Short Term Selection to Diflubenzuron and Bacillus Thuringiensis Subsp. Israelensis Differentially Affects the Winter Survival of Culex Pipiens F. Pipiens and Culex Pipiens F. Molestus (Diptera: Culicidae)

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Research

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Abstract

Background

The Culex pipiens mosquito consists of two forms named pipiens and molestus that exhibit substantial differences in their biology including overwintering behavior. Diflubenzuron (DFB) and Bacillus thuringiensis subsp. israelensis (Bti) are among the most widely used larvicides for controlling Cx. pipiens populations. The high dependency on these two larvicides, pose major concerns for resistance development. The evolution and stability of tolerance/resistance to insecticides has been associated with fitness costs that may be manifested under stressful conditions such as the winter period. This study aimed to investigate how short term selection of pipiens and molestus forms to both larvicides affect their susceptibility and their overwintering success.

Methods

Colonies of both forms of Cx. pipiens were established from the same area. Following standard protocols, the efficacy of both larvicides was determined for each mosquito population. Then, larvae from each form, were selected for three successive generations by applying fixed doses corresponding to IE$_{80}$ (Inhibition of adult Emergence) and LC$_{80}$ (Lethal Concentration) for DFB and Bti, respectively. At the end of this process, the susceptibility levels and the winter survival of the selected populations relative to controls (colonies that received no selection) were determined.

Results

Contrary to Bti, selection with DFB induced different susceptibility levels between the two forms of Cx. pipiens. The selected populations of Cx. pipiens f. molestus to both larvicides exhibited a high fitness cost in terms of reduced winter larval survival rates relative to control. Moreover, the obtained adults of the Bti selected population experienced significantly shorter lifespan compared to control and DFB selected population. On the other hand, selection with both DFB and Bti had no apparent effects on Cx. pipiens f. pipiens female winter survival rates relative to control. Furthermore, the reproductive parameters and the longevity of the overwintered females were similar between the selected populations and the control.

Conclusions

Our findings reveal that sort term selection to DFB and Bti induces a high fitness cost on the winter survival of Cx. pipiens f. molestus but not of pipiens form, suggesting potential differences on the persistence of tolerant individuals from year to year between the two forms.

Background

The common house mosquito, Culex pipiens (L.) is a widespread insect pest of extremely high medical and veterinary importance as it is considered effective vector of several human and animal diseases,
including filarial nematodes and arbovirus such as West Nile virus (WNV), Sindbis virus, Rift Valley fever and Japanese encephalitis virus [1, 2]. *Culex pipiens* includes two distinct forms (usually referred as biotypes), *pipiens* and *molestus*, which are morphologically identical but differ in several behavioral and physiological aspects [3, 4]. In particular, the *molestus* form prefers to colonize underground breeding sites, while *pipiens* is commonly found in above ground habitats. Moreover, *Cx. pipiens f. molestus* is stenogamous (copulation can occur in confined spaces), autogenous (ability to develop a first batch of eggs without a blood meal), and mammophilic (prefers to feed on mammals, including humans). On the other hand, *Cx. pipiens f. pipiens* is eurygamous (copulation occur outdoors in swarms), anautogenous (blood feeding is necessary for eggs development), and rather ornithophilic (prefers to feed on birds) [3]. Another major difference between the two forms is lying on their winter biology at temperate regions. Contrary to the *molestus* form which remain active and reproduce during winter, the *pipiens* form undergoes diapause as inseminated females with arrested ovariole development and elevated fat body reserves that serve as energy source [3, 5, 6]. Short day length and relative low temperatures perceived in larval and pupal stage during autumn are responsible for triggering the physiological changes underling diapause induction [7, 8]. The two forms often co-occur and can hybridize, while the hybridization rates may reach up to 31.8% as has recently demonstrated [9]. Hybrids are considered to play a key role on WNV transmission, since they may exhibit a more opportunistic biting behavior and therefore act as effective bridge vectors between the avian/WNV reservoirs and humans [10].

Although nowadays several non-chemical methods for mosquito control are under development and evaluation such as the Sterile Insect Technique (SIT), the Release of Insects carrying a Dominant Lethal (RIDL) and the release of *Wolbachia*-infected mosquitoes [11], insecticide applications still remain the principal tool for tackling mosquitoes related problems. This is mostly because several medically important mosquito species, including *Cx. pipiens f. pipiens*, tend to colonize fixed, open breeding sites that can concentrate very large populations. In such situations, the use of insecticides can be very effective since both mapping and treating is feasible. However, this is not the case for the artificial container-breeding mosquitoes, such as *Aedes aegypti* and *Ae. albopictus* which distribute individual eggs in several, ephemeral oviposition sites [12]. Among the insecticides that are employed, larvicides are considered as the most important means for the prevention of mosquito-borne diseases, as they target immature stages (larvae and pupae) and thus prevent females' emergence which are responsible for the pathogens transmission. Despite the high importance of this approach, under the current European Union biocide legislation and the prohibition of organophosphates (OPs) such as temephos, larval control relies almost exclusively on two main categories of biocides, the Insect Growth Regulators (IGRs) and the microbial ones [13]. Diflubenzuron (DFB) and *Bacillus thuringiensis* subsp. *israelensis* (*Bti*) are the most widely used larvicides in each category, as they combine some very desirable features such as the high efficacy against mosquito larvae and the very low toxicity to vertebrates. DFB is a member of the Benzoylurea insecticide family that inhibits the chitin biosynthesis process causing abnormal molting during the immature development preventing adult emergence [14], while contrary to other IGRs like methoprene it has a remarkable ability to control all mosquito larval instars [15]. Following its introduction in the early 1970s, DFB has been extensively used in agricultural, forestry and more recently
in medical insect pest larvae [14]. On the other hand, over the last years the microbial larvicide *Bti* has increasingly used as an alternative to chemical insecticides as it displays high selective action against mosquito larvae and minimal impacts on the environment and non-target organisms including human [16]. During sporulation *Bti* produces a parasporal crystalline body composed of four major (Cry11Aa, Cry 4Ba, Cry 4Aa and Cyt1Aa) and at least two minor (Cry10Aa and Cyt2Ba) protoxins. Upon ingestion, the crystals are solubilized under the alkaline conditions in the larval midgut and the protoxins are proteolytically converted into active toxins [17]. The Cry toxins bind to specific midgut membrane receptors, while the Cyt toxins appear to act synergistically with the Cry toxins functioning as surrogate receptors that improve their capacity to bind on the available target sites (receptors) [18, 19]. After the toxins binding on the midgut receptors, they insert in the plasma membrane forming lytic pores that disturb the cell's osmotic balance, resulting in cell lysis and consequently death of the larvae [20]. The synergistic interactions between the Cyt and the Cry toxins have attracted particular attention due to the drastic enhancement of their larvicidal activity against mosquitoes [21]. Moreover, the synergism between the Cyt1Aa and the Cry toxins is considered the key factor for the low potential of resistance development following extensive selection with *Bti* [17, 22].

