Electrophysiological, Behavioural and Biochemical Effect of Ocimum Basilicum Oil and Its Constituents Methyl Chavicol and Linalool on Musca Domestica L.

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Abstract

The *Ocimum basilicum* essential oil (EO) was evaluated for its biological effect on *M. domestica*. Characterization of *O. basilicum* EO revealed the presence of methyl chavicol (70.93%), linalool (9.34%), epia-cadinol (3.69 %), methyl eugenol (2.48%), γ-cadinene (1.67%), 1,8-cineole (1.30%) and (E)-β-ocimene (1.11%). The basil EO and its constituents methyl chavicol and linalool caused the neuronal response in female adults of *M. domestica*. Adult female flies showed reduced preference to food source laced with basil EO and methyl chavicol. Substrate treated with EO and methyl chavicol at 0.25% caused an oviposition deterrence of over 80%. The ovicidal effect was high in *O. basilicum* EO (EC$_{50}$ 9.74mg/dm$^3$) followed by methyl chavicol (EC$_{50}$ 10.67mg/dm$^3$) and linalool (EC$_{50}$ 13.57mg/dm$^3$). On contact toxicity, adults exposed to EO (LD$_{50}$ 10.01 μg/adult) were more susceptible than to methyl chavicol and linalool (LD$_{50}$ 13.62 μg/adult and LD$_{50}$ 43.12 μg/adult respectively). EO and its constituents methyl chavicol and linalool induced the detoxifying enzymes Carboxyl esterase (Car E) and Glutathione S – transferases (GST)

Introduction

The cosmopolitan housefly, *Musca domestica* L. (Diptera: Muscidae) is synanotrophic and thrives in close association with humans and livestock animals causing annoyance, food spoilage and transmitting pathogenic organisms (causing enteric disease, typhoid and shigellosis) (Sasaki et al. 2000; Kumar et al. 2012; Khamesipour et al. 2018; Moon 2019). The transmission of microbes is facilitated through dislodgement from exoskeleton, feeding regurgitation and feces (Fotedar 2001; Meerberg et al. 2007; Wanaratana et al. 2013). The ability of housefly to acquire avian influenza H9N2 aid in spreading the virus to humans and poultry birds (Salamatian et al. 2020). Reports from Wuhan confirmed 2 – 10 % of COVID-19 patients had diarrhoea and abdominal pain, a symptom that could have occurred due to faeco oral transmission (Dawei wang et al. 2020). In such a situation, vectoral role of flies carrying virus from faecal visits to human settlements cannot be ruled out (Dehghani and Kassiri 2020).

*M. domestica* menace in human settlements, animal and poultry sheds are managed by treating with insecticides or raising the birds on feed premix of growth regulators (Macovei et al. 2008; Ghosh and Zurek 2015). The excessive dependence on chemical insecticides leads to development of insecticide resistance to pyrethroids, organophosphates, spinosad, indoxacarb, spiromesifen (Khan et al. 2015; Alam et al. 2020) and also build-up of xenobiotics that are toxic to humans and nontarget animals. Botanical based insecticides are an alternative to overcome the problem as they pose minimal safety risk with potential bioaction and can be used in tandem with biocontrol agents (Kaufman et al. 2010; Khan et al. 2015; Scott 2017; Saeed et al. 2018; Shi et al. 2020). This necessitates development of green insecticide and repellents with varied mode of action (Pavela and Benelli 2016; Benelli and Pavela 2018a; Pavela et al. 2019; Isman 2020) that aids to mitigate the problems posed to human and environment (Isman 2017; Pandiyan et al. 2019; Ikbal and Pavela 2019).

Sweet basil, *Ocimum basilicum* belonging to lamiaceae is a herbaceous and perennial plant grown in Asia, Africa, Central and South America (Simon et al. 1999). *O. basilicum* EO is widely used in preparation of food, cosmetics and medicine (Moura et al. 2020). The major constituents of *O. basilicum* EO viz., methyl
chavicol and linalool possess antioxidant, anesthetic, anti-inflammatory, antimicrobial (Radulovic et al. 2013; Varga et al. 2017). In addition they are reported to be cidal (Rice and Coats 1994; Palacios et al. 2009; Tarelli et al. 2009; Gallardo et al. 2015; Yuexun Tian 2017) and insect repellent (Delille 2007; Yuexun Tian 2017).

Previous studies on basil EO and its constituent's linalool and methyl chavicol were limited to assessing its fumigant and contact toxicity to adults and larval stages (Palacios et al. 2009; Yuexun Tian 2017). In this investigation, we have reported the electrophysiological and olfactory response of *O. basilicum* EO and its major constituents linalool and methyl chavicol to adult females of *M. domestica*. The ovicidal, adulticidal and the activity of detoxification enzyme in adult exposed to *O. basilicum* EO, linalool and methyl chavicol are investigated.

**Materials And Methods**

**House fly rearing**

House fly, *M. domestica* adults were collected using a sweep net from suburbs of Bengaluru, India. The collected adult flies were kept in 30×30×30 cm cage (aluminium frame fitted with acrylic sheets). The flies were fed with diluted honey solution (10%) in water. An oviposition substrate (250 gm) prepared by mixing wheat bran + milk powder + egg yolk powder (10: 2: 1) with water was placed in the cage to facilitate the mated flies to lay their eggs. The larvae on hatching continued to feed on wheat bran medium. When the larvae in the medium reached third instar, ragi (*Elusine coracona*) husk was added to the dry larval medium to facilitate pupation. The fly rearing unit was maintained at 28±2°C, RH 65±5 %. The bio stages of the flies collected from the rearing chamber were used in experiments.

