Monitoring resistance to *Bacillus thuringiensis* subsp. *israelensis* in the field by performing bioassays with each Cry toxin separately

Guillaume Tetreau*, Renaud Stalinski, Jean-Philippe David, Laurence Després

Laboratoire d’Écologie Alpine, Unité Mixte de Recherche 5553, Université de Grenoble 1, Grenoble, France

*Bacillus thuringiensis* subsp. *israelensis* (Bti) is increasingly used worldwide for mosquito control and is the only larvicide used in the French Rhône-Alpes region since decades. The artificial selection of mosquitoes with field-persistent Bti collected in breeding sites from this region led to a moderate level of resistance to Bti, but to relatively high levels of resistance to individual Bti Cry toxins. Based on this observation, we developed a bioassay procedure using each Bti Cry toxin separately to detect cryptic Bti-resistance evolving in field mosquito populations. Although no resistance to Bti was detected in none of the three mosquito species tested (Aedes rusticus, Aedes sticticus and Aedes vexans), an increased tolerance to Cry4Aa (3.5-fold) and Cry11Aa toxins (8-fold) was found in one Ae. sticticus population compared to other populations of the same species, suggesting that resistance to Bti may be arising in this population. This study confirms previous works showing a lack of Bti resistance in field mosquito populations treated for decades with this bioinsecticide. It also provides a first panorama of their susceptibility status to individual Bti Cry toxins. In combination with bioassays with Bti, bioassays with separate Cry toxins allow a more sensitive monitoring of Bti-resistance in the field.

Key words: *Bacillus thuringiensis* subsp. *israelensis* - bioassays - field resistance - Cry toxins - mosquito

Numerous insect species represent major economic and sanitary issues for human societies as they can strongly impact crop cultures or transmit severe human diseases such as malaria, leishmaniasis or trypanosomiasis (Medlock et al. 2012, Savary et al. 2012, van Griensven & Diro 2012). The control of pest populations is therefore of high importance and relies mainly on the use of chemical and biological insecticides (Meissle et al. 2011, van den Berg 2011). The main threat for an efficient pest control is the ability of insects to quickly develop resistance to nearly all the insecticides used, leading to a strong decrease of insecticides efficiency (Tabashnik et al. 2008, Rivero et al. 2010). To bypass these resistances and adapt the insecticide treatment strategies, detecting incipient resistance in treated populations as early as possible is required (Carriere et al. 2010).

The most obvious approach to detect resistance in the environment is to perform bioassays on field-sampled individuals, which consist in exposing insects to increasing amount of insecticide to calculate their relative tolerance to this insecticide (Sivasupramaniam et al. 2007). Such basic approach is often preliminary to further studies exploring in-depth the molecular bases of the resistance (Tabashnik et al. 2006, Marcombe et al. 2009). Bioassays have been performed on numerous field populations of insects with different chemical insecticides (Ocampo et al. 2011, Shin et al. 2011, Silva et al. 2011, Springate & Colvin 2012), but also with biological insecticides such as *Bacillus sphaericus* (Su & Mulla 2004, Silva et al. 2008, Akiner et al. 2009) and *Bacillus thuringiensis* subspecies (Tabashnik et al. 1990, Gao et al. 2010, Tabashnik & Carriere 2010, Alcantara et al. 2011, Gassmann et al. 2011). *B. thuringiensis* subsp. *israelensis* (Bti) is increasingly used worldwide for mosquito control and investigating potential *Bti*-resistance in the environment is essential to ensure the long term use of this insecticide. Resistance to *Bti* has been extensively searched in natural populations of mosquitoes (Liu et al. 2004, Vasquez et al. 2009, Loke et al. 2010, Kamgang et al. 2011) and generally concluded in a lack of *Bti* resistance in field populations even after decades of *Bti* treatments, although a few studies described a decreased susceptibility in some populations (Zhang et al. 2004, Paul et al. 2005, Boyer et al. 2007, 2012). Nevertheless, these studies may suffer from technical biases, notably on the choice of the reference strain.

Usually, when bioassays are performed, a laboratory strain from the same species, susceptible to the insecticide tested, is used as a reference to calculate the resistance ratio of the strains/populations tested. Although this approach is widely used, it is adapted only if the reference strain has been sampled recently in populations geographically close to those tested. If this is not the case, results can reflect a basal difference of tolerance between the field populations and the reference strain, independent from *Bti*-treatments, rather than a selected resistance evolving in the populations treated with *Bti*. 