The high dependency on both DFB and *Bti* for the suppression of mosquito populations, including *Cx. pipiens*, pose major concerns for resistance development which may jeopardize the control efforts and increase the risk of diseases transmission such as the WNV. Indeed, striking DFB resistance levels associated with specific mutations have been already detected in *Cx. pipiens* natural populations from Italy and Turkey in areas with intense use of this insecticide [23–26]. On the other hand, there is only a single record of high (33-fold) *Bti* resistance levels in *Cx. pipiens* wild populations with a background of previous exposure to support for potential resistance development [27]. Interestingly, evaluation of field populations of *Cx. pipiens* without history of *Bti* exposure has shown variations of resistance ratios ranging from less than 3- to 10-fold [28]. Therefore, this inherent variability in *Cx. pipiens* populations’ susceptibility to *Bti* may be of high importance since it is possible to affect their response to selection pressure. Despite the prominence of the above findings regarding both DFB and *Bti* resistance in *Cx. pipiens* populations, no efforts have been made to separate potential differences between the form *pipiens* and *molestus* given their divergent biology.

Insecticide resistance emergence and evolution in the wild is a dynamic process that depends among others, on the genetic background and the biology of mosquito species, the intensity of biocides selection pressure and the resistance mechanisms that are involved [29]. However, resistance development is often associated with significant reductions in fitness parameters of the resistant individuals as a result of increased metabolic costs of physiological/biochemical resistance mechanisms. The reason underlying this phenomenon is because there is a trade-off of energetic resources from primary biological functions associated with fitness to secondary ones that contribute to the function of resistance mechanisms [30]. In *Cx. pipiens* resistance development against organophosphates (OPs) has been associated with male mating competition cost [31], predation avoidance cost [32] and decreases in preimaginal survival [33]. Moreover, adverse abiotic conditions can also affect the impact of insecticide resistance on key life history traits. Winter in temperate areas represents a very challenging and stressful period for the survival
of many insects including mosquitoes and therefore potential fitness costs associated with insecticide resistance may reduce their overwintering success. Indeed, the frequency of two genetic loci in *Cx. pipiens* f. *pipiens* females associated with OPs resistance decreased over winter, indicating reduced survival for resistant individuals [34]. This is of great importance since it can determine the persistence of resistant mosquitoes from year to year affecting both the evolution and stability of acquired resistance in the wild. However, it remains completely unknown if a similar phenomenon holds in the case of DFB and *Bti* given their different mode of action relative to OPs.

Considering the importance of *Cx. pipiens* as an effective vector of several diseases, the fundamental differences in biology between its two forms and the high dependency on DFB and *Bti* to tackle its populations, the aim of the present study was to explore how short term exposure of *pipiens* and *molestus* forms to both larvicides affect their susceptibility and most importantly their overwintering success. For this purpose, we established colonies of both *Cx. pipiens* f. *pipiens* and *Cx. pipiens* f. *molestus* originated from the same area and subjected them to the same selective pressure (80% population mortality) for three successive generations with DFB and *Bti*. The susceptibility levels and the overwintering survival of the selected populations relative to controls (colonies that received no selection) were determined.