**Extraction and characterization of essential oil**

Fresh leaves of *O. basilicum* (300 g) was extracted by hydro-distillation in a Clevenger type apparatus to obtain EO. The leaves along with 500 ml of water were loaded into a round bottom flask placed over a heating mantle. The contents in the flask were heated to 100°C. The oil along with water collected in the receiver tube was phase separated using a separating funnel. The collected essential oil was dried by passing over anhydrous sodium sulphate to remove the moisture trace and stored in amber vials at 4°C until use.

The *O. basilicum* oil was characterized using GC-MS (Agilent GC- 7890A (G3440A) and MS- G3171A 5975) as suggested by Ravindran et al. (2019). One µl of 0.01 % essential oil diluted in dichloromethane was introduced by microsyringe into injector port attached to HP-5 MS Phenylmethylsilox capillary column (30 m × 250 µm i.d. × 0.25 µm film thickness, Agilent Technologies, USA) through a glass liner. The oven and column temperature were maintained at 40°C for one minute and then raised at the rate of 20°C per minute to 280°C. The temperature during the post run was held at 300°C for 10 min. The temperature in injector and detector was maintained at 250°C. The total run was for 23 min. In MS, the ion source temperature was set at 250°C. The flow rate of the carrier gas Helium was maintained at 1 ml per minute. An electron ionization (EI) mode with 70 eV ionization energy was used for the GC–MS identification. Identification of
the components in EO was done by taking into account the relative peak percent area, retention time and mass fragmentation pattern using the NIST library. Major constituents in oil were verified by co-injecting the compounds.

**Chemicals**

Methyl chavicol and linalool used for bioassays was purchased from Sigma Aldrich. HPLC grade dichloromethane and acetone were purchased from Merck. Dimethyl-2,2-dichlorovinyl phosphate (DDVP) and Imidacloprid (PESTANAL®) analytical standard were purchased from Sigma Aldrich. Eserine and Fast Blue RR salt were procured from Fluka (Sigma Aldrich). α-Napthyl acetate (α-NA), 2,4-dinitrochlorobenzene (CDNB), Coomassie brilliant blue G-250, pyrogallol, guaiacol, and H₂O₂ were purchased from Shanghai Chemical Industry Co., Ltd, China.

**Electroantennography (EAG)**

The response of *M. domestica* adult female antennae (3 days old) to sweet basil EO, methyl chavicol, linalool and neem oil was recorded using an electroantennographic system (Syntech). The dual electrode probe was used for mounting the antennal prep. The head of the adult female fly was decapitated and mounted on one electrode and the proximal tip funiculus to another electrode using a conductive gel (Spectra 360 Parker, Orange, New Jersey). The clean air (activated charcoal filtered) was continuously flushed over the antennae. The EO and its constituent's methyl chavicol, linalool and neem oil were diluted in HPLC grade dichloromethane at concentration of 1 µg µL⁻¹. Dichloromethane alone was used as a control. One µl of the aliquot amounting to 1 µg of the test compound placed on Whatman filter paper strips (Advantec 5C (110 mm) Japan of 2 cm length and 4 mm diameter) was dried for 5 min in fume hood and then it was inserted into the Pasteur pipettes. This setup was connected to stimulus controller (CS 05 Syntech) by Tygon silicone tube. The first puff was blown off after 30 seconds of loading filter paper. After sixty seconds, the antennae were exposed to vapour phase of the stimulus through pipette placed 15 mm upstream from the antennae that had continuous air stream (pulse time 0.5 seconds, continuous flow 25ml/s, pulse flow 21 ml/s) as suggested by Vibina et al. (2019). Between the stimulus puffs a time delay of 20 seconds was maintained. The antennal responses were recorded through a high impedance probe that was in turn connected to amplifier (IDAC-4, Syntech) and the signals were recorded with EAG software (Syntech). Responses were expressed as a summated response of neurons, sorted according to shape and amplitude, emitted during 1 sec after the onset of the stimulation. The control stimulus was at the beginning, middle and end of each session. EO and its constituents and neem oil were tested on six fly antennae with four replications of per stimuli per antennae in randomized manner.

**Y olfactometer assay**

The olfactory response of 2 – 3 days old adult females to sweet basil EO its constituent methyl chavicol, linalool and neem oil was evaluated using a glass Y - tube olfactometer having a main arm of 17 cm and choice arms of 17 cm with an inner diameter of 4 cm. Atmospheric air pumped using an air sampler was allowed to flow out through activated charcoal cartridge with a flow rate of 0.5 L / min. The purified air was then let into the arms of the Y - tube with steady flow rate. Whatman No 1 filter paper stripes (3 cm length
and 0.5 cm width) were treated with 200 µL of 20 % sugar solution and dried for 30 minutes. To this 10 µL of 100 ppm of odorants diluted in dichloromethane was loaded and the paper strips were allowed to dry in room temperature for 10 minutes to permit the solvent to evaporate. The paper strips prepared as mentioned above without the odorants was used as control. Both treated and control paper strips were inserted into an odor tube that was connected between the air flow tube and the Y-tube arm. The entire Y-tube setup was placed inclined at 20º. Pairwise comparison was made between odorants and control. The adult female flies were starved for 4 hours with water satiation prior to test. A total of 100 female flies (N=100) were introduced into the main arm. The choice made by the flies in an arm was considered if they crossed the halfway mark made in the arm in three minutes of start of the test. Those flies that failed to participate in the test were considered non respondents. The odorants were switched between the arms to avoid position effect. The test was conducted at room temperature of 25 ± 2°C under red light with slight modification as suggested by Ravindran et al. (2019).

