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A highly expressed odorant receptor from the yellow fever mosquito, AaegOR11, responds to (+)- and (–)-fenchone and a phenolic repellent

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Abstract

The cornerstone of the reverse chemical ecology approach is the identification of odorant receptors (OR) sensitive to compounds in a large panel of odorants. In this approach, we de-orphanize ORs and, subsequently, measure behaviors elicited by these semiochemicals. After that, we evaluate behaviorally active compounds for applications in insect vector management. Intriguingly, multiple ORs encoded by genes highly expressed in mosquito antennae do not respond to any test odorant. One such case is CquiOR125 from the southern house mosquito, Culex quinquefasciatus Say. To better understand CquiOR125's role in *Culex* mosquito olfaction, we have cloned a CquiOR125 orthologue in the genome of the yellow fever mosquito, Aedes aegypti (L.), AaegOR11. Unlike the unresponsive nature of the orthologue in *Cx. quinquefasciatus*, oocytes co-expressing AaegOR11 and AaegOrco elicited robust responses when challenged with fenchone, 2,3-dimethylphenol, 3,4-dimethylphenol, 4-methycyclohexanol, and acetophenone. Interestingly, AaegOR11 responded strongly and equally to (+)- and (-)-fenchone, with no chiral discrimination. Contrary to reports in the literature, fenchone did not show any repellency activity against Ae. aegypti or Cx. quinquefasciatus. Laboratory and field tests did not show significant increases in egg captures in cups filled with fenchone solutions compared to control cups. The second most potent ligand, 2,3-dimethylphenol, showed repellency activity stronger than that elicited by DEET at the same dose. We, therefore, concluded that AaegOR11 is a mosquito repellent sensor. It is feasible that CquiOR125 responds to repellents that remain elusive.

Keywords

Culex quinquefasciatus; CquiOR125; AaegOR84; AaegOR113; *Aedes aegypti*; 2,3-dimethylphenol

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Author contributions

W.L., W.S.L., K.K.B., S.A. and A.J.C. performed research; W.S.L. designed research; W.L., WS.L. and A.J.C. analyzed the data; W.S.L. wrote the paper; all authors read and approved the final manuscript.

Competing financial interest

The authors declare no conflict of interest.

1. Introduction

The explosive advancement in our understanding of the molecular basis of insect olfaction in the last two decades paved the way for the application of reverse chemical ecology (Leal, 2005). In conventional chemical ecology approaches, attractants, repellents, and other semiochemicals are chemically characterized after bioassay-guided isolation from natural sources. In reverse chemical ecology, active compounds are identified first by their interactions with olfactory proteins, such as odorant receptors (ORs), and subsequent behavioral assays (Leal, 2020).

Semiochemicals must first activate receptors housed in olfactory neurons in the peripheral sensory system of insects (e.g., antennae, maxillary palps, and proboscis) (Leal, 2013) to elicit behavior. Highly expressed odorant receptors are intuitive targets for reverse chemical ecology because high expression levels imply that they are involved in the reception of crucial semiochemicals. Therefore, de-orphanizing highly expressed ORs may lead to the discovery of novel attractants or repellents, provided the test ORs are adequately expressed and challenged with appropriate compounds.

The two-electrode voltage-clamp (TEVC) recording system has been widely used to de-orphanize ORs (Pask, 2020-Wang et al., 2020). With this technique, a test OR is co-expressed with the co-receptor Orco in *Xenopus laevis* oocytes and challenged with test compounds. Co-expression is necessary because, in more evolved insect species, the binding units of ORs form ion channels (pathways) with the obligatory odorant receptor co-receptor Orco. In more primitive insects, tetramers of the binding unit *per se* form ion channels (Del Marmol et al., 2021). Previously, we have de-orphanized ORs in the genome of the Southern house mosquito, *Culex quinquefasciatus* (Hughes et al., 2010; Pelletier et al., 2010; Xu et al., 2014; Xu et al., 2013; Xu et al., 2015; Zhu et al., 2013), as part of our reverse chemical ecology approach. For example, we expressed CquiOR36/CquiOrco in *Xenopus* oocytes, challenged them with a panel of 230 odorants, and identified acetaldehyde as the most potent ligand (Choo et al., 2018). Subsequent behavioral tests showed that acetaldehyde is an oviposition attractant.

Considering the metabolic resources used for their synthesis, it is conceivable that highly expressed ORs play crucial roles in the ecology of the insect. Intriguingly, however, many ORs from the Southern house mosquito were found to be "silent," i.e., no compounds in our extensive panel of odorants elicit measurable currents in these CquiORx/CquiOrco-expressing oocytes. One such case is CquiOR125, which is the second-most expressed OR in *Cx. quinquefasciatus* antennae (Choo et al., 2018; Leal et al., 2013; Xu et al., 2014). We surmised that testing an ortholog from the yellow fever mosquito, *Aedes aegypti*, AaegOR11 (Matthews et al., 2018), might shed some light on the function of the highly expressed CquiOR125. AaegOR84, AaegOR113, and AaegOR11 are three of the most expressed ORs in the female yellow fever mosquito antennae (Matthews et al., 2018). While AaegOR84 and AaegOR113 have no orthologs in the *Cx. quinquefasciatus* genome, AaegOR11 shares 73% amino acid identity with CquiOR125.