**Methods**

**Mosquito colonies and rearing methods**

All mosquito colonies were established during early September to late October of 2017 from eggs that were collected at the vicinity of Volos and Larissa city, Thessaly province, Greece. Earlier surveys in the study area revealed *Cx. pipiens* as the most abundant mosquito species, followed by *Aedes caspius* [35]. Since 2010 WNV outbreak in Greece, there is a regular presence of human cases in the area, with a peak of 24 incidents in Larissa in 2019 [36]. Mosquito control is performed routinely since 2010 by private enterprises and involves mainly the use of DFB in urban and suburban breeding sites, while *Bti* is applied in protected wetlands such as rivers, streams and lakes [25, 37]. In total 74 and 59 egg rafts were recovered for *Cx. pipiens* form *pipiens* and *molestus* respectively. Separation of *Cx. pipiens* forms was confirmed by matrix-assisted laser desorption/ionization time-of-flight mass spectrometry (MALDI-TOF MS) protein profiling [38]. Colonization took place within the insectary facilities of the laboratory of Entomology and Agricultural Zoology at the University of Thessaly. The insectary walk-in chamber was maintained at 25 ± 1 °C, 65 ± 5% relative humidity and a photoperiod of L14 : D10 with a simulated dusk and dawn for 45 min. Photophase initiation was set at 00:00 hr and termination at 14:00 hr. Larvae were reared in 42 × 30 × 10 cm white plastic containers in 3 l of bottled table water (Table Water, Epirotic Bottling Industry S.A. Ioannina, Greece) fed a total amount of 2 g of ground cat food (Friskies Adult, Purina, Italy) and held at a density of approximately 1000 individuals per container. Adults were kept in 32 × 32 × 32 cm screened cages at a density of 400–500 individuals and fed with 10% sugar solution that was renewed every week. Females of *Cx. pipiens* f. *pipiens* were fed on certified human blood derived from samples that were provided by a blood analysis laboratory. Blood temperature was set at 38 °C.
using two custom made, cylindrical (7.5 cm in diameter and 10 cm in height) feeding apparatus operated with circulated water from a warm bath via 12V DC mini water pumps. The apparatus were placed on the top of the holding cages for 1 h, and females had accesses to feed via a stretched Parafilm M (Bemis, USA) membrane. Depending on the experimental needs, colonies received two to three blood meals per month. In general, colonies of *Cx. pipiens f. molestus* were kept without access to blood. Only in a few cases they were provided with a blood meal after the deposition of their first autogenus egg raft in order to reinforce colonies population during the selection process (see below). Both mosquito populations were reared for 3 generations in order to establish a uniform genetic background before the initiation of the experiments.

**Larval bioassays**

Standard World Health Organization (WHO) guidelines [39] were adopted to evaluate the susceptibility of each collected population against DFB and *Bti*. Larval susceptibility was evaluated against analytical standard DFB (Purity ≥ 99.8%, Pestanal®, Sigma-Aldrich, Germany) and formulated *Bti* (Vectobac® 12AS, 11.61% w/w *Bti* serotype H-14, strain AM65-52, 1200 ITU/mg, Valent BioSciences Corporation, USA). Stock solutions were prepared in 99.5% acetone for DFB and distilled water for *Bti* and stored at -22 °C until use for up to two weeks. For DFB, six concentrations ranging from 0.001 to 0.01 mg/l and for *Bti* five doses ranging from 0.02 to 0.04 mg/l were used yielding larval control between 10 and 95%. Six replicates were performed per concentration and an equal number of controls, each involving twenty five 3rd and early 4th instar larvae for DFB and *Bti*, respectively. Bioassays were run three times on different days using new batches of larvae and larvicide solutions. Larval mortality for *Bti* and adult emergence inhibition for DFB were recorded according to WHO recommended exposure times [39]. For each mosquito population, dose mortality responses were used to calculate IE$_{50}$, IE$_{80}$ and IE$_{90}$ values (IE: Adult emergence inhibition) for DFB and LC$_{50}$, LC$_{80}$ and LC$_{90}$ values (LC: Lethal concentration) for *Bti* using Probit Analysis [40].

**Larval selection**

Larvae from each population were exposed for three successive generations to fixed concentrations of DFB and *Bti* corresponding to IE$_{80}$ and LC$_{80}$, respectively. During the selection process, six to eight groups of ≈ 1.000 larvae were placed into rearing containers (see above) provided with 3 l of table water and the fixed dose of each larvicide. Selection against DFB involved 3rd instar larvae, while 1.2 g of cat food was added in the containers to allow development (pupation). Selection against *Bti* involved the exposure of early 4th instar larvae to fixed doses for 24 h without any accesses to food. Surviving larvae were placed at a maximum density of ≈ 1.000 individuals into rearing containers with 3L of clean table water and offered 1 g of food to complete development. Resulting pupae from both DFB & *Bti* selection process, were transferred daily into cages and reared following the standard procedures described above. Additionally, for each population, two larvae groups (≈ 1.000 individuals each) were maintained under identical conditions but in the absence of DFB & *Bti* exposure serving as controls. There was at no point exchange of individuals among treatments and controls during the above mentioned procedures. After the completion of the final (third) selection process, new dose-response bioassays were performed for
each population to establish the new susceptibility levels against both DFB & Bti. A range of five concentrations was tested for each larvicide (DFB: 0.002–0.015 mg/l; Bti: 0.03–0.08 mg/l) using the same procedures described previously. Moreover, the winter survival of the descendants of both selected and control populations was evaluated.

**Winter survival**

Since *Cx. pipiens* f. *molestus* remains active and reproduce during the winter period, both immature and adult survival were assessed. For this purpose, 1,000 first instar larvae 3–5 hours after their eclosion from both selected populations and the control were equally apportioned into five white, plastic, containers (30 × 20 × 10 cm) (200 larvae/container) with 1.5 l of table water and 2 mg of cat food per larvae. Then, all containers were randomly placed side by side in a humid, unheated warehouse located at the outdoor facilities of the laboratory that simulated the winter breeding sites of the species. Natural daylight was provided through a 70 × 70 cm window in the east side of the warehouse. To prevent water evaporation, each container was shield with a well fitted lid bearing a 2.5 cm hole at the centre covered with mesh. Larval exposure took place on December 23, 2018. Containers were inspected every five days until the appearance of the first 4th instar larvae and since then daily. The initiation of larvae pupation took place on February 5 and completed on March 3. Resulting pupae from each population were transferred into transparent, plastic bowls filled with 200 ml of table water kept at the same conditions. A maximum number of 15 pupae was placed in each bowl, while a lid prevented emerging adults from escaping. First adult emergence was observed on February 17 and the last on March 5. Upon adult appearance, pairs consisting of a male and a female from each population were placed into individual cages. Each cage comprised a 0.4 l capacity transparent plastic cup (height 12.5 cm high, upper diameter 6.5 cm, base diameter 9.2 cm) fitted into a 9.2 cm in diameter plastic Petri dish lid. On the side of each cup, an opening of 25 cm² covered with nylon mesh was formed for ventilation. Each individual cage was placed upon a 9 cm in diameter Petri dish provided with 5% sugar solution while adults had access to feed via a small piece of wick made of sponge cloth (Wettest® classic, Freudenberg, Sweden). Depending on the adult emergence rates, 30, 47 and 50 replications (pairs) were established for DFB and Bti selected and the control population respectively. Adults were kept at the same place where larval development took place (warehouse). Their survival was monitored daily from February 18 until the death of the last individual on June 27, while the sugar solution in the individual cages was renewed every week. Prevailing temperature and RH conditions inside the warehouse during the experiments were recorded by an indoor data logger (HOBO UX100-011, ONSET, USA) set to receive 4 recordings per 24 h. For each selected population and the control, both larval and pupal developmental duration and survival were assessed as well as adult longevity.