**Oviposition repellence**

Gravid *M. domestica* females (10 Nos.) housed in cage (30 L × 30 W × 30 H cm) made of aluminium frame having acrylic sheets were exposed to 2 gms of oviposition substrate (wheat bran + milk powder + egg yolk powder (10: 2: 1) treated with EO, methyl chavicol, linalool and neem oil dissolved in acetone so as to achieve 0.05, 0.15 and 0.25 % concentrations. Oviposition substrate treated with acetone alone was maintained as control. In both the cases, the solvent was allowed to evaporate from the oviposition substrate prior to their placement in treated and control dish that were placed at diagonal ends in the floor of the cage. Four replications were maintained (One cage per replicate). The number of eggs laid were counted after 24 hrs.

Oviposition Activity Index (OAI) was calculated using the formula

\[ OAI = \frac{(NT - NC)}{(NT + NC)} \]

where NT = total number of eggs on the treated substrate and NC = total number of eggs on the control substrate (Cheah et al. 2013). For OAI values ≤-0.3, the EO was considered as repellent (Kramer and Mulla 1979).

The percent effective repellency (ER%) for EO, linalool, methyl chavicol and neem oil were calculated using the following formula:

\[ ER% = \frac{NC - NT}{NC} \times 100 \]

as suggested by (Phasomkusolsil and Soonwera 2012).

**Ovicidal effect**

Freshly laid *M. domestica* eggs (0 - 3 hrs), were collected using a soft paint brush. Twenty eggs of *M. domestica* were placed in the base of the 50 ml plastic sample container (Tarsons) having screw cap lid. Sweet basil EO, methyl chavicol and linalool of varied concentration ranging from (0.5 - 79mg/dm³) were applied to filter paper attached to the base of the lid. Acetone alone was maintained as control and DDVP was maintained as positive control. This setup was sealed tightly using a parafilm and placed in incubator...
at 28 ± 2ºC and RH 65 ± 5%. The number of hatched and unhatched eggs were counted after 48 hrs. The experiments were replicated for five times.

Contact toxicity

Topical bioassay:

Acute toxicity of EO, methyl chavicol and linalool to female *M. domestica* was assessed by topical application. Acetone was used as a solvent to prepare varying concentration of EO, methyl chavicol and linalool. Acetone alone was maintained as negative control and imidacloprid was used as positive control. Preliminary range finding test was done to fix the doses. Two-day old adult female flies were immobilized using CO$_2$. The flies were transferred to flat surface and 1µl of the test compounds were placed on the pronota using a micro applicator. The treated adults were transferred to an insect breeding dish (Himedia) having ventilation in the lid. The adults had access to 10 % honey solution soaked in absorbent cotton. Four replications were maintained per treatment. Mortality was assessed 24 hrs after the treatment. Those flies that ceased to move its appendages in mild pin prick were considered dead.

Enzyme assays

The surviving flies after treatment with LC$_{50}$ doses of EO, methyl chavicol and linalool (9.98, 13.62 and 43.13 µg / adult respectively) were used for the extraction of whole-body homogenate. Acetone treated adults were used as control. The flies were homogenised in ice cold 50 mM phosphate buffer (pH 7.4). Homogenates were centrifuged at 4 ºC for 20 minutes at 10,000 rpm. The supernatant was subjected to protein estimation using Bradford's method (Bradford 1976) and used as an enzyme source for the estimation of carboxylesterase (CarE) and glutathione S-transferase (GST) activities.

Glutathione S-transferase (GST) activity was determined as described by Habig et al. (1974) and Zhang et al. (2007). The components in reaction mixture include 30µl of enzyme solution, 10 µl each of reduced glutathione (GSH) and 1-chloro-2,4-dinitrobenzene (CDNB) (dissolved in ethanol and prepared with 50 mM po4 buffer, PH 7.4) and 950 µl phosphate buffer. Control mixture had 10 µl each of GSH and1-chloro-2,4-dinitrobenzene and 950 or 980 µl of phosphate buffer. The rate of change in absorbance at 340 nm was measured for 5 minutes in 96-well microplate reader and converted to specific activity using extinction coefficient of 9.6 mM$^{-1}$ after necessary path length correction.

The activity of carboxylesterases (Car E) was determined using the method suggested by Li et al. (2007). The reaction mixture consisting of 15µl of enzyme homogenate made upto 500 ul with sodium phosphate buffer (50 mM, pH 7.4) and 800 µl α-naphthyl acetate (dissolved in acetone, prepared in PO$_4$ buffer) was incubated in dark condition for 20 minutes. The control mix had 800µl of substrate and 500µl of phosphate buffer (50mM, pH 7.4). Formation of blue colour on addition of 200 µl of staining solution indicated the production of α-napthol. The absorbance was measured at 595 nm using micro plate reader (iMark, BioRad). Enzyme activity in the sample was calibrated using α-napthol standard curve.
For both the enzyme estimation, three biological replicates were maintained per treatment. The specific activity of the enzyme was expressed as nmol/min/mg protein.