We cloned these three ORs and challenged AegORx/AaegOrco-expressing oocytes with a panel of 286 compounds. AaegOR84/AaegOrco- and AaegOR113/AaegOrco-expressing oocytes were not activated by any compound in our panel of odorants but elicited small inward currents when challenged with the Orco ligand candidate, OLC 12 (Chen and Luetje, 2012). By contrast, AaegOR11/AaegOrco-expressing oocytes generated strong inward currents when stimulated with fenchone, 2,3-dimethylphenol, and other compounds. Interestingly, AaegOR11 does not discriminate fenchone stereoisomers. In-door and field tests suggest that fenchone is neither an oviposition attractant nor a repellent for the yellow fever mosquito, and 2,3-dimethylphenol is a potent repellent.

2. Material and Methods

2.1. Mosquitoes

Culex quinquefasciatus mosquitoes used in this study originated from the CQ1 colony, established in the 1950s, from specimens collected in Merced County, CA. The CQ1 colony has been used as a standard insecticide-susceptible strain in insecticide bioassays (McAbee et al., 2004). In Davis, mosquitoes were maintained at $27\pm1^{\circ}$ C, $75\pm5\%$ relative humidity, and under a photoperiod of 12:12 h.

Aedes aegypti mosquitoes used for molecular studies and blood feeding repellency choice assays originated from second-filial generation individuals originating from adults collected in Clovis, (Fresno County, CA). The first two of the three laboratory oviposition repellency/ attractiveness trials were conducted using females reared from *Ae. aegypti* eggs that were laid by wild females in oviposition cups placed at multiple properties in Reedley. In the third repellency/attractiveness trial, females reared from the eggs laid by females in the first two trials were used.

2.2. Cloning and de-orphanization

The p-GEMHE plasmids for AaegOR11, AaegOR84, and AaegOR113 were obtained by using *Ae. aegypti* female antennae cDNA (Zeng et al., 2019) as a template and the infusion primers below. The resulting PCR products were purified with gel electrophoresis, QIAquick gel extraction kit (Qiagen, Germantown, MD), and then cloned into pGEMHE using In-Fusion HD Cloning Kit according to the manufacturer's instructions (Clontech).

PCR In-Fusion primers were designed according to the user manual.

InFuAaOR11F primer:

5'- GATCAATTCCCCGGGACCATGCAGCTGAAAGACGAATGG -3' and

InFuAaOR11R primer:

5'- CAAGCTTGCTCTAGATTAGCCGGCAGCTTGTTTC -3';

InFuAaOR84F primer:

5'- GATCAATTCCCCGGGCACATCATGAACGCGATTGT -3' and

InFuAaOR84R primer:

5'- CAAGCTTGCTCTAGAGCCATTGCATTATTCCGATT -3'.

InFuAaOR113F primer:

5'- GATCAATTCCCCGGGACCATGTTTGCGGAAATTCGTGGC -3' and

InFuAaOR113R primer:

5'- CAAGCTTGCTCTAGACTAAACCATATTGATCAGAAATGTTAACACTG -3'.

The colonies from transformations were verified by regular PCR. The positive clones were cultured and subsequently extracted using the QIAprep Spin Miniprep kit (Qiagen) and sequenced by ABI 3730 automated DNA sequencer.

TEVC was performed as previously described (Xu et al., 2022). In brief, linearized pGEMHE-AaegORs were used as templates to transcribe into capped cRNA with poly(A) using an mMESSAGE mMACHINE T7 kit (Ambion, Austin, TX) following the manufacturer's protocol. The cRNAs were dissolved in RNase-free water and adjusted to a concentration of 200 µg/mL by UV spectrophotometry (NanoDropTM Lite Spectrophotometer). 9.2 nl of a 50/50 mixture of a test OR and AaegOrco [25] cRNAs were microinjected into V or VI Xenopus oocytes (purchased from EcoCyte Bioscience, Austin, TX). Then, injected oocytes were incubated at 18°C for 3–7 days in modified Barth's solution [in mM: 88 NaCl, 1 KCl, 2.4 NaHCO₃, 0.82 MgSO₄, 0.33 Ca(NO₃)₂, 0.41 CaCl2, 10 HEPES, pH 7.4] supplemented with 10 µg/mL of gentamycin, 10 µg/mL of streptomycin, and 1.8 mM sodium pyruvate. Test oocytes were placed in a perfusion chamber and challenged with a panel of odorants (see below). Currents were amplified with an OC-725C amplifier (Warner Instruments, Hamden, CT) holding the voltage at -80 mV and a low-pass filter at 50 Hz, and digitized at 1 kHz. Data acquisition and analysis were carried out with Digidata 1440A and pClamp10 software (Molecular Devices, LLC, Sunnyvale, CA). To standardize the measurement, 6-7 days-old eggs were used for TEVC measurement.

2.3. Panel of odorants

Oocytes were challenged with the following compounds: 1-butanol, 1-pentanol, 1hexanol, 1-heptanol, 1-octanol, 1-nonanol, 2,3-butanediol, 2-butoxyethanol, 3-methyl-1butanol, 2-hexen-1-ol, 3-hexen-1-ol, 1-hexen-3-ol, 1-hepten-3-ol, 3-octanol, 1-octen-3ol, 2-octanol, 2-butanol, 2-nonen-1-ol, 2-pentanol, 4-methylcyclohexanol, 1-hexadecanol, menthyl acetate, methyl acetate, ethyl acetate, propyl acetate, butyl acetate, pentylacetate, hexyl acetate, heptyl acetate, octyl acetate, nonyl acetate, decyl acetate, methyl propionate, ethyl propionate, methyl butyrate, ethyl butanoate, methyl hexanoate, ethyl hexanoate, ethyl 3-hydroxyhexanoate, ethyl 3-hydroxybutanoate, (*E*)-2-hexenyl acetate, (\mathbb{Z}) -3-hexenyl acetate, α -terpinene, γ -terpinene, ethyl lactate, methylsalicylate, 1-octen-3-yl acetate, isopentyl acetate, m-tolyl acetate, ethyl phenylacetate, geranyl acetate, octadecyl acetate, propanal, butanal, penatanal, hexanal, (*E*)-2-methyl-2-butenal, heptanal, octanal, nonanal, decanal, undecanal, 1-dodecanal, (*E*)-2-hexenal, (*Z*)-8-undecenal, (*E*)-2-heptenal,

