Mutagenesis of the odorant receptor co-receptor (Orco) reveals severe olfactory defects in the crop pest moth Helicoverpa armigera

Xiao-Bin Fan¹,², Bao-Tong Mo¹,², Guo-Cheng Li¹,², Ling-Qiao Huang¹, Hao Guo¹,², Xin-Lin Gong¹,² and Chen-Zhu Wang¹,²*

Abstract

Background: Odorant receptors (ORs) as odorant-gated ion channels play a crucial role in insect olfaction. They are formed by a heteromultimeric complex of the odorant receptor co-receptor (Orco) and a ligand-selective Or. Other types of olfactory receptor proteins, such as ionotropic receptors (IRs) and some gustatory receptors (GRs), are also involved in the olfactory system of insects. Orco as an obligatory subunit of ORs is highly conserved, providing an opportunity to systematically evaluate OR-dependent olfactory responses.

Results: Herein, we successfully established a homozygous mutant (Orco⁻/⁻) of Helicoverpa armigera, a notorious crop pest, using the CRISPR/Cas9 gene editing technique. We then compared the olfactory response characteristics of wild type (WT) and Orco⁻/⁻ adults and larvae. Orco⁻/⁻ males were infertile, while Orco⁻/⁻ females were fertile. The lifespan of Orco⁻/⁻ females was longer than that of WT females. The expressions of most Ors, Irs, and other olfaction-related genes in adult antennae of Orco⁻/⁻ moths were not obviously affected, but some of them were up- or down-regulated. In addition, there was no change in the neuroanatomical phenotype of Orco⁻/⁻ moths at the level of the antennal lobe (including the macroglomerular complex region of the male). Using EAG and SSR techniques, we discovered that electrophysiological responses of Orco⁻/⁻ moths to sex pheromone components and many host plant odorants were absent. The upwind flight behaviors toward sex pheromones of Orco⁻/⁻ males were severely reduced in a wind tunnel experiment. The oviposition selectivity of Orco⁻/⁻ females to the host plant (green pepper) has completely disappeared, and the chemotaxis toward green pepper was also lost in Orco⁻/⁻ larvae.

Conclusions: Our study indicates that OR-mediated olfaction is essential for pheromone communication, oviposition selection, and larval chemotaxis of H. armigera, suggesting a strategy in which mate searching and host-seeking behaviors of moth pests could be disrupted by inhibiting or silencing Orco expression.

Keywords: Olfaction, Odorant receptor co-receptor (Orco), CRISPR/Cas9, Helicoverpa armigera, EAG, SSR, Chemotaxis

Background

Insects live in an ever-changing chemical world comprising a variety of odors [1]. Olfaction plays a significant role in the insect life cycle and regulates many important behaviors, such as host plant location [2], mate choice [3], oviposition site selection [4], predator avoidance [5],...
and behavioral division [6]. Insect olfaction has, thus, attracted considerable research attention.

Insects have an extremely complex olfactory system, which is responsible for the detection of different types of odors. The antennae and maxillary palps, two major olfactory organs in insects, are covered with different types of olfactory sensilla (trichoid, basiconic, and coeloconica sensilla), which contain olfactory sensory neurons (OSNs) [7, 8]. The antennal lobe (AL), which contains many gloubule-shaped neuropils known as glomeruli, is the primary olfactory processing center responsible for the integration of peripheral olfactory information [9]. Odorant receptors (ORs) expressed in the dendrite membranes of OSNs are important in the insect olfactory signal transduction pathway and have been studied in depth [10]. ORs are sensitive to many different types of compounds, including insect pheromone components [11] and host plant compounds, such as esters, alcohols, and ketones [12, 13]. Other types of chemosensory receptors, such as ionotropic receptors (IRs) and gustatory receptors (GRs), have also been identified in the insect peripheral olfactory system [14]. IRs are mainly for responding to volatile amines and acids [12, 13, 15, 16]. For example, in Drosophila melanogaster, DmIr92a is involved in sensing amines [17], and DmIr8a and DmIr64a are required for response to carboxylic acids (such as acetic acid and HCl) [16]. In addition, 2–3 GRs are involved in the detection of CO₂ in insects [18, 19]. DmGr21a and DmGr63a are involved in the rejection of CO₂ in Drosophila [20]. Using these different types of chemosensory receptors, insects can complete important behaviors such as mating, oviposition, and host selection in their natural habitats.

Insect ORs are odorant-gated ion channels that are formed by a heteromultimeric complex of the odorant receptor co-receptor (Orco) and a tuning Or [21–27]. Both Orco and ORs possess seven inverted transmembrane domains (TMDs), with an intracellular N-terminus and an extracellular C-terminus, which is opposite to the topology of vertebrate olfactory receptors that are conventional G-protein coupled receptors (GPCRs) [25]. The number of ORs varies greatly among insect species. For example, up to 400 CfloOrs are expressed in the social insect Camponotus floridanus [28], while only 10 PhumOrs are expressed in Pediculus humanus [29]. The sequence similarity of ORs is very low (the highest similarity is only 20%), whereas Orco (as an obligate partner) is highly conserved among insect species, especially toward the C terminal, and there is a single Orco gene in each insect species [30, 31].

Studies of ORs have mainly focused on the characterization of tuning profiles of each Or in Xenopus oocytes [32] and Drosophila OSNs [33]. Some ORs are narrowly tuned, while others are broadly tuned to diverse odorants and coding is combinatorial [34]. Pheromone receptors (PRs) are a subset of ORs responsible for olfactory detection of sex pheromone compounds. In general, the tuning spectrum of PRs is specific [35, 36]. For example, BmOr1 and BmOr3 are specifically tuned to bombykol and bombykal, respectively, in Bombyx mori [37, 38]. Conventional ORs are usually tuned to host plant volatiles and help insects to locate and select host plants. In B. mori, BmOr56 is specifically tuned to the compound cis-jas-mone emitted by mulberry leaves, which strongly attracts silkworm larvae [39]. Two PxylOrs (PxylOr35 and PxylOr49) in Platella xylostella are specifically tuned to isothiocyanates in cruciferous plants, guiding the adults to these plants [40]. HassOr31 is highly expressed in the ovipositor of Helicoverpa assulta and widely tuned to 12 plant odorants including Z-3-hexenyl butyrate, acting as a key chemical cue for locating oviposition sites [41]. However, in Manduca sexta, plant-seeking and oviposition behaviors of the Orco knocking-out adults were sustained although foraging and pollination behaviors were disrupted [42]. This proves that the OR-mediated olfactory responses are not the whole olfactory responses of insects. Orco, as an obligate partner, is essential for each functional Or. When Orco is knocked out, the Or functional repertoire will be abolished. Thus, Orco is a prime candidate to study OR-dependent olfactory responses in the insect olfactory system. It has been showed that the Orco family can form functional ion channels in the absence of a tuning Or, and its agonist VUAA1 is capable of gating orthologues across multiple insect taxa [43].