**Statistical analysis**

The egg and adult mortality data were subjected to probit analysis to determine median lethal dose LD$_{50}$ and the corresponding 95% CI values and chi-square test were calculated. Pooled EAG response of antennae to EO and its constituents, variations in effective repellence (ER), OAI and variation in enzyme activity was compared using one-way analysis of variance (ANOVA) followed by Tukey's Post hoc test (P<0.05) using SPSS. In case of Y tube olfactory assay, the binomial test was used to compare the orientation of the flies to the arms with odorant and control.

**Results**

**GC-MS analysis for Ocimum basilicum EO**

The extracted *O. basilicum* EO was characterized by gas chromatography coupled to mass spectroscopy. The compounds were identified by the reported retention index. The chemical formula, molecular weight and their chemical structures are presented in Table-1 and Figure 1. The major components present in *O. basilicum* EO were methyl chavicol (70.93%), linalool (9.34%), epi-α-cadinol (3.69%), methyl eugenol (2.48%), γ-cadinene (1.67%), 1,8-cineole (1.30%) and (E)-β-ocimene (1.11%).

**Table 1 Chemical composition of the *O. basilicum* essential oil**
| No | Component            | Chemical formulae | Molecular weight g/mol | Reported RRI | Experimental RRI | Percentage Composition |
|----|----------------------|-------------------|------------------------|--------------|-----------------|------------------------|
| 1  | β-Pinene             | C$_{10}$H$_{16}$  | 136                    | 974          | 978             | 0.12                   |
| 2  | Myrcene              | C$_{10}$H$_{16}$  | 136                    | 988          | 991             | 0.48                   |
| 3  | Limonene             | C$_{10}$H$_{16}$  | 136                    | 1024         | 1028            | 0.14                   |
| 4  | 1,8-Cineole          | C$_{10}$H$_{18}$O | 154                    | 1026         | 1032            | 1.3                    |
| 5  | (E)-β-ocimene        | C$_{10}$H$_{16}$  | 136                    | 1044         | 1047            | 1.11                   |
| 6  | Linalool             | C$_{10}$H$_{18}$O | 154                    | 1095         | 1101            | 9.34                   |
| 7  | Camphor              | C$_{10}$H$_{16}$O | 152                    | 1141         | 1140            | 1.44                   |
| 8  | Methyl chavicol      | C$_{10}$H$_{12}$O | 148                    | 1195         | 1208            | 70.93                  |
| 9  | p-Anisaldehyde       | C$_{8}$H$_{6}$O$_{2}$ | 137              | 1247         | 1257            | 0.28                   |
| 10 | Bornyl acetate       | C$_{12}$H$_{20}$O$_{2}$ | 196              | 1287         | 1288            | 0.1                    |
| 11 | β-Bourbonene         | C$_{15}$H$_{24}$  | 204                    | 1387         | 1387            | 0.09                   |
| 12 | β-Elemene            | C$_{15}$H$_{24}$  | 204                    | 1389         | 1394            | 0.62                   |
| 13 | Methyl eugenol       | C$_{11}$H$_{14}$O$_{2}$ | 178            | 1403         | 1408            | 2.48                   |
| 14 | β-Caryophyllene      | C$_{15}$H$_{24}$  | 204                    | 1417         | 1422            | 0.45                   |
| 15 | α-Guaiene            | C$_{15}$H$_{24}$  | 204                    | 1437         | 1441            | 0.29                   |
| 16 | α-Humulene           | C$_{15}$H$_{24}$  | 204                    | 1452         | 1456            | 0.24                   |
| 17 | Germacrene D         | C$_{15}$H$_{24}$  | 204                    | 1484         | 1484            | 0.16                   |
| 18 | Bicyclogermacrene    | C$_{15}$H$_{24}$  | 204                    | 1500         | 1499            | 0.35                   |
| 19 | α-Bulnesene          | C$_{15}$H$_{24}$  | 204                    | 1509         | 1508            | 0.63                   |
| 20 | γ-Cadinene           | C$_{15}$H$_{24}$  | 204                    | 1513         | 1517            | 1.67                   |
| 21 | trans-Calamenene     | C$_{15}$H$_{22}$  | 202                    | 1521         | 1526            | 0.12                   |
| 22 | (E)-Nerolidol        | C$_{15}$H$_{26}$O | 222                    | 1561         | 1566            | 0.11                   |
| 23 | 4-Methoxycinnamaldehyde | C$_{10}$H$_{10}$O$_{2}$ | 162          | 1562         | 1571            | 0.67                   |
Electroantennograph (EAG) response

The sweet basil EO and its constituents methyl chavicol, linalool and positive control neem oil at 1 µg treatment caused the neuronal response in female adults (Fig. 2)\((t = 2.71; P< 0.05)\). The mean EAG response to control was less than 0.5 mV. Sweet basil EO and its major constituent caused highest mean antennal response of 2.51 and 2.38 mV respectively and were at par. The response of female adults to linalool (1.69 mV) was lower than the methyl chavicol but higher than neem oil (1.37 mV).

Olfactometer bioassay

There was significant reduction in adult female flies selecting the arm having food source with sweet basil EO \((P< 0.05)\) and methyl chavicol \((P<0.05)\) as compared to arm having food source alone (Control). However, there was no significant difference between the flies choosing the arms having food source with linalool \((P = 0.22)\) and neem oil \((P = 0.7)\) and control having food source alone (Fig. 3).