Diapausing females of *Cx. pipiens* f. *pipiens* from both selected populations and the control were reared from 1st instar larvae, using the standard procedures described above, in an environmental chamber set at 20 °C, 8L:16D, and 70% RH [6]. Adults were provided constant access to 10% sugar solution 10–14 days post eclosion to allow female copulation and lipid accumulation reserves. To confirm diapause induction, 10 randomly selected females from each population were dissected and the primary
foll./germa. length ratio was used as criterion of ovarian diapause [41]. After the lipid accumulation period, two hundred females from each population were equally apportioned into five 20 × 20 × 20 cm screened cages (40 females/cage) having access only to water and transferred to the same warehouse described before to simulate winter conditions. Female exposure took place on January 15, 2019. Cages were inspected at two week intervals and female survival for each population was recorded. Dead mosquitos were removed with an aspirator through a 2 cm hole in the cages door shielded with a cork. Female winter survival was terminated at the end of March (26/3), following temperature rise, by providing the remaining individuals with 10% sugar solution without been removed from their winter shelter. Two weeks later, on April 8, survived females from each population (25–32 individuals, see results) were merged in a 20 × 20 × 20 cm screened cage and were given the opportunity to receive a blood meal for an hour over three successive nights. All three cages (one per population) were transferred inside the walk-in chamber between 20:00 and 22:00 and females were allowed to feed through the apparatus described above. After each blood feeding trial, cages were transferred back to the warehouse. One week following the last blood meal, a white, plastic cylindrical (10 cm in diameter and 5 cm in height) bowl with 200 mL of table water and 0.05 g of cat food was placed in each cage to allow oviposition. Bowls remained inside cages for 7 days. During that period, cages were daily inspected and deposited egg rafts were collected and pictured under a binocular stereoscope (ZEISS, SteREO, Discovery.V12) equipped with a digital camera (ZEISS, AxioCam, ERc 5 s) to facilitate eggs counting. Then, each egg raft was transferred individually into a white bowl (same as those that described just above) covered with mesh to assess larval hatch rates under the warehouse conditions. The females of each population remained in the warehouse under ambient conditions and their survival was monitored every two days until the death of the last individual on July 28, while the sugar solution in each cage was renewed every week. Climatic data (temperature and RH) during the course of the above experiments were recorded as described previously. For post-overwintering females of each population we estimated a) the percentage of individuals that received a blood meal b) preoviposition period c) mean number of eggs per raft d) mean larval hatch rate per egg raft and e) lifespan under the ambient conditions inside the warehouse.

Data analysis

The effect of mosquito population on larval and female (in each sampling date) winter survival rates of *Cx. pipiens* f. *molestus* and *Cx. pipiens* f. *pipiens* respectively were assessed using One-way analysis of variance, after appropriate transformations for normality and homoscedasticity when necessary, followed by Tukey’s HSD post hoc to separate means. The same analysis was also performed to assess the effect of mosquito population on the number of eggs per raft and the larval hatch rates per egg raft of the survived *Cx. pipiens* f. *pipiens* females. The proportions of pupae survived in each population of *Cx. pipiens* f. *molestus*, their sex ratio as well as the proportions of female of *Cx. pipiens* f. *pipiens* that received a blood meal were analyzed using the Chi-square test. The effects of mosquito population and sex on larval and pupal developmental duration as well as on obtained adult lifespan of *Cx. pipiens* f. *molestus* were assessed using the Cox proportional hazards model. This model is commonly applied to assess the effects of one or more predictors on time to event incidents such as time to pupation, adult emergence/death etc. Pairwise comparisons were conducted using the log rank (Mantel–Cox) test. The
same analysis was also performed to assess the effect of mosquito population on both the preoviposition period and the lifespan of the survived *Cx. pipiens* f. *pipiens* females. Data analysis was performed using IBM SPSS 25 (IBM Corp., Armonk, NY).

**Results**

Dose-response larval bioassays results are given in Tables 1 & 2 for DFB and *Bti* respectively. The EI$_{50}$ values for DFB were similar between the two forms of *Cx. pipiens* before the initiation of the selection trials, however the EI$_{90}$ value of *Cx. pipiens* f. *pipiens* was twice as much as that of *molestus* suggesting a lower inherent susceptibility. The selection process for three successive generations by applying fixed DFB doses corresponded to the EI$_{80}$ for each population resulted in 3.7 and 3.1 resistance ratio values for EI$_{50}$ and EI$_{90}$ respectively for *Cx. pipiens* f. *pipiens*, and 1.7 and 2.9 for the *Cx. pipiens* f. *molestus* (Table 1). The LC$_{50}$ and LC$_{90}$ values for *Bti* were almost identical between the two populations before selection. The selection processes against *Bti* had minor effects on the susceptibility levels of both *Cx. pipiens* forms (Table 2).