(E)-2-nonenal, phenylacetaldehyde, furfural, 2-butanone, 2-heptanone, geranylacetone, 6methyl-5-hepten-2-one, 5-methyl-2-hexanone, 2,3-butanedione, 3-hydroxy-2-butanone, 2undecanone, (-)-menthone, 2-tridecanone, 2-nonanone, 5-isobutyl-2,3-dimethylpyrazine, (S)-(-)-perillaldehyde, (1S,4R)-fenchone, (1R,4S)-fenchone, cyclohexanone, acetophenone, $(\alpha+\beta)$ -thujone, p-coumaric acid, isovaleric acid, 1-dodecanol, dodecanoic acid, (\pm) -lactic acid, ethanoic acid, propanoic acid, butanoic acid, isobutyric acid, 2-oxobutyric acid, pentanoic acid, 2-oxovaleric acid, palmitoleic acid, oleic acid, hexanoic acid, (E)-2-hexenoic acid, 5-hexanoic acid, (E)-3-hexenoic acid, heptanoic acid, octanoic acid, nonanoic acid, decanoic acid, trimethylamine, n-tridecanoic acid, linoleic acid, ammonia, trimethylamine, propylamine, butylamine, pentylamine, hexylamine, heptylamine, camphor, octylamine, 1,4-diaminobutane, cadaverine, 1,5-diaminopentane, benzaldehyde, phenol, 2-methylphenol, 3-methylphenol, 4-methylphenol, 4-ethylphenol, 3,5-dimethylphenol, 2,3-dimethylphenol, guaiacol, 2-methoxy-4-propylphenol, 2-phenoxyethanol, 1,2-dimethoxybenzene, benzyl alcohol, 2-phenylethanol, 1-phenylethanol, phenyl ether, isoprene, limonene, linalyl acetate, α -humulene, linalool oxide, geraniol, nerol, thymol, (±)- linalool, eucalyptol, citral, eugenol, a-pinene, ocimene, (±)-citronellal, indole, permethrin, acetaldehvde, 3methylindole, 3-pentanol, 3-methyl-2-butanol, 3-methyl-2-buten-1-ol, 2-methyl-3-buten-2ol, γ -valerolactone, γ -hexalactone, γ -octalactone, γ -decalactone, α -phellandrene, nerolidol, γ -dodecalactone, jasmine, 1-octen-3-one, dibutyl phthalate, dimethyl phthalate, menthol, benzyl formate, 2-acetylthiophene, phenethyl formate, carvone, isovaleraldehyde, n-methylbenzamide, 3-methylbenzamide, ethyl stearate, tetradecanoic acid, methyl myristate, p-cymene, sabinene, 2-tridecanone, terpinolene, valencene, 2,4-pentanedione, (R)-(+)-pulegone, β -myrcene, 5-isobutyl-2,3-dimethylpyrazine, phenyl propanoate, phenyl isobutyrate, eugenyl acetate, (+)- δ -cadinene, (+)-limonene oxide, 2ethyltoluol, 2,4-hexadienal, 2-methyl-2-thiazoline, phenethyl propionate, ethyl (2E,4Z)decadienoate, methyl anthranilate, α-hexylcinnamaldehyde, 2-pentanone, 2-hexanone, 2-octanone, 4,5-dimethylthiazole, (E,E)-farnesol, (E,E)-farnesyl acetate, farnesene, 4dimethylamino-1-naphthaldehyde, α -methylcinnamaldehyde, cinnamyl alcohol, α -terpineol, citronellol, (*E*)-cinnamaldehyde, (–)-caryophyllene oxide, β -caryophyllene, carvacrol, terpinen-4-ol, neo-alloocimene, palmitic acid methyl ester, 2-ethylhexanoic acid, pentadecanoic acid, geranyl isovalerate, geranyl butyrate, α -cetone, 7-hydroxcitronellal, isopropyl myristate, methyl N,N-dimethylanthranilate, pyridine, pyrrolidine, pyrrolidinone, amiloride, allethrin, allopurinol, methyl-p-benzoquinone, lilial, styrene, indole-3-aldehyde, dimethyl trisulfide, methyl disulfide, cyclohexanol, 2-methylbutyl 2-methylbutanoate, sabinene, citronellyl acetate, methyl geranate, 3-methylbutyl 2-methylbutanoate, tertbutanol, 2,4-dimethylphenol, 2,5-dimethylphenol, 2,6-dimethylphenol, 3,4-dimethylphenol, ethyl undecanoate, neryl acetate, isobutyl acetate, methyl sulfide, 2-hexanol, (+)-3-carene, ethyl butyrate, α -ionone, β -ionone, formic acid, linalyl formate, diethyl sebacate, 3,5-di-tert-butyl-4-hydroxyacetophenone, 2,4-di-tert-butylphenol, 2nonanol, N-methylpiperidine, 5-ehtyl-2-methylpyridine, piperine, 1-methylpiperidine, 2-methylpiperidine, 3-methylpiperidine, 4-methylpiperidine, dimethyl carbonate, 5ethyl-2-methylpyridine, 3-pentyn-1-ol, 4-methylthiazole, 4,5-dimethylthiazole, tryptamine, dicyclohexyl disulfide, and isoamyl alcohol. The Orco ligand candidate 2-{[4-Ethyl-5-(4pyridinyl)-4H-1,2,4-triazol-3-yl]sulfanyl}-N-(4-isopropylphenyl)acetamide (OLC 12 (Chen

and Luetje, 2012) = VUAA-3 (Taylor et al., 2012)) was used for confirming receptor protein expression.