Ovipositional repellence

Effective repellence (ER%) and oviposition activity index (OAI) of *O. basilicum* EO and its constituent’s methyl chavicol and linalool at three concentration (0.05, 0.15 and 0.25%) were determined. The ER% and OAI for EO and its constituents was concentration dependant and it increased with rise in concentration. Among the samples tested, higher oviposition deterrence to *M. domestica* adults was observed in EO \((89.52\pm1.00)\), methyl chavicol \((80.12\pm 2.40)\) and linalool \((78.99\pm2.93)\) at a dose of 0.25 % which was superior to neem oil 0.25%. EO and methyl chavicol at 0.15 % were at par in causing oviposition deterrence to *M. domestica* adults. At a lower concentration of 0.05% the EO, methyl chavicol and linalool caused a lowest deterrence ranging from 27.83 - 35.86 % \((F_{11,36} = 50.07 \ P < 0.001)\). Across the concentration, neem oil caused lower oviposition deterrence.
O. basilicum EO, methyl chavicol, linalool and neem oil caused significant difference on OAI. O. basilicum EO, methyl chavicol at 0.25% caused an OAI -0.81 ± 0.01 and -0.67 ± 0.03, respectively. All the samples tested at 0.15% and 0.05% recorded OAI value below 0.5. An OAI value from -0.5 to -1 an indication of better oviposition repellence. In case of neem oil all the doses tested had OAI less than -0.5 (F_{11,36} = 53.87 P < 0.001) (Table 2).

Table 2: Effective repellence of O. basilicum EO, methyl chavicol, linalool and neem oil against M. domestica

| Test sample     | Concentration (%) | Effective Repellency (ER) | OAI      |
|-----------------|-------------------|---------------------------|----------|
| O. basilicum EO | 0.05              | 35.86 ± 0.88^e            | -0.21 ± 0^d |
|                 | 0.15              | 71.44 ± 4.24^bc           | -0.50 ± 0.02^c |
|                 | 0.25              | 89.52 ± 1.00^a            | -0.81 ± 0.01^a |
| Methyl chavicol | 0.05              | 33.27 ± 0.95^e            | -0.19 ± 0.0^d |
|                 | 0.15              | 59.81 ± 4.36^cd           | -0.43 ± 0.04^c |
|                 | 0.25              | 80.12 ± 2.40^ab           | -0.67 ± 0.03^ab |
| Linalool        | 0.05              | 31.28 ± 2.07^e            | -0.18 ± 0.01^d |
|                 | 0.15              | 56.54 ± 3.75^d            | -0.39 ± 0.03^c |
|                 | 0.25              | 78.99 ± 2.93^ab           | -0.65 ± 0.03^b |
| Neem oil        | 0.05              | 27.83 ± 3.57^e            | -0.16 ± 0.02^d |
|                 | 0.15              | 53.95 ± 1.70^d            | -0.37 ± 0.01^c |
|                 | 0.25              | 62.31 ± 4.18^cd           | -0.45 ± 0.04^c |

The data are given as Mean ± SE. *Denote significant different at P < 0.05 compared with the control.

Means followed by same alphabet in a column do not differ significantly by Tukeys test p<0.05

Ovicidal effect

O. basilicum EO and its constituent methyl chavicol and linalool were evaluated for ovicidal activity. O. basilicum EO had 5.8-fold higher ovicidal activity (EC_{50} 9.74 mg/dm\(^3\)) on M. domestica eggs than linalool (EC_{50} 13.57 mg/dm\(^3\)). Among the constituent’s methyl chavicol was 1.27-fold higher toxicity than linalool (EC_{50} 10.67 mg/dm\(^3\)). The ovicidal activity of DDVP was superior to (EC_{50} 390.37 mg/dm\(^3\)) EO and its constituents (Table 3).

Table 3. Ovicidal activity of O. basilicum EO, methyl chavicol and linalool on M. domestica eggs
| Test sample        | Concentration (mg/dm$^3$) | 48 h% mortality ± SE | EC$_{50}$ (mg/dm$^3$) | 95% CL | df | Chi-square | p value |
|-------------------|---------------------------|----------------------|------------------------|--------|----|------------|---------|
| O. basilicum EO   | 45.59                     | 100 ± 0              |                        |        |    |            |         |
|                   | 30                        | 86 ± 2.91            | 9.74                   | 7.36 – 12.75 | 5  | 10.79      | 0.058   |
|                   | 22.79                     | 68 ± 2.54            |                        |        |    |            |         |
|                   | 11.4                      | 51 ± 4.30            |                        |        |    |            |         |
|                   | 5.7                       | 28 ± 2.54            |                        |        |    |            |         |
|                   | 0.3                       | 16 ± 1.87            |                        |        |    |            |         |
|                   | 0.15                      | 6 ± 1.0              |                        |        |    |            |         |
| Methyl chavicol   | 45.59                     | 99 ± 0.00            |                        |        |    |            |         |
|                   | 30                        | 85 ± 1.58            |                        |        |    |            |         |
|                   | 22.79                     | 72 ± 2.54            |                        |        |    |            |         |
|                   | 11.39                     | 48 ± 2.54            |                        |        |    |            |         |
|                   | 5.6                       | 29 ± 2.91            |                        |        |    |            |         |
|                   | 2.8                       | 8 ± 1.22             |                        |        |    |            |         |
|                   | 1.5                       | 3 ± 1.22             |                        |        |    |            |         |
| Linalool          | 79.33                     | 100 ± 0.00           |                        |        |    |            |         |
|                   | 63.47                     | 94 ± 2.44            |                        |        |    |            |         |
|                   | 39.66                     | 84 ± 4.30            |                        |        |    |            |         |
|                   | 23.8                      | 74 ± 1.87            |                        |        |    |            |         |
|                   | 15.86                     | 63 ± 2.54            |                        |        |    |            |         |
|                   | 11.9                      | 44 ± 2.91            |                        |        |    |            |         |
|                   | 7.93                      | 20 ± 2.23            |                        |        |    |            |         |
|                   | 3.96                      | 12 ± 2.54            |                        |        |    |            |         |
| DDVP              | 1.2                       | 95 ± 2.73            | 0.15                   | 0.13 – 0.18 | 5  | 1.43       | 0.92    |
Each value represents the mean of five replicates, and each set-up had 20 individuals (n=100).

