



Molecular characterization of the gene profile of *Bacillus thuringiensis* Berliner isolated from Brazilian ecosystems and showing pathogenic activity against mosquito larvae of medical importance



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ABSTRACT

The occurrence of *Aedes aegypti*, *Culex quinquefasciatus*, and mosquitoes of the genus *Anopheles* potentiate the spread of several diseases, such as dengue, Zika, chikungunya, urban yellow fever, filariasis, and malaria, a situation currently existing in Brazil and in Latin America. Control of the disease vectors is the most effective tool for containing the transmission of the pathogens causing these diseases, and the bacterium *Bacillus thuringiensis* var. *israelensis* has been widely used and has shown efficacy over many years. However, new *B. thuringiensis* (*Bt*) strains with different gene combinations should be sought for use as an alternative to *Bti* and to prevent the resistant insects selected. Aiming to identify diversity in the *Bt* in different Brazilian ecosystems and to assess the pathogenicity of this bacterium to larvae of *Ae. aegypti*, *C. quinquefasciatus*, and *Anopheles darlingi*, *Bt* strains were obtained from the Amazon, Caatinga (semi-arid region), and Cerrado (Brazilian savanna) biomes and tested in pathogenicity bioassays in third-instar larvae of *Ae. aegypti* under controlled conditions in the laboratory. The isolates with larvicidal activity to larvae of *Ae. aegypti* were used in bioassays with the larvae of *C. quinquefasciatus* and *An. darlingi* and characterized according to the presence of 14 *cry* genes (*cry1*, *cry2*, *cry4*, *cry10*, *cry11*, *cry24*, *cry32*, *cry44Aa*, *cry1Ab*, *cry4Aa*, *cry4Ba*, *cry10Aa*, *cry11Aa*, and *cry11Ba*), six *cyt* genes (*cyt1*, *cyt2*, *cyt1Aa*, *cyt1Ab*, *cyt2Aa* and *cyt2Ba*), and the *chi* gene. Four hundred strains of *Bt* were isolated: 244 from insects, 85 from Amazon soil, and 71 from the Caatinga biome. These strains, in addition to the 153 strains isolated from Cerrado soil and obtained from the Entomopathogenic Bacillus Bank of Maranhão, were tested in bioassays with *Ae. aegypti* larvae. A total of 37 (6.7%) strains showed larvicidal activity, with positive amplification of the *cry*, *cyt*, and *chi* genes. The most frequently amplified genes were *cry4Aa* and *cry4Ba*, both occurring in 59.4% in these strains, followed by *cyt1Aa* and *cyt2Aa*, with 56.7% and 48% occurrence, respectively. Twelve (2.2%) strains that presented 100% mortality within 24 h were used in bioassays to estimate the median lethal concentration (LC₅₀) for *Ae. aegypti* larvae. Two strains (BtMA-690 and BtMA-1114) showed toxicity equal to that of the *Bti* standard strain, and the same LC₅₀ value (0.003 mg/L) was recorded for the three bacteria after 48 h of exposure. Detection of the presence of the *Bt* strains that showed pathogenicity for mosquito larvae in the three biomes studied was possible. Therefore, these strains are promising for the control of insect vectors, particularly the BtMA-1114 strain, which presents a gene profile different from that of *Bti* but with the same toxic effect.

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1. Introduction

Diseases whose etiological agents are transmitted by mosquito vectors are among the major diseases affecting humans (WHO, 2017). Considering Latin America, among the species of greater epidemiological importance is *Aedes (Stegomyia) aegypti* (Diptera: Culicidae) (Linnaeus, 1762), it is the main vector of the Zika, dengue, chikungunya, and urban yellow fever viruses (Honório et al., 2015; Ebi and Nealon, 2016; Ferreira-de-Brito et al., 2016).

Culex quinquefasciatus Say, 1823 (Diptera: Culicidae) is another species of mosquitoes of importance to public health in the Americas. It is the vector of lymphatic filariasis, a disease of a chronic nature that mainly affects populations of low socioeconomic level (Brasil, 2016; Rebollo and Bockarie, 2017).

The species *Anopheles darlingi* Root 1926 is the main vector of malaria in America, mainly in the Amazon region, which recorded more than 83% of the cases (Siqueira et al., 2017; Tadei et al., 2017).

There are epidemic cycles of these diseases in the Latin America, it is necessary to seek ways to control. Several obstacles to mosquito control exist, with resistance to chemical agents being noted as one of the main challenges to the current vector control program (Moyes et al., 2017; Seixas et al., 2017).