As a typical polyphagous insect, the cotton bollworm, Helicoverpa armigera, is an important agricultural pest globally. A total of 84 candidate OR genes have been identified in the published genomic data of H. armigera [44] and at least 65 candidate OR transcripts (1 Orco, 7 PRs, at least 55 conventional ORs), and at least 21 Irs have been identified in the transcriptome data of H. armigera [45–49]. Seven PRs are involved in sex pheromone communication between male and female H. armigera, which are mainly expressed in three types of trichoid sensilla in male antennae. Among them, type A sensilla respond to the sex pheromone component Z11-16: Ald, type B sensilla to the behavior antagonist Z9-14: Ald, and type C sensilla to another sex pheromone component Z9-16: Ald and the behavior antagonists Z9-14: Ald and Z9-16: OH [50]. To date, the functions of most PRs have been studied. HarmOr13 is tuned to Z11-16: Ald and Z9-14: Ald [51, 52], HarmOr6 to Z9-16: Ald and Z9-14: Ald [51], and Z9-16: OH [47, 53], HarmOr14b to Z9-14: Ald [11, 47, 52], and HarmOr16 to Z9-14: Ald and Z11-16: OH [52]. The functions of 29 conventional ORs have been identified, and they are mainly responsible for detecting the green leaf volatiles,
terpenes, aliphatic, and aromatic compounds of plants involved in searching for nectar and the selection of host plants in *H. armigera* [54, 55]. Most Ors have a broad tuning spectrum. For example, HarmOr60 expressed in larval antennae can respond to 25 different compounds, with cis-3-hexen-ol-1 being the most effective ligand. Some Ors are narrowly tuned, such as HarmOr42 expressed in adult antennae, which is specifically tuned to phenylacetaldehyde [54, 55]. However, the functions of more than half of the Or repertoire in *H. armigera* are still unknown, and the role of the OR-mediated olfactory pathway in the olfactory system is still not fully understood. In recent years, the CRISPR/Cas9 system has been used for functional studies of HarmOr14 [3], HarmOr16 [56], HarmOr42 [55], PBP1 [57], and SNMP1 [58] of *H. armigera*. In this study, we employed CRISPR/Cas9 gene editing to generate the Orco mutant of *H. armigera*. Next, we compared the antennal transcriptional profiles, electrophysiology, and neuroanatomical phenotypes of the mutant and WT adults. Moreover, we examined the behaviors of their adults and larvae. Taken together, our results reveal the OR-mediated olfaction play an essential role in mate searching and host plant selection of *H. armigera*.

Results

Generation of the Orco homozygous mutant of *H. armigera*

To obtain Orco mutant lines of *H. armigera*, we employed the CRISPR/Cas9 gene editing system, which has been successfully developed in many different organisms [59–61]. A single-guide RNA (sgRNA) targeting the third exon of the Orco (Fig. 1A) was designed using the

![Fig. 1 CRISPR/Cas9 directed heritable mutagenesis of Orco in Helicoverpa armigera. A Schematic of the Orco structure. Exons are shown as boxes and introns as bent lines. Shaded boxes represent the coding sequences of the Orco gene. An sgRNA targeting the sequence of exon 3 is indicated (PAM: red). A representation of mutations with a 2-bp deletion (−2 bp) is shown (Orco−2bp). B DNA sequencing chromatograms of Orco in the wild type (WT) (top), heterozygous mutant (Orco+/−) (middle), and homozygous mutant (Orco−/−) (bottom). Targeted sequences are underlined and PAM sequences (CGG) are in boxes (inverted triangles indicate the expected cleavage sites of the Cas9 protein). C Predicted secondary structure of Orco in the WT (top) and Orco in the homozygous mutant (Orco−/−) (bottom). These models were predicted using the software TOPO2 (http://www.sacs.ucsf.edu/TOPO2/) based on the methods [11].](image)
tool CHOPCHOP [62]. A mixture of sgRNA (200 ng/µL) and Cas9 protein (100 ng/µL) was co-injected into nearly 711 embryos. Then 77 injected embryos were successfully hatched, and finally, 48 G0 adults were obtained. To screen for targeted mutations, genomic DNA was extracted from one of the hind legs of each G0 adult, then confirmed by both PCR and DNA sequencing. Among 48 G0 adults, 36 G0 harbored multiple targeted mutations (Table 1), indicating a high mutation efficiency (75%) in G0 adults. Each mutated G0 was mated with three wild type (WT) heterosexual adults to produce G1 offspring. From the 36 crosses, 13 (36.1%) G0 adults produced heterozygous mutant progeny in G1, suggesting that G0 adults carried stable and heritable mutagenesis (Table 1, Additional file 1: Figure S1A). One of the Orco mutations with a 2-bp deletion at exon 3 was inferred to produce a nonfunctional, truncated Orco protein and was selected for subsequent crosses (Fig. 1B). Interestingly, no offspring were produced when homozygous mutant males were mated with homozygous mutant females or WT females but fertile offspring were produced from homozygous mutant females mated with WT males (Fig. 2). Hence, in order to obtain sufficient Orco−/− for the experiments, Orco+/− were maintained for self-crossing to screen Orco−/− in subsequent generations (Fig. 2). There were no distinct morphological differences between Orco−/− and WT adults (Additional file 1: Figure S1B). Therefore, each adult used was genotyped by sequencing prior to the experiments.

It has been reported that olfaction could regulate the longevity of Drosophila and ants [63, 64]. Thus, we compared the lifespans of Orco−/− and WT adults (Fig. 3A, B). Orco−/− females exhibited a 48% increase in median life span compared with WT females (Fig. 3A), but the lifespan of Orco−/− males was indistinguishable from that of WT males (Fig. 3B).

**Antennal transcriptome sequencing of Orco−/− and WT adults**

In order to detect transcriptional changes induced by the loss of Orco, antennal transcriptome sequencing of Orco−/− and WT adults was performed. Differentially expressed genes (DEGs) were identified with the thresholds (a log2 fold change > 1 and P value < 0.05) (Fig. 4A, B). The results clearly showed that, compared with genes from WT adults, the numbers of upregulated genes in the antennae of female and Orco−/− males were 98 and 199, respectively, and the numbers of downregulated genes were 134 and 70 (Fig. 4A, B). Gene ontology (GO) enrichment analyses were conducted for males and females. Among the 20 significantly enriched pathways in Orco−/− females, there were five related to negative chemotaxis (“induction of negative chemotaxis,”

| Table 1 Summary of the CRISPR/Cas9 directed mutations from G0 to G2 |
|---------------------------------------------------------------|
| **Total number** | **Mutated number** | **Germline mutation** |
|------------------|-------------------|----------------------|
| G0 Injected embryo | 711 | / | / |
| Hatched larvae | 77 | / | / |
| Adults | 48 | 36 (75%) | 13 (36.1%) |
| Wild type | 49 | 21 (−2 bp) | 0 |
| Heterozygotes | 5 | 9 (−5 bp) | 0 |
| Homozygous | 24 | 17 (−1, +4 bp) | 0 |
| G1 Adult | 51 | 8 (−7 bp) | 0 |
| 20 | 17 (+3 bp) | 0 |
| 34 | 41 (−12 bp) | 0 |
| 23 | 36 (−2, +7 bp) | 0 |
| 12 | 7 (−4 bp) | 0 |
| 9 | 8 (−1, +5 bp) | 0 |
| 21 | 19 (−1 bp) | 0 |
| 7 | 10 (−1, +3 bp) | 0 |
| 11 | 13 (−3 bp) | 0 |
| 33 | 21 (−8 bp) | 0 |
| Total | 299 (56.84%) | 227 (43.15%) | 0 |
| Wild type | 27 (24.54%) | 48 (43.63%) | 35 (31.82%) |
| Heterozygotes | Homozygous (−2 bp) |
“positive regulation of negative chemotaxis,” “regulation of negative chemotaxis,” “negative chemotaxis,” and “negative regulation of chemotaxis”) and three related to axon orientation (“negative regulation of axon guidance,” “negative regulation of axon extension involved in axon guidance,” and “axon midline choice point recognition”) (Fig. 4C). While in Orco−/− males, there were two allelopathy-related pathways (“olfactory receptor activity” and “neuron projection membrane”) (Fig. 4D).

We also compared the expression of 61 Ors (including 7 PRs and 54 conventional Ors), 27 Ir3s, and 9 other olfaction-related genes (including Gr1, Gr2, Gr3, TRPA1, PBP1, PBP2, PBP3, GOBP1, and GOBP2) between the antennae of Orco−/− and WT adults (Additional file 2: Table S1). For the seven PRs, only Or11 and Or6 were highly expressed in WT females. The expression level of Or11 in the antennae of Orco−/− females was 73.41% higher than that in WT females and there was no significant difference in the expression level of Or6 (Fig. 5A). The expression levels of Or6, Or11, Or13, and Or15 in the antennae of Orco−/− males were 51.55%, 31.27%, 12.87%, and 22.70% lower than those in the WT males, respectively, and the expression levels of Or14, Or14b, and Or16 were not significantly different (Fig. 5B).

