Efficacy of used coffee grounds as larvicide against *Aedes albopictus* (Skuse, 1894) and *Ae. aegypti* Linné, 1762 (Diptera: Culicidae)

A. Drago¹*, S. Vettore¹, S. Martini¹ and M. Dutto²

¹Entostudio S.r.l., Viale del Lavoro 66, 35020 Ponte San Nicolò (PD), Italy; ²Studio di Entomologia e Fitopatologia, Via Papò 4, 12039 Verzuolo (CN), Italy; drago@entostudio.com

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### Abstract

*Aedes albopictus* and *Aedes aegypti* are two synanthropic, anthropophilic container-breeding mosquitoes. These species are very annoying, but are also vectors of dengue, chikungunya, yellow fever, Zika, and Usutu viruses, and other pathogens. Because these mosquitoes breed very close to humans, cheap homemade methods, as alternatives to commercial insecticides, could be important for their control. Coffee being a very common beverage, the grounds extracted from used coffee capsules have been tested for their larvicidal efficacy. The grounds were extracted with either 30 ml or 70 ml of 65-70 °C water. The content of one capsule was used as a unit dose to treat the quantity of water contained in a medium sized flowerpot tray. The test provided a clear indication that at this dosage, the used coffee capsules were completely ineffective at killing the larvae of *Aedes* species, so this method cannot be suggested to control these mosquitoes.

### Keywords: mosquitoes, coffee, larvicide, *Aedes albopictus*, *Aedes aegypti*

### 1. Introduction

*Aedes albopictus* (Skuse, 1894) and *Aedes aegypti* Linné, 1762 are two species of relevant health importance due to their arbovirus vectorial capacity (Tsai and Teng, 2016). Only *Ae. albopictus* is to date widely and permanently established in Europe, where it is spreading by about 100 km/year (Kraemer *et al.*, 2019). *Ae. aegypti* is currently present in Europe only on Madeira and in the coastal region of the Black Sea in southern Russia, Abkhazia and Georgia (Medlock *et al.*, 2012), although it has been sporadically found in various European states following imports (Medlock *et al.*, 2012). The ongoing climate change (increase in temperatures) together with the increasing of global trade could favour the expansion of these species. *Ae. aegypti* in particular, thereby increasing the risk of vector-borne disease outbreaks on the European continent (Liu-Helmersson *et al.*, 2019).

The role of vectors, but above all the nuisance, makes the fight against these domestic mosquitoes necessary. The human population, in situations of non-infectious emergency, seems to give preference to the use of popular insecticide methods (authors’ observation) the real efficacy of which is, in most cases, unknown. The internet has, surely, contributed to the wide circulation of information regarding possible alternative and biological methods for fighting mosquitoes. This led to the testing of different substances and materials including metallic copper (Della Torre *et al.*, 1993), coffee grounds (Dieng *et al.*, 2018; Ellias *et al.*, 2015; Guirado and Bicudo, 2007; Laranja *et al.*, 2003; Wiwanitkit, 2018) and tea extracts (Dieng *et al.*, 2016). One advantage of the use of alternative insecticides would be to overcome the onset of resistance phenomena (Dieng *et al.*, 2016).
Given that coffee is a very common beverage in Europe (Landais et al., 2018) and therefore coffee grounds are widely available, we wanted to test their effect on the larval development of two species of *Aedes* known for their vectorial importance. The possibility of a mosquito control material / substance at very low cost, with little / no environmental impact, safe for humans and easily available, would represent a strong incentive for the implementation of control measures on private properties.

### 2. Material and methods

The study was performed using larvae of *Ae. albopictus* (colony originated from field eggs collection in Padua during years from 2011 to 2017) and *Ae. aegypti* (colony originated in 2015 by eggs from Biogents AG, Regensburg Germany) bred in a laboratory. These species breed in small containers in or near dwellings and are therefore commonly faced by inhabitants. Unlike other species that breed in sites like marshes, channels or lakes, the larval habitats of *Ae. albopictus* and *Ae. aegypti* can be treated by private citizens on their ground.

The insects used for the test were bred in the laboratory, leaving them to lay their eggs on filter paper, which were then placed to hatch in dechlorinated water in the same room where the adults were reared. The larvae were fed with Altromin International® (Lage, Germany) rat chow until the pupal stage is reached, then were transferred into small containers and put into cages to complete development. *Ae. albopictus* was reared in 50 cm side cubic cages at 25±1 °C and 50±5% of RH while *Ae. aegypti* was reared in identical cages but at 27±1 °C and 80±5% of RH. The photoperiod lasted 12 hours at a solar spectrum artificial light of 6,000 K and 300 lux intensity. The adults were fed with a 10% sugar solution and twice a week bovine blood was supplied. The adults were fed 0.04 g of rat food per day.

To perform the test capsules of the brand Lavazza Bourbon Gustoso, a blend of a mix of Robusta coffee (*Coffea canephora*) and Arabica coffee (*Coffea arabica*) from Africa and South-East Asia, were used. They were extracted with a domestic coffee machine (‘A Modo Mio’ – Lavazza, Turin, Italy), operating at 65-70 °C. Because across Europe coffee is made ‘strong’ or ‘light’, two different extractions were applied, making 30 or 70 ml of coffee beverage with one capsule. Because with a longer extraction, less caffeine remains in the coffee grounds, the dosage ‘70 ml’ has to be considered as lower than the dosage ‘30 ml’. As soon as the capsules were used, they were stored in a fridge at 3-4 °C until the test was performed, no more than 7 days later.

