory sensitivity of the palps to foraging- or oviposition-associated volatiles in females following mating. For the proboscis, expression of *MsexORCo* in male *M. sexta* has been demonstrated previously using RT-PCR and immunohistochemistry methods (Haverkamp et al., 2016). In our study, however, we detected *MsexORCo* transcripts only in one out of three samples each of virgin and mated males, thus not fulfilling our criterion for gene expression (see section “Materials and methods” for details). *MsexORCo* is expressed only in a single neuronal cell in a single sensillum



at the tip of the proboscis (Haverkamp et al., 2016). As we extracted RNA from the entire length of the proboscis and not only from the tip, such cell-specific expression might have gone undetected. This sensillum responds strongly to benzyl

acetone (the major component of *N. attenuata* flowers) but in addition detects other typical floral odors such as cis-jasmone, linalool, geraniol, nerol and benzyl alcohol (Haverkamp et al., 2016; Bisch-Knaden et al., 2022). *MsexOR84/89*, *MsexOR64* and

MsexOR77 were highly expressed in the proboscis and might be candidate receptors involved in the detection of these floral odors. Another intriguing feature was expression of one of the male-specific antennal pheromone receptors, *MsexOR51*, in the proboscis of female virgin *M. sexta*. This suggests that females might have multiple mechanisms of detecting their own pheromone, and that *MsexOR51* might not really be “male-specific.”

IRs. All IRs expressed in the labial palps were also detected in the proboscis (Figure 2B). In both mouthparts, the IRs with the highest counts were those encoding co-receptors, *MsexIR25a* and *MsexIR76b*, and the Lepidoptera-specific IR *MsexIR7d.3*, which were expressed in each of the tested tissues (Table 1). The IR-co-receptor *MsexIR8a*, on the other hand, was not detected in the mouthparts. This is contrary to a previous report where both *MsexIR8a* and *MsexIR25a* were shown to be expressed in the proboscis by using RT-PCR (Haverkamp et al., 2016). We detected *MsexIR8a* transcripts in only one sample of a virgin female, indicating that *MsexIR8a* is expressed in very low levels in the proboscis, and PCR amplification would be a better technique for detecting this expression. The proboscis expression of genes of three IRs from the “expanded acid-sensing cluster” (*MsexIR1.2*, *MsexIR4* and *MsexIR75q.1*) suggests that the proboscis could detect volatile carboxylic acids (Hou et al., 2022); however, based on the functional characterization of IRs from this cluster in the turnip moth *A. segetum*, such acid-sensing would require expression of the co-receptor IR8a.

Several IRs that were not detected in the antennae, were expressed in the mouthparts. These are *MsexIR7d.2*, a duplicate of *MsexIR7d.3* expressed in the antenna, and *MsexIR68a*, whose *Drosophila* ortholog is involved in hygrosensation (Knecht et al., 2017). Additionally, the proboscis expressed three more Lepidoptera-specific IRs (*MsexIR100a*, *MsexIR100f*, and *MsexIR143*; Liu et al., 2018) with unknown functions.

GRs. Four GRs were commonly expressed between labial palps and proboscis (Figure 2C). High counts of the ubiquitously expressed *MsexGR41* (Table 1) were found in both mouthparts. A distinct and high expression of all three putative CO₂ receptor genes, *MsexGR1*, 2 and 3, was observed only in the labial palps, the organ known for CO₂ detection in *M. sexta* (Guerenstein et al., 2004a). In the noctuid moth *H. armigera*, a similar high expression of the orthologous GRs, *HarmGR1*, 2, and 3, was reported for the labial palps (Guo et al., 2018). In the proboscis, like in the antennae, transcripts of *MsexGR2* and *MsexGR3*, but not of *MsexGR1* were detected. Therefore, CO₂-detection in *M. sexta* seems to be restricted to the labial palps.

All other GRs expressed in the mouthparts belong to the clade of bitter receptors (Koenig et al., 2015) and, surprisingly, no sugar or fructose receptors were detected. However, as the proboscis is considered the major gustatory organ, we expected to find expression of sugar or fructose receptors. In *H. armigera*,

for example, sugar receptors are expressed in the proboscis (Guo et al., 2018). Reiter et al. (2015) showed that gustatory neurons in the proboscis of *M. sexta* can discriminate various tastants including sucrose and other sugars, strongly suggesting the presence of sugar receptors in the proboscis. However, gustatory neurons are mainly found in the distal one-third of the proboscis (Reiter et al., 2015) and the expression level of these sugar receptors might be very low.

Legs

We detected 19, 18, and 9 OR transcripts in fore-, mid- and hindlegs respectively (Figure 3). In addition, 11, 10, and 7 IRs, and 9, 10, and 5 GRs were expressed in fore-, mid- and hindlegs, respectively.