95% CL=confidence interval at 95% confidence level.

**Topical bioassay**

The toxicity assay carried out by topical application showed that *O. basilicum* EO caused higher mortality over methyl chavicol and linalool to adult stages. Adults exposed to EO (LD$_{50}$ 10.01 μg/adult) were more susceptible than those exposed to methyl chavicol and linalool (LD$_{50}$ 13.62 μg/adult and LD$_{50}$ 43.12 μg/adult respectively). Imidacloprid was highly toxic (1.41 μg/adult) than EO and its constituents (Table 4).

**Table 4: Acute toxicity of *O. basilicum* EO, methyl chavicol and linalool on *M. domestica*- Topical application**

| Test sample       | Stage | Period (Hours) | LD$_{50}$ * | 95% CL     | df | Chi-square | P value |
|-------------------|-------|----------------|-------------|------------|----|------------|---------|
| *O. basilicum* EO | Adult | 24             | 10.01       | 9.19 – 10.94 | 6  | 11.83      | 0.066   |
| Methyl chavicol   | Adult | 24             | 13.62       | 10.76 – 16.78 | 4  | 9.19       | 0.05    |
| Linalool          | Adult | 24             | 43.12       | 23.80 – 64.39 | 2  | 5.93       | 0.05    |
| Imidacloprid      | Adult | 24             | 1.41        | 1.19 – 1.68  | 5  | 2.8        | 0.718   |

*(μg/adult)*

**Enzyme assay**

The effect LD$_{50}$ dose of *O. basilicum* EO, methyl chavicol and linalool on *M. domestica* adult detoxifying enzymes carboxyl esterase and GST are reported in the Table 5.
Adults exposed to linalool at LD$_{50}$ value caused inhibition of GST but it was on par with control. Whilst LD$_{50}$ values of EO and methyl chavicol caused induction of GST levels with EO having a higher ER (4.61) than methyl chavicol (3.09) ($F_{6,14} = 224.49$ P<.005). Topical application of EO, methyl chavicol and linalool induced the enzymes Car E in adults. In case of Car E levels in adults, those exposed to methyl chavicol at LD$_{50}$ had Car E levels at par with control samples. Linalool had higher level of induction of Car E in adults than $O. basilicum$ EO ($F_{6,14} = 224.49$ P<.005).

Table 5. Activities of carboxylesterase and glutathione S-transferase in $M. domestica$ adult

| Treatment                  | GST        | ER    | Car E   | ER    |
|----------------------------|------------|-------|---------|-------|
| Control                    | 15.58 ± 0.49$^{e}$ | 1.35 ± 0.04$^{e}$ |          |       |
| $O. basilicum$ EO LD$_{50}$ | 71.83 ±1.92$^{a}$ | 4.61  | 2.21 ±0.00$^{b}$ | 1.63  |
| Methyl chavicol LD$_{50}$  | 48.16 ±2.76$^{b}$ | 3.09  | 1.53 ±0.03$^{de}$ | 1.13  |
| Linalool LD$_{50}$         | 13.86 ±0.37$^{e}$ | 0.88  | 2.84 ±0.07$^{a}$ | 2.10  |

$^{a}$ Means within a column followed by the same letter are not significantly different (One-way ANOVA).

$^{b}$ Unit of enzymes: GST - Glutathione S-transferase and Car E - Carboxylesterase = n moles/min/mg protein/min.

The enzyme activities were expressed as enzyme ratio (ER, mean activity of enzyme in different treatments/mean activity of enzyme in control group).

Discussion

Indiscriminate use of chemicals has led to development of insecticide resistance and negative effect to consumers. To tide over the ill effects caused by chemical insecticides there is a growing demand for environmentally safe pesticides to manage $M. domestica$. Essential oils (EO) derived from plant parts are an alternative as they possess bioactive compounds that can be used in isolation or in combination for managing the pests of agriculture, medical and veterinary importance (Isman 2006; Pavela 2011; Pavela and Benelli 2016; Khater and Geden 2019). Biodegradable nature and safety to nontargets puts EOs as an alternative to synthetic insecticides (Poorjavad et al. 2014). The potency of essential oil and its constituents against $M. domestica$ has been reported earlier (Sinthusiri and Soonwera 2014; Benelli et al. 2018b).
Insecticidal, ovicidal, deterrence/repellence and growth regulating effect of *O. basilicum* crude extracts and EO against *M. domestica* was reported earlier (Pavela 2008a; El Zayyat et al. 2015; Chowdhary et al. 2018).

