Insecticide susceptibility of *Aedes albopictus* and *Ae. aegypti* from Brazil and the Swiss-Italian border region

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**Abstract**

**Background:** *Aedes aegypti* and *Ae. albopictus* are two highly invasive mosquito species, both vectors of several viruses, including dengue, chikungunya and Zika. While *Ae. aegypti* is the primary vector in the tropics and sub-tropics, *Ae. albopictus* is increasingly under the public health watch as it has been implicated in arbovirus-transmission in more temperate regions, including continental Europe. Vector control using insecticides is the pillar of most control programmes; hence development of insecticide resistance is of great concern. As part of a Brazilian-Swiss Joint Research Programme we set out to assess whether there are any signs of existing or incipient insecticide resistance primarily against the larvicide *Bacillus thuringiensis* sv. *israelensis* (*Bti*), but also against currently applied and potentially alternative insecticides in our areas, Recife (Brazil) and the Swiss-Italian border region.

**Methods:** Following World Health Organization guidelines, dose-response curves for a range of insecticides were established for both colonized and field caught *Ae. aegypti* and *Ae. albopictus*. The larvicides included *Bti*, two of its toxins, Cry11Aa and Cry4Ba, *Lysinibacillus sphaericus*, Vectomax CG\(^*\), a formulated combination of *Bti* and *L. sphaericus*, and diflubenzuron. In addition to the larvicides, the Swiss-Italian *Ae. albopictus* populations were also tested against five adulticides (bendiocarb, dichlorodiphenyltrichloroethane, malathion, permethrin and \(\lambda\)-cyhalothrin).

**Results:** Showing a similar dose-response, all mosquito populations were fully susceptible to the larvicides tested and, in particular, to *Bti* which is currently used both in Brazil and Switzerland. In addition, there were no signs of incipient resistance against *Bti* as larvae were equally susceptible to the individual toxins, Cry11Aa and Cry4Ba. The field-caught Swiss-Italian populations were susceptible to the adulticides tested but DDT mortality rates showed signs of reduced susceptibility.

**Conclusions:** The insecticides currently used for mosquito control in Switzerland and Brazil are still effective against the target populations. The present study provides an important reference as relatively few insecticide susceptibility surveys have been carried out with *Ae. albopictus*.

**Keywords:** Vector control, Insecticide resistance, Biolarvicides, Insect growth regulator
Background

Dengue (DENV), chikungunya (CHIKV) and Zika virus (ZIKV) are mosquito-borne viruses of medical importance in most tropical regions but also emerging in more temperate regions including continental Europe. Dengue fever is the most prevalent mosquito-borne disease worldwide with an estimated 390 million cases per year [1]. In the Americas, most dengue cases have been reported from Brazil, which has been affected by several epidemics since the 1990s [2]. In 2016, around 1.5 million cases were reported from all states of the country [3]. In Brazil Aedes aegypti is the major vector of dengue and all four DENV serotypes are co-circulating in the country [4]. In 2014, the first autochthonous chikungunya cases have been detected in Brazil and in 2016 a total of 271,824 confirmed cases have been reported from several states, including Pernambuco [3]. In 2015, autochthonous cases of ZIKV were also reported from Brazil for the first time [5] and 125,319 cases were confirmed in 2016 [3]. All the above cases have been linked to *Ae. aegypti*, a highly competent vector of arboviruses.

In continental Europe, the most prominent example is the chikungunya outbreak in Ravenna, Italy in 2007 with over 200 confirmed cases and one death. The outbreak was linked to a transmission by the invasive mosquito *Ae. albopictus* and a single viraemic person that returned with chikungunya from India [6]. CHIKV is a mosquito-borne alphavirus indigenous to African countries, the Indian subcontinent and Southeast Asia where it causes endemic and epidemic fever outbreaks [7]. The outbreak in Italy demonstrates the vector capacity of the local *Ae. albopictus* population to transmit CHIKV. Following this outbreak, additional cases of autochthonous chikungunya were recorded in mainland France [8] as well as dengue cases in both Croatia and France (e.g. [9–11]). These outbreaks show that continental Europe is vulnerable to the transmission of "tropical" arboviruses, particularly in regions where *Ae. albopictus* and *Ae. aegypti* are present.

* Ae. aegypti and * Ae. albopictus* are the main vectors of DENV and CHIKV worldwide. Both mosquito species have recently shown a large geographical expansion. *Ae. albopictus*, known as the Asian tiger mosquito is among the 100 of the world’s most invasive alien species [11] and is currently present in many regions in the Americas, Africa, Australia and Europe, where its presence has been reported in several European countries, including Switzerland [12–14]. In Brazil, *Ae. albopictus* was recorded for the first time in 1986 and has since spread throughout the country [15]. The rapid worldwide expansion of *Ae. albopictus* is attributed to its eggs that resist desiccation and may undergo diapause, an adaptation to lower temperatures. Eggs are passively dispersed across the globe through container shipments of used tyres and wet plants [16, 17].

As a reaction to the dengue epidemics in Brazil, a national programme with the aim to eliminate *Ae. aegypti* (Programa para Erradicação do *Aedes aegypti*) was launched in 1996, and in 2002 it became the National Programme for Dengue Control (Programa Nacional de Controle de Dengue). The main goal of the programme is to fight dengue through integrated vector control strategies, including the use of larvicides [18]. Until 2008 the main larvicide used in the programme was the organophosphate temephos. Due to increasing resistance observed in several *Ae. aegypti* populations [19–21] temephos was replaced by the biological larvicide *Bacillus thuringiensis* svar. *israelensis* (*Bti*) in some municipalities in 2002. Then, in 2009, it was replaced by the insect growth regulators (i.e. chitin synthesis inhibitors), diflubenzuron and novaluron [19, 22] and finally, between 2014 and 2015, by pyriproxyfen, a juvenile hormone analogue [23]. Already in 2001, the health secretary of the city of Recife had decided to use *Bti* as the sole larvicide to fight *Ae. aegypti*. *Bti* equally targets *Ae. albopictus* larvae that share the same breeding sites with *Ae. aegypti* in many urban areas [24, 25] and, therefore, monitoring of *Bti* susceptibility to both *Aedes* species is needed.