ORs. Nine ORs were common among all three leg pairs (Figure 3A). The ORs with the highest counts in all leg pairs were *MsexOR6* and the ubiquitous *MsexOR84/89* (Table 1). The putative linalool receptor gene *MsexOR6*, which was highly expressed and female-specific in the antennae, was also highly expressed and female-specific in all three leg pairs. Transcripts of its paralog *MsexOR5*, however, was not detected in the legs. Based on our criteria (see section “Materials and methods”), the obligatory co-receptor *MsexORCo* was only expressed in forelegs of mated females. *MsexORCo* was, however, detected in one foreleg sample each of virgin females and mated males. We also note that there are 15 ORs expressed in the mated female versus only eight in virgin females (see discussion of a similar trend in female labial palps). It could be possible that expression of ORs including *MsexORCo* is upregulated in female forelegs after mating and that the forelegs might be involved in the assessment of oviposition substrates via olfaction. Transcripts of putative pheromone receptor gene *MsexOR83*, which is male-specific in the antennae, were expressed in the virgin male midlegs. This suggests that in addition to the antennae, legs may also detect pheromones.

Our results indicate a potential olfactory function of the legs of *M. sexta*. A class of chemosensory sensilla found on the *M. sexta* legs, the scattered blunt sensilla, has been hypothesized to have olfactory function due to the presence of wall pores (Kent and Griffin, 1990). Both male and female legs possess an equal number of these sensilla; however, their olfactory function still needs to be tested.

IRs. Six IRs were common among all leg pairs (Figure 3B). The transcripts of IR co-receptor genes *MsexIR25a*, *MsexIR76b* were detected in all leg pairs, whereas expression of acid-sensing co-receptor *MsexIR8a* seemed to be absent in leg tissues. However, we detected expression of IRs that potentially code for IRs tuned to acids based on their homolog’s functions: *MsexIR4*, *MsexIR75p.2*, and *Msex75q.1* (Ai et al., 2010; Rytz et al., 2013; Hou et al., 2022). This suggests that like in the mouthparts, *MsexIR8a* expression might be very low in