2.4. Chemical analysis

Gas chromatography analyses were performed on an Agilent 6890 Series GC System equipped with a flame ionization detector (FID) and a chiral capillary column (HP-Chiral-20B, 30 m x 0.25 mm; 0.25 μ m). The injection port and the FID detector were operated at 250°C. The oven temperature was set at 85°C for 70 min, with a 5 min post-run at 210°C.

2.5. Repellent assays

Repellency was measured using the surface landing and feeding assays, previously described in detail (Leal et al., 2017; Xu et al., 2014). In short, two Duddle bubbling tubes were inserted inside a mosquito cage (arena) through a wooden board attached to the back of the cage. The tubes were covered with non-lubricated condoms to avoid contamination; they were replaced every five tests (Xu et al., 2022). Water circulated inside the tubes to maintain the surface at 37°C. A needle was inserted through the wooden board above each tube to deliver CO₂ (50 ml/min). A dental cotton roll holding defibrinated sheep blood (100 µl) was inserted between a Duddle tube and a needle. One filter paper cylinder was impregnated with solvent only (control; 200 µl of hexane) and the other with a solution of the test compound (200 µl of a 1% solution in hexane). After the solvent evaporated, the cylinders were placed around each tube and held in place by insect pins. Repellency was measured using cages with one hundred non-blood-fed (two weeks old) female mosquitoes deprived of water and sugar for at least one hour. Responses were recorded for 5 min; the filter paper rings were swapped or renewed to start another trial. DEET 1% was used as a positive control. Behavioral responses were expressed in percent protection, $P \approx (C-T)/2$ C)X100 (EPA, 2010; WHO, 2009).

2.6. Oviposition assays

Three trials were performed. In the first trial, 6 concentrations of fenchone were tested. One hundred µl from each of the primary concentrations of 0.0001, 0.001, 0.01, 0.1, 1, and 10% of fenchone were added to 6 white plastic cups (old-fashioned tumblers, Finelinesettings.com) filled with 100 ml of tap water which would represent in effect a further final 1000x dilution of each primary concentration in each cup. Before adding water and fenchone into each cup, the inside walls were lined with two slightly overlapping 5.8 x 11.4 cm strips of seed germination paper (Regular weight brown seed germination or toweling paper, Seedburo Equipment Company, Des Plaines, IL, USA). The seed germination paper was held against the sides with forceps while pouring the tap water into them to ensure the seed germination paper remained hugged against the cup walls. Control cups contained 100 µl of 100% ethanol added to 100 ml of tap water. Two cups (one with treatment and the other the control cup) were placed in the opposite corners of each of the 6 cages (30.5 cm x 30.5 cm x 30.5 cm; Cat # 1450B Bioquip Products Inc, Rancho Dominguez, CA, USA; discontinued). Eighteen to 21 gravid Ae. aegypti females were released into each of the cages. The cages with the mosquitoes and treatment and control cups in them were spaced 128 cm apart in the insectary (80% RH with lighting set at 11 h daylight, 1 h crepuscular, and 12 h

dark) for 7 days. Females had constant access to 10% sucrose solution in each cage. On each of the 7 days the dead mosquitoes were counted and removed, and the minimum and maximum temperatures were recorded. The light/dark conditions we simulated in our insectary are conducive for maximum egg-laying activity in *Ae. aegypti* (Farnesi et al., 2018). The concentration of the fenchone remained consistent in each cup for the 7 days as very little evaporation was noted. The second trial with 6 replicates was set up like the first one, except for the tested doses. The control and treatment cups were placed in opposite corners of the cage. The third trial was set up the same as the first trial but used 24 cages, each with 18-25 gravid females, to test a single dose.

2.7. Field tests

The entire inside walls of black plastic cups (16-oz, 473 ml black stadium cups, csbdstore.com) Charlotte, NC, USA) were lined with rough brown paper (Regular weight seed germination or toweling paper, Seedburo Equipment Company). In each of the six properties in Reedley (Fresno County, CA), traps were placed in shady areas 30 m apart from each other. The control cups were filled with 2.5 ml of ethanol in 247.5 ml of tap water; the infusion cups were prepared with 250 ml 7-day-old Orchard grass infusion; and fenchone cups had 2.5 ml of 1% (+/-)-fenchone solution in ethanol plus 247.5 ml tap water, Seven-day-old infusions were prepared by adding 30 g of Orchard grass to 11.3 L of tap water and left outdoors to ferment in a plastic drum while mixing once a day. The cups were spaced 30 m apart to allow a single female oviposition choice in a corner area of a property. Orchard grass infusion in black cups lined with rough germination paper is used by the local mosquito abatement district for Ae. aegypti abundance surveillance purposes. After 7 days, the oviposition paper was removed carefully, and the number of eggs attached to the oviposition paper was counted. This was repeated three times, comprising a field evaluation of 3 x 6 replicates. In each of the three replicates, the cups containing different solutions were placed in alternate positions to avoid possible position effects. In Reedley, Ae. aegypti is the only Aedes mosquito species that lay eggs in these cups.