Table 1 Effective doses of diflubenzuron against the two forms of *Culex pipiens* after the selection process for three successive generations by applying fixed doses corresponding to EI$_{80}$ of the control populations.
| Population          | N*  | El$_{50}$     | RR$_{50}$ | El$_{90}$     | RR$_{90}$ | Slope | $\chi^2$ (df) |
|---------------------|-----|---------------|-----------|---------------|-----------|-------|---------------|
|                     |     | (95% CL)$^a$ | (95% CL)$^a$ |               |           |       |               |
| **Cx. pipiens f. pipiens** |     |               |           |               |           |       |               |
| Control             | 3150| 0.0025        | -         | 0.0081        | -         | 2.47  | 177.57$^b$ (105) |
| Selected            | 2700| 0.0093        | 3.7       | 0.0252        | 3.1       | 2.96  | 70.98 (87)    |
|                     |     | (0.0076–0.0106)|          | (0.0218–0.0321)|          |       |               |
| **Cx. pipiens f. molestus** |     |               |           |               |           |       |               |
| Control             | 3150| 0.0022        | -         | 0.0040        | -         | 4.49  | 201.99$^b$ (105) |
| Selected            | 2700| 0.0037        | 1.7       | 0.0116        | 2.9       | 2.60  | 57.94 (87)    |
|                     |     | (0.0028–0.0046)|          | (0.0101–0.0132)|          |       |               |

$^a$ Number of larvae tested

$^a$ El values are expressed in milligrams per liter, and they are considered significantly different when 95% of confidence limits (CL) fail to overlap

$^b$ Since goodness-of-fit test is significant ($P<0.05$), a heterogeneity factor was used in the calculation of confidence limits (CL)

Table 2. Effective doses of *Bti* against the two forms of *Culex pipiens* after the selection process for three successive generations by applying fixed doses corresponding to LC$_{80}$ of the control populations.
| Population       | N*  | LC<sub>50</sub> (95% CL)<sup>a</sup> | RR<sub>50</sub> | LC<sub>90</sub> (95% CL)<sup>a</sup> | RR<sub>90</sub> | Slope | χ² (df) |
|------------------|-----|-------------------------------------|----------------|-------------------------------------|----------------|-------|--------|
| *Cx. pipiens f.* |     |                                     |                |                                     |                |       |        |
| Control          | 2700| 0.031 (0.029–0.033)                 | -              | 0.047 (0.044–0.051)                 | -              | 7.52  | 120.73<sup>b</sup> (87) |
| Selected         | 2700| 0.042 (0.035–0.048)                 | 1.3            | 0.069 (0.065–0.074)                 | 1.5            | 6.12  | 47.99  |

| *Cx. pipiens f.* |     |                                     |                |                                     |                |       |        |
| molestus         |     |                                     |                |                                     |                |       |        |
| Control          | 2700| 0.032 (0.029–0.034)                 | -              | 0.047 (0.044–0.051)                 | -              | 7.62  | 126.68<sup>b</sup> (87) |
| Selected         | 2700| 0.045 (0.034–0.059)                 | 1.4            | 0.081 (0.065–0.089)                 | 1.7            | 4.98  | 117.21<sup>b</sup> (87) |

* Number of larvae tested

- LC values are expressed in milligrams per liter, and they are considered significantly different when 95% of confidence limits (CL) fail to overlap

- Since goodness-of-fit test is significant (P < 0.05), a heterogeneity factor was used in the calculation of confidence limits (CL)

Ambient temperature and relative humidity conditions inside the warehouse from the beginning of the exposure of 1st instar larvae of *Cx. pipiens f. molestus* until the death of the last adult are depicted in Fig. 1. Temperatures ranged between 3.9–13.6 °C with a mean value of 10.2 °C from larvae exposure (on December 23) until the formation of the last pupae (on March 3). Corresponding values regarding the first adult emergence until the death of the last individual ranged between 7.7–26.8 °C with a mean value of 17.4 °C. Population differentially affected larva-to-pupa winter survival rates of *Cx. pipiens f. molestus* (F<sub>2,12</sub> = 25.26, P < 0.001). Selection to both DFB & *Bti* significantly reduced larva-to-pupa survival relative to control, while significant differences were also found between the two selected populations (Fig. 2). Cox regression analysis revealed mosquito population as a significant predictor (Wald test, χ²<sub>2</sub> = 30.26, P < 0.001) of the larval developmental duration considering the total number (viable and dead) of pupae formed. Larval developmental duration was significantly longer in both DFB & *Bti* selected populations relative to control, while significant differences were also observed between the two selected populations (Fig. 3). Cox regression analysis revealed mosquito population (Wald test, χ²<sub>2</sub> = 7.87, P = 0.020) and sex (Wald test, χ²<sub>1</sub> = 11.49, P = 0.001) as significant predictors of the larval developmental duration considering the viable pupae yielded. Larval developmental duration of control male larvae was significantly shorter than the *Bti* selected population but not than the DFB (Fig. 4). On the other hand, no
significant differences were observed between the control and the two selected populations as far as the female larvae developmental duration is regarded. Pupal survival rates were 50.7%, 45.0% and 44.4% for control, DBF and $Bti$ selected population respectively and did not differ significantly among the three populations (Chi-square test, $X^2_2 = 2.10$, $P = 0.349$). Neither mosquito population (Wald test, $X^2 = 4.00$, $P = 0.135$) nor sex (Wald test, $X^2_1 = 1.20$, $P = 0.272$) were significant predictors of the pupae developmental duration of $Cx. pipiens f. molestus$ (Fig. 5). Sex ratio rates (females/males) were 52.3%, 58.3% and 44.3% for control, DBF and $Bti$ selected population respectively, though no significant differences were observed (Chi-square test, $X^2_2 = 3.54$, $P = 0.170$). Cox regression analysis revealed mosquito population (Wald test, $X^2_2 = 13.29$, $P = 0.001$) and sex (Wald test, $X^2_1 = 4.52$, $P = 0.033$) as significant predictors of adult lifespan. Both males and females of the $Bti$ selected population exhibited significantly shorter longevity relative to control. Females of the DFB selected population also outlived those of $Bti$ (Fig. 6, Table 3).