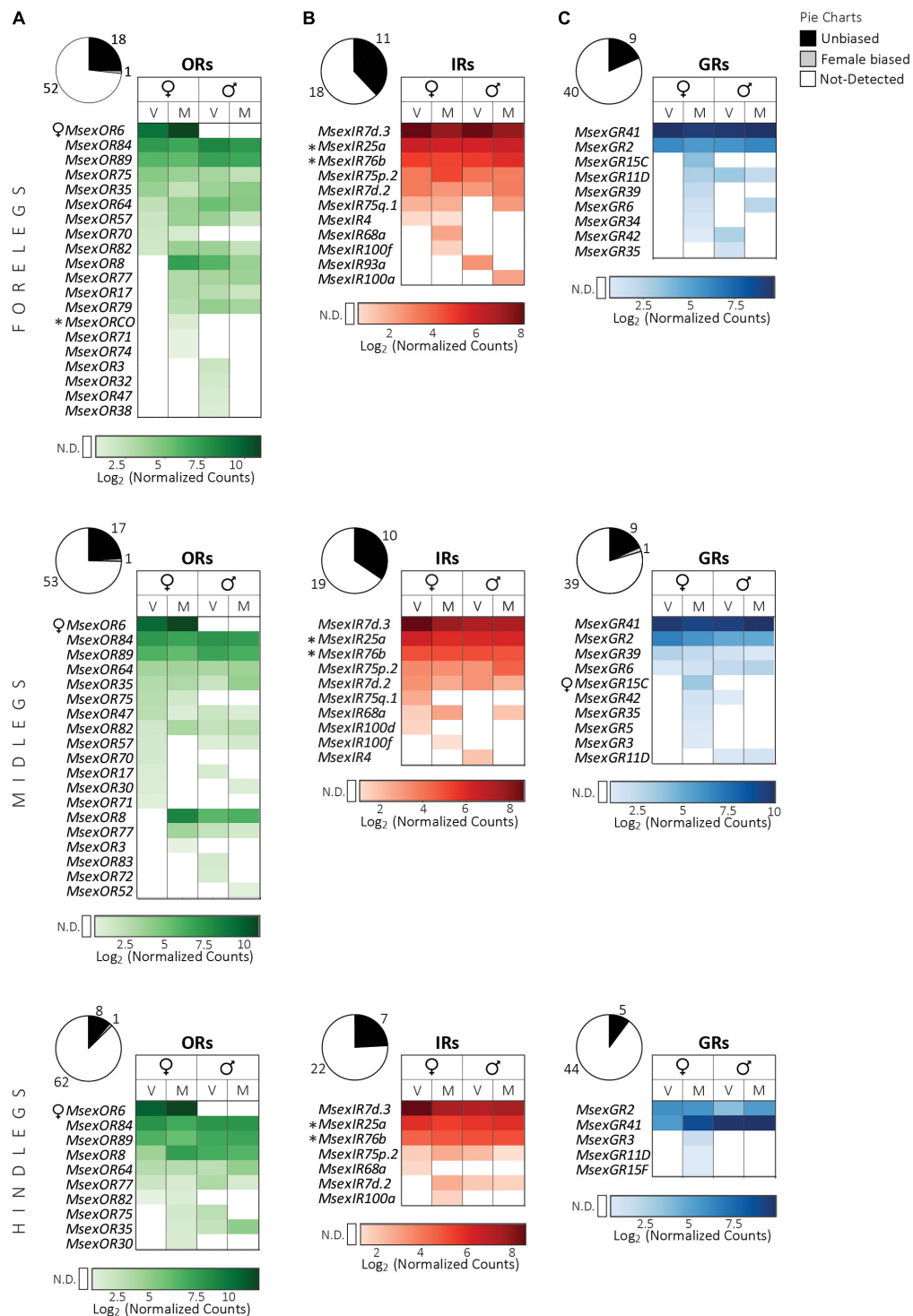


FIGURE 3
 Expression of chemosensory receptor genes in the legs of adult *M. sexta* (top panel, forelegs; middle panel, midlegs; bottom panel, hindlegs). **(A)** Odorant receptors, **(B)** ionotropic receptors, **(C)** gustatory receptors. Pie charts depict the percentage of expressed genes (absolute numbers next to the charts) of each receptor family (see legend on top right). Heatmaps were created separately for each receptor family using the \log_2 of the geometric mean ($n = 3$ biological replicates) of normalized counts obtained from the NanoString assay (see section “Materials and methods”). Light to dark shades indicate low to high counts (see color bar at bottom of each heatmap); white cells indicate no detection (N.D.) of transcripts (for a definition see section “Materials and methods”). Columns represent female and male moths of different mating states (V: virgin, M: mated). Rows represent receptors that were expressed in the legs of at least one of the four groups and are sorted first according to values in virgin females, followed by mated females, virgin males and mated males. Symbols next to receptor gene names: *, OR or IR co-receptors; ♀, female-biased receptors (FDR corrected $p < 0.05$, nested ANOVA). Mating states of males and females did not have a significant effect on receptor gene expression (FDR corrected $p > 0.05$, nested ANOVA).

the legs. Alternatively, these receptors might have a different function in the legs of *M. sexta*. In fact, some *A. segetum* homologs AsegIR75p.2 and AsegIR75q.2 did not respond to any acid, aldehyde or alcohol stimulation when co-expressed with AsegIR8a in *Xenopus* oocytes (Hou et al., 2022), either indicating that their volatile ligands were not tested, or that they have a non-olfactory function. Like in the labial palps and proboscis, the Lepidoptera-specific *MsexIR7d.2* and the putative gene for humidity sensor *MsexIR68a* are expressed in all leg pairs (Knecht et al., 2017). *MsexIR93a*, which could have both hygro- and thermosensory roles, was detected only in the forelegs (Knecht et al., 2016).

GRs. Among the three leg pairs, four GRs were commonly expressed (Figure 3C). Once again, *MsexGR41* had the highest counts in each of them. Due to the crucial gustatory function of a moth's legs, especially for oviposition, we expected high expression of several GRs. However, we detected transcripts of fewer GRs in the legs than in the antennae. A similar, unexpectedly low GR expression was also seen in the legs of *S. littoralis* (Koutroumpa et al., 2021). Interestingly, we found expression of more GRs in the legs of mated females, than in virgin females or in males. Expression of a bitter receptor transcript, *MsexGR15C*, was detected only in the fore- and midlegs of mated females, and in the midlegs we found a statistically significant, female-biased expression of this GR. This is interesting as there is evidence of sexual dimorphism in the chemosensory sensilla on the legs of *M. sexta*. A class of putative contact chemosensilla, the “spine-associated clustered sensilla”, is more abundant on the females' fore-, mid- and hindlegs than on the males' legs (Kent and Griffin, 1990). The female might have subclasses of these sensilla that express female specific receptors. Transcripts of sugar receptors *MsexGR5* and *MsexGR6* (Koenig et al., 2015) were detected in the fore- and midlegs of both sexes but not in the hindleg samples.

As in the antennae and proboscis, GRs from the CO₂ receptor group, *MsexGR2* and *MsexGR3*, were expressed. While *MsexGR2* had a consistent and high expression in males and females, *MsexGR3* was only detected in mated female mid- and hindlegs. As these receptors are known for detecting odorants rather than tastants, they could be associated with the scattered blunt sensilla on the legs.

Overall, fore- and midlegs expressed about two times more GRs than the hindlegs. After arriving at a potential hostplant via multisensory cues such as olfaction, vision, CO₂, humidity, and temperature, a gravid female *M. sexta* touches the leaves with its fore- and midlegs while still hovering in front of the plant. Such direct contact is important for the stimulation of egg laying behavior in hawkmoths (Yamamoto et al., 1969; Mechaber et al., 2002) as well as in butterflies (Staedler et al., 1995), and allows the assessment of primary and secondary plant metabolites to decide whether the plant is suitable for oviposition. Both the expressed sugar and bitter receptors might be involved in this behavior. The spines on the moth's legs may help them to

pierce through the leaf surface to allow these metabolites to come in contact with the gustatory sensilla, such as the spine-associated clustered sensilla (Kent and Griffin, 1990). In another Lepidopteran species, *Cydia pomonella*, primary metabolites, like plant surface sugars can influence egg laying decisions (Lombarkia and Derridj, 2008).