The field evaluation was conducted in late summer when the daily average maximum and minimum temperatures were 37 and 19°C. After 7 days when the cups were removed, there were still about 50 ml of infusion or water remaining in all the cups, which meant for the entire 7 days females were able to lay eggs on moist germination paper above the water or infusion line. Cups were left for 7 days in the field because that is the procedure the mosquito abatement district follow for *Ae. aegypti* egg surveillance purposes.

3. Results and Discussion

3.1. Cloning and de-orphanization of ORs

We cloned AaegOR11, 84, and 113 using cDNA obtained from *Ae. aegypti* mosquitoes collected in Clovis, CA, as a template. Four clones of AaegOR11 gave identical sequences (GenBank OM568708), but they differed in five residues compared to the AaegOR11 amino acid sequence in VectorBase (AAEL011583). Specifically, Ser163Thr, Met167Ile, Asn240His, Met249Ile, and Met338Ile. In short, three methionine residues in the VectoBase sequence are Ile in the Clovis strain, a His replaces an Asn, and a Ser changed to a Thr. Our

AaegOR84 and AaegO113 clones showed sequences identical to those in VectorBase, i.e., AAEL017043 and AAEL017123, respectively.

We co-expressed each OR with the odorant receptor co-receptor Orco in *Xenopus* oocytes and challenged them with a large panel of odorants. AaegOR113/AaegOrco-expressing oocytes did not respond to any test compound except for a small current elicited by the Orco ligand OLC 12 (Chen and Luetje, 2012). Likewise, no currents were recorded when AaegOR84/AaegOrco-expressing oocytes were challenged with individual compounds in our panel of odorants, except for OLC 12-induced currents. By contrast, AaegOR11/AaegOrco-expressing oocytes responded to twelve compounds in our panel, specifically, fenchone, 2,3-dimethylphenol, 2,6-dimethylphenol, 3,4-dimethylphenol, 2-methylphenol, 4-methylcyclohexanol, acetophenone, guiacol, 2-acetylthiophene, indole, 1,2-dimethoxybenzene, and (*Z*)-3-hexenyl acetate. We then challenged AaegOR11/AaegOrco-expressing oocytes at least six times with each active compound (Fig 1). Fenchone and 2,3-dimethylphenol, 3,4-dimethylphenol, 4-methylcyclohexanol, and acetophenone gave the most robust responses (Fig 1).

Concentration-dependent relationships for the five ligands that produced the most robust responses confirm that fenchone was the most potent ligand, followed by 2,3-dimethylphenol, and 3,4-dimethylphenol (Fig 2).

We then asked whether this receptor could discriminate fenchone stereoisomers. Interestingly, AaegOR11/AaegOrco-expressing oocytes responded equally to (1R,4S)-fenchone (=(-)-fenchone, PubChem CID #82229) and (1S,4R)-fenchone (=(+)-fenchone, PubChem CID #1201521) (Fig 3A). A dose-dependent curve confirmed that the receptor responds to the two stereoisomers equally (Fig 3B)

3.2. Functional studies

(+)-Fenchone has been reported as a potent repellent against *Ae. aegypti* (Kim et al., 2002) and *Ae. albopictus* (Yan et al., 2022) comparable to DEET, but a moderate repellent 30 min after sample preparation and about 50% activity one hour later (Kim et al., 2002). We tested both (+)-fenchone and (–)-fenchone with our surface-landing and feeding assay (Leal et al., 2017; Xu et al., 2022; Xu et al., 2014), and using freshly prepared samples. In our hands, neither (+)-fenchone, nor (–)-fenchone showed significant repellency activity against *Ae. aeypti* (Fig 4).

The (+)-fenchone sample reported being a potent repellent against *Ae. aegypti* was extracted and isolated from fruits of the common fennel *Foeniculum vulgare* Miller (Kim et al., 2002). The reported specific rotation (Kim et al., 2002) suggests an enantiopure compound was used in the previous work. In any case, we analyzed our sample to rule out a possible problem with enantiomeric impurities. Gas chromatography analysis by separation on a capillary column with a chiral stationary phase showed that our (+)-fenchone sample was enantiopure, whereas the (–)-fenchone sample contained a small percent of (+)-fenchone (Fig 5).

Given the discrepancy between the literature (Kim et al., 2002; Yan et al., 2022) and our findings, we tested whether the (+)- and (–)-fenchone repel *Cx. quinquefasciatus*, but neither isomer showed repellency activity. The protection elicited by (+)-fenchone at 1% was $-52\pm27.5\%$ (n = 8), whereas DEET showed 92.2 $\pm2.5\%$ (n = 4). Similarly, (–)-fenchone gave low protection $-18\pm20.2\%$ (n = 8); DEET, 100% (n = 4). In unusual cases (Borden et al., 1976), semiochemicals are active only when presented as a racemic mixture. We, therefore, tested a (+/–) mixture of fenchone and observed no activity: 9.5 \pm 8.0 % protection (n = 8); DEET 90.1 \pm 2.4% (n = 8).