For the 54 conventional Ors, the expression levels of 10 Ors were higher in the antennae of Orco−/− females than those in the WT, the expression levels of 9 Ors were lower, and there were no significant differences in the expression levels of the other 35 Ors (Fig. 5C). In Orco−/− males, the expression levels of 4 Ors were higher than those of the WT males, the expression levels of 23 Ors were lower, and the expression levels of the other 27 Ors were not changed (Fig. 5D).

For the 27 identified Ir3s, the expression level of one Ir in the antennae of Orco−/− females was higher than that in the WT females. The expression levels of 6 Ir3s were lower in the antennae of Orco−/− females, and the expression levels of the other 20 Ir3s were not significantly different (Additional file 3: Figure S2A). Compared with WT males, Orco−/− males had 5 Ir3s with higher expression levels, 6 Ir3s with lower expression levels, and 16 Ir3s with
Fig. 4 Transcriptional profiling revealed changes in gene expression after the gene *Orco* was knocked out in adult *Helicoverpa armigera* antennae. A Up-regulated and B down-regulated differentially expressed genes (DEGs) after *Orco* was knocked out in females and males. The top 20 gene ontology terms with significant enrichment in females (C) and males (D).
no difference in expression levels (Additional file 3: Figure S2B).

In addition, we also detected the expression levels of nine olfaction-related genes (Gr1, Gr2, Gr3, TRPA1, PBP1, PBP2, PBP3, GOBP1, and GOBP2). In Orco−/− females, both Gr1 and Gr2 showed lower expression levels than in WT females; TRPA1, PBP2, PBP3, GOBP1, and GOBP2 showed higher expression levels than in WT females, while the expression levels of Gr3 and PBP1 did not differ (Additional file 3: Figure S2C, D). In Orco−/− males, four genes (Gr1, PBP2, PBP3, and GOBP1) had lower expression levels than in WT males. One gene (PBP1) showed higher expression levels in WT than mutant females but the expression levels of other
genes (\textit{Gr2}, \textit{Gr3}, \textit{TRPA1}, and \textit{GOBP2}) did not differ (Additional file 3: Figure S2E, F).

In sum, the expression levels of some PRs in males were affected, while the expression levels of most Ors, Ir or, and other olfaction-related genes were not affected when \textit{Orco} was knocked out.

\textbf{Morphology of the antennal lobes in \textit{Orco}^{-/-} and WT adults}

To detect if there were changes in the neuroanatomical phenotype of the antennal lobe (AL) in \textit{H. armigera} \textit{Orco}^{-/-} adults, we compared the total volume and total number of glomeruli in the AL of \textit{Orco}^{-/-} and WT adults (Fig. 6). In both females and males, the total volume and total number of glomeruli in a single AL of \textit{Orco}^{-/-} adults did not differ from those of WT adults. The volume and number of macroglomerular complex (MGC) regions also showed no difference (Fig. 6).

\textbf{EAG responses to pheromone compounds and other odorants of \textit{Orco}^{-/-} and WT adults}

To evaluate the role of \textit{Orco} in the olfactory pathway, antennal EAG analysis was conducted on \textit{Orco}^{-/-} and WT adults. In total, 27 different types of chemical compounds were used for EAG analysis (Fig. 7; Additional file 4: Table S2). For female moths, our results clearly showed that WT females had robust antennal EAG responses to 20 plant odorants; however, limited responses to these compounds were observed in \textit{Orco}^{-/-} females (Fig. 7A). For male moths, strong antennal EAG responses to 23 compounds (including four sex pheromone-related components) were observed in WT males, while \textit{Orco}^{-/-} males were anosmic to these plant odorants and sex pheromone-related components (Fig. 7B). Furthermore, we found that both sexes of \textit{Orco}^{-/-} and WT showed no significant response to propanoic acid, acetic acid, and spermine.

\textbf{Electrophysiological responses of trichoid sensilla to pheromone-related compounds in \textit{Orco}^{-/-} and WT males}

Single sensillum recordings (SSRs) were further performed to compare antennal sensilla responses to sex pheromone compounds between WT and \textit{Orco}^{-/-} males. Following the method reported for mosquitoes [4], we processed the SSR assay with a

---

**Fig. 6** Neuroanatomical phenotypes of wild type (WT) and \textit{Orco}^{-/-} \textit{Helicoverpa armigera}. **A** Confocal images of the female WT and mutant antennal lobe (AL). **B** Volume of all glomeruli from WT (\(n = 4\)) and mutant female (\(n = 4\)) ALs. **C** Number of glomeruli in the ALs of WT (\(n = 4\)) and mutant females (\(n = 4\)). **D** Confocal images of the ALs of WT and mutant males. Areas of the macroglomerular complex (MGC) in the ALs are indicated by dotted lines. **E** Volume of all glomeruli and volume of the MGC area in ALs of WT (\(n = 4\)) and mutant males (\(n = 4\)). **F** Number of glomeruli in the ALs of WT (\(n = 4\)) and mutant males (\(n = 4\)). Confocal images were taken using 20 × 0.6 objective. Scale bars = 100 µm. Error bars, mean ± SD.
random sampling of antennal sensilla from WT and Orco−/− males. In total, responses were recorded from 72 and 70 antennal sensilla from WT and Orco−/− males, respectively. We readily located and detected three types of sensilla in WT males: 72.22% type A sensilla responded to Z11-16: Ald, 4.17% type B sensilla responded to Z9-14: Ald, 8.33% type C sensilla responded to Z9-16: Ald and Z9-14: Ald. In contrast, none of the recorded antennal sensilla responded to these sex pheromone components in Orco−/− males (Fig. 8, Additional file 5: Figure S3).
Behavioral responses to sex pheromones in Orco<sup>−/−</sup> and WT males

A wind tunnel assay was performed to detect differences in attractiveness to a blend of the main sex pheromones [Z11-16: Ald and Z9-16: Ald (97:3)] between Orco<sup>−/−</sup> and WT males. The WT males showed a robust response to the main pheromone blend, exhibiting male sexual behaviors, including flight, upwind, close, landing, and copulate (Fig. 9, Additional file 6: Video S1). The Orco<sup>−/−</sup> males also took off in the wind tunnel, but only 13% of adults exhibited upwind flight behavior. Specifically, the behaviors of close flight, landing, and copulate were completely absent in mutant males (Fig. 9, Additional file 7: Video S2).
Oviposition preferences of Orco−/− and WT females
A simple dual-choice oviposition assay was performed using Orco−/− and WT females (Additional file 8: Figure S4). The mean number of eggs was 2.6 times higher in areas exposed to fresh green pepper fruit discs than in areas with no fruit discs (Fig. 10A). The preference index was 0.44±0.07 (mean±SD) (Fig. 10B), suggesting a significant preference for oviposition on green pepper fruit discs. When Orco−/− females were tested, there was no obvious difference in the numbers of eggs between treatments (Fig. 10A); thus, the preference index was almost zero (Fig. 10B). Our results clearly suggested that Orco−/− females lost the oviposition preference for green pepper fruit discs.

Chemotaxis of Orco−/− and WT larvae
A larval chemotaxis test (Additional file 9: Figure S5) was performed to detect whether Orco is necessary for food selection by H. armigera larvae. Our results showed that eight WT larvae chose the green pepper (Fig. 11A) and the mean time to make a choice was 6±3 min (mean±SD) (Fig. 11B). However, only one Orco−/− larva chose the green pepper, four larvae chose the control, and the other three larvae remained still for 30 min (Fig. 11A). The mean time for choices was longer at nearly 21±9 min (mean±SD) (Fig. 11B).