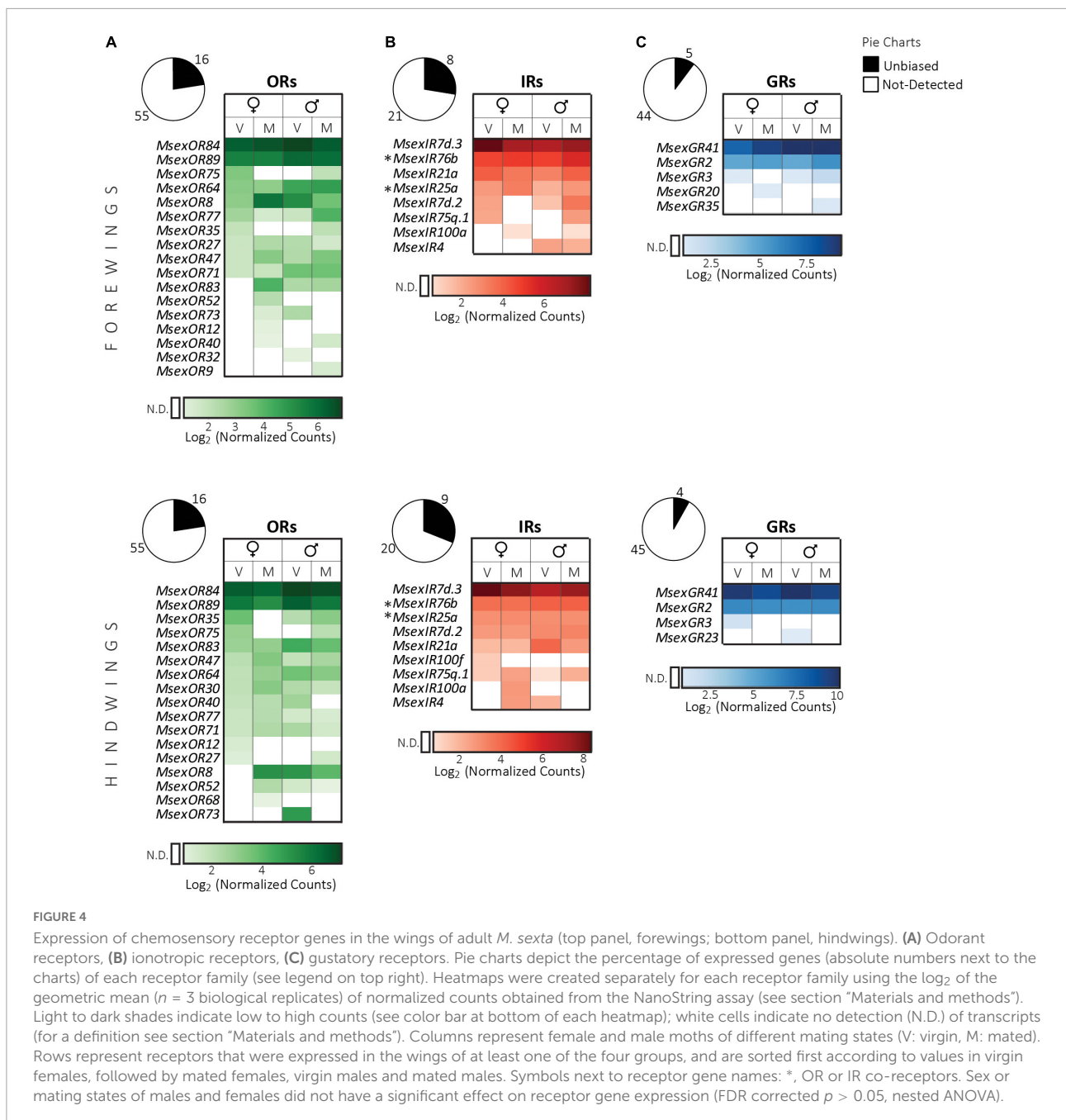
Wings

We detected expression of 16 ORs, eight IRs and five GRs in forewings and 16 ORs, nine IRs and four GRs in hindwings (Figure 4).

ORs. Of the 16 expressed ORs, 14 were common in fore- and hindwings (Figure 4A). In both wing pairs, *MsexOR84/89* transcripts had the highest counts. However, *MsexORCo* transcripts were not detected in any of the samples. Similar expression of ORs but not *ORCo* was reported for the wings of the geometrid moth *Ectropis obliqua* (Ma et al., 2016). Such expression of ORs without that of *ORCo* in the wings, suggests that ORs may have more functions than the detection of odor molecules. Alternatively, some ORs might function without *ORCo* as in the case of basal insect species that lack the *ORCo* gene (del Mármol et al., 2021). *MsexOR83*, whose transcripts were specifically detected in antennae and midlegs of males, was in addition expressed in wings of both males and females. Therefore, moth wings might detect female pheromones. For females, this information might be useful to regulate pheromone production or release. In moths, pheromone perception with wings has not been reported to the best of our knowledge. However, an IR in gustatory sensilla on the wings of *Drosophila* is required for normal mating behavior in both males and females and is thought to detect pheromonal cuticular hydrocarbons (He et al., 2019).

IRs. Except for one additional IR (*MsexIR100f*) in the hindwings, we detected expression of the same set of IR transcripts in both wings (Figure 4B). *MsexIR7d.3* transcripts had the highest counts in the wings like in other tissues. In both pairs of wings, two IR co-receptor genes, *MsexIR25a* and *MsexIR76b*, were expressed, but the third one *MsexIR8a* was not. The expression of *MsexIR25a* and *MsexIR76b* might indicate olfactory sensing of amines, or in case of *MsexIR76b*, gustatory detection of salts, sour acids and amino acids (Zhang et al., 2013; Hussain et al., 2016; Knecht et al., 2016, 2017; Chen and Amrein, 2017). Of special interest is the expression of *MsexIR21a* in fore- and hindwings of both sexes. The only other tissue where this gene was expressed was the antennae. *MsexIR21a* is potentially involved in thermosensation (cool sensing) as reported for its ortholog *DmelIR21a* (Knecht et al., 2016; Ni et al., 2016).