In Switzerland, *Ae. albopictus* was found for the first time in the Canton of Ticino in the southernmost tip of the country in 2003 [13]; its surveillance has since been continuously expanded [26]. Today, the monitoring system consists of more than 1000 ovitraps that are analysed bi-weekly and the trapping data are used to coordinate targeted applications of insecticides [26]. In addition, public information campaigns are carried out in order to reduce breeding sites on private grounds. On public grounds, where larval breeding sites may not be removed (e.g. water drains), the authorities mainly apply *Bti* and diflubenzuron for larval control. In addition, focal spraying of permethrin to target adult mosquitoes is implemented if there is a risk of autochthonous transmission due to imported fever cases. Despite these efforts, *Ae. albopictus* has expanded its range in the Canton of Ticino over the last years [12, 26], requiring careful monitoring of the current insecticides’ efficacy.

*Bti* formulations are widely used and shown to be effective in controlling mosquitoes [27–29]. The toxicity of *Bti* against mosquito larvae is linked to crystals, produced during bacterial sporulation, that contain mainly four protoxins Cry11Aa, Cry4Aa, Cry4Ba and Cyt1Aa [30]. When ingested by mosquito larvae the crystals are dissolved in the alkaline milieu of the midgut and the released protoxins are then activated into toxins by gut proteases. The toxins bind to the receptors on the midgut cell membranes, leading to the formation of pores causing cell lysis, septicaemia and finally larval death [31, 32]. *Bti* toxins are highly active due to their
synergistic effects to the target species, while showing low toxicity for other organisms due to their specificity [33]. Bti may be used in combination with Lysinibacillus sphaericus, another entomopathogen that also produces insecticidal crystals. In combination, the toxins from both bacteria display synergistic efficacy in a wide range of mosquito species, including Aedes spp. [29].

This study was part of a Brazilian-Swiss Joint Research Programme and aimed to examine whether the Ae. albopictus populations in the Swiss-Italian border region and in Recife are still fully susceptible to the insecticides currently applied despite their use over many years, in particular to Bti that has been widely employed in both study areas. While Ae. albopictus is the only potential vector of DENV, CHIKV and ZIKV in Switzerland, the main vector in Brazil is Ae. aegypti and, therefore, insecticide susceptibility assays were done for both Aedes species in Brazil. As there are no data available on adulticides in Switzerland, additional WHO insecticide susceptibility bioassays were carried out. In Ticino, Switzerland, a surveillance and control programme targets Ae. albopictus, such a programme does not exist in the neighbouring Lombardy region in Italy; hence mosquitoes were collected from both areas and their insecticide susceptibility compared.

Methods

Aedes susceptible reference colonies

Three Aedes colonies were used as susceptible reference colonies for all the compounds tested in this study: (i) Rockefeller, an international standard Ae. aegypti colony; (ii) RecL, an Ae. aegypti colony established from eggs collected from the Recife Metropolitan Region (RMR) in 1996 [34]; and (iii) RecLalb, an Ae. albopictus colony established from eggs collected from the same area as the RecL. The colonies were maintained in the insectary of the Instituto Aggeu Magalhães in Recife, Brazil as previously described [34]. Briefly, insects were maintained under controlled conditions at 26 ± 1 °C, 70% relative humidity and a 14:10 h light:dark photoperiod. Larvae were reared in de-chlorinated tap water and fed with cat food (Whiskas®, Brazil). Adults were fed on a 10% sucrose solution and females were provided with chicken blood twice per week.

Establishment of Aedes field colonies

The field colonies were set up from eggs collected in the Canton of Ticino in southern Switzerland (TICINO), the Province of Como Lombardy in northern Italy (COMO) and from the Recife Metropolitan Region in Brazil, Sítio dos Pintos (SP) and Recife “field” (RF). The eggs for the TICINO and COMO colonies were provided by an already existing network of 280 ovitraps that were set across the Swiss-Italian border region [35]. The eggs were collected from the wooden slats in the ovitraps every other week between July and August 2013. To hatch out the eggs the slats were transferred to trays filled with de-chlorinated tap water. First-instar larvae were then split into equally sized batches, transferred to plastic trays and provided with TetraMin® fish food (Tetra, Melle, Germany). The larval trays were kept in a climate chamber (KBWF 720 E5.2, Binder GmbH, Tuttlingen, Germany) at 28 °C, 70% relative humidity and a 16:8 h light:dark photoperiod until pupation occurred. Adults that emerged from the pupae were transferred to a 30 × 30 × 30 cm Bugdorm-1® insect cage (Bugdorm, Taichung, Taiwan). Adults were allowed to mate and had access to water and 10% sucrose solution ad libitum. The founder populations of the TICINO and COMO colonies consisted of 520 (380 females and 140 males) and 610 (330 females, 280 males) adult mosquitoes, respectively. The females were blood-fed twice per week and their eggs collected on filter papers inside the cage to produce the test population. In Brazil, eggs were collected in 60 ovitraps distributed across Sítio dos Pintos (SP), a district of Recife city and used to establish the Ae. albopictus (SPalb) and Ae. aegypti (SPAeg) colonies, as described in Regis et al. [36]. Upon eclosion, larvae were maintained at the insectary of IAM-FIOCRUZ (Recife, Brazil) as described above. SPalb and SPAeg colonies were founded by 1774 (887 females and 887 males) and 3129 (1536 females and 1593 males) adult mosquitoes, respectively. Recife Field (RF), another Ae. aegypti colony representing 45 Recife districts was established from eggs sampled using ovitraps that were set according to the protocol previously described [37]. At least 1000 adults from these field collections were used to set up the RF colony. Bioassays were performed using larvae from the first (F₁), the second (F₂) or, in exceptional cases, also the third filial generation (F₃).

Larval bioassays

In the larval bioassays, the microbacteria Bti, individual Bti toxins and L. sphaericus as well as the chitin inhibiting diflubenzuron were tested against the above Aedes laboratory and field colonies (Table 1). Bti was prepared from the lyophilized reference powder IPS82 (Pasteur Institute, Paris, France), serotype H-14, as an aqueous suspensions at 5 g/l, and stored at -20 °C until use. In addition to Bti, individual Bti toxins, Cry11Aa and Cry4Ba, were produced with the Bt acrystalliferous strain 4Q2-81 that was transformed with plasmids carrying the respective protoxin genes [38]. Cry11Aa and Cry4Ba were chosen because they show the highest larval toxicity among the major protoxins of the crystal [39]. Spore-crystal biomass from each recombinant strain was produced and then lyophilized according to Barros et al. [40]. Similar to Bti an aqueous suspensions of
**Table 1** *Aedes* spp. colonies and evaluated insecticides

| Species | Colony | Source | Bti H-14 | Cry11Aa | Cry4Ba | Vectomax* | L. sphaericus | Diflubenzuron | Adulticides* |
|---------|--------|--------|----------|---------|---------|------------|---------------|--------------|-------------|
| Ae. albopictus | RecLalb | Lab. Brazil | x | x | x | x | x | - | - |
| TICINO | Field Switzerland | x | x | x | - | x | - | - |
| COMO | Field Italy | x | x | x | - | - | x |
| SPalb | Field Brazil | x | x | x | - | - | - |
| Ae. aegypti | Rockefeller | Lab. Brazil | x | x | x | - | - | x |
| Recl | Lab. Brazil | x | - | - | - | - | - |
| SPAeg | Field Brazil | x | x | x | - | - | - |
| RF | Field Brazil | x | - | - | - | - | - |