Discussion
Insects sense a variety of volatile molecules using different types of olfactory receptor proteins (Ors, Irs, Grs, and TRPA1). As a notorious worldwide agricultural pest, H. armigera uses its complex olfactory system to find mates and nectar and select host plants. In this study, we explored the role of the OR-mediated olfactory pathway in the foraging and reproductive behavior of H. armigera,
using an Orco homozygous mutant, established using CRISPR/Cas9, in which OR-mediated olfaction was non-functional and demonstrated that OR-mediated olfaction is essential for pheromone communication, oviposition selection, and larval chemotaxis of H. armigera.

The type II CRISPR/Cas system (only one Cas9 protein required) was used in our study owing to its simplicity and precision [65, 66]. Through direct injection of synthesized sgRNA and Cas9 protein into newly laid embryos, a high mutation efficiency (75%) was obtained in G0 adults, which was similar to efficiencies reported in other insects [60, 67, 68]. After screening in the next generations, a homozygous Orco mutant line with a 2-bp deletion was obtained. There were no obvious changes in the appearance of mutant moths, so it was necessary to identify the genotype of each adult prior to the experiments using DNA sequencing of the targeted genomic region by PCR amplification. However, this method was time consuming and labor intensive. In Anopheles coluzzii and silkworm Orco mutants, this obstacle was overcome by distinguishing mutants using a DsRed visible eye color marker and fluorescent proteins, respectively [4, 69]. In future studies with H. armigera, CRISPR-based applications, such as knock-ins and transgenesis with the aid of a piggyBac transposon [70], should be explored.

Although there were no apparent morphological differences between the Orco−/− and WT adults, no offspring were produced when Orco−/− males were mated with Orco+/− females or WT females (Fig. 2). Thus, we could not maintain homozygotes and had to use another strategy (HET × HET) in subsequent generations to propagate the mutant. This phenomenon also exists in other Orco mutant insects, for example Spodoptera littoralis [71] and M. sexta [42]. There are two possible reasons for this phenomenon. Sperm activity might be impaired when Orco is knocked out in male moths. In mosquitoes and some mammalian species, some Ors and Orco are involved in the activation and (possibly) orientation of spermatozoa in male germ cells [72–74]. Another possible explanation is that Orco−/− males could not perceive sex pheromones released by female moths. At the same time, we found that Orco−/− females were able to mate with WT males in our oviposition test and healthy progeny were obtained (Fig. 2, Fig. 10). In contrast with these results, a previous study found that Orco−/− females had significantly impaired fecundity (e.g., delayed oviposition and fewer eggs). In Harpegnathos saltator ants, Orco−/− males had an equivalent mating ability to WT males [64]. An interesting question for future studies is how Orco affects the fecundity of the different sexes in these two insects.

Orco seems indispensable in detection of sex pheromone and plant odor in moth species. In B. mori, the males of BmOrco mutant displayed a significantly impaired mating selection behavior in response to the sex pheromone mixture of bombykol and bombykal (11:1), and the mutant larvae displayed defective selection for mulberry leaves and different concentrations of the volatile compound cis-jasmonate [69]. In S. littoralis, Orco knockout caused defects in plant odor and sex pheromone olfactory detection in homozygous individuals [71]. In H. armigera, it has been showed that OR-mediated olfaction is involved in plant volatile detection by adults and larvae [54, 55]. The present study shows that the antennal electrophysiological responses of the Orco−/− to sex pheromone components and plant volatile substances were abolished, the upwind flight behaviors of Orco−/− males to sex pheromones were severely reduced, the oviposition selectivity of the Orco−/− females to the host plant green pepper has completely disappeared, and the chemotaxis of larvae to the green pepper was also lost in the Orco−/− larvae. However, in M. sexta, a specialist insect with a classical interaction with Datura wrightii, the oviposition behavior was maintained in Orco−/− females, which is mainly determined by the carboxylic acids 3-methylvaleric acid and hexanoic acid from feces [42]. These acids are sensed through IR-mediated olfaction in M. sexta [75].

The heteromultimeric complex of Orco and Orx expressed in OSNs plays a crucial role in insect olfaction, but little is known about the effect of Orco knockout on the expression of Ors and other chemosensory-related proteins. In Orco−/− adults of B. mori, the expression levels of two PRs (Or1 and Or3), PBP1, PBP2, and PBP3 were decreased significantly in silkworm [69]. RNA sequencing revealed that Orco knockout caused differential expression of Or genes in the antenna of honey bees, but the expression of other types of chemoreceptor genes was generally unaffected [76]. We used transcriptional profiling to detect expression changes of Ors, IRS, IRS, TRPA1, PBP5, and GOBP3 in antennae induced by the loss of Orco. There was no significant effect on the expression levels of most olfaction-related genes, but the expression of some genes increased or decreased. For example, the expression level of Or11 and Or13 were decreased in Orco−/− males, while that of Or13 was increased in Orco−/− females. Or13 as the receptor of the major pheromone component, Z11-16: Ald is specifically expressed in male antennae, while Or11 is unbiasedly expressed in male and female antennae [11, 51]. Up to now, the function of Or11 is unknown [77]. It was reported that the OSN expressing Or13 are colocalized with the OSN expressing Or11 in the A type sensilla of males [78, 79], which may explain why the expression level of both OR11 and OR13 were decreased in males.
However, why does the opposite effects of Orco knock-out on OR11 expression in two sexes deserves further investigation. In addition, the trafficking and dendritic localization of tuning Ors were strikingly impaired in Orco Or83b mutants of D. melanogaster [25]. Whether the transportation, trafficking, and localization of these Ors from cell body to neuronal dendrites in H. armigera Orco−/− adults are scrambled or not remains unclear.

As the primary olfactory center in the insect brain, the AL has drawn a lot of attention in insect olfaction studies [80, 81]. There are numerous glomeruli (functional units of olfactory information processing) in the AL and ORNs expressing the same Or will converge on the same glomeruli [9]. The results of previous studies on changes in AL morphology caused by defective Orco were inconsistent [64]. In D. melanogaster, when Orco was knocked out, the neuroanatomy of the AL remained unchanged [27, 82]; however, in ants with expanded Ors, the number of glomeruli was severely reduced in two dependent Orco mutant alleles [64, 83]. In lepidopteran insects, the AL structure is sexually dimorphic. The MGC is specialized for processing sex pheromones in the ALs of males [84]. In M. sexta, the activity of ORNs in the MGC region was lost, the glomeruli existed, and the volume of the MGC region was reduced after Orco was knocked out [42]. In this study, we found that the number and volume of glomeruli in the AL and MGC region were very similar between the WT and Orco−/− adults (Fig. 6). These results indicate that the development of glomeruli is independent of the development of OSN in H. armigera, which is in accordance with findings for Drosophila [85]. In Drosophila, OR gene expression begins only after AL patterning, and Orco is just localized exclusively to dendrites and OSN cell bodies [27].

**Conclusions**

In summary, we established an Orco mutant using the CRISPR/Cas9 system and investigated the function of Orco in H. armigera using transcriptomics, neuroanatomy, electrophysiological recordings, and behavioral tests. Severe olfactory defects were observed in Orco−/− moths, demonstrating a crucial role of Orco (OR-dependent olfactory responses) in survival, mating, and the location of host plants. This study not only demonstrated the important multifaceted role of Orco in the H. armigera olfactory system, but also highlights potential avenues for the development of novel pest control strategies that combine knock-in or even gene drive.

**Methods**

**Insects rearing**

H. armigera was originally obtained from Zhengzhou, Henan province of China, and have been reared in Institute of Zoology, Chinese Academy of Sciences, Beijing, for successive generations. The climate chambers were maintained under a 16:8 h (light: dark) photoperiod, at 26±1 °C and 55–65% relative humidity. As previously reported [11], the larvae of H. armigera were fed on artificial diet in glass tubes. Pupae were sexed and placed in cages for eclosion. Adults were fed on 10% honey water.