**Chemical characterization**

In our study, the sweet basil EO had methyl chavicol (70.93 %) and linalool (9.34%) as major constituents. This agrees with earlier reports mentioning methyl chavicol, linalool, β-elemene and camphor as major constituents of basil oil (Sajjadi 2006; Sonmezdag et al. 2018). The proportion of the constituents in basil oil vary according to the chemotypes of plant (Telci et al. 2006; Ogendo et al. 2008) viz., methyl chavicol, linalool, methyl eugenol and methyl cinnamate types (Lawrence 1998). The EO of *O. basilicum* had thirty components, accounting for 99.31% of the total composition. As the EO used in our study had higher load of methyl chavicol (70.93%) it could be considered as methyl chavicol chemotype as proposed by Varga et al. (2017).

In our study, EAG assay revealed that summated response of the olfactory neurons in the antennae of adult female *M. domestica* was higher to sweet basil EO and methyl chavicol followed by linalool and neem oil. As EAG response is linked to insect olfaction (Ghabbari et al. 2018) higher amplitude of response trace to EO and methyl chavicol may be due to higher number of receptors in antennal neuron to these stimuli that may influence the fly’s behaviour. Basil oil when presented at 100 ng caused mean antennal response of 0.99 mv in coconut rhinoceros beetle, *Oryctes rhinoceros* (Ravindran et al. 2019). White and Hobson (1993) reported that methyl chavicol caused EAG response in mountain pine beetle. Screening of the odorants for the antennal response using EAG provides a lead to select the compounds for behavioural assay (Cosse et al. 1995; Zito et al. 2013; Ruschioni et al. 2015) as the EAG active compounds in the essential oils may elicit attractive or repulsive behavioural response (Meza et al. 2020)

Compounds that cause physiological response in antennae need not be behaviorally active, hence an olfactory assay must be done to establish the behavioral response (Ravindran et al. 2019). In Y– tube olfactory assay, the adults of *M. domestica* when provided a choice preferred the control arm having food source alone over the arm having food source laced with sweet basil EO and methyl chavicol. This aversive response of *M. domestica* adults to EO and methyl chavicol may be due to its ability to perceive the odorants in antennae. The peripheral response of antennal neurons to EO and methyl chavicol in EAG provides a physiological basis for the aversive olfactory mediated behaviour in Y- tube assay. The essential oils of *Ocimum gratissimum*, *Thymus serpyllum*, *Myristica fragrans* and *Curcuma amada* caused 100% repellence to housefly for a duration of 5 h. (Singh and Singh 1991). Methyl chavicol a constituent in *Tagetes filifolia* and *O. selloi* were found to be repellent to *Aedes aegypti* (Diptera) (Gleiser et al. 2011) and *Anopheles braziliensis* (Diptera) (Padiha Paula et al. 2003). Sadeh et al. (2018) reported that rosemary variety 11 having 0.6 – 0.9 % methyl chavicol caused repellency to whitefly, *Bemisia tabaci*. Salom and Kenneth (1995) reported that methyl chavicol occurring in the pine trees by its repellent action reduced 40-68 % aggregation of *Dendroctonus* beetles. Repellents prevent the orientation / alighting of housefly on treated surface there by scaling down the possibilities of spread of diseases in locations where the population of housefly is high (Haselton et al. 2015). Identifying the repellents of house fly is an paramount important because its pestiferous status in agriculture and human health (Malik et al. 2007).
In the present study *O. basilicum* EO, methyl chavicol and linalool at 0.25% showed highest ovipositional repellence against *M. domestica* ranging from 79 – 89 % and OAI of - 0.65 to - 0.67. Essential of *O. gratissimum* at 2 % caused 100 % repellent activity against *M. domestica* (Singh and Singh 1991). Repellence of essential oil against *M. domestica* has impact on the population build up (Maganga et al. 1996). The toxic effects of *O. basilicum*, EO and its constituent's methyl chavicol and linalool to larval, pupal and adult stages of *M. domestica* have been documented (Pavela 2008b; Tarelli et al. 2009; Scalerandi et al. 2018). The success of pest management is at high when the vulnerable bio stage of insects is targeted. Eggs of *M. domestica* being a non-motile stage are easier target than motile larval and adult stages. Identifying the plant derived parts like essential oil having the ovicidal activity will aid to contain the pest buildup (Hong et al. 2018). *O. basilicum* EO had 5.8-fold higher ovicidal activity on *M. domestica* eggs than its constituent’s linalool and methyl chavicol. Eggs of mosquito vectors, *Aedes aegypti*, *Anopheles dirus* and *Culex quinquefasciatus* exposed to *O. basilicum* oil had extremely low hatchability (EC50<1.9%) (Phasomkusolsil and Soonwera 2012). *O. basilicum* oil at 1.0 ml/ 38.5 ml of air caused 100% mortality of *Callosobruchus chinensis* (L.) three days after exposure period (Abd El salam 2010). The ovicidal activity of linalool against the eggs of insecticide resistant lice BR-HL females proved to be a novel ovicide (Yang et al. 2009). Transgenic *Nicotiana tabacum* producing higher level of linalool were not preferred for oviposition by *Helicoverpa armigera* as linalool was reported to cause repellence (McCallum et al. 2011).