*Bendiocarb, DDT, malathion, permethrin (25:75 cis:trans ratio) and λ-cyhalothrin* 

*L. sphaericus* was prepared from lyophilized reference powder SPH88 (Pasteur Institute, Paris, France), serotype H5a5b strain 2362. Vectomax CG* (Valent Biosciences Corporation, Libertyville, IL, USA) is a commercial product available as water-soluble pouches containing a granular formulation that combines 4.5% *Bti* (serotype H-14, strain AM65-52) and 2.7% *L. sphaericus* (2362, serotype H5a5b, strain ABTS 1743) spores and insecticidal crystals as active ingredients (AIs). To prepare the stock suspension of 70 g/l (i.e. 5 g/l of AI) pouches (batch 179,654 N8) were incubated at 25 °C for 72 h in order to allow the release of crystals into the suspension. Aliquots of this suspension were then stored at -20 °C until use. Dose-response curves for *Bti*, Cry11Aa and Cry4Ba toxins, *L. sphaericus* and Vectomax® were estimated following the WHO guidelines for testing larvicides [41]. Briefly, batches of 20 third-instar larvae were exposed to serial dilutions of lyophilized spore-crystal powder in cups containing 100 ml bacterial suspensions in distilled water, without adding food. Five to seven concentrations for each compound were tested in each bioassay alongside a negative control group in three replicates. The negative control group was only exposed to distilled water. Each bioassay was repeated at least three times on different days. For *Bti*, Cry11Aa and Cry4Ba mortality rates were recorded after a 24 h exposure and for Vectomax* and *L. sphaericus* after a 48 h exposure time.

In addition to the above larvicides, the efficacy of diflubenzuron was assessed against the TICINO colony because the compound is being used in the Ticino surveillance and control programme [26]. The efficacy of diflubenzuron to prevent adult emergence was assessed in third-instar larvae as it inhibits the production of chitin. For the bioassays diflubenzuron analytical standard powder (Sigma-Aldrich: St. Louis, MO, USA, code 45446) was dissolved in acetone to make a 0.3% (w/v) stock solution and aliquots were stored at -20 °C until use. The bioassays then followed the protocols described in Martins et al. [42]. Briefly, 8–12 concentrations, between 0.2 and 4.0 μg/l, were tested alongside a negative control containing a 0.2% acetone in water solution. To avoid starvation effects during the long term assay period food was added to the test cups. Larvae were exposed in 8 batches of 10, yielding 80 individuals at each diflubenzuron concentration and in the negative control. Any dead larva or pupa was removed from the bioassay cups every other day and adult emergence was observed up to 30 days. The assays were then repeated on different days up to four times (Additional file 1: Tables S1-S3).

The mortality and inhibition rates were the basis to estimate dose-response curves in order to predict the average concentrations, and 95% confidence intervals, at which 50% and 90% of the larval population would be killed (i.e. LC50 and LC90) or, in the case of diflubenzuron, prevented from reaching the adult stage (i.e. EI50 and EI90). The dose-response curves were estimated using generalised linear models with a binomial distribution and a “probit” link function. The models were computed using the statistical software IBM SPSS 10.0 for Windows.

On the basis of the estimated LC50s and LC90s a resistance ratio (RR) was calculated, where RR is the ratio between the LC for the test colony and the LC of the reference colony. For chemical insecticides, Mazzarri & Georghiou [43] proposed the following classifications: low resistance for an RR below 5, moderate resistance for an RR between 5 and 10, and high resistance for an RR above 10. However, for biological compounds such as *Bti*, RR values lower than 10-fold are considered as natural variations [19, 44].

**Adult bioassays**

In addition to larvicides, the *Ae. albopictus* TICINO and COMO colonies sampled from the Swiss-Italian border region were also tested for their susceptibility against the four insecticide classes of WHO recommended adulticides. The insecticides evaluated were bendiocarb,
dichlorodiphenyltrichloroethane (DDT), malathion, the non-alpha-cyano pyrethroid permethrin (25:75 cis:trans ratio) and the alpha-cyano pyrethroid λ-cyhalothrin (Table 1). The pyrethroids were kindly provided by Syngenta Crop Protection (Basel, Switzerland), while the other insecticides were purchased as technical grades from Sigma-Aldrich. The bioassays were performed on females of the F2 generation raised from the field-sampled eggs following the WHO guidelines for testing adulticides [45]. Using a series of insecticide-impregnated filter papers, dose-response curves were estimated to determine the lethal dosage (LD) that would kill 50% (LD50) and 90% (LD90) of the TICINO and COMO colonies. The filter papers (Whatman no. 1) were impregnated with insecticide in acetone solutions mixed with silicon oil (Dow Corning 556 Silicon) according to the WHO test procedures [46]. The insecticide solutions were serial dilutions with five to six concentrations that would yield mortality rates between 0 and 100%. In the test, batches of 17–25 non-blood-fed Ae. albopictus females, aged 2–5 days were introduced into the exposure tubes lined with the insecticide-treated filter papers. The mosquitoes were exposed for 1 h, then gently blown back into the holding tube and provided with 10% sucrose solution. Following a 24 h recovery period, the numbers of dead and alive mosquitoes were recorded. Mosquitoes were considered to be alive if they were able to fly. Any knocked-down mosquito, with or without legs and wings, were considered moribund and were recorded as dead [45]. Tests were repeated aiming at 100 mosquitoes exposed per insecticide and concentration, including a negative control.

For the dose-response curves the 24 h mortality rates were the basis to estimate the dosage at which 50% and 90% of the adult population would be killed (i.e. LD50 and LD90). The dose-response curves were estimated using generalised linear models (GLM) with a binomial distribution and a “logit” link function, predicting mortality as a function of the log-transformed concentration. The models were computed in the freely available software package R, 3.3.2 [47] and the graph was produced with the R package “ggplot2” [48].