**CRISPRCas9 gene editing methods**

The sequence of HarmOrco was downloaded in NCBI (https://www.ncbi.nlm.nih.gov) and confirmed by our PCR analysis. With the help of online tool CHOPCHOP (http://ch.opchop.cbu.uib.no/), single-guide RNA (sgRNA) targeting the sequence in exon 3 of HarmOrco was designed. And sgRNA was synthesized according to the manufacturer’s protocol (GeneArtTM Precision gRNA Synthesis Kit, Invitrogen). Cas9 protein was purchased from the company (Thermo Fisher Scientific, Shanghai, China).

Embryo microinjection was performed according to a reported method [4] with minor modifications. Briefly, embryos used for microinjection were collected within 1–2 h old after laid. The newly collected embryos were washed thoroughly with 10% sodium chloride solution for 1 min, followed by 5 min distilled water three times. After these washes, embryos were dried on filter paper and arranged on a coverslip using double-sided tape. A mixture of sgRNA (200 ng/µL) and Cas9 protein (100 ng/µL) was co-injected into embryos. The microinjection was performed under a Zeiss microscope. After microinjection, embryos were covered with a little flour and thereafter reared in a climate chamber to develop. Newly hatched larvae were timely transferred to artificial diet.

The genomic DNA was extracted from the hind leg of each G0 adult using the TransDirect® Animal Tissue PCR Kit (Beijing, China) as template for PCR. The gene-specific primers (forward primer; 5′-CGTAAAAAGTTATTATCG AATGGCA-3′; reverse primer; 5′-AGAAAGCAAGCT TGATGATA-3′) were designed by NCBI Primer-BLAST (http://www.ncbi.nlm.nih.gov/tools/primer-blast/). The PCR reaction consisted of 12.5 µL Premix Taq, 6.5 µL ddH2O, 2 µL DNA template, 2 µL forward primer, and 2 µL reverse primer. Reaction conditions were as follows: denaturation for 5 min at 98 °C and 35 cycles of denaturation at 98 °C for 10 s, annealing at 55–60 °C for 30 s, and extension at 72 °C for 40 s. PCR products encompassing the target Orco sequence were sequenced to determine genotypes.
Comparisons of lifespan between the Orco−/− and WT of H. armigera
There were totally four groups used for lifespan statistics experiments: WT females, WT males, mutant females, and mutant males. In each group, thirty adults that emerged on the same day were placed in a cage and fed on 10% honey water. The number of surviving individuals in each group was counted daily until all individuals died. Three biological repetitions were performed.

Antennal transcriptome sequencing from the Orco−/− and WT of H. armigera
Antennae from males and females of the Orco−/− and WT of H. armigera were collected separately for RNA-seq. All the insects were reared in the same condition, and the unmated three-day-old adults were used. In each group, antennae were dissected from nearly 30 moths and quickly stored at 80 °C until RNA extraction. Three biological replicates were conducted in each group. Total RNA was firstly extracted according to a previous work [86], then evaluated with Agilent 2100 BioAnalyzer (Agilent, USA) and an ND-2000 spectrophotometer (Nanodrop, Wilmington, DE, USA). High-quality RNA was used for cDNA library construction. And these libraries were sequenced on an Illumina HiSeq2000 platform with the Nova-PE150 mode. Clean data were de novo assembled with software Trinity [87]. Blast nucleotide database was created from the assembled Trinity.fasta file and were queried by the protein sequence file of OR genes annotation repertoire of H. armigera [44]. The putative OR transcripts were obtained with BLAST, and expression abundances were calculated with the RSEM package [88]. Differentially expressed genes (DEGs) were screened by threshold (a log2 fold change > 1 and p value <0.05) though software DESeq2 [89]. The GO function enrichment of DEGs was completed by Omicshare tool (https://www.omicshare.com/).

Comparisons of antennal lobe between Orco−/− and WT adults
The neuroanatomy of antennal lobes from homozygous mutant (Orco−/−) and wild type moths were examined following a protocol previously published [90]. Brains were firstly dissected in phosphate-buffered saline (PBS) and then fixed in 4% paraformaldehyde overnight at 4 °C. Glomeruli were marked using the antibody SYN-ORF1 (Developmental Studies Hybridoma Bank, IA, USA) antibody and visualized with Alexa Fluor 488 goat anti-mouse secondary antibody (Invitrogen). Photos were obtained with a Zeiss LSM710 Meta laser scanning microscope (Zeiss, Oberkochen, Germany). The software AMERA 6.0 (ZIB, Germany) was used for construct brain atlas. Four adults were observed in each group.

Chemicals and electrophysiology analysis
According to our previous method [91], antennal EAG (electroantennograms) responses using antennae from 3- to 5-day-old unmated wild type and mutant adults (females and males) were conducted. A total of 27 odors including four previously identified pheromone gland components (Z11-16:Ald, Z9-16:Ald, Z9-14:Ald, Z11-16:OH) [50, 92], two green leaf volatiles (hexanal, trans-2-hexenal), four terpenoids (geraniol, farnesene, citral, linalool), seven aliphatic compounds (nonanal, 6-methyl-5-hepten-2-one, 2-octanone, acetoin, allyl-isothiocyanate, acetonphene, and 1-octanol), seven aromatic compounds (methyl salicylate, benzyl acetate, methyl phenylacetate, methyl benzoate, eugenol, ethyl benzoate, and benzaldehdye), two acids (acetic acid, propionic acid), and one amine (spermine) (Additional file 4: Table S2). All chemicals were diluted with paraffin oil, and the concentrations were set up as µg/µL. Briefly, 10 µL of each chemical was added into a filter paper (0.2 cm × 1 cm) in a pastre tube, and 10 µL paraffin oil was used for negative control. Ten repeats were conducted for each odorant. A computer was coupled with an IDAC-2 amplifier and used for record data, and all EAG data were analyzed with software EAG2000 (Syn-tech, Hilversum, the Netherlands).

Single sensillum recordings (SSRs) were conducted only using 3- to 5-day-old, unmated males from homozygous mutant (Orco−/−) and wild type. Following the previously described methods in H. armigera [93], six previously identified pheromone gland components (Z11-16: Ald, Z9-16: Ald, Z9-14: Ald, Z11-16: OH, Z9-16: OH, Z11-16: Ac) [93] were detected in our experiments. Similar with EAG responses experiments, odorant cartridges were made by loading 10 µL of stimulus (10 µg/µL) onto a filter paper. Briefly, a moth was fixed in a 1 mL disposable Eppendorf pipette tip with a narrow end cut. The head and antenna of the moth were protruded from the narrow end, then immobilized by the dental wax. An electrolytically sharpened tungsten electrode was inserted into the base of a single sensillum in the antenna and used for record signals. Another tungsten electrode was inserted into a compound eye of the moth as a reference electrode. For odorant delivery, a flow of humidified and
purified air continuously blew to the antenna (12.5 mL/s) from a 15-cm-long steel tube. When tested, a 200-ms air pulse was generated after a stimulus was added into the air stream using a stimulus flow controller (CS-55, Syntech, Hilversum, the Netherlands) and the air pulse was delivered with a stable flow rate of 10 mL/s. The recorded signals were displayed through the IDAC interface amplifier (IDAC-4, Syntech, Hilversum, the Netherlands) and further analyzed with the software Autospike, version 3.4 (Syntech, Hilversum, the Netherlands).

Wind tunnel assay of male adults
A mixture of Z11-16: Ald and Z9-16: Ald at a ratio of 97:3 (the principal sex pheromone components in *H. armigera* [92]) was used in our wind tunnel assay. Different from above electrophysiology analysis, the binary sex pheromone components (Z11-16: Ald, Z9-16: Ald) were dissolved in hexane and the concentration was 0.1 µg/µL. In order to adapt experiment conditions (0.45 lx of red light, 22–25 °C, and 20–40% RH), each virgin male (4-day-old) was placed in a wind tunnel (size: 2.5 m × 1 m × 1 m (L × W × H)) for at least 30 min prior to the experiment.