Next, we tested whether fenchone could act as an oviposition attractant for the yellow fever mosquito. First, we conducted in-door assays and focused on (–)-fenchone at doses ranging from 10^{-2} % to 10^{-7} % (Fig 6A). For each dose, 100 µl of a concentrated fenchone solution in ethanol was added to a cup filled with 100 ml of tap water. One hundred µl of ethanol was added to each control cup with 100 ml tap water. There were no significant difference in the number of eggs laid in treatments at doses lower than 0.01% (10^{-2} %) compared with their respective controls. However, the number of eggs laid in the 0.01% (–)-fenchone cups was significantly higher than those laid in control cups (P=0.0244, n = 6, two-tailed, paired t-test). Considering that activity was observed at the highest dose tested, we repeated these experiments with 4 doses, 0.001, 0.005, 0.01, and 0.05% (Fig 6B). We found no significant difference between treatment and control in all doses tested, including 0.01% (10^{-2} %).

Given this discrepancy in the two in-door trials regarding the activity of (–)-fenchone at 10^{-2} % (Fig 6), we conducted a third trial with a single dose of 0.01% that included a larger number of replicates (Fig 7). Again, there was no significant difference between treatment and control (P=0.1882, n = 22, two-tailed paired t-test).

Lastly, we performed field tests comparing the deposition of eggs in cups containing Orchard grass infusion (Ponnusamy et al., 2010) (positive control), racemic fenchone, or water (control) (Fig 8). Traps were placed in seven properties in Reedley (Fresno County, California), with three replicates of one week each. The total number of eggs laid in control, infusion, and fenchone cups, respectively, were: 224, 741, and 259 (week 1); 297, 797, and 546 (week 2), and 276, 984, and 567 (week 3). While cups loaded with infusion had significantly more eggs than control cups (145.7 ± 22.7 vs. 44.3 ± 7.9 ; adjusted P=0.0002, n = 18, one-way ANOVA), egg counts in cups holding racemic fenchone did not significantly differ from egg counts in control traps (77.3 ± 16.7 vs. 44.3 ± 7.9 ; adjusted P=0.2969, n = 18, one-way ANOVA). This led us to conclude that fenchone solution was no more or less attractive than water but confirmed that grass infusion offered higher egg-laying attractive properties. For now, mosquito control agencies should still have cups filled with Orchard grass infusion for *Ae. aegypti* oviposition field surveillance purposes.

Surprisingly, the most potent ligand for a highly expressed receptor did not show any activity as either a repellent or an oviposition attractant. We cannot rule out that fenchone plays a crucial role in the chemical ecology of the yellow fever mosquito, particularly for behaviors not measured in our study, including, but not limited to, mating behavior, attraction to flowers, and pollination. Additionally, fenchone may elicit repellency or oviposition attraction in synergy with other semiochemicals. However, our studies

demonstrate that fenchone is neither a repellent nor an oviposition attractant when used as a single compound.

Lastly, we tested whether the second most potent ligand, 2,3-dimethylphenol, would repel mosquitoes. In contrast to fenchone, 2,3-dimethylphenol at 1% showed significantly stronger (P = 0.0133, Wilcoxon matched-pairs signed rank test) repellency activity against *Cx. quinquefasciatus* mosquitoes: 94.3 \pm 0.8 % protection (n = 32), than DEET at 1%: 81.5 \pm 2.6 % (n = 22). These findings suggest that AaegOR11, one of the ORs most expressed in the female yellow fever mosquito antennae (Matthews et al., 2018), is involved in the reception of repellents.

AaegOR11 is 73% identical to CquiOR125. Of note, there is a proposition to rename CquiOR125 to CquiOR11 (Dr. Carolyn McBride, personal communication). Previous work showed that none of the ligands for AaegOR11 identified in this study activated CquiOR125 in an earlier study (Xu et al., 2014). Indeed, CquiOR125 did not respond to any compound in a large panel of odorants (Xu et al., 2014).

3.3. Overall conclusions

In summary, we have de-orphanized an OR highly expressed in the antennae of the female yellow fever mosquito, AaegOR11, which shared high amino acid identity with CquiOR125, a "silent" receptor from the southern house mosquito. In the *Xenopus* oocyte recording system, AaegOR11, co-expressed with AaegOrco, generated robust responses when challenged with fenchone, 2,3-dimethylphenol, 3,4-dimethylphenol, 4-methylcyclohexanol, and acetophenone. Fenchone did not elicit blood-feeding repellency or oviposition activity. By contrast, behavioral measurement showed that 2,3-dimethylphenol is a potent blood-feeding repellent. Considering that AaegOR11 is a repellent-detecting OR, it is conceivable that AaegOR11 orthologue in the genome of *Cx. quinquefasciatus*, CquiOR125, is also a "repellent receptor," but the active ligand is still elusive.

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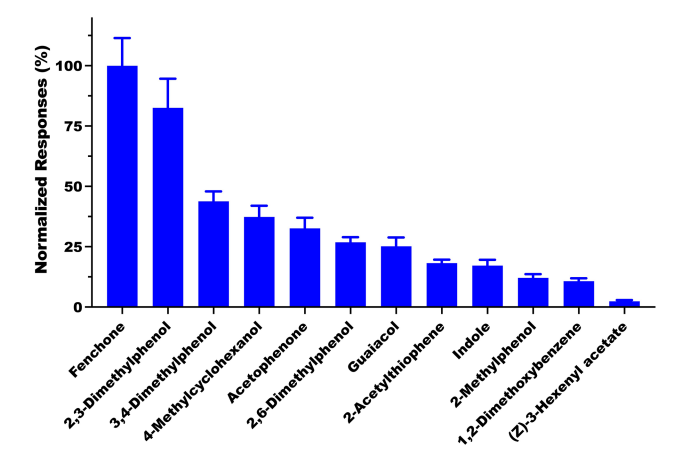


Fig 1. Quantification of current responses recorded from oocytes co-expressing AaegOR11 and AaegOrco.