**Results**

The susceptibility to Bti and its toxins was assessed for each Aedes population as they are all being exposed to Bti in the study areas. The Ae. albopictus populations from the Canton of Ticino in southern Switzerland, the Como area in northern Italy and Recife, Brazil were all still susceptible to Bti. The LC50 values were similar with a concentration of 0.015 mg/l, while the LC90 values varied between 0.030–0.036 mg/l (Table 2). RRs between the Ae. albopictus field colonies and RecL Lab reference colony were all below two-fold, showing that the field populations remain fully susceptible to Bti. Likewise, the LC values for SPaeg and RF, the two Ae. aegypti populations from Brazil were close to those observed for the reference susceptible Rockefeller and the RecL colonies (Table 2). The LC values for Bti across Ae. albopictus and Ae. aegypti suggest that both species display a similar level of susceptibility to this agent.

In order to detect early development of resistance to individual Bti toxins, lyophilized powders containing Cry11Aa or Cry4Ba toxins were tested separately. Here, the LC50 values of Cry11Aa and Cry4Ba against the Ae. albopictus and Ae. aegypti field colonies were close to those found for the corresponding reference colonies (Table 3). The LC values of the selected Cry toxins were an order of magnitude higher than those of the overall Bti crystal, corroborating the Bti cocktail to be more effective than individual toxins.

The efficacy of L. sphaericus, another entomopathogenic bacterium, was tested against Ae. albopictus for which the susceptibility status has been poorly documented, in contrast to Ae. aegypti that is well known to be refractory. The reference powder SPH88 that contains crystals of the binary (Bin) toxin from the 2362 strain gave an LC50 of 0.084 mg/l and an LC90 of 0.336 mg/l in the RecL Lab reference colony (Table 4).

The activity of Vectomax®, a mixture of Bti and L. sphaericus crystals, was also investigated in order to evaluate if this combination of insecticidal components could be an effective alternative to control Ae. albopictus. Data from our evaluation showed similar LC values for TICINO, COMO and SPalb Ae. albopictus colonies (Table 4) and RRs below 2, suggesting Vectomax® to be effective.

The third control agent that was tested against immature mosquito stages was diflubenzuron, a compound used in Ticino to control Ae. albopictus [26] but neither in Recife nor Como; hence the efficacy of diflubenzuron was only evaluated against the TICINO colony. The EL50 preventing 50% larvae from developing into the adult stage was 0.376 mg/l (95% confidence interval; 95% CI: 0.289–0.462 mg/l) and the EL90 was 1.197 mg/l (95% CI: 1.033–1.448 mg/l). These concentrations were similar to those observed for the Ae. aegypti Rockefeller colony that was used as a reference. Here, the EL50 was 0.456 mg/l (95% CI: 0.352–0.549 mg/l) and the EL90 was 1.655 mg/l (95% CI: 1.322–2.249 mg/l).

Finally, the two Ae. albopictus populations, COMO and TICINO from the Swiss-Italian border region were also tested against five insecticides, representing the four available classes of adulticides for which the susceptibility status was unknown. Among the five insecticides λ-cyhalothrin showed the lowest LC50 followed by bendiocarb, permethrin, malathion and DDT (Table 5). The two populations showed very similar dose-response profiles (Fig. 1). Although there are no diagnostic concentrations...
available for *Ae. albopictus*, adult mortalities after a 1 h exposure and 24 h holding period to bendiocarb, malathion, permethrin and λ-cyhalothrin suggest that both *Ae. albopictus* field populations are susceptible given that the mortality rates were close to 100% at the diagnostic concentration for other mosquito species (Fig. 1) and below those reported for permethrin and λ-thion, permethrin and *Ae. albopictus* LC50 (95% CI) are estimated to be 96% for the TICINO and 96.9% for the COMO population. However, assuming *Ae. aegypti* to be a reference species, the COMO and TICINO colonies showed decreased sensitivity to DDT as mortality rates at the *Ae. aegypti* diagnostic concentration of 4% are estimated to be 96% for the TICINO and 96.9% for the COMO population, even at the extended exposure time of 1 h. Note that, according to WHO, the exposure time for *Ae. aegypti* against DDT would only be 30 min [51].

The original data used in the statistical analysis are provided in the Additional file 1: Tables S1-S3.

### Discussion

In the absence of commercially available vaccines or treatments, dengue, chikungunya, Zika and other arbovirus transmissions may only be averted through vector control. However, vector control heavily relies on insecticides, raising concerns over the development of insecticide resistance beside adverse effects on the environment and human health [52]. Knowing the insecticide susceptibility status of a local mosquito population is, therefore, crucial [53]. Still, many programmes have been implemented without previously evaluating the susceptibility profiles of the target field populations to the intended control agents. In some cases, laboratory colonies have been used as surrogates to establish the susceptibility status, yet such colonies may underestimate the existence of resistance alleles in the field due to founder and bottle neck effects when maintaining laboratory colonies [44].

The biological larvicide *Bti* is known to be effective in reducing mosquito densities in control programmes and has a high toxicity to the target species without causing unwanted side-effects to the environment [28, 54]. The specificity of *Bti* is particularly important for the control of mosquito species that breed in ecologically sensitive areas where broad-spectrum insecticides may not be used. Likewise, in urban settings the control of day-active mosquito species like *Ae. albopictus* and *Ae. aegypti* by adulticides is critical because of human exposure to the insecticides. With the exception of one case in *Culex quinquefasciatus* in New York, USA [55], to our knowledge, no resistance to *Bti* has been reported.