Climatic conditions during the experiments of $Cx. pipiens f. pipiens$ are shown in Fig. 1. Temperatures ranged between 7.7–16.5 °C with a mean value of 11.9 °C from the beginning of female exposure until the termination of the overwintering period on March 26. Corresponding values regarding the post-overwintering period until the death of the last female ranged between 13.5–27.1 °C with a mean value of 21.2 °C. Analyses in each sampling date revealed no significant differences on female survival rates among the three populations ($F_{2, 12} = 0.48$ to 1.05, $P = 0.379$ to 0.631) (Fig. 7). Additionally, no significant differences were observed in female reproductive parameters (Table 4). Finally, Cox regression analysis showed that mosquito population was not a significant predictor (Wald test, $X^2_2 = 3.35$, $P = 0.187$) of female post-overwintering survival. Average survival was approximately three months for all three populations demonstrating that overwintered females of $Cx. pipiens f. pipiens$ may experience extensive remaining lifespans (Fig. 8, Table 5).

**Table 3** Longevity parameters of *Culex pipiens f. molestus* populations that either were selected against diflubenzuron (DFB) and $Bti$ for three successive generations or not (control). Adults were maintained at the same place (warehouse) where they developed as larvae during winter having access to 5% sugar solution. Within sex, different letters indicate significant differences (pairwise comparisons log-rank test, $P < 0.05$).
Table 4 Reproductive parameters of post-overwintering *Culex pipiens* f. *pipiens* females. Females originated from populations that either were selected against diflubenzuron (DFB) and *Bti* for three successive generations or not (control).

| Population       | n   | Blood meal acceptance % | Preoviposition period days (± SE) | Eggs per raft (± SE) | Larval hatch rate per egg raft (± SE) |
|------------------|-----|-------------------------|-----------------------------------|----------------------|--------------------------------------|
| Control          | 31  | 35.71                   | 10.00 ± 0.76                      | 61.10 ± 5.43         | 89.10 ± 3.42                         |
| DFB selected     | 32  | 34.38                   | 11.72 ± 0.61                      | 51.54 ± 4.50         | 92.58 ± 3.04                         |
| *Bti* selected   | 25  | 46.43                   | 11.53 ± 0.50                      | 51.46 ± 3.70         | 90.68 ± 3.81                         |
| *F*              | -   | -                       | -                                 | 1.43                 | 0.22                                 |
| *X*^2^           | -   | 1.06 ^a                 | 2.64 ^b                          | -                    | -                                   |
| df               | -   | 2                       | 2                                 | 2.31                 | 2.31                                 |
| *P*              | -   | 0.587                   | 0.266                             | 0.254                | 0.798                                |

^a^ Chi-square test

^b^ Wald test

Table 5. Post-overwintering longevity parameters of *Culex pipiens* f. *pipiens* populations that either were selected against diflubenzuron (DFB) and *Bti* for three successive generations or not (control). Females
were maintained at the same place (warehouse) were they overwintered having access to 10% sugar solution. Different letters indicate significant differences (pairwise comparisons log-rank test, \( P<0.05 \)).

| Longevity parameters in days ± SE |
|-----------------------------------|
| Average | Quartiles |
|        | 25 | 50 | 75 |
| Population |
| Control (n = 31) | 81.45 ± 6.02 a | 109 ± 6.59 | 89 ± 5.00 | 68 ± 15.02 |
| DFB selected (n = 32) | 92.40 ± 5.84 a | 115 ± 1.23 | 105 ± 2.80 | 83 ± 22.04 |
| Bti selected (n = 25) | 99.84 ± 3.19 a | 115 ± 1.62 | 97 ± 1.48 | 92 ± 2.91 |

**Discussion**

Our study revealed that the implementation of equivalent selective pressure for three successive generations with both DFB and *Bti* against the form *pipiens* and *molestus* of *Cx. pipiens* induced differential susceptibility levels only in the first case. The selected populations of *Cx. pipiens f. molestus* to both larvicides exhibited a high fitness cost in terms of reduced winter larval survival rates accompanied with increased larval developmental times. Moreover, the obtained adults of the *Bti* selected population appeared to suffer an additional cost in terms of shorter lifespan compared to the other two (control and DFB selected). On the other hand, the selection process had no apparent effect on *Cx. pipiens f. pipiens* female winter survival rates relative to control. Additionally, overwintered females showed similar reproductive parameters among populations. Interestingly, these females, irrespective of population origin, experienced considerable post-overwintering longevity periods.