All compounds were tested at 1 mM (source dose, n = 6). For clarity, bars are displayed in decreasing order of response (mean \pm SEM).

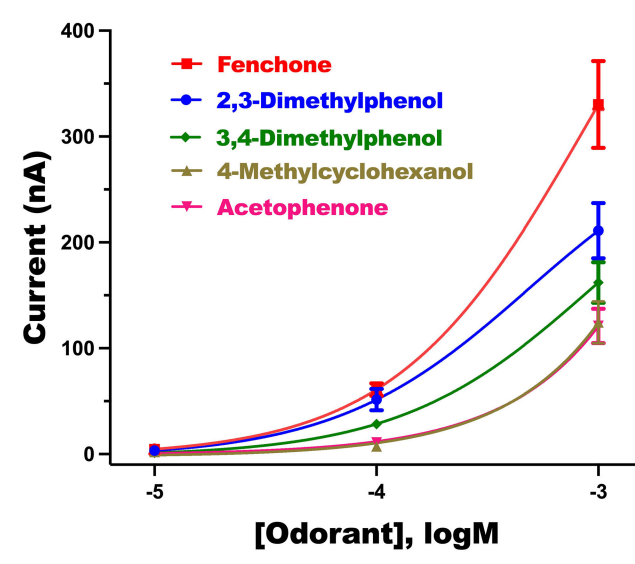


Fig 2. Concentration-dependent responses were recorded from AaegOR11/AaegOrco-expressing oocytes.

Currents were recorded from seven oocytes challenged at 0.01, 0.1, and 1 mM, with seven replicates (n = 7) per stimulus per dose.

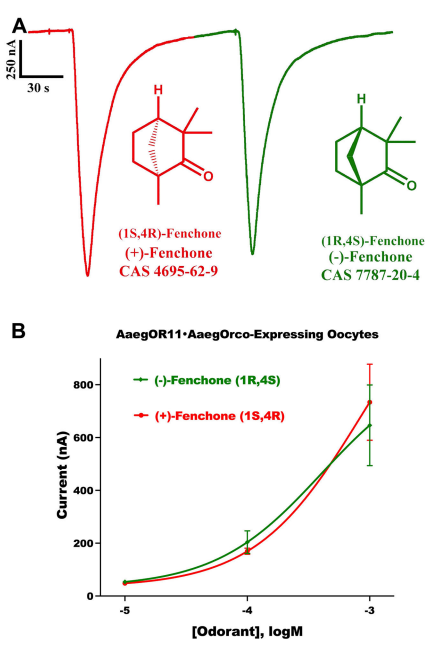


Fig 3. Responses were recorded from oocytes co-expressing AaegOR11 and AaegOrco. (A) A representative trace with recordings from the same oocyte challenged with (+)- and (-)-fenchone at 1 mM. Similar results were obtained when oocytes were stimulated first with (-)- and then (+)-fenchone. (B) Concentration-dependent relationships obtained with oocytes (n = 3) challenged with the two isomers at three different doses (n = 6-9 replicates/ dose).



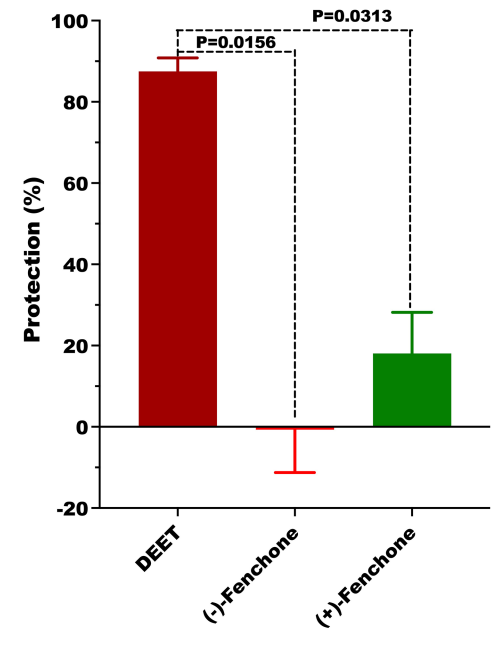


Fig 4. Fenchone repellency measured in the surface-landing and feeding assay. Repellency is expressed in percent protection for DEET (n = 7), (–)-fenchone (n = 8) and (+)-fenchone (n = 6). Tests were conducted in tandem by measuring the responses to DEET (2 replicates, one on each side of the arena) and two replicates of (+)- or (–)-fenchone and repeating this cycle until the response to fresh samples of DEET started to decrease. Data were analyzed by the Wilcoxon matched-pairs signed rank test.