### Table 2 Toxicity of *Bacillus thuringiensis* svar. *israelensis* (IPS82) against third-instar Aedes spp. larvae

| Species         | Colony     | Number | LC50 (95% CI)a | RRb | LC90 (95% CI) |
|-----------------|------------|--------|----------------|-----|--------------|
| *Ae. albopictus*| RecLalb    | 1080   | 0.009 (0.008–0.011) | –   | 0.028 (0.023–0.037) |
|                 | TICINO     | 1440   | 0.015 (0.012–0.018) | 1.7 | 0.036 (0.030–0.060) |
|                 | COMO       | 1120   | 0.015 (0.012–0.016) | 1.7 | 0.030 (0.026–0.036) |
|                 | SPalb      | 1560   | 0.015 (0.011–0.020) | 1.7 | 0.036 (0.027–0.098) |
| *Ae. aegypti*   | Rockefeller| 1320   | 0.008 (0.007–0.009) | –   | 0.026 (0.021–0.036) |
|                 | RecL       | 1080   | 0.013 (0.011–0.015) | 1.6 | 0.032 (0.027–0.039) |
|                 | SPaeg      | 1140   | 0.014 (0.012–0.016) | 1.7 | 0.029 (0.025–0.035) |
|                 | RF         | 1860   | 0.013 (0.012–0.016) | 1.6 | 0.037 (0.030–0.050) |

aConcentration (mg/l) that is lethal to 50% or 90% of the larvae over a 24 h exposure, mean and 95% confidence interval

bResistance ratio (RR) between the LCs of the test colony and the reference colony

### Table 3 Toxicity of Cry11Aa and Cry4Ba against third-instar Aedes spp. larvae

| Species         | Colony     | Cry11Aa | Cry4Ba |
|-----------------|------------|---------|--------|
| *Ae. albopictus*| RecLalb    | 1500    | 1380   |
|                 | TICINO     | 1020    | 1440   |
|                 | COMO       | 1120    | 1060   |
|                 | SPalb      | 1560    | 980    |
| *Ae. aegypti*   | Rockefeller| 1080    | 1120   |
|                 | SPaeg      | 1140    | 1140   |

| Species         | Colony     | Cry11Aa n | LC50 (95% CI)a | RRb | Cry4Ba n | LC50 (95% CI)a | RRc |
|-----------------|------------|------------|----------------|-----|----------|----------------|-----|
| *Ae. albopictus*| RecLalb    | 1500       | 0.410 (0.311–0.514) | –   | 1380     | 0.595 (0.431–0.787) | –   |
|                 | TICINO     | 1020       | 0.539 (0.437–0.648) | 1.3 | 1440     | 0.483 (0.213–0.839) | 0.8 |
|                 | COMO       | 1120       | 0.650 (0.517–0.798) | 1.6 | 1060     | 0.782 (0.589–1.042) | 1.3 |
|                 | SPalb      | 1500       | 0.432 (0.335–0.530) | 1.1 | 980      | 0.830 (0.622–1.095) | 1.4 |
| *Ae. aegypti*   | Rockefeller| 1080       | 0.162 (0.121–0.210) | –   | 1120     | 0.331 (0.209–0.492) | –   |
|                 | SPaeg      | 1140       | 0.266 (0.207–0.339) | 1.6 | 1140     | 0.685 (0.482–0.969) | 2.1 |

aConcentration (mg/l) that is lethal to 50% or 90% of the larvae over a 24 h exposure, mean and 95% confidence interval

bResistance ratio (RR) between the LCs of the test colony and the reference colony

cReference colony
from mosquito field populations [56–59] and decreased larval susceptibility to \textit{Bti} is also rare [60–62]. Under laboratory conditions resistance has been found to single \textit{Bti} toxins in selection experiments [63–65] but not to \textit{Bti}. With regards to the case of \textit{Bti} resistance reported from New York, it is inconclusive whether the observed resistance is linked to the application of \textit{Bti} as there are neither data available from the pre-treatment period nor has the finding been confirmed in a follow up study.

Here, we performed larval bioassays with \textit{Bti} reference powder IPS82 and two \textit{Bti} toxins, Cry11Aa and Cry4Ba. Our study showed no increased tolerance in any of the \textit{Aedes} populations and susceptibility was also similar between the intervention and the non-intervention areas in the Swiss-Italian border. Also, the results from the \textit{Aedes} populations in Recife, Brazil suggest the exposure to \textit{Bti} for several years had not selected for insecticide resistance. Comparing our results to the findings from other studies it appears that variations in \textit{Bti} susceptibility in \textit{Aedes} spp. are narrow [19, 61, 66, 67]. Likewise, our data show that \textit{Ae. albopictus} and \textit{Ae. aegypti} are equally susceptible to \textit{Bti}, suggesting that the same application rates may be used where both species co-exist. This is an important finding since \textit{Ae. albopictus} can be found in many urban environments together with \textit{Ae. aegypti}, and presence of both species in these areas is being increasingly reported [24].

Tetreau et al. [65] stated that one of the main reasons why no resistance to \textit{Bti} has yet been detected in the field is due to the synergistic effect of the individual toxins which may mask failure of individual toxins. Previous laboratory studies have shown that exposure to single \textit{Bti} toxins selects for resistance but not when the toxins are combined [64, 67–69]. In this study the approach of Tetreau et al. [65] was followed and bioassays with two individual \textit{Bti} toxins were performed with larvae in order to have a more sensitive assay that may detect early development of resistance. However, the mosquito test populations were still fully susceptible even to the individual Cry11Aa and Cry4Ba toxins. We, therefore, conclude that \textit{Bti} treatments in both Ticino, Switzerland and Recife, Brazil have not exerted a selection pressure strong enough to cause a differential larval response to these individual toxins.

Like \textit{Bti}, \textit{L. sphaericus} is a naturally occurring soil bacterium that produces a larvicidal toxin [29]. The efficacy of \textit{L. sphaericus} against \textit{Ae. albopictus} has not been well investigated; and was also assessed here in order to account for the wide variations of \textit{L. sphaericus} toxicity generally observed in this genus [70–72]. For

### Table 4 Lethal concentrations for Vectomax® and \textit{Lysinibacillus sphaericus} (SPH88) against third-instar \textit{Aedes albopictus} larvae

| Larvicide                | Colony | Number | LC50<sup>a</sup> Mean (95% CI) | RR<sup>b</sup> | LC90<sup>a</sup> Mean (95% CI) | RR<sup>b</sup> |
|--------------------------|--------|--------|-------------------------------|---------------|-------------------------------|---------------|
| Vectomax®                | RecLalb| 1140   | 0.087 (0.080–0.094)            | –             | 0.163 (0.145–0.190)           | –             |
| TICINO                   | 1440   | 0.131 (0.118–0.144)            | 1.5            | 0.221 (0.194–0.228)           | 1.4           |
| COMO                     | 1120   | 0.076 (0.069–0.083)            | 0.9            | 0.145 (0.130–0.169)           | 0.9           |
| SPalb                    | 1260   | 0.092 (0.077–0.105)            | 1.1            | 0.191 (0.159–0.305)           | 1.2           |
| \textit{L. sphaericus}   | RecLalb| 1080   | 0.084 (0.070–0.099)            | –             | 0.336 (0.239–0.630)           | –             |