When tested, 10 µL solution was added to the filter paper and the wind speed was at 0.5 m/s. One virgin male was transferred into a cylindrical mesh cage, and their behaviors were observed for 5 min. Five different behavioral responses were recorded: (1) Flight: male moth quit out of the cage and took off; (2) Upwind: male moth flew against the wind or shown a zigzag pursuing flight model; (3) Close: male moth flew close to the pheromone source less than 10 cm; (4) Landing: male moth contacted and landed on the filter paper; (5) Copulate: male moth showed a series of behaviors such as curling abdomen, stretching the hair-pencils from the abdominal cavity. Thirty repetitions were carried out for both homozygous mutant (*Orco−/−*) and wild type.

Oviposition choice analysis of female adults
In order to detect whether oviposition preference of female moth was impaired in homozygous mutant (*Orco−/−*), a dual-choice assay was conducted following the method described [4]. At the beginning, newly emerged mutant females or wild type females were respectively mated with wild type males for 3 days (sex ratio = 1:3). Then 3 gravid females were transferred into a cylinder cage (size: diameter 24 cm, height 26 cm) to lay eggs. A piece of gauze covered on the top side of cage as an oviposition substrate and was equally divided into four sections. When scotophase begins, two 1.5-cm-diameter fresh green pepper fruit discs were placed on the opposite corners of cage, no fruit discs were placed on another two corners as control. To avoid the contact clues, a stainless net shelf was placed between gauze and green pepper fruit discs (Additional file 8: Figure S4). After 24 h, numbers of eggs were manually counted in different oviposition areas and a new piece of gauze was placed on the cage. This dual-oviposition assay was conducted for 3 days, and the mean number of eggs was recored. Five repetitions were conducted for both mutant females and wild type females. According to the formula ((T−C)/(T+C)) [4], preference index was calculated.

Larval chemotaxis analysis
Larval chemotaxis experiments were performed according to the previously reported work in silkworm [39] with some modifications. Four 1.5-cm-diameter fresh green pepper fruit discs and four 1.5-cm-diameter filter papers were arranged at opposite ends in a closed box (30 cm × 20 cm × 10 cm). A single fifth-instar larva was starved for 2 h prior to the onset of the experiment and was placed in the middle of the box. A video camera was used to track the behavior trajectory of larva (Additional file 9: Figure S5). Once the larva touched the green pepper or filter papers, the video recording was stopped. Each larva was recorded and observed up for 30 min. Videos were analyzed with the software EthoVision XT (v.11.5, Noldus Information Technology). Eight biological replications were conducted for each genotype.

Statistical analysis
Multiple *t*-test was used to analyze the below experiments (comparisons of lifespan between wild types and *Orco−/−* adults, the comparisons of genes expression levels between wild types and *Orco−/−* adults, comparisons of antennal lobe between wild types and *Orco−/−* adults, SSR data, wind tunnel assay of male adults, oviposition choice analysis of female adults, and larval chemotaxis analysis) with *, *P* < 0.05; **, *P* < 0.01; ***, *P* < 0.001. ANOVA followed by Tukey’s test was used for the comparisons of antennal EAG responses between wild types and *Orco−/−* adults. All statistical analyses were conducted by the software GraphPad Prism 7.

**Abbreviations**
Orco: Odorant receptor co-receptor; Ors: Odorant receptors; IrIs: Ionotopic receptors; Grs: Gustatory receptors; CRISPR: Clustered regularly interspaced short palindromic repeats; Cas9: CRISPR-associated protein 9; *H. armigera*: Helicoverpa armigera; AL: Antennal lobe; MGC: Macroglomerular complex; Z11-16: Ald: (Z)-11-hexadecenyl acetate; Z9-16: Ald: (Z)-9-hexadecenyl acetate; Z19-16: Ald: (Z)-19-hexadecenyl acetate; Z9-16:OH: (Z)-9-hexadecenol; Z11-16:OH: (Z)-11-hexadecenol; Z11-16:Ac: (Z)-11-hexadecenyl acetate; EAG: Electroantennogram; SSR: Single sensillum recording; ORNs: Odorant receptor neurons; GPCRs: G-protein coupled receptors; PRs: Pheromone receptors; sgRNA: Single-guide RNA; WT: Wild type; DEGs: Differentially expressed genes; GO: Gene ontology; TPM: Transcripts per million; TRPA1: Transient receptor potential cation channel subfamily A member 1; PBP: Pheromone-binding protein; GOBP: General odorant-binding protein; ANOVA: Analysis of variance.
Supplementary Information

The online version contains supplementary material available at https://doi.org/10.1186/s12195-022-01411-2.

Additional file 1. Figure S1. (A) Sequences of different mutations in G1 from G0 (germinal mutations occurred in these G0 individuals). The top sequence represented wild type. The target sequences were in bold and PAM sequences (CGG) was highlighted with red. Expected cut sites were marked by a red triangle. Among these mutations, deletions were indicated by dotted lines and insertions by red lower case letters. (B) Phenotypes of homozygous mutant (Orco−−) and wild type adults.

Additional file 2. Table S1. The TPM value of main olfactory related genes in Orco−− and wild type.

Additional file 3. Figure S2. (A, B) TPM of irs in females and males. (C, D, E, F) TPM of other olfactory related genes in females and males. Multiple t test was used with *, P < 0.05; **, P < 0.01; ***, P < 0.001 as significant differences. Error bars, mean ± SD.

Additional file 4. Table S2. Summary of the chemicals used in our experiments.

Additional file 5. Figure S3. Summary of the recorded sensilla types according to the response profiles (SO) in wild type (n = 72) (A) and homozygous mutant males (n = 70) (B). Type A sensilla only responded to Z11-16:Ac and while type B sensilla responded to Z9-14:Ac. Type A sensilla responding to Z9-16:Ac and Z9-14:Ac were classified as type C. The remaining sensilla were considered as “others” in pie chart.

Additional file 6. Video S1. A typical wind tunnel trajectory of a wild type male.

Additional file 7. Video S2. A typical wind tunnel trajectory of an Orco−− male.

Additional file 8. Figure S4. The set-up of oviposition choice tests (A, B) and the spread of eggs laid by mated females: (C) wild type females, (D) Orco−− females.

Additional file 9. Figure S5. Setup for behavioral tracing of a larva in a dual-choice arena system.

Acknowledgements

We thank our colleagues Hao Yin for his assistance in insect rearing, Feng-Ming Wu for her helps in dissection of antennal lobes, Ke Yang for his suggestion in bioassays, and Jun Yang for his helps in visualization.

Authors’ contributions

XBF: Conceptualization; Data curation; Software; Formal analysis; Validation; Investigation; Visualization; Methodology; Writing—original draft. BTM: Conceptualization; Resources; Data curation; Supervision; Funding acquisition; Validation; Methodology; Project administration; Writing—original draft; Writing—review and editing. All authors read and approved the final manuscript.

Funding

This work was supported by the National Natural Science Foundation of China (Grant No. 31830088 and 32130090) and Strategic Priority Research Program of Chinese Academy of Sciences (Grant No. XDBB16).

Availability of data and materials

All data generated or analyzed during this study are included in this published article, its supplementary information files and publicly available repositories. Sequencing data generated in this study has been uploaded to SRA database of NCBI under PRJNA813394.

Declarations

Ethics approval and consent to participate

Not applicable.

Consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

Author details

1 State Key Laboratory of Integrated Management of Pest Insects and Rodents, Institute of Zoology, Chinese Academy of Sciences, 1 Beichen West Road, Chaoyang District, Beijing 100101, People’s Republic of China. 2 CAS Center for Excellence in Biotic Interactions, University of Chinese Academy of Sciences, Beijing, People’s Republic of China.