The EI_{50} values estimated in the current study for the control populations of both forms were found to be almost identical with that of the Benaki *Cx. pipiens f. molestus* laboratory reference strain [23], suggesting high susceptibility to DFB. Similarly, earlier studies conducted in different regions of Greece using the WHO diagnostic dose protocols demonstrated high susceptibility of most *Cx. pipiens* populations tested [37]. Moreover, recent surveys failed to detect any specific mutations in the chitin synthase gene of *Cx. pipiens* sampled from Greece that are associated with high levels of resistance against DFB. On the contrary, two and three different mutations at amino acid I1043 of the chitin synthase gene have been reported in neighboring Turkey and Italy, respectively with phenotypes exhibiting up to 128 fold Resistance Ratios (RR) relative to the Benaki reference strain [23–26]. The different selection pressure regimes imposed on these populations over the past years relative to the Greek ones, have been proposed as a possible explanation. Interestingly, in our study by imposing the same short term selection pressure with DFB we observed differential response on the susceptibility levels between the form *pipiens* and *molestus*, especially as EI_{50} RR values are concerned. This tendency was found to be even more pronounced after additional generations of continuous selection to DFB (F6,
F9 & F12 generation) in the laboratory (Ioannou et al. in preparation). The reasons underlying this
differential response to DFB selection of the two forms of *Cx. pipiens* remain largely unknown. A possible
explanation could lay on their different biology. The tendency of the *molestus* form to reproduce below
ground may largely decrease both its exposure to insecticides as well as the gene flow rates among
different populations. On the other hand there are no such limitations for the above ground free living
*pipiens* form. Therefore, it is anticipated that the *molestus* form populations may lack the genetic
background for a rapid apparition and evolution of tolerance. This argument is also supported by the fact
that in the current study the El$_{90}$ value of the control *Cx. pipiens* f. *pipiens* population (without selection)
was twice as much as of the *molestus* form. Another possible explanation could be the fact that the DFB
selected populations of each form may encounter differential levels of fitness costs in the wild such as
winter survival rates as evidenced by the present study.

Sort term selection with *Bti* had a minor impact on the susceptibility levels of both biotypes. This is an
expected outcome since there are no consistent recordings of mosquito resistance development to the
full crystal. Despite some sporadic reports describing considerable resistance levels of wild mosquito
populations against *Bti* [27, 42–44], long-term studies under both laboratory and field conditions provide
no support to these data [45–51]. The low tolerance against *Bti* after long periods of applications in
natural settings or intensive laboratory selection have been primarily attributed to the synergistic action
between the Cry and the Cyt toxins as previously mentioned. In contrast, selection with single, purified Cry
toxins can rapidly lead to considerable resistance levels [45, 52, 53]. Another factor that seems to
promote the lack of *Bti* resistance in mosquitoes is the fact that in the absence of selection pressure
within only a few generations (3–5), any acquired tolerance disappears almost completely [46, 48]. The
high fitness costs following extensive selection to this microbial larvicide have been proposed as the
most convincing explanation of this phenomenon.

Both DFB and *Bti* selection against *Cx. pipiens* f. *molestus* conferred a high fitness cost in terms of larvae
winter survival as mortality rates relative to control increased more than 50% and 30% respectively. When
we repeated the same experimental procedure after three additional generations of selection (F6) under
optimum (laboratory) conditions we found no apparent differences in larval survival rates among tested
populations. This suggests that the observed costs are manifested only under stressful conditions.
Indeed, the mean prevailing temperature during larval development reached the lowest developmental
thresholds described for the species [54] shaping a very challenging environment for survival. In general,
the fitness costs determined in optimal conditions are not always representative of that experienced in
the wild. This is because stressful environments and/or limited resources might be more deleterious for
tolerant/resistant individuals. Similarly to our findings, the prolongation of larval developmental times in
OPs resistant *Cx. pipiens* populations was found to emerge only under stressful crowding conditions in
natural breeding sites [33]. Therefore, both biotic and abiotic factors as in our case, may shape the
expression of fitness costs related to insecticides’ selection. The observed differences on the reduction of
larval survival rates between DFB and *Bti* selected populations suggest deferential levels of fitness costs.
A possible explanation may lay on the different mode of action of the two larvicides. An interesting
finding that emerged from the current study is the fact that while short selection against \textit{Bti} has minor effects on the tolerance levels, it may confer a high fitness cost under stressful conditions.

Reduction on winter larvae survival rates following selection against DFB and \textit{Bti} in \textit{Cx. p. molestus} was also accompanied with a discrete increment of their developmental duration relative to control. Interestingly, the longest larvae developmental times were recorded in the \textit{Bti} selected population. Moreover, the adults obtained exhibited significantly shorter lifespan compared to DFB selected and control population. This is probably because a higher tolerance to \textit{Bti} has been associated with a modified microbiota of larvae, which may affect their midgut proper function and therefore the nutrients assimilation, developmental processes and ultimately adult performance as it is well documented that nutrition during the mosquito larval stage may shape important fitness elements of the emerging adults [55, 56]. Similarly to our results, \textit{Bti} selection against \textit{Ae. aegypti} also resulted in significant prolongation of larvae developmental times relative to control [48]. No effects were found regarding adult survival in the same study. However, it should be stressed that in this case, adults had access to only water and therefore no direct comparisons can be made with our results. A moderate reduction on both male and female longevity following selection with \textit{Bti} was observed in \textit{Cx. p. p.} compared to untreated control [46]. The optimal laboratory conditions in this study may account for not detecting significant differences as in our case.