<sup>a</sup>Concentration (mg/l) that is lethal for 50% or 90% of larvae over a 48 h exposure, mean and 95% confidence limits

<sup>b</sup>Resistance ratio (RR) between the test colonies and the RecLalb reference colony

### Table 5 Lethal concentrations for adulticides in \textit{Aedes albopictus} from the Swiss-Italian border region

| Insecticide | Population | Number<sup>a</sup> | LC50<sup>b</sup> Mean (95% CI) | LC90<sup>b</sup> Mean (95% CI) |
|-------------|------------|--------------------|-------------------------------|-------------------------------|
| Bendiocarb  | TICINO     | 463                | 0.015 (0.014–0.017)            | 0.021 (0.019–0.024)            |
|             | COMO       | 470                | 0.017 (0.016–0.019)            | 0.027 (0.024–0.031)            |
| DDT         | TICINO     | 523                | 1.359 (1.220–1.514)            | 3.048 (2.557–3.635)            |
|             | COMO       | 470                | 1.126 (1.003–1.263)            | 2.807 (2.309–3.413)            |
| \textit{λ-cyhalothrin} | TICINO | 440                | 0.007 (0.006–0.007)            | 0.012 (0.010–0.014)            |
|             | COMO       | 426                | 0.006 (0.006–0.007)            | 0.011 (0.010–0.013)            |
| Malathion   | TICINO     | 489                | 0.116 (0.104–0.128)            | 0.262 (0.222–0.310)            |
|             | COMO       | 486                | 0.120 (0.108–0.133)            | 0.284 (0.239–0.338)            |
| Permethrin  | TICINO     | 481                | 0.046 (0.042–0.051)            | 0.094 (0.081–0.110)            |
|             | COMO       | 430                | 0.051 (0.047–0.056)            | 0.092 (0.079–0.106)            |

<sup>a</sup>Total number of mosquitoes exposed across 5–6 concentrations

<sup>b</sup>Concentrations are expressed as % insecticide on the filter paper in the WHO insecticide susceptibility assay to kill 50% (LC<sub>50</sub>) and 90% (LC<sub>90</sub>) of the mosquito population over a 24 h holding period
**Ae. albopictus** we found LC values that were 8 to 13-fold higher than that reported for C. quinquefasciatus [73]. That is far better than the LC values for *Ae. aegypti*, which are 100 to 1000-fold higher and, for this reason, it is considered a refractory species [71]. When comparing LC values of both bacteria towards *Ae. albopictus*, *L. sphaericus* showed good activity since this species was only 11-fold less susceptible compared to Bti.

The biological larvicide Vectomax®, containing a mixture of Bti and *L. sphaericus* crystals, was also evaluated against *Ae. albopictus* since *L. sphaericus* activity can be enhanced by Bti [74]. Vectomax® combines Bti’s advantage of resistance-blocking together with *L. sphaericus*’ advantage of longer residuality in a single formulation and a broader spectrum of action. Our evaluation shows that Vectomax® is effective against *Ae. albopictus*, making it an alternative for the control of immature stages in our study areas. Conjugated products such as Vectomax® offer a mixture of 5 toxins that can target a wider range of medically important insect species, while showing a low risk of selection for resistance due to their complex mode of action. Other studies evaluating of Vectomax® in various environments have also shown its efficacy [75–77]. In conclusion, Vectomax® is a promising candidate to replace the actual microbial larvicides in our study areas.

Diflubenzuron showed to be another viable alternative to control *Ae. albopictus* larvae given the low inhibitory concentration found in the Ticino population, regardless of its history of utilisation in this area. Nevertheless, application of diflubenzuron needs careful assessment because it may harm non-target organisms.

In Ticino, the control of *Ae. albopictus* is mainly based on larval source reduction, either by removing breeding sites or by applying Bti or diflubenzuron if the larval sources cannot be removed [26]. Adulticides are rarely used. Only in exceptional cases is permethrin sprayed on vegetation when mosquito densities cause nuisance to residents in a confined area, or in surroundings from where symptomatic patients with arboviral disease have been reported [26]. Nevertheless, the susceptibility status of the Swiss *Ae. albopictus* population has never been investigated; and hence it is useful to know that permethrin shows good activity against the local *Ae. albopictus* population. While the results for permethrin, bendiocarb, λ-cyhalothrin and malathion suggest that the *Ae. albopictus* populations across the Swiss-Italian border region may be considered susceptible to these insecticides, there are some indications that the population shows decreased susceptibility to DDT, or perhaps even resistance. Alteration of susceptibility to DDT has also been recorded in *Aedes* spp. in Thailand, Japan, Malaysia, Cameroon and the Central African Republic [49, 56, 78–80]; the underlying mechanisms remain unclear.

In Brazil, although the use of adulticides have decreased due to their toxicity to humans, resistance to most used compounds has already been widely documented in *Aedes* spp. populations [81, 82] and was not further investigated in the present study.

Although we made some inference about adulticide susceptibility in *Ae. albopictus*, we lack discriminating concentrations for this mosquito species. However, with its increasing importance for public health it would be helpful to have explicit discriminating concentrations also established for adulticides against *Ae. albopictus*. 

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**Fig. 1** Dose-response effects of adulticides in *Aedes albopictus* from the Swiss-Italian border region. The curves show the estimated dose-response relationship between the 24 h mortality and the percentage insecticide on the filter paper in WHO insecticide susceptibility assays. The symbols represent the summaries of the actual measurements in the bioassays, while the curves are the predicted estimates of the mean and the shaded areas the 95% confidence intervals around the means. The vertical lines indicate the discriminating concentrations in *Ae. aegypti* for DDT, λ-cyhalothrin, malathion and permethrin, and in *Anopheles gambiae* for bendiocarb, as there are no discriminating concentrations established for *Ae. albopictus*.
In summary, the *Aedes* populations evaluated in this study were equally susceptible to the insecticides evaluated. The study implies that the currently applied mosquito larvicides in Ticino, southern Switzerland as well as in Recife, Brazil and adulticides in Ticino are still effective for the control of *Ae. albopictus* and *Ae. aegypti*. The larvicides tested have distinct modes of action and this feature is important to avoid the onset of resistance. Besides the use of insecticides, other strategies show promising results in decreasing vector densities and should be considered as part of integrated mosquito control programmes [83–87]. Highly productive artificial breeding sites are often found on private properties [83] and information campaigns encouraging the elimination of these water containers, alongside a correct use of biological insecticides, can significantly decrease the local mosquito population. The cost-effectiveness of such approaches and their long-term success should be evaluated when compared with conventional control methods [88]. Our results demonstrate the importance of research on the susceptibility status of mosquito populations to insecticides to prevent the spread of resistance in these important vectors of human diseases.