The use of the bacterium *Bacillus thuringiensis* (Bt) Berliner, 1915 is one of the biological control strategies that has been showing better efficacy (Bravo et al., 2011; Lacey et al., 2015). *Bacillus thuringiensis* var. *israelensis* (Bti) is a natural enemy of several species of mosquitoes of the genera *Culex* Linnaeus, 1758; *Aedes* Meigen, 1818; and *Anopheles* Meigen, 1818. It is the microbial agent more commonly used in the control of these insects, and its continued use for more than 30 years has not resulted in the evolution of resistance in mosquito populations treated in different regions of the world (Bravo et al., 2011; Stalinski et al., 2016).

The lack of resistance is attributed to the complex mechanism of action of Bti, which has toxins known as Cry toxins (4Aa, 4Ba, 11Aa, 11Ba, and 10Aa) that are capable of interacting with the intestinal epithelium of the mosquito larvae and also has cytolytic (Cyt) toxins that are less specific but facilitate the insertion of Cry toxins into the intestinal epithelium and may thus increase insecticidal activity (Bravo et al., 2007; Ben-Dov, 2014; Zhang et al., 2016). The synergism between the Cry and Cyt toxins is fundamentally important for the efficacy of the bacterium (Frankenhuyzen, 2013; Zhang et al., 2016).

Some Bt strains also produce vegetative insecticidal proteins (Vips) produced in vegetative phase and chitinolytic toxins (Chi), the toxins chitinolytics are another group of toxins that may contribute to larval mosquitoes mortality by destroying the peritrophic matrix of insects (Sampson and Gooday, 1998; Djenane et al., 2017).

Although no records exist of resistance to Bti, the possibility of the select of resistant populations cannot be discounted. This bacterium is an important biological agent; therefore, the use of other strains with different combinations of cry and cyt genes is necessary as a form of management and prevention of resistance to Bti (Cánton et al., 2015; Peralta and Palma, 2017).

The diversity of the Cry toxins already found and described in the literature demonstrates the possibility of discovery of different combinations of Bti with different insecticidal potentials (Crickmore, 2017).

Several studies have sought to obtain more *B. thuringiensis* isolates with insecticidal potential for mosquitoes, which is done by isolating native strains from substrates such as soils from different ecosystems, dead insects, plants, and other sources. In addition, these strains are investigated at the molecular level by detecting the genes encoding the Cry and Cyt toxins present in the toxic crystal, which makes predictions of their insecticidal activity possible (Bravo et al., 1998; Jozani et al., 2008; Costa et al., 2010; El-kersh et al., 2016).

The present study investigated the diversity of Bt strains isolated from soils of different Brazilian biomes and from dead insects, and showing larvicidal activity against mosquitoes *Ae. aegypti*, *Cx*

quinquefasciatus and *An. darlingi* in the laboratory which are the main mosquitoes of medical importance in Latin America.; in addition, the gene profiles of the strains pathogenic.

2. Methods

2.1. Sampling and isolation of *Bacillus thuringiensis*

A total of 37 soil samples from two biomes (15 from the Caatinga biome and 22 from the Amazon biome) and 44 samples from dead insects were processed according to the World Health Organization (1985) protocol for the isolation of Bt strains.

The soil samples consisted of 10 g of soil, which was collected at a depth of 5 cm, placed in sterile flasks, and sent to the Laboratory of Medical Entomology (LABEM) at the Universidade Estadual do Maranhão – UEMA.

The insect samples consisted of 44 dead insects collected in the Cerrado biome. The insects were identified as belonging to the orders Coleoptera (22), Hymenoptera (15), and Hemiptera (07). All samples were collected in the state of Maranhão, Brazil, which contains the three biomes (SISBIO/59840; IBGE, 2017).

2.2. Morphological identification of *Bacillus thuringiensis* isolates

The strains were cultured in nutrient agar (peptic digest of animal tissue 5 g/L, sodium chloride 5 g/L, meat extract 1.5 g/L, and yeast extract 1.5 g/L pH 7.4 ± 2) containing penicillin G (100 mg/L) for 48 h; then, and viewed at 1000 x magnification under an Axio Scope A.1 (Zeiss) microscope by phase-contrast to detect the presence of crystal for differentiation from *Bacillus cereus*. The Bt strains were submitted the gram-staining test (Jung et al., 1998).

The strains the Bt were stored at 4 °C in filter-paper strips, impregnated with spore suspension, immersed in autoclaved distilled water, and stored at 4 °C in triplicate. The strains were individually identified with BtMA (*Bacillus thuringiensis* from Maranhão) followed by the number corresponding to the order of isolation and deposited in the Entomopathogenic Bacillus Bank of Maranhão (BBENMA), located in the municipality of Caxias, Maranhão, Brazil.