GRs. Three GRs that were detected in almost all tissues described so far, *MsexGR2* and *MsexGR3* of the CO₂ subfamily, and *MsexGR41*, were expressed in both wing pairs, with



MsexGR41 having the highest counts (Figure 4C). Three more bitter receptor transcripts were detected. No putative sugar or fructose receptors (Koenig et al., 2015) were found in the wings. So far, no chemosensory sensilla have been reported on the wings of *M. sexta*. GR expression was also found in the wings of *E. obliqua* including a putative sugar receptor gene (Ma et al., 2016). In *Drosophila*, functional gustatory sensilla on the wings are important for sugar and bitter detection eliciting food exploration behavior and grooming (Raad et al., 2016; Yanagawa et al., 2019).

Ovipositor

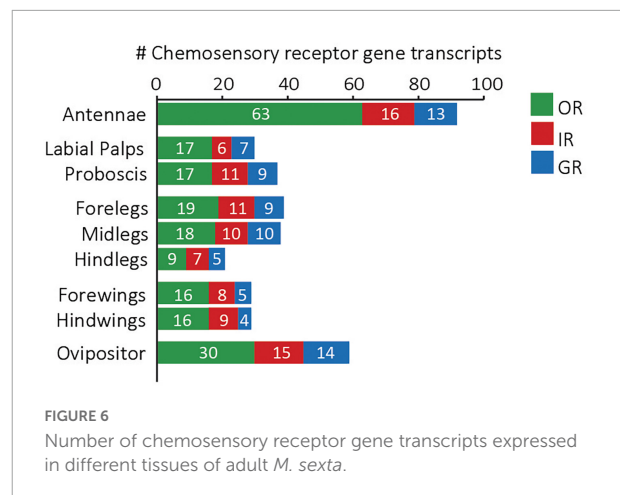
We detected 30 OR, 15 IR and 14 GR transcripts in the ovipositor (Figure 5). This is a higher number of receptor genes than was found to be expressed previously (Klinner et al., 2016). This difference might be due to different sample collection methods applied in both studies. As opposed to the previous study, where the last four abdominal segments were taken, our sample consisted only of the anal papillae at the last abdominal segment that bears the sensilla. Hence, it is possible that we were

able to detect more genes expressed in chemosensory neurons housed in these sensilla because potential receptor transcripts were more concentrated in our samples.

ORs. The highest counts were found for *MsexOR84/89* transcripts, followed by *MsexOR35* and *MsexOR75* (Figure 5A). Our results furthermore, included the three ORs that were previously reported to be expressed in the ovipositor using RNA-sequencing, *MsexOR9*, *MsexOR26*, and *MsexORCo* (Klinner et al., 2016). The expression of *MsexORCo* in our NanoString assay was detected only in the virgin female ovipositor as it was the case for the RNA sequencing data in our previous study (Klinner et al., 2016). However, *MsexORCo* expression was confirmed using RT-PCR and western blot analysis for both mating states (Klinner et al., 2016). Hence, *MsexORCo* might be expressed in both mating states, but the expression level is most probably lower in mated than in virgin females. This indicates that OR-mediated olfactory function of the ovipositor might be involved in courtship rather than in oviposition behavior. *MsexOR17*, *MsexOR26*, and *MsexOR87* that were female-biased on the antennae, were expressed in the ovipositor as well and might be involved in detection of mating or oviposition related cues.

Our result of low expression of *ORCo* and higher expression of several ORs is similar to reports from other moth species. Transcriptome sequencing together of pheromone gland and ovipositor of *Spodoptera frugiperda* revealed the expression of *SfruORCo* and 11 *SfruORs*, of which the transcript level of *SfruORCo* was the lowest (Sun et al., 2022). Furthermore, in the ovipositor of *Helicoverpa assulta*, the expression level of a tuning OR gene, *HassOR31*, is higher than that of *HassORCo* (Li et al., 2020). *HassOR31* is co-expressed with *HassORCo* in some, but not all sensilla on the ovipositor. Heterologous expression in *Xenopus* oocyte show that *HassORCo* is necessary for olfactory detection of host-plant volatiles by *HassOR31*, therefore cells where *HassORCo* is not co-expressed with *HassOR31* might use this OR in a non-olfactory function. One such example is *CpomOR1*, the pheromone receptor of *C. pomonella*, whose expression in the ovipositor has been suggested to play a role in the process of egg production and maturation (Garczynski et al., 2017). In *H. virescens*, *HvirOR13*, the receptor for its major pheromone component (Z)-11-hexadecenal was detected in the ovipositor, along with the pheromone binding protein PBP2, which is needed for the function of the pheromone receptor. Pheromone receptor and binding protein were proposed to be involved in a potential feedback mechanism (Widmayer et al., 2009). As the putative pheromone receptor gene *MsexOR83* was expressed in the ovipositor of *M. sexta*, it might accordingly be involved in the regulation of pheromone emission or in other non-chemosensory cellular functions.