**Conclusions**

Currently used larvicides (i.e. *Bti*, diflubenzuron) and adulticides (permethrin) used for mosquito control in the Ticino and Recife control programmes remain effective against the local *Ae. albopictus* and *Ae. aegypti* populations. The susceptibility profiles of the different mosquito populations were similar, despite distinct differences in the deployed interventions and geographical context. In addition, *Ae. albopictus* and *Ae. aegypti* display similar susceptibility levels to *Bti*, suggesting that this biolarvicide may target both species where they co-exist.

**Additional files**

Additional file 1: Original data that were the basis for the statistical analysis. Table S1. Larvicide mortalities recorded in the bioassays with diflubenzuron. Table S2. Mortalities recorded in the bioassays with *Bti* and individual *Bti* toxins. Table S3. Mortalities recorded in the bioassays with adulticides. (XLSX 122 kb)

**Abbreviations**

95% CI: 95% confidence interval; *Bti*: *Bacillus thuringiensis* svar. *israelensis*; CHIKV: Chikungunya virus; DDT: Dichlorodiphenyltrichloroethane; DENV: Dengue virus; EI: Emergence inhibition; LC: Lethal concentration; OP: Organophosphate; RR: Resistance ratio; ZIKV: Zika virus

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11. Global Invasive Species Database. http://www.isg.org/database/species/search.asp?st=100ss&n=1&lan=en-EN. Accessed 7 Dec 2012.
12. Flacio E, Engeler L, Tonolla M, Müllner P. Spread and establishment of Aedes albopictus in southern Switzerland between 2003 and 2014: an analysis of oviposition data and weather conditions. Parasit Vectors. 2016;9:304.
13. Flacio E, Lüthy P, Patocchi N, Guidotti F, Tonolla M, Peduzzi R. Primo ritrovamento di Aedes albopictus in Svizzera. Boll Della Soc Tisicene Sci Nat. 2004;92:141–2.
14. ECDC. VBORNET Mosquito maps. European Centre for Disease Control and Prevention. 2017. http://www.ecdc.europa.eu/en/healthtopics/vectors/vector-maps/Pages/VBORNET_maps.aspx. Accessed 5 Jul 2017.
15. dos Santos RL. Updating of the distribution of Aedes albopictus in Brazi (1997). Rev Saúde Pública. 2003;37:5.
16. Scholte EJ, Jacobs F, Linton Y-M, Dijkstra E, Fransen J, Taekken W. First record of Aedes (Stegomyia) albopictus in the Netherlands. Eur Mosq Bull. 2007;22:5–9.
17. Reiter P, Sprenger D. The used tire trade: a mechanism for the worldwide dispersal of container breeding mosquitoes. J Am Mosq Control Assoc. 1999;15:494–501.
18. Braga IA, Valle D. Aedes aegypti: histórico do controle no Brasil. Epidemiol Serv Saúdes. 2007;16:113–8.
19. Araújo AP, Araújo Diniz DF, Helvecio E, de Barros RA, de Oliveira CMF, CFJ A, et al. The susceptibility of Aedes aegypti populations displaying temephos resistance to Bacillus thuringiensis israelensis: a basis for management. Parasit Vectors. 2011;6:297.
20. Lima EP, Santos Paiva MH, de Araújo AP, Gomes da Silva EV, da Silva UM, de Oliveira LN, et al. Insecticide resistance in Aedes aegypti populations from Ceará, Brazil. Parasit Vectors. 2011;4:5.
21. Macoris Mdle LG, MTM A, Takaku L, Glasser CM, Garbeloto VC, Bracco JE. Resistance of Aedes aegypti from the state of São Paulo, Brazil, to organophosphates insecticides. Mem Inst Oswaldo Cruz. 2003;98:703–8.
22. Secretaria de Vigilância em Saúde. Nota Técnica no 015/2009/CGPNCD/Aedes aegypti – var. Brasil: Ministério da Saúde; 2009.
23. Secretaria de Vigilância em Saúde. Nota Técnica no 123/2014/GAB/SVS/MS. Brasília: Ministério da Saúde; 2014.
24. Carvalho RG, Lourenço-de-Oliveira R, Braga IA. Updating the geographical distribution and frequency of Aedes albopictus in Brazil with remarks regarding its range in the Americas. Mem Inst Oswaldo Cruz. 2014;109:787–96.
25. de Albuquerque CM, Melo-Santos MAV, Bezerra MAS, Barbosa RM, Silva DF, da Silva E. First report of Aedes albopictus in areas of rain forest in Brazil. Rev Saúde Pública. 2000;34:314–5.
26. Flacio E, Engeler L, Tonolla M, Lüthy P, Patocchi N. Strategies of a thirteen year surveillance programme on Aedes albopictus (Stegomyia albopicta) in southern Switzerland. Parasit Vectors. 2015;8:208.
27. Becker N. Management of mosquitoes: management of the Upper Rhine mosquito population as a model programme. Parasitol Today. 1997;13:498–7.
28. Guidi V, Patocchi N, Lüthy P, Tonolla M. Distribution of Bacillus thuringiensis subsp. israelensis in Soil of a Swiss Wetland reserve after 22 years of mosquito control. Appl Environ Microbiol. 2011;77:3663–8.
29. Lacey IA. Bacillus thuringiensis serovar israelensis and Bacillus sphaericus for mosquito control. J Am Mosq Control Assoc. 2007;23:133–63.
30. Becker N, O'Neill S, Ben-Dov E, Jones AF, Murphy L, Quail MA, et al. Complete sequence and organization of ptoxins, the toxin-coding plasmid of Bacillus thuringiensis subsp. israelensis. Appl Environ Microbiol. 2002;68:5082–95.
31. de Maagd RA, Bravo A, Crickmore N. How Bacillus thuringiensis has evolved specific toxins to colonize the insect world. Trends Genet. 2001;17:193–9.
32. Gómez I, Pardo-López L, Muñoz-Garay C, Fernandez LE, Pérez C, Sánchez J, et al. Role of receptor interaction in the mode of action of insecticidal Cry and Cyt toxins produced by Bacillus thuringiensis. Peptides. 2007;28:169–73.
33. Thomas WE, Ellar DJ. Mechanism of action of Bacillus thuringiensis var israelensis insecticidal delta-endotoxin. FEBS Lett. 1983;154:362–8.
34. de Melo-Santos MAV, de Araújo AP, Rios EMM, Regis L. Long lasting persistence of Bacillus thuringiensis serovar. israelensis larvicidal activity in Aedes aegypti (Diptera: Culicidae) breeding places is associated to bacteria recycling. Biol Control. 2009;49:1896–91.
35. Suter TT, Flacio E, Farlha BF, Engeler L, Tonolla M, Regis LN, et al. Surveillance and control of Aedes albopictus in the Swiss-Italian border region: differences in egg densities between intervention and non-intervention areas. PLoS Neg Trop Dis. 2016;10:e0004515.
61. Boyer S, Tsiquin M, Ravani P. Differential sensitivity to Bacillus thuringiensis var. israelensis and temephos in field mosquito populations of Culex tritaeniorhynchus (Diptera: Culicidae): toward resistance? Environ Toxicol Chem. 2007;26:157–62.