Received: 3 April 2022 Accepted: 16 September 2022

Published online: 30 September 2022

References

1. George J, Robbins PS, Alessandro RT, Stelinski LL, Rapinote SL. Formic and acetic acids in degradation products of plant volatiles elicit olfactory and behavioral responses from an insect vector. Chem Senses. 2016;41(4):325–38. https://doi.org/10.1093/chemse/bjw005.

2. Riffell JA, Alkon R. Multimodal floral signals and moth foraging decisions. PLoS ONE. 2013;8(8):e72809. https://doi.org/10.1371/journal.pone.0072809.

3. Cao S, Huang T, Shu J, Liu Y, Wang G. An orphan pheromone receptor affects the mating behavior of Helicoverpa armigera. Front Physiol. 2020;11:413. https://doi.org/10.3389/fphys.2020.00413.

4. Sun H, Liu F, Ye Z, Baker A, Zwiebel LJ. Mutagenesis of the orco odorant receptor co-receptor impairs olfactory function in the malaria vector Anopheles coluzzi. Insect Biochem Mol Biol. 2020;127: 103497. https://doi.org/10.1016/j.ijimbmb.2020.103497.

5. Leonhardt SD, Menzel F, Nehring V, Schmitt T. Ecology and evolution of communication in social insects. Cell. 2016;164(6):1277–87. https://doi.org/10.1016/j.cell.2016.01.035.

6. Liebig J, Peeters C, Oldham NJ, Markstadter C, Hohloobler B. Are variations in cuticular hydrocarbons of queens and workers a reliable signal of fertility in the ant Harpegnathos saltator? Proc Natl Acad Sci U S A. 2000;97(8):4124–31. https://doi.org/10.1073/pnas.97.8.4124.

7. Su CY, Menuz K, Carlson JR. Olfactory perception: receptors, cells, and circuits. Cell. 2009;139(1):45–59. https://doi.org/10.1016/j.cell.2009.09.015.

8. Vosshall LB, Stocker RF. Molecular architecture of smell and taste in Drosophila. Annu Rev Neurosci. 2007;30:505–33. https://doi.org/10.1146/annurev.neuro.30.051606.094306.

9. Vosshall LB, Wang AM, Axel R. An olfactory sensory map in the fly brain. Cell. 2000;102(2):147–59. https://doi.org/10.1016/s0092-8674(00)00021-0.

10. Vosshall LB, Amrein H, Morozov PS, Rzhetsky A, Axel R. A spatial map of olfactory receptor expression in the Drosophila antenna. Cell. 1999;96(5):725–36. https://doi.org/10.1016/s0092-8674(00)80582-6.

11. Yang K, Huang LQ, Ning C, Wang CZ. Two single-point mutations shift the ligand selectivity of a pheromone receptor between two closely related moth species. eLife. 2017;6. https://doi.org/10.7554/eLife.21910.

12. Silbering AF, Ryttz R, Grosjean Y, Albuin L, Ramdya P, Jefferis GS, Benton R. Complementary function and integrated wiring of the evolutionarily distinct Drosophila olfactory subsystems. J Neurosci. 2011;31(38):13357–75. https://doi.org/10.1523/JNEUROSCI.2360-11.2011.

13. Yao CA, Ignell R, Carlson JR. Chemosensory coding by neurons in the coelomic sensilla of the Drosophila antenna. J Neurosci. 2003;23(3):8359–67. https://doi.org/10.1523/JNEUROSCI.2432-02.2003.

14. Montell C. Drosophila sensory receptors-a set of molecular Swiss Army Knives. Genetics. 2021;217(1):1–34. https://doi.org/10.1039/genetics/aya01171.

15. Hussain A, Zhang M, Ucpunar HK, Svensson T, Quillery E, Gompel N, Ignell R, Grunwald Kadow IC. Ionotropic chemosensory receptors mediate the taste and smell of polyamines. PLoS Biol. 2016;14(5): e1002454. https://doi.org/10.1371/journal.pbio.1002454.

16. AI M, Min S, Grosjean Y, Leblanc B, Bell R, Benton R, Suh GS. Acid sensing by the Drosophila olfactory system. Nature. 2010;468(7324):691–5. https://doi.org/10.1038/nature09537.
17. Min S, Ai M, Shin SA, Suh GS. Dedicated olfactory neurons mediating attraction behavior to ammonia and amines in Drosophila. Proc Natl Acad Sci U S A. 2013;110(14):E1321–23. https://doi.org/10.1073/pnas.1215880110.

18. Jones WD, Caygillou P, Kadog IW, Vosshall LB. Two chemosensory receptors together mediate carbon dioxide detection in Drosophila. Nature. 2007;445(7123):86–90. https://doi.org/10.1038/nature05466.

19. Kwon JY, Dahanukar A, Weiss LA, Carlson JR. The molecular basis of CO2 reception in Drosophila. Proc Natl Acad Sci U S A. 2007;104(9):3574–8. https://doi.org/10.1073/pnas.0700791104.

20. Faucher C, Forstreuter M, Hilker M, de Bruyne M. Behavioral responses of Drosophila to pheromonal levels of carbon dioxide depend on life-stage, sex and olfactory context. J Exp Biol. 2006;209(14):2739–48. https://doi.org/10.1242/jeb.02297.

21. Butterwick JA, Del Marmol J, Kim KH, Kahlos MA, Rogaw JW, Wala T, Ruta V. Cyto-EM structure of the insect olfactory receptor Orco. Nature. 2018;560(7719):447–52. https://doi.org/10.1038/s41586-018-0420-z.

22. Del Marmol J, Yedlin MA, Ruta V. The structural basis of odorant recognition in insect olfactory receptors. Nature. 2021;570(7764):126–31. https://doi.org/10.1038/s41586-021-03794-8.

23. Neuhaus EM, Gisselmann G, Zhang W, Dooley R, Stortkuhl K, Hatt H. Odorant signals mediated by specific combinations of olfactory receptors. Science. 2011;334(6059):1365–2583. 2010. 01045.x.

24. Sato K, Pellegrino M, Nakagawa T, Vosshall LB, Touhara K. Sensory receptors mediate one-way signaling in Drosophila. Proc Natl Acad Sci U S A. 2007;104(22):9125–30. https://doi.org/10.1073/pnas.0702120104.

25. Benton R, Sachse S, Michnick SW, Vosshall LB. Atypical membrane topology and heteromeric function of Drosophila odorant receptors in vivo. PLoS Biol. 2006;4(2):e20. https://doi.org/10.1371/journal.pbio.0040020.

26. Wicher D, Schafer R, Bauernfeind R, Stensmyr MC, Heller R, Heinemann SF. Insect olfactory receptors are heterodimers of ligand-gated ion channels. Nature. 2008;452(7190):1002–6. https://doi.org/10.1038/nature06850.

27. Del Marmol J, Yedlin MA, Ruta V. The structural basis of odorant recognition in insect olfactory receptors. Nature. 2021;570(7764):126–31. https://doi.org/10.1038/s41586-021-03794-8.

28. Zhang J, Wang B, Dong S, Cao D, Dong J, Walker WB, Liu Y, Wang G. Antennal transcriptome analysis and comparison of chemosensory gene families in two closely related noctuid moths, Helicoverpa armigera and H. assulta. PLoS ONE. 2015;10(2):e0117054. https://doi.org/10.1371/journal.pone.0117054.

29. Liu NY, Xu W, Papanicolaou A, Dong SL, Anderson A. Identification and characterization of three chemosensory receptor families in the cotton bollworm Helicoverpa armigera. BMC Genomics. 2014;15:597. https://doi.org/10.1186/1471-2164-15-597.