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GT current address: Department of Entomology, Cornell University, New York State Agricultural Experiment Station, Geneva, NY, USA  
* Corresponding author: guillaume.tetreau@gmail.com  
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Production of individual Bti Cry toxins - A crystal negative strain of Bti (4Q2-81) transformed with the plasmids pHT606, pHT618 or pWF53 was used for the production of Cry4Aa, Cry4Ba or Cry11Aa toxins respectively. These strains were obtained from the Pasteur Institute (Paris, France) or from Prof. B Federici (University of California, Riverside, CA, USA). Each toxin was produced, purified and quantified as previously described (Tetreau et al. 2012b). Toxin suspensions were stored at -20°C until use.

Bioassays procedure and data analyses - To avoid any influence of the larval stage on bioassays, larvae were maintained in the water from the breeding site where they were sampled until the third instar. Considering that the species studied cannot reproduce in laboratory conditions, we could not perform bioassays on F1 larvae, but directly used field-collected larvae. In order to limit the impact of environmental parameters in the sampled sites (e.g. pH of the water, pollutants etc.) on bioassay results, field collected larvae were kept in fresh water for 24 h before performing the bioassays.

When more than 2,000 larvae of the same species were sampled, standard bioassays were conducted by exposing larvae to five-seven concentrations of Cry4Aa, Cry4Ba, Cry11Aa and commercial Bti (Vectobac WG, 3500 ITU/mg) for 48 h in order to obtain 5-95% mortality. When 1,000-2,000 larvae were sampled, high sensitive standard bioassays could be performed, but not for all the Cry toxins; bioassays with Bti and Cry4Aa were performed in priority since Cry4Aa is the main toxin persisting in the field after Bti treatment (Tetreau et al. 2012a) and with the highest resistance ratio in the Bti-

Materials and methods

Mosquito sampling and species identification - Mosquito larvae were sampled in 17 different breeding sites of the French Rhône-Alpes region in close collaboration with the Entente Interdépartementale pour la Démosuccion (EID) (Fig. 1). Among them, 12 sites were regularly treated with Bti for decades and five had never been treated with Bti. Each sampling site is identified by the letter “P” followed by a number. Each site can be associated to different “P” numbers corresponding to different sampling dates. As mosquito breeding sites often contained more than one mosquito species, different species sampled at the same time in the same site were associated to the same “P” number.

As numerous different species can share the same breeding site, larvae were individually identified before bioassays using the identification key provided in Becker et al. (2010). Bioassays with different species were performed independently.

Fig. 1: localisation of the 32 mosquito breeding sites visited in the French Rhône-Alpes region. The 22 treated (triangles) and the 10 untreated (circles) sites visited are shown. Sampled sites used for bioassays (12 for treated and 5 for untreated sites) are indicated by a filled item and by the name of the population in italic (untreated site) or not (treated site).
resistant strain described in Tetreau et al. (2012b). Lethal concentrations for 50% individuals (LC\textsubscript{50}) were calculated for each population and each toxin using a probit statistical model with the module “dose” of XLSTAT v.2009.4.06 (Addinsoft).

When only a limited number of larvae of the same species were sampled, standard bioassays could not be performed, as a reduced number of replicates and/or concentrations performed would lead to a high variability in larval mortality and to a low reliability of the LC\textsubscript{50} values. In this case, a “diagnostic dose” approach was used, consisting in using one concentration of Cry toxin or Bti chosen in order to obtain 60-80% mortality. These doses were designed for each species and each toxin from the bioassays data from the first population sampled. For each toxin and Bti, statistical differences in larval mortality between two populations from the same species were measured by a Wilcoxon test performed with R 2.8.1 software (R Development Core Team 2007).

For each toxin tested, four replicates per concentration was performed on 20 third-instar larvae in 50 mL of tap water containing the insecticide according to the standard bioassay procedure described by the World Health Organization (WHO 2005). For each population tested, five negative controls, consisting in 20 larvae exposed to tap water only, were performed. All bioassays were performed at 17ºC. For each population, the mean larval size was calculated from more than 15 larvae.

Significance of larval size and mortality differences between the three species (Aedes rusticus, Aedes sticticus and Aedes vexans) were estimated by performing a Wilcoxon test using R 2.8.1 software (R Development Core Team 2007).

Considering that the species studied cannot be maintained in laboratory, we were unable to compare the tolerance levels of the sampled populations to a susceptible laboratory strain. Therefore, for each species we compared populations of mosquitoes with each other. Then, we calculated “tolerance ratios” which represent an increased or decreased susceptibility of a population relative to the others.