62. Hongyu Z, Changju Y, Jingle H, Lin L. Susceptibility of field populations of Anopheles sinensis (Diptera: Culicidae) to Bacillus thuringiensis subsp. israelensis. Biocontrol Sci Tech. 2004;14:321–5.

63. Cadavid-Restrepo G, Sahaza J, Orend S. Treatment of an Ae. aegypti colony with the Cry11Aa toxin for 54 generations results in the development of resistance. Mem Inst Oswaldo Cruz. 2012;107:74–9.

64. Georgiou GP, Wirth MC. Influence of exposure to single versus multiple toxins of Bacillus thuringiensis subsp. israelensis on development of resistance in the mosquito Culex quinquesquaxius (Diptera: Culicidae). Appl Environ Microbiol. 1997;63:1095–101.

65. Tetreau G, Stallinski R, David J-P, Després L. Monitoring resistance to Bacillus thuringiensis subsp. israelensis in the field by performing bioassays with each Cry toxin separately. Mem Inst Oswaldo Cruz. 2013;108:894–900.

66. Marcombe S, Farajollahi A, Healy SP, Clark GG, Fonseca DM. Insecticide resistance status of United States populations of Aedes albopictus and mechanisms involved. PLoS One. 2014;9:e101992.

67. Wirth MC. Mosquito resistance to bacterial larvicidal toxins. Open Toxinology J. 2010;3:126–40.

68. Paris M, Tetreau G, Laurent F, Lelu M, Despres L, David J-P. Persistence of Bacillus thuringiensis israelensis (Bti) in the environment induces resistance to multiple Bti toxins in mosquitoes. Pest Manag Sci. 2011;67:122–8.

69. Wirth MC, Walton WE, Federici BA. Inheritance patterns, dominance, stability, and allelism of insecticide resistance and cross-resistance in two colonies of Culex quinquesquaxius (Diptera: Culicidae) selected with Cry toxins from Bacillus thuringiensis subsp. israelensis. J Med Entomol. 2012;49:886–94.

70. Berry C, Hindley J, Erhardt AF, Grounds T, De Souza I, Davidson EW. Genetic determinants of host ranges of Bacillus sphaericus mosquito larvicidal toxins. J Bacteriol. 1993;175:510–8.

71. Thiery I, de Barjac H. Selection of the most potent Bacillus sphaericus strains based on activity ratios determined on three mosquito species. Appl Microbiol Biotechnol. 1989;31:577–81.

72. Wright SP, Molloy DP, Singer S. Studies on the culicine mosquito host range of Bacillus sphaericus and Bacillus thuringiensis var. israelensis with notes on the effects of temperature and instar on bacterial efficacy. J Invertebr Pathol. 1987;49:291–302.

73. Chalorgo KDM, Romao TP, Amorim LB, Anastacio DB, de Barros RA, de Oliveira CMF, et al. Detection of an allele conferring resistance to Bacillus sphaericus binary toxin in Culex quinquesquaxius populations by molecular screening. Appl Environ Microbiol. 2009;75:1044–9.

74. Wirth MC, Jiannino JA, Federici BA, Walton WE. Synergy between toxins of Bacillus thuringiensis subsp. israelensis and Bacillus sphaericus. J Med Entomol. 2004;41:935–41.

75. Anderson JF, Ferrandino CJ, Dingman DW, Main AJ, Andreidis TIG, Becnel JJ. Control of mosquitoes in catch basins in Connecticut with Bacillus thuringiensis israelensis, Bacillus sphaericus, and spinosad. J Am Mosq Control Assoc. 2011;27:45–55.

76. Dritz DA, Lawler SP, Evkhanian C, Graham P, Baracosa V, Dula G. Control of mosquito larvae in seasonal wetlands on a wildlife refuge using Vectomax FX. Curr. J Am Mosq Control Assoc. 2011;27:498–403.

77. Eritja R. Laboratory tests on the efficacy of V8C60035, a combined larvicidal formulation of Bacillus thuringiensis israelensis (Strain AM65-52) and Bacillus sphaericus (Strain 2362) against Aedes aegypti in simulated catch basins. J Am Mosq Control Assoc. 2013;29:280–3.

78. Kawada H, Maekawa Y, Abe M, Ohashi K, Ohba S, Takagi M. Spatial distribution and pyrethroid susceptibility of mosquito larva collected from catch basins in parks in Nagasaki City, Nagasaki, Japan. Jpn J Infect Dis. 2010;63:19–24.

79. Somboon P, Prapantadhara L, Suwonkerd W. Insecticide susceptibility tests of Anopheles minimus s.l., Aedes aegypti, Aedes albopictus, and Culex quinquesquaxius in northern Thailand. Southeast Asian J Trop Med Public Health. 2003;34:87–93.

80. Ngaoguoni C, Kamgang B, Brengues C, Yahouedo G, Paupy C, Nkoué E, et al. Susceptibility profile and metabolic mechanisms involved in Aedes aegypti and Aedes albopictus resistant to DDT and deltamethrin in the Central African Republic. Parasit Vectors. 2016;9:599.

81. Maceiros MLG, Andrighetti MTM, Wandeler DMV, Ribolla PEM. Impact of insecticide resistance on the field control of Aedes aegypti in the State of São Paulo. Rev Soc Bras Med Trop. 2014;47:573–8.