Groups of 20 larvae (L1), taken 24 hours after hatching, were placed in food grade polypropylene (PP) containers (Ø 86 mm, h 120 mm), with 430 ml of tap water that had previously been stored in the open for 24 hours to permit chlorination to evaporate. As soon pupae appeared, a mosquito mesh was used to cover the container to avoid emerging adults flying away. This quantity of water was chosen because it is the capacity of a medium sized flower pot tray, the most typical domestic breeding site. The larvae were fed 0.04 g of rat food per day.

Treated replicates (n=3) and control replicates (n=3) were set up for each dosage and each target species. For each replicate 14.0 g of wet used coffee grounds (UCG) (corresponding to 31.5 mg/ml) were added immediately before the introduction of the larvae. Coffee capsules contain around 7 grams of dry coffee but they almost double in weight because of the water adsorbed during extraction. The 14 grams of coffee grounds used per replication, therefore correspond to the quantity of coffee contained in a used capsule or pod. The coffee powder was taken from the capsules and inserted into filter bags (like the ones used for tea) before placing in the water containers, to facilitate observation of the larvae. An empty tea filter was inserted in each control replicate.

The test was performed in a room at 27.0±1 °C and 80±5% of R.H., the temperature of the water in the containers was 25.5±1 °C. The containers were checked every 24 hours. At each assessment the emerged adults were counted and then removed by means of an entomological aspirator.

The test was concluded when live larvae or pupae were no longer present. Mortality was calculated using Abbott’s formula (Abbott, 1987). Because the different dosages were tested at different times during 2020, for each species and dosage, specific control replicates were run contemporarily.

### 3. Results

The results are expressed as percentage of larval mortality or emerged adults (complementary value). The results are shown in Table 1.

| Species          | Dosage  | Mean larval mortality (%) | Emerging adults (%) |
|------------------|---------|---------------------------|---------------------|
| *Ae. albopictus* | 30 ml (control) | 5 | 95 |
|                  | 30 ml (treated) | 3.3 | 96.7 |
|                  | 70 ml (control) | 1.7 | 98.3 |
|                  | 70 ml (treated) | 6.7 | 93.3 |
| *Ae. aegypti*    | 30 ml (control) | 1.7 | 98.3 |
|                  | 30 ml (treated) | 1.7 | 98.3 |
|                  | 70 ml (control) | 1.7 | 98.3 |
|                  | 70 ml (treated) | 2.5 | 97.5 |

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The two-sample Mann-Whitney U test was used to compare values of emerged adults from the treated and control replicates (Table 2). This test is equivalent to a two-sample Wilcoxon rank-sum test. The test assumes that the observations are independent. There was no difference between adults emerged from control and treated samples, independently of the species or concentration.

4. Discussion and conclusion

The repellence of coffee versus Ae. albopictus females from the breeding sites and the inhibition of the embryonic development was shown by Satho et al. (2015). The capability of coffee extracts to reduce embryonic maturation as well as the shorter adult life span in dengue vectors were also proved (Dieng et al., 2016). Regarding the efficacy of the used coffee grounds as larvicide, the literature shows very different results. Guirado and Bicudo (2007) recorded a complete mortality of Ae. aegypti larvae but using a very high quantity of dry UCG (300 mg/ml) while Laranja et al. (2003) completely killed the larvae of the same species using only 50 mg/ml.

The insecticidal action of coffee extracts or UCG is mainly driven by the caffeine, the dosage of 1.0 mg/ml of caffeine is sufficient to kill 100% of Aedes larvae (Laranja et al., 2003). The larvicidal efficacy of the exhausted coffee grounds is therefore strictly connected to their content of this alkaloid. Because caffeine is water-soluble the more water was used to produce the coffee drink, the less caffeine is present in the exhausted coffee grounds and therefore the preparation method of the beverage is very important. Brewed coffee, cold brew, instant coffee and espresso content have very different quantities of caffeine and therefore leave UCG with very different larvicidal efficacy. The plant species from where the coffee comes from is also very important. In general C. canephora (also known as ‘Robusta’) contains twice as much caffeine as C. arabica but this amount also depends on the different geographical origin (Jeszka-Skowron et al., 2016). The very different results found in the literature are therefore dependent on the many variables that affect the caffeine content.

As Laranja et al. (2003) and Derraik and Stanley (2005) showed, the use of UCG in nutrient-poor water can even create a more suitable environment for larval development thanks the availability of fatty acids, amino acids and other nutrients, as well as an increased growth of algae and bacteria.

The difficulty of standardisation and the very different results in terms of larval control suggest that the use of UCG shouldn’t be advised because of the uncertainty of its efficacy. The use of partially effective methods can be very damaging when proved effective tools are available, while in specific conditions, when no better solutions are available, they could be helpful.

Conflict of interest

The authors declare no conflict of interest.

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Table 2. Mann-Whitney U test values for emerged adults of Aedes albopictus and Aedes aegypti with the two coffee ground dosages.

| Species     | Dosage      | Mean ± sd   | P-value |
|-------------|-------------|-------------|---------|
| Ae. albopictus | 30 ml (control) | 2.46±4.39  | 0.653   |
|              | 30 ml (treated) | 2.42±5.67  |         |
|              | 70 ml (control) | 1.90±2.11  | 0.1633  |
|              | 70 ml (treated) | 1.87±3.70  |         |
| Ae. aegypti  | 30 ml (control) | 2.46±4.31  | 0.9898  |
|              | 30 ml (treated) | 2.46±4.89  |         |
|              | 70 ml (control) | 2.19±4.52  | 0.5792  |
|              | 70 ml (treated) | 2.15±3.16  |         |
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