**RESULTS AND DISCUSSION**

**Limits of performing bioassays on field populations** - Among the 71 natural mosquito populations sampled, data from 59 populations (83%) were not analysable due to insufficient larvae to perform bioassays (69%), to mortality in controls (11%) or to high rate of pupation during bioassays (3%) (Table I). No bioassay data could be obtained for Aedes cantans, Aedes cataphylla, Aedes cinererus and Culex pipiens field populations in the present study, while sufficient data were obtained for Ae. rusticus, Ae. sticticus and Ae. vexans populations (Table I). Among the three Ae. rusticus populations tested in bioassays, two were sampled in treated sites (P3 and P5) and one in an untreated site (P14) while Ae. sticticus (P16, P20 and P52) and Ae. vexans (P18, P20, P30, P42 and P44) populations were all sampled in treated sites. Only one population of Culiseta annulata (P51) was sampled in sufficient amount to perform a diagnostic dose approach and showed comparable mortality rates as Ae. vexans populations at the same doses (data not shown).

The fact that only less than 17% of the populations sampled and only four out of the eight species sampled gave analysable results highlights the difficulty to find sites producing enough calibrated larvae of the same species to perform bioassays. This limit is mainly due to typical features of the Rhône-Alpes region, constituted of numerous sites of moderate size largely scattered in woodlands.

**Interspecific comparisons -** Ae. sticticus and Ae. vexans larvae exhibited the same patterns of tolerance (Wilcoxon test, p > 0.05) except for Cry4Aa for which

### Table I

Summary of mosquito breeding sites sampled where at least one of the eight studied species were found

| Species            | Sampled | Bioassays | Not analysable populations |
|--------------------|---------|-----------|----------------------------|
|                    |         | LC\textsubscript{50} | Diagnostic dose | Low sample size | High mortality in controls | Pupation |
| Aedes cantans      | 11      | 0         | 0                          | 9              | 2                          | 0        |
| Aedes cataphylla   | 5       | 0         | 0                          | 5              | 0                          | 0        |
| Aedes cinererus     | 3       | 0         | 0                          | 1              | 1                          | 1        |
| Aedes rusticus      | 15      | 3         | 0                          | 12             | 0                          | 0        |
| Aedes sticticus     | 14      | 3         | 0                          | 8              | 3                          | 0        |
| Aedes vexans        | 12      | 3         | 2                          | 6              | 0                          | 1        |
| Culex pipiens       | 6       | 0         | 0                          | 4              | 2                          | 0        |
| Culiseta annulata   | 5       | 0         | 1                          | 4              | 0                          | 0        |
| Total              | 71      | 9         | 3                          | 49             | 8                          | 2        |

for each species, the number of populations for which bioassays were performed is indicated. The main reason why bioassays were not performed for the other populations is also indicated. LC\textsubscript{50}: lethal concentrations for 50% individuals.
Aedes vexans larvae were 4.6-fold more tolerant than Aedes sticticus larvae (Wilcoxon test, p < 0.05) (Table II). Aedes rusticicus was significantly 3.9-fold–18.5-fold more tolerant than Aedes sticticus and Aedes vexans species when tested with Cry toxins and commercial Bti (Wilcoxon test, p < 0.05) (Table II). Aedes sticticus and Aedes vexans had similar larval size (6.3 and 6.2 mm; Wilcoxon test, p > 0.05) while Aedes rusticicus larvae were significantly 70% larger than larvae from other species (10.6 mm; Wilcoxon test, p < 0.05) (Table II). Correlating larval size with Bti toxicity tolerance indicated that the higher tolerance of Aedes rusticicus larvae compared to the two other species is due to the size differences (Supplementary data), linked to its particular life cycle containing a long winter larval period. This induces a slow larval development and metabolism that could lead to an alteration of feeding behaviour, therefore reducing the quantity of Bti ingested that could partly explain the lower susceptibility to Bti observed for this species. These results are congruent with observations made by the local mosquito control agency EID that Aedes rusticus requires a higher dose of Bti than other Aedes species for efficient control.

**Intraspecific comparisons** - Among the populations analysed, no increased tolerance to Bti was found in Aedes rusticus, Aedes sticticus and Aedes vexans (Figs 2-5). This result is congruent with the observations of the EID who reported no decreased efficacy of Bti treatments in the treated sites over time and also with previous works conducted in other Bti-treated areas (Liu et al. 2004, Vasquez et al. 2009, Loke et al. 2010, Kamgang et al. 2011).

The untreated P14 population of Aedes rusticus was significantly 1.77-fold more tolerant than treated P3 and P5 populations to Cry4Aa, but 1.72-fold less tolerant than P5 to Cry11Aa (Fig. 2). No significant differences between these populations were found when tested with Cry4Ba and Bti (Fig. 2).