30. Kirkness EF, Haas BJ, Sun W, Braig HR, Perotti MA, Clark JM, Lee SH, Roberts TP. Helicoverpa armigera and H. assulta: considerations for pest management and control. J Insect Physiol. 2018;101:21–33. https://doi.org/10.1016/j.jinsphys.2018.01.020.

31. Liu Y, Gu SH, Zhang YJ, Guo YY, Wang GR. Candidate olfaction genes in the cotton bollworm Helicoverpa armigera. Insect Biochem Mol Biol. 2018;560:447–52. https://doi.org/10.1038/s41586-018-0420-z.

32. Liu Y, Gu SH, Zhang YJ, Guo YY, Wang GR. Candidate olfaction genes in the cotton bollworm Helicoverpa armigera. Insect Biochem Mol Biol. 2018;560:447–52. https://doi.org/10.1038/s41586-018-0420-z.

33. Liu Y, Gu SH, Zhang YJ, Guo YY, Wang GR. Candidate olfaction genes in the cotton bollworm Helicoverpa armigera. Insect Biochem Mol Biol. 2018;560:447–52. https://doi.org/10.1038/s41586-018-0420-z.

34. Bisch-Knaden S, Dahake A, Sachse S, Knaden M, Hansson BS. Spatial representation of feeding and oviposition odors in the brain of a hawkmoth. Cell Rep. 2018;22(2):2482–92. https://doi.org/10.1016/j.celrep.2018.01.082.

35. Wang G, Vazquez GM, Schal C, Zwiebel LJ, Gould F. Functional characterization of chemosensory genes in Drosophila and the tobacco budworm Heliothis virescens. Insect Mol Biol. 2011;20(1):125–33. https://doi.org/10.1111/j.1365-2918.2010.00454.x.

36. Cao S, Liu Y, Wang B, Wang G. A single point mutation causes one-way alteration of chemosensory receptor function in two Heliotris species. iScience. 2021;24(9):103293. https://doi.org/10.1016/j.isci.2021.103293.

37. Nakagawa T, Sakurai T, Nishihata T, Touhara K. Sensory information mediated by specific combinations of olfactory receptors. Science. 2005;307(5715):1638–42. https://doi.org/10.1126/science.1106267.
58. Ye ZF, Liu XL, Han Q, Liao H, Dong XT, Zhu GH, Dong SL. Functional characterization of PBP1 gene in Helicoverpa armigera (Lepidoptera: Noctuidae) by using the CRISPR/Cas9 system. Sci Rep. 2017;7(1):8470. https://doi.org/10.1038/s41598-017-08769-2.

59. Liu S, Chang H, Liu W, Cui W, Liu Y, Wang Y, Ren B, Wang G. Essential role for SNMP1 in detection of sex pheromones in males. Insect Biochem Mol Biol. 2020;127:105845. https://doi.org/10.1016/j.ibmb.2020.105845.

60. Li X, Fan D, Zhang W, Liu G, Zhang L, Zhao L, Fang X, Chen L, Dong Y, Chen Y, et al. Outbred genome sequencing and CRISPR/Cas9 gene editing in butterflies. Nat Commun. 2015;6:2812. https://doi.org/10.1038/ncomms12912.

61. Li Y, Zhang J, Chen D, Yang P, Jiang F, Wang X, Kang L. CRISPR/Cas9 in locusts: Successful establishment of an olfactory deficiency line by targeting the mutation of an odorant co-receptor (Orco). Insect Biochem Mol Biol. 2016;79:27–35. https://doi.org/10.1016/j.ibmb.2016.10.003.

62. Shalem O, Sanjana NE, Hartenstein E, Shi X, Scott DA, Mikkelsen T, Heckl D, Ebert BL, Root DE, Doench JG, et al. Genome-scale CRISPR-Cas9 knockouts screening in human cells. Science. 2013;343(6166):84–7. https://doi.org/10.1126/science.1247055.

63. Libert S, Zwiener J, Chu X, Vanvoorhies W, Roman G, Pletcher SD. Regulation of Drosophila life span by olfaction and food-derived odors. Science. 2007;318(5853):1133–7. https://doi.org/10.1126/science.1143660.

64. Yan H, Opalchaleomphan C, Mancini G, Yang H, Galillo M, Miejsnek J, Leitbol H, Haight K, Ghaninia M, Hoo L, et al. An engineered orco mutation produces aberrant social behavior and defective neural development in ants. Cell. 2017; 170(4):736–747 e739. https://doi.org/10.1016/j.cell.2017.06.051.

65. Hsu PD, Lander ES, Zhang F. Development and applications of CRISPR-Cas9 for genome engineering. Cell. 2014;157(6):1262–78. https://doi.org/10.1016/j.cell.2014.06.002.

66. Jiang W, Bikard D, Cox D, Zhang F, Marraffini LA. RNA-guided editing of bacterial genomes using CRISPR-Cas systems. Nat Biotechnol. 2013;31(3):233–9. https://doi.org/10.1038/nbt.2508.

67. Wei W, Xin H, Yang R, Bai J, Miao Y, Gao G. Heritable genome editing with CRISPR/Cas9 in the silkmoth, Bombyx mori. PLoS ONE. 2014;9(7):e101210. https://doi.org/10.1371/journal.pone.0101210.

68. Yu Z, Ren M, Wang Z, Zhang B, Rong Y, Jiao R, Sun Y, Zeng Q, et al. Full-length transcriptome assembly of the odorant receptor co-receptor (Orco) in the silkworm, Bombyx mori. Insect Biochem Mol Biol. 2016;73:12–9. https://doi.org/10.1016/j.ibmb.2016.05.007.

69. Huang Y, Liu Z, Zeng Q, Zhao L, Fang X, Chen L, Dong Y, Chen Y, et al. Olfactory perception and Sensillar expression and regulation of optimal mating time in the moth Helicoverpa armigera. Chem Senses. 2009;34(6):469–77. https://doi.org/10.1093/chemse/bkp012.

70. Chang H, Guo M, Wang B, Liu Y, Dong S, Wang G. Sensillar expression and responses of olfactory receptors reveal different peripheral coding in two Helicoverpa species using the same pheromone components. Science. 2016;356:18742. https://doi.org/10.1038/srep18742.

71. Hansson BS, Anton S. Function and morphology of the antennal lobe: new developments. Annu Rev Entomol. 2000;45:203–31. https://doi.org/10.1146/annurev.ent.45.1.203.

72. Zhao XC, Ma BW, Berg BG, Xie YG, Tang QG, Guo XR. A global-wide search for sexual dimorphism of glomeruli in the antennal lobe of female and male Helicoverpa armigera. Sci Rep. 2016;6:35204. https://doi.org/10.1038/srep35204.

73. Chiang A, Priya R, Ramaswami M, Vijayraghavan K, Rodrigues V. Neuronal activity and Wnt signaling act through Gsk3-beta to regulate axonal integrity in mature Drosophila olfactory sensory neurons. Development. 2009;136(8):1273–82. https://doi.org/10.1242/dev.031377.

74. Tribble W, Olivos-Cisneros L, McKenzie SK, Saragosti J, Chang NC, Matthews B, Dixley PR, Kronauer DRC. orco mutation causes loss of antennal lobe glomeruli and impaired social behavior in ants. Cell. 2017;170(4):727–735 e710. https://doi.org/10.1016/j.cell.2017.07.001.

75. Kirsch K, Koo J. Olfactory receptors in non-chemosensory tissues. BMB Rev. 2012;54(4):155–167. https://doi.org/10.5422/Bmbre.2012.04.001.

76. Chen Z, Tanielian IM, Rana S, Cash-Ahmed AC, Sankey AL, Yang C, Robinson GE. Neurodevelopmental and transcriptomic effects of CRISPR/Cas9-induced somatic mutation in honey bees. J Neurogenet. 2021;35(3):320–32. https://doi.org/10.1080/10707706.2021.1887173.