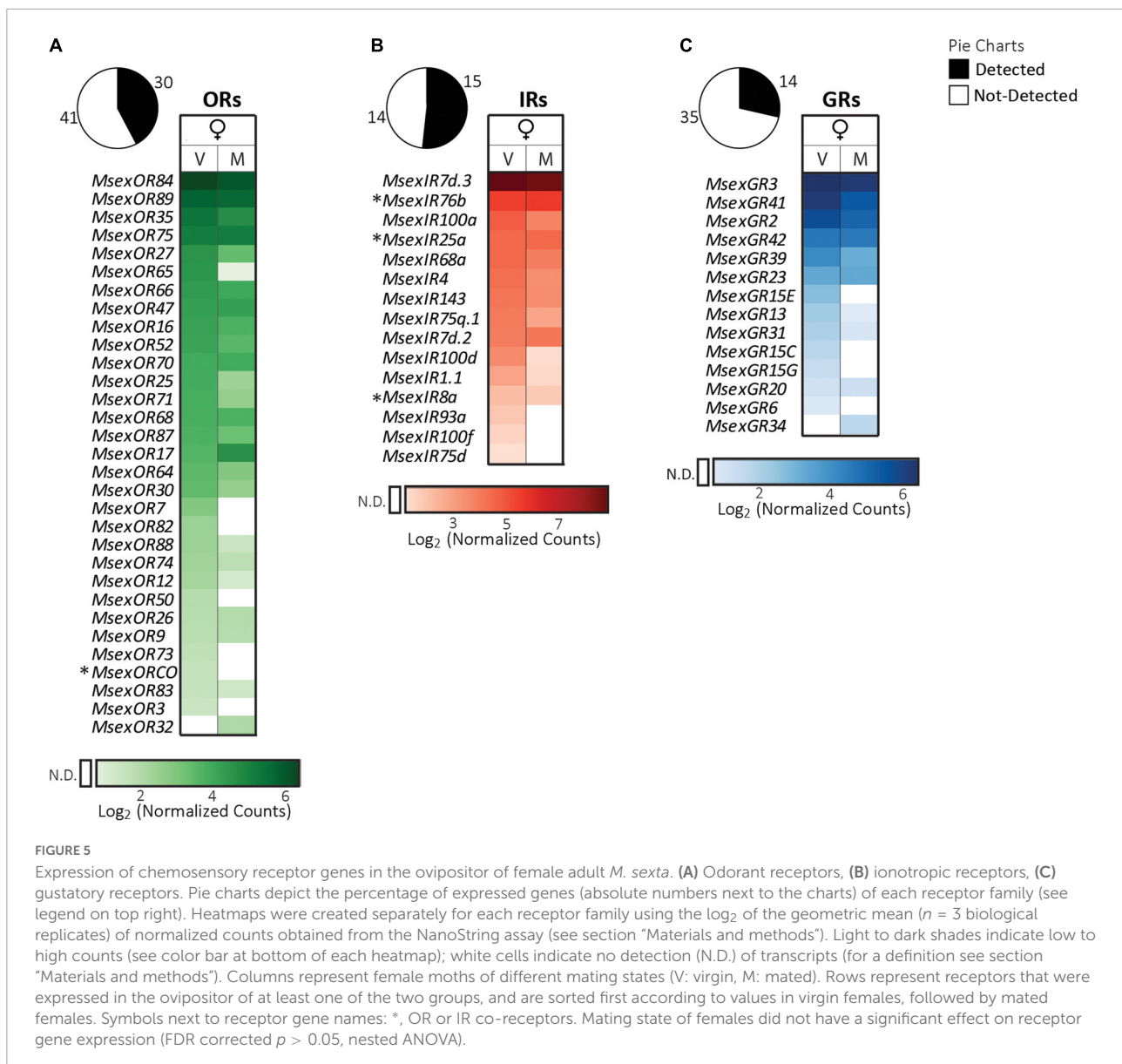
IRs. Similar to the proboscis, legs and wings, among the expressed IRs, *MsexIR7d.3* transcripts again had the highest counts (Figure 5B). Expression of all known IR co-receptors (*MsexIR8a*, *25a*, and *76b*) were detected in the ovipositor, and



also of most previously reported IRs (*MsexIR4*, *7d*, *68a*, and *75d*; Klinner et al., 2016). In fact, *MsexIR75d* expression was found exclusively in the ovipositor in our study. Based on homology with *Drosophila*, we predict that *MsexIR75d* might be involved in the detection of pyrrolidine or other amines (Silbering et al., 2011; Rytz et al., 2013), in line with the strong activation of sensilla on the ovipositor upon stimulation with pyrrolidine (Klinner et al., 2016). *MsexIR8a* is involved in acid detection (Zhang J. et al., 2019) and sensilla at the ovipositor of *M. sexta* also detect acids, especially hexanoic acid (Klinner et al., 2016). Acids play a key role in oviposition deterrence of *M. sexta* (Zhang J. et al., 2019). As the ovipositor is the only tissue apart from the antennae where we find *MsexIR8a* expression, the role of the ovipositor in acid-induced oviposition deterrence would be worth investigating. We also found expression of other IR8a-associated putatively acid-sensing IR genes: *MsexIR1.1*, *MsexIR4*, and *MsexIR75q.1* (Hou et al., 2022). Non-chemosensory IRs such as *MsexIR68a* and *MsexIR93a* were also expressed in the ovipositor. We found a very clear expression of a Lepidoptera-specific IR, *MsexIR143*, in the ovipositor. The only other tissue where this IR was expressed is the male proboscis.