The tolerance of Aedes vexans populations for which full bioassays were performed did not show significant variations for Cry4Aa, Cry4Ba and Bti while the P44 population was 2.5-fold more tolerant than the P20 population to Cry11Aa (Fig. 3). P30 and P42 populations, for which the diagnostic doses approach was used, did not show any significant differential tolerance to Cry4Ba, Cry11Aa and Bti (Fig. 4).

Bioassays with individual Cry toxins being more sensitive than bioassays with the whole Bti toxins mixture, slight changes in tolerance were observed to Cry4Aa and Cry11Aa in the P14 population of Aedes rusticus and to Cry11Aa in the P44 population of Aedes vexans.
Even if these changes were significant, tolerance ratios were low (less than 2.5-fold) possibly due to natural tolerance variations between populations rather than to an increased resistance selected by insecticide treatments.

The *Aedes sticticus* population P52 was significantly more tolerant to Cry4Aa and Cry11Aa toxins compared to P16 and P20 populations (mean of the two populations), while no significant difference was found with *Bti* (Fig. 5). This trend is similar to the resistance pattern observed for the resistant *Aedes aegypti* strain selected in the laboratory with field-persistent *Bti* sampled in the Rhône-Alpes region which exhibited a moderate level of resistance to *Bti* (3.5-fold), but a higher resistance to individual Cry toxins (up to 60-fold) (Tetreau et al. 2012b). It is therefore possible that even if no difference of tolerance to *Bti* was detectable in the three populations of *Ae. sticticus*, a cryptic resistance may be developing in the P52 population, only detectable with bioassays using each Cry toxins separately. Moreover, these fluctuations could be partly explained by the environmental parameters in the collected sites (e.g. pH of the water, pollutants etc.) that could affect the susceptibility of the larvae, for example by inducing the production of detoxification enzymes or metabolites that may interfere with larval response to *Bti* toxins (Riaz et al. 2009, Poupardin et al. 2012). These effects were limited by keeping larvae 24 h in fresh water before performing the bioassays. This regularly *Bti*-sprayed breeding site was located behind a deserted windmill, close to both treated and untreated sites. No obvious characteristic, in terms of vegetation, ecological corridors and insecticide treatments strategy could explain the increased tolerance to *Bti* toxins in this particular site. Regular bioassays with *Bti* and individual toxins will confirm if tolerance of this population is increasing along with *Bti* treatments.

**Fig. 4:** differential mortality of P30 and P42 *Aedes vexans* populations to Cry4Ba (16.3 ng/mL), Cry11Aa (32.5 ng/mL) and commercial *Bacillus thuringiensis* subsp. *israelensis* (*Bti*). Error: standard error. 600 larvae were sampled from P30 and P42 populations. Approximate distance between the sites: P30-P42, 66 km. NS: not significant (Wilcoxon test).

**Fig. 5:** lethal concentrations for 50% individuals (LC$_{50}$) of P16, P20 and P52 *Aedes sticticus* populations to Cry4Aa, Cry4Ba, Cry11Aa toxins and commercial *Bacillus thuringiensis* subsp. *israelensis* (*Bti*). Error: 95% confidence intervals calculated with the probit statistical analysis. 1060, 1120 and 2280 larvae were sampled from P16, P20 and P52 populations, respectively. Approximate distance between the sites: P16-P20, 45 km, P16-P52, 58 km and P20-P52, 14 km. Asterisks mean p < 0.05. NS: not significant.
could be detected with classical bioassays with Bti. This hypothesis has to be confirmed by monitoring the evolution of the tolerance of the populations tested in this study for several years. In combination with bioassays with Bti, bioassays with separate Cry toxins allow a more sensitive monitoring of the susceptibility of field-mosquito populations to Bti. Therefore, we believe that such approach should be used in further studies looking for Bti-resistance in other species from the Rhône-Alpes region (e.g. Aedes cantans), from the Mediterranean region (e.g. Aedes detritus, Aedes caspius) and should also be applied to human pathogens vector species such as Aedes aegypti, Aedes albopictus and Cx. pipiens exposed to regular Bti treatments.

Bti is widely used for mosquito control. Based on observations of the resistance phenotype of a Bti-resistant laboratory strain, we developed a new tool: bioassays with individual Bti Cry toxins. This highly sensitive tool could be used in combination with bioassays with Bti in order to improve the monitoring of the susceptibility of field-populations of mosquitoes regularly treated with Bti.

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