Contrary to \textit{Cx. p. molestus}, selection against DFB and \textit{Bti} had no apparent effects on the winter survival of the \textit{p.} form. The differential overwintering developmental stages (larva vs adult) in these two forms may account for the observed outcome. Mosquito larvae require a minimum amount of nutrition to fully mature and pupate. Furthermore, larval developmental completion takes place within specific time limits which are endogenously defined. This dynamic process may be more prone to selection costs relative to the adult stage where full development have been already attained. In contrast to our findings, resistance of \textit{Cx. p. p.} against OPs have been associated with reduced overwintering survival [34]. Among others, the differential exhaustion of fat reserves was proposed as the basic proximate cause. Indeed, a later study confirmed that the presence of resistance alleles against OPs in this species is negatively correlated with female lipid reserves [57]. The fact that female winter survival patterns in our study were almost identical between the control and the selected populations suggest no differences on the physiological process of fat reserves accumulation and/or exploitation. But most importantly, it means that these females have equal probabilities of survival as the untreated ones, providing the base for the building of higher tolerance levels in the next year. This argument seems to be supported by the fact that the resistance ratio against DFB of \textit{Cx. p.} field populations in Italy, from 32.5 fold in 2015 reached 128.5 fold in 2016 [23], which backs the idea of the persistence of resistant mosquitoes in the wild from year to year. However, this is something that needs to be demonstrated for the specific populations, as in contrast to our case, they carry mutations of the chitin synthase gene associated with striking DFB resistance levels. Interestingly, further studies in the same country reveled a high focal distribution of DFB resistant \textit{Cx. p.} mosquitoes which was attributed to the differential selection pressure imposed by both agricultural and mosquito control applications with DFB in the tested areas [24]. Since no separation between the two forms of \textit{Cx. p.} took place in this study, the
following explanation is possible. The high focal distribution of DFB resistant mosquitoes may only reflect differences in the mosquito populations’ composition regarding the two forms, with the predominance of *pipiens* form to account for the observed outcome. For instance, analysis of population structure from different areas of Greece, a country with identical climate, revealed extreme variations between the two forms on the composition of *Cx. pipiens* populations [9]. Nevertheless, as mentioned above, the validity of this hypothesis has to be confirmed for the specific populations.

Winter survival of diapausing *Cx. pipiens* f. *pipiens* females may largely vary depending on the hibernacula conditions [3, 58]. It has been observed that females may abandon their overwintering sites and actively search for new ones, a behavior described as an adaptive response associated with increased survival. It seems that the quality of each overwintering site, as experienced by females, is depending on multiple parameters such as the prevailing temperature and humidity levels, predator density, parasite frequency and human disturbance and therefore this quality may change during winter [34]. Under our experimental design, mosquitoes were not able to select the most optimal environment for maximizing their survival as they were forced to overwinter in a given place (warehouse). Nevertheless, this does not diminish the reliability of our results since females from all populations experienced exactly the same conditions. Female survival rates observed in the current study are comparable with that found by Koenraadt et al. [58] considering similar overwintering sites such as unheated house rooms. Interestingly, they found that non-diapausing females kept under the same conditions, died within four days, suggesting that in the absence of nutritional resources such females have a very limited ability to survive but also see Rinehart et al. [59].

Reproductive parameters of overwintered females of *Cx. pipiens* f. *pipiens* were similar between the two selected populations and the control. Blood meal acceptance percentages ranged between 34.4–46.4% suggesting that females had partially terminated their reproductive diapause by the time that trials took place. Contrary to other studies that used artificial conditions to terminate diapause in order to induce females to either respond to host stimuli [60] or receive a blood meal [58] we intentionally preferred to simulate as much as possible the natural conditions in an attempt to detect any potential variations in feeding activity. Although, no significant differences were observed regarding both the preoviposition period and the average number of eggs per egg raft, the better performance of the control population in respect to these parameters may reflect an early form of fitness cost in the selected ones. The short term selection to both larvicides may have acted against more pronounced differences as indicated by other studies. For example, *Bti* selection against *Cx. pipiens* f. *pipiens* for 20 generations resulted in 44.8% decrease of female fecundity [46] while Belinato & Valle [61] by applying the same experimental protocol as we, found that DFB selection against *Ae. aegypti* for 6 generations also conferred a significant reduction on this parameter. Finally, an interesting finding that emerged from the current study is the fact that females, irrespective of population origin (selected or not), experienced extensive post-overwintering longevity periods, which appeared to even exceed the total lifespans of *Cx. pipiens* f. *molestus* females (Tables 3 & 5). This observation is in accordance with a previous study documented that the physiological changes that take place during the pre-hibernation transition of these females confer a considerable increase in their longevity potential [62].
Conclusions

By imposing the same short term selection with both DFB and Bti against Cx. pipines form papiens and molestus we found differences on the acquired tolerance in the first case which is probably associated with their divergent biology. Moreover, the selection process with both larvicides had a significant negative impact on the winter survival of the molestus form but not that of the papiens. These findings are of high importance since provide insights on the early phase of tolerance development between the two forms but mostly on its prevalence in the wild form year to year. The incorporation of such information on a properly developed model it is anticipated to contribute on the better understanding of the two Cx. papiens forms performance against these two important larvicides over time.

Abbreviations

DFB: Diubenzuron; Bti: Bacillus thuringiensis subsp. israelensis; IE: Inhibition of adult Emergence; LC: Lethal Concentration; SIT: Sterile Insect Technique; RIDL: Release of Insects carrying a Dominant Lethal; WNV: West Nile virus; IGRs: Insect growth regulators; WHO: World Health Organization; OPs: Organophosphates; 95% CL: 95% confidence limits; RR: Resistance Ratio.

Declarations

Ethics approval and consent to participate

Not applicable.

Consent for publication

Not applicable.

Availability of data and materials

The datasets used and/or analysed during the current study are available from the corresponding author on reasonable request.

Competing interests

The authors declare that they have no competing interests.

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Authors' contributions
CSI, CH and NTP conceived and designed the study. CSI and MAK performed the experiments. CSI and NTP analyzed the data. CSI, CH, MAK and NTP wrote the manuscript. All authors read and approved the final manuscript.

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