GRs. *MsexGR3* and *MsexGR41* transcripts had the highest counts among all GRs in the ovipositor (Figure 5C). In agreement with a previous report, the putative genes for CO₂ receptors *MsexGR2* and *MsexGR3* were expressed in the ovipositor (Klinner et al., 2016). The gene for sugar receptor *MsexGR6* and several bitter receptors (Koenig et al., 2015) were expressed in addition, and might be involved in detecting sugars and other plant metabolites that can be used for evaluating leaf quality for oviposition.

Of the chemosensory tissues we examined, the ovipositor expressed the highest number of GRs. At the anal papillae of the *M. sexta* ovipositor, there are blunt tipped sensilla, which have both a terminal pore and wall pores, indicating olfactory and gustatory functions of the neurons residing in



them (Klinner et al., 2016). Single sensillum recordings (Klinner et al., 2016) have demonstrated only an olfactory function so far. In the moth *S. littoralis*, however, gustatory responses of ovipositor sensilla to salts, sugar and bitter compounds have been shown (Seada et al., 2016). It is possible, that gravid female moths, after landing on a potential host plant, use the spines on their legs to scratch the leaf surface and expose essential sugars and bitter secondary plant metabolites. Sensing these substances with sensilla on both the leg and the ovipositor might help the females to make an oviposition choice. As more GRs are present in the ovipositor than in the legs (Figure 6), gustation by the ovipositor might have the most important contribution to this decision-making in *M. sexta*.

Comparison among different tissues

In our results, the antennae and the ovipositor of *M. sexta* expressed every co-receptor known so far: *MsexORCO*, *MsexIR8a*, *MsexIR25a*, and *MsexIR76b* (Table 3), indicating that all pathways of odor detection take place in these organs. The labial palps and the forelegs express all co-receptors except *MsexIR8a*. The striking expression of *MsexORCO* in the labial palps suggests that these might be the second most important olfactory organs in *Manduca*, analogous to the maxillary palps in *Drosophila*. In the noctuid moth *S. frugiperda* labial palps detect other volatiles in addition to CO_2 (Chen et al., 2021), although the response profile indicates IR-mediated olfaction. Transcripts of two IR co-receptor (*MsexIR25a* and *MsexIR76b*)

TABLE 3 Expression of co-receptor genes in different tissues.

Co-receptors	No. of tissues	Tissues
<i>ORCo</i>	4	Antennae, Labial palps, Forelegs, Ovipositor
<i>IR8a</i>	2	Antennae, Ovipositor
<i>IR25a, IR76b</i>	9	Antennae, Labial Palps, Proboscis, Forelegs, Midlegs, Hindlegs, Forewings, Hindwings, Ovipositor

were detected in all nine tissues, similar to the expression patterns observed in other insects and even mollusks. These receptors are expressed in neurons of gustatory organs all over the body (Croset et al., 2010; van Schooten et al., 2016; Liu et al., 2018; Zhu et al., 2018). However, the expression of the *MsexIR25a* and *MsexIR76b* in peripheral tissues may not only indicate chemosensory activity as these genes could be involved in hygro- and thermosensation.

Apart from these two IR-coreceptors, 13 more receptors were expressed ubiquitously in all or almost all tissues (Table 1). The expression of three of these receptors (*MsexOR64*, *MsexOR75*, and *MsexOR77*) would require further validation due to possible cross-hybridization of the probes with duplicate receptor transcripts. For the remaining 12 receptors our probes did not have any off-targets. *MsexOR84/89* was broadly expressed even in tissues where no *ORCo* was detected, suggesting a possible non-olfactory function of this gene. The ortholog of this gene in the silkworm, *BmorOR23*, is expressed only in the larvae, but its function remains unknown (Tanaka et al., 2009; Koenig et al., 2015). However, expression of some ORs in several tissues including non-chemosensory ones was also detected in other moth species like *Spodoptera littoralis* (Walker et al., 2019). *MsexIR7d.2* and *MsexIR7d.3* belong to the Lepidoptera-specific IR7d group and homologs of their genes in other moth species have a broad expression in several tissues in adult and larval stages (Liu et al., 2018; Zhu et al., 2018). However, their paralog *MsexIR7d.4* was not detected in any of the tissues investigated here. *MsexGR41*, similar to its ortholog *BmorGR63*, is expressed in different chemosensory tissues (Guo et al., 2017).