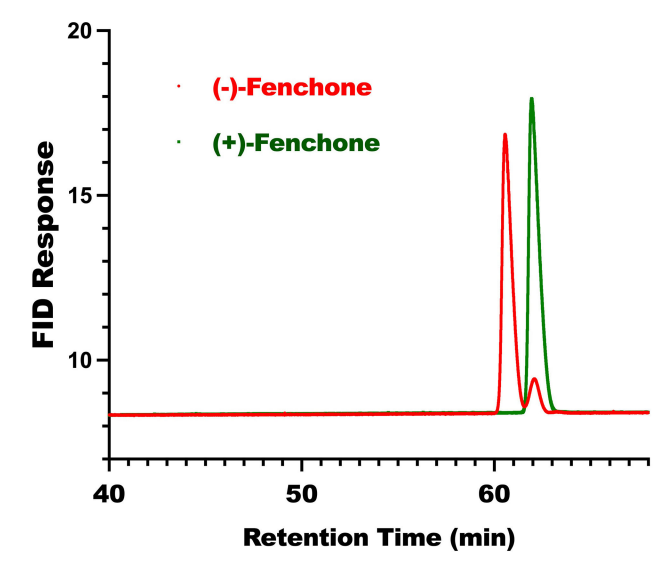


Fig 5. Separation of fenchone isomers on a gas chromatograph equipped with a capillary column with a chiral stationary phase.

Baseline separation of (–)- and (+)-fenchone was achieved with an HP-CHIRAL-20B column. For clarity, the traces were reconstructed by Prism using the FID data imported from the Agilent software.

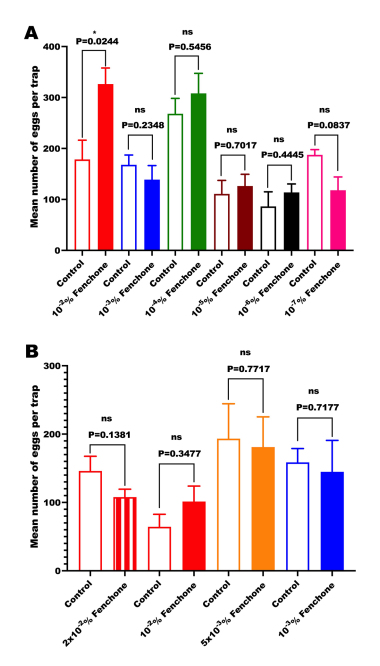


Fig 6. Results of in-door oviposition assays with fenchone at various doses.

(A) The first screening of (–)-fenchone activity at multiple doses, and (B) a closer examination of the responses to concentrations closer to 10^{-2} %. All tests were performed with six replicates (n = 6 per dose). Data passed the Shapiro-Wilk normality test and were, therefore, analyzed by paired, two-tailed t-test.

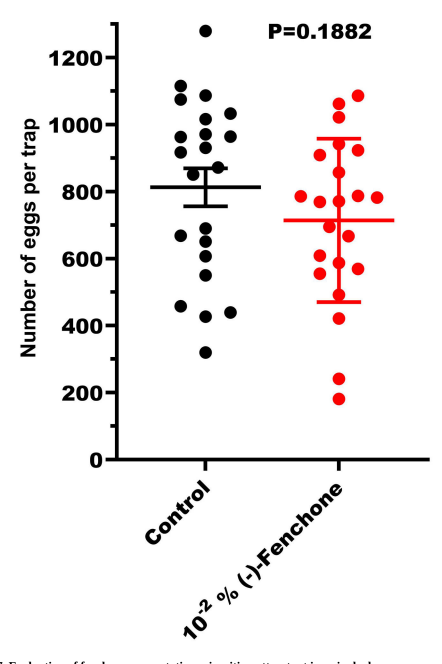


Fig 7. Evaluation of fenchone as a putative oviposition attractant in a single dose. Each of the six cages used had control and treatment; the experiment was replicated four times. Two outliers were omitted for clarity (cage 2: control, 731; treatment, 1477; cage 3: control, 1952; treatment, 959). The complete dataset, including the two outliers, did not pass the Shapiro-Wilk normality test and was analyzed by the Wilcoxon matched-pairs signed rank test (p=0.2768, n = 24).



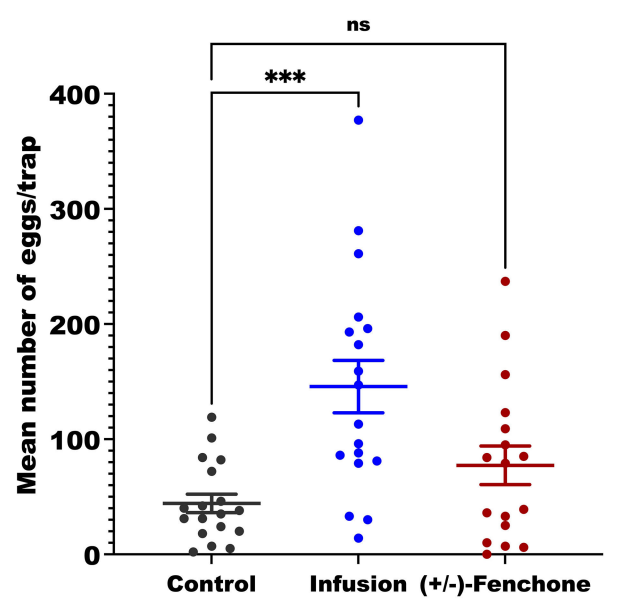


Fig 8. Results of field tests in Reedley (Fresno Co.) in late summer.

Control traps were baited with a 7-day-old Orchard grass (*Dactylis glomerata* L.)-based infusion; fenchone traps were filled with 0.01% racemic fenchone. A set of control-, infusion- and fenchone-trap, separated by 30 m, was placed in seven properties in Reedley. Eggs were counted after seven days. These experiments were repeated three times late in summer with average maximal and minimal temperatures of 37 and 19°C, respectively.