We observed the topical application of *O. basilicum* EO to be more toxic than methyl chavicol and linalool to adult stages of *M. domestica*. Increased toxicity in EO may be due to the combined effect of the compounds in natural oil as reported by Cheng et al. (2009). In contrary to this, methyl chavicol (11.01 ppm) and linalool (35.17 ppm) in isolation had higher larvicidal activity than *Clausena anisate* EO as a whole (LC50 119.59 ppm) (Govindarajan 2010). The structure and functional group of terpenoids facilitates their entry into the insect cuticle and attach to the target site to bring in desirable bioaction (Rice and Coats 1994).

Resistance in insects to xenobiotics and phytochemicals is due to metabolic detoxification mediated by the action of enzymes like, glutathione S-transferase (GST) and esterase (Castaneda et al. 2010; Waliwitiya et al. 2012). GST is involved in metabolism of endogenous compounds and acts by conjugation to make them water soluble and less toxic (Yu 2004). Esterases act by binding, sequestering and detoxification of toxic chemicals (Hegeto et al. 2015).

Our investigations reveal that GST and Car E activity in adult stage of *M. domestica* increased on exposure to *O. basilicum* EO and its constituent’s methyl chavicol and linalool. *M. domestica* adults exposed to EO at LD50 values induced a higher level of GST as compared to control and its constituents. Treating with LD50 doses of linalool caused significant increase in Car E levels in adults .This agreement with earlier studies that *C.pipiens* larvae exposed *O. basilicum* EO at LC50 values caused a stimulation of the detoxification GST activity. Increase in degrading enzymes may be increase in active compounds in insect body so as to bring in toxicity (Zibaee and Bandani 2010). Previous reports state that plant derived products having mixture of compounds that include triterpenoids and phenols inhibit the activity of GST and Car E (Tak et al. 2017; Yang et al. 2018). Increase in the degrading enzymes Car E and GST on exposure of insect to natural
products have been reported earlier (Kumrungsee et al. 2014). Mosquitoes, *C. quinquefasciatus*, *A. stephensi* exposed to *Lantana camara* root and *Anacardium occidentale* caused a rise in GST activity in larvae. This scenario of rise in enzymes does not limit to insecticide resistance alone, this could be due to degrading enzymes generated by reactive oxygen species (ROS) (Vontas et al. 2001). Induction of detoxifying enzymes due to exposure to EO may be due to synergy caused by the constituents in EO (Miresmailli et al. 2006; Isman et al. 2008; Jiang et al. 2009; Senthil-Nathan 2013). Increase in degrading enzyme levels is clear indication that EO and its constituents possess cidal effect in *M. domestica* and the fall out of which is the possibilities of resistance development, which needs further investigation.

The work in this study has demonstrated the bioactive potential of *O. basilicum* EO and its constituent's methyl chavicol and linalool in causing toxicity to eggs and adult stage. The electrophysiological and the behavioral response of adult female flies to EO and its constituents treated substrate supports their impact by exhibiting repellence and ovipositional deterrence there by serving as an olfactory barrier to avoid the files from settling in human settlements and animal sheds . EO as whole possess a higher ovicidal, adulticidal and repellency to flies and these traits make it fit to be harmoniously blended in integrated pest management (IPM) of *M. domestica*. Further studies are needed to develop a formulation of EOs of sustained spatio temporal release rates as they would be techno economically feasible to use as there is a market acceptance for plant derived parts for managing the pest like *M. domestica* that are common in dwelling areas of humans, birds and animals.

**Declarations**

**Ethics approval and consent to participate:** Not applicable.

**Consent for publication:** Not applicable

**Availability of data and materials:** Data is available by request to the corresponding author

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**Authorship contribution**

**Rajendran Senthooraraja:** Investigation, Data generation. **Kesavan Subaharan:** Conceptualization, Writing - Original Draft, Data Curation. **Sowmya Manjunath:** Investigation on biochemical aspects. **Muthu Gounder Mohan:** Resources. **Vppalayam Shanmugam Pragadheesh:** Investigation, Validation. **Nandagopal Bakthavatsalam:** Writing - Review & Editing. **Sekarappa Basavarajappa:** Writing - Review & Editing. **Sengottayan Senthil-Nathan:** Writing - Original Draft and editing.
All authors read and approved the final manuscript.

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Figures

Figure 1

Structures of chemical constituents in O. basilicum essential oil
Figure 2

EAG response of M. domestica against EO and its constituents Mean (+ SE) antennal EAG responses of female M. domestica to the assay performed using 1 μL of 1000 ppm of sweet basil EO, methyl chavicol, linalool and neem oil. Bars having same letter do not differ significantly at P < 0.05 (ANOVA followed by Tukey test)
Figure 3

Y-tube olfactometer response of female M. domestica against EO and its constituents Behavioural response of adult female M. domestica in a two choice Y- olfactometer (Per cent repelled n = 100). Starved adult female flies were offered a choice between treated arm (Food source with sweet basil EO and its constituents’ methyl chavicol, linalool and neem oil) and control arm (Food source alone). Both bars represent the per cent flies orienting to treated and control arm. The non-responding adult female flies were presented in right hand side (No choice %). Asterisks shows a preference that was significantly different (binomial test) from a 50:50 distribution: * = P < 0.05. The flies that failed to respond were excluded from the statistical analysis.