In contrast to the broad expression of several genes, expression of 25 GRs were not detected in any of the tissues under investigation. We hypothesize that the low expression levels of GRs underlay the underrepresentation of this gene family in our expression data which has also hampered the detection of these genes in transcriptomes and RNA *in situ* hybridizations (Clyne et al., 2000; Koutroumpa et al., 2021). Another aspect is the spatially restricted distribution of cells expressing different receptors, for example, the segregation of *MsexORCo* and *MsexIR8a* expression in the proboscis (Haverkamp et al., 2016). Extracting RNA from smaller segments of each organ and increasing the amount of input RNA could also aid in detection of lowly expressed genes. Alternatively, these genes could be expressed in internal organs such as the brain or gut, where they might be involved in

cell signaling acting as transmembrane receptors for various molecules. In addition, many of these not expressed GRs in adults could be larvae-specific, as larval feeding behaviors are strongly influenced by gustation (Glendinning et al., 2000; Zhang Z. J. et al., 2019).

Among all the tissues investigated, we found the highest number of expressed chemosensory receptors in the antennae (92), followed by the ovipositor (59, Figure 6). The hindlegs, in contrast, showed the lowest number of chemosensory receptor genes, particularly the least number of ORs compared to all other tissues, indicating that hindlegs might have a minor role in chemoreception, a phenomenon also seen in mosquitoes (Yang et al., 2020). In female *Ae. aegypti*, electrophysiological responses from different tissues (antennae, mouthparts, wings and tarsal segments of legs) to seven insect repellents and attractants showed that all these tissues can detect odors with different capacities. The hindlegs were the least chemoreceptive as they responded only to one odor, triethylamine. This amine response is likely due to an IR25a or IR76b pathway, as these receptors are known to be broadly expressed and amine-detecting in many insects.

Furthermore, the antennae had the highest number of exclusively expressed genes, with 24 ORs, 4 IRs and 3 GRs (Table 1). One antennal GR is a putative inositol receptor gene, *MsexGR8* (Zhang et al., 2011). In *M. sexta* caterpillars inositol, an alcohol sugar that occurs in host plants (Nelson and Bernays, 1998), promotes feeding and helps overcome the inhibitory effects of bitter compounds (Glendinning et al., 2000), whereas, in adults, this GR may help identify host plants. Remarkably, *MsexGR1* showed specific expression in the labial palps, which differs from the broad expression patterns of the other two candidate CO₂ receptor genes *MsexGR2* and *MsexGR3*. If *MsexGR1* is necessary for CO₂ perception, then no tissue other than the labial palps might be able to detect it.

Sex-dependent expression

We found sex-dependent expression of chemosensory receptor genes only in the antennae (12 ORs) and legs (1 OR and 1 GR), the two organs for which sexual dimorphism has been described. In *M. sexta*, similar to other moth species, the antenna is sexually dimorphic with respect to its shape as well as the size, number and type of its sensilla (Sanes and Hildebrand, 1976; Keil, 1989; Shields and Hildebrand, 1999). For all three pairs

of legs, females have more spine-associated sensilla than males (Kent and Griffin, 1990). No sexual dimorphism is observed in the palps (Kent et al., 1986), and none has been reported for proboscis or wings.

However, sex-biased receptors in one tissue may have a sex-independent expression in other tissues. Examples are the expression of the “male-specific” putative pheromone receptor gene *MsexOR51* in the proboscis of virgin females and the expression of the “female-biased” *MsexOR17* in the legs of males. A receptor could thus have multiple functions, being involved in certain sex-specific roles when expressed in one tissue and a role common for both sexes when expressed in another tissue.

Mating-status dependent expression

Few studies have shown plasticity in insect chemosensory receptor gene expression dependent on mating state. In female *D. sukikii*, for example, several *ORs* are upregulated after mating (Crava et al., 2019). In the moth *Dendrolimus punctatus*, both up- and down-regulation have been reported for *ORs* and other chemosensory genes in both males and females (Zhang et al., 2017). In *H. armigera*, two antennal *IRs* are downregulated after mating in females (Liu et al., 2018). Such differential expression of chemosensory receptors can modulate sensitivity to essential cues and lead to changes in behavior and physiology.

For *M. sexta*, post-mating changes of odor-evoked behavioral and physiological responses have been described. Host plant volatiles induce more often a directed upwind flight and abdomen curling in mated than in virgin female *M. sexta* (Mechaber et al., 2002). In addition, some plant odor-evoked activation patterns in the antennal lobe of female *M. sexta* change after mating (Bisch-Knaden et al., 2022). However, we did not find significant differences between virgin and mated moths in the expression levels of *ORs* in the antennae or other tissues, suggesting that other mechanisms are responsible for the observed state-dependent changes in physiology and behavior.

Conclusion

We provide an extensive data set of the expression pattern of 149 chemosensory receptor genes across nine peripheral chemosensory tissues and both sexes in different mating states in adults of the hawkmoth *M. sexta*. Our data suggest that all peripheral tissues under investigation are potentially multimodal, having both olfactory and gustatory functions. Based on this information, we can predict functions or discover potential new functions for the repertoire of chemosensory receptors in different tissues of this insect species.

Data availability statement

The datasets presented in this study can be found in online repositories. The name of the repository and accession number can be found below: Edmond, <https://edmond.mpdl.mpg.de/>, doi: 10.17617/3.AEVVZV.

Author contributions

SB-K and BH: study conception and editing the manuscript. MT, LC, SB-K, and BH: study design. MT and SB: sample and data collection. MT, LC, and SB-K: data analysis. MT: writing of first draft. All authors revised the manuscript and approved the submitted version.

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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.2022.976521/full#supplementary-material>

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