2.3. Selection of strains for mosquito pathogenicity assays

For preliminary screening of strain pathogenicity to *Ae. aegypti*, 553 strains of Bt, of which 400 were obtained in this study and 153 were isolated from Cerrado soil and kept at BBENMA, were used after pre-selection of the strains for larvicidal activity. The bioassay was performed in triplicate: three plastic cups containing 10 mL of drinking distilled water and 10 third-instar *Ae. aegypti* larvae were used, with 1 mL of the total bacterial culture being added to each. The negative control consisted of 10 larvae placed in a plastic cup with water, but without inoculation with the bacterium; for the positive control, Bti T04001 lyophilized was used under the same conditions. The bioassays were conducted at the LABEM, UEMA, with a temperature of 26 ± 2 °C, relative humidity of 80% and photoperiod of 12 h light followed by 12 h dark (12L:12D) (WHO, 2005).

The strains that reached 100% mortality in up to 48 h, along with the Bti standard strain, were grown in NYSM medium incubated at 28 °C for 5 days for complete sporulation and release of the crystal proteins. After, the samples were centrifuged at 1700xg for 15 min at 4 °C, and the pellet was recovered and transferred to Falcon tubes with 10 mL of autoclaved distilled water and 0.01% Triton® X-100. The spores were then counted using a Neubauer chamber in an Axio Scope A.1 (Zeiss) phase-contrast optical microscope (Alves and Moraes, 1998). The strains were tested again for *Ae. aegypti* larvae at the standard concentration of 1.5 × 10⁷ spores/mL under the same abovementioned conditions.

Strains that killed 100% of the larvae in 48 h were cultured in

600 mL of nutrient yeast extract salt medium (NYSM) (Yousten, 1984) for 5 days at 28 °C at 180 rpm. The culture obtained was centrifuged at 10.000 x g for 30 min at 4 °C, washed with autoclaved distilled water, frozen, and lyophilized for approximately 16 h (Santos et al., 2012).

2.4. Pathogenicity bioassays in *Culex quinquefasciatus* and *Anopheles darlingi* larvae

Strains that killed 100% of the *Ae. aegypti* were selectively tested at a concentration of 10 mg/L against *C. quinquefasciatus* and *An. darlingi* larvae. The larvae of the F1 generation, obtained from adults collected in the field and eggs reared in the laboratory (SISBIO/21264-3). Collection was carried out in the city of Caxias, Maranhão for *C. quinquefasciatus* and in Manaus, Amazonas for *An. darlingi*. The bioassays followed the same conditions of the tests described above for *Ae. aegypti*.

2.5. Bioassays for estimating the lethal concentration (LC₅₀) and (LC₉₀) for *Aedes aegypti* larvae

These bioassays were performed with the 12 strains that showed 100% mortality in 24 h for *Ae. aegypti*. Toxicity bioassays were performed in the laboratory with *Ae. aegypti* due to the ease of obtaining this species.

For each strain, six concentrations (0.04, 0.03, 0.02, 0.01, 0.008 and 0.005 mg/L) were initially tested, from which more concentrations (0.09–0.001 mg/L) were established to obtain a mortality of 100% and 5% for each strain.

Each concentration was tested in three replicates, was performed in the different days. Each replicate used five plastic cups containing a final volume of 150 mL and 20 third-instar larvae, and the amount corresponded to each concentration of the isolates. For each bioassay, a negative control group was added, which consisted of a cup with larvae and without inoculation with the bacillus. The bioassays were monitored at intervals of 24, 48, and 72 h after the application of the bacillus, with dead larvae being counted at each interval.

To compare the larvicidal activity of the isolates, *Bti* T04001 was used. The bioassay was performed under the same conditions described for the other isolates and in accordance with the recommendations Guidelines For Laboratory And Field Testing Of Mosquito Larvicides (WHO, 2005).

2.6. Lethal concentration (LC₅₀) and (LC₉₀) and statistical analyses

The mortality data obtained in the toxicity bioassays were submitted to Probit analysis at p = 0.05 (Finney, 1971), with the statistical software POLO PLUS (LeOra Software, 2003) being used to estimate the LC₅₀ and LC₉₀.

The toxicity of each isolate was compared to that of the standard strain *Bti* T04001 using Student's *t*-test when the data were parametric or the Mann-Whitney test for non-normal data; the level of significance was set as 5% (α = 0.05). The statistical program used was BioEstat 5.0 (summer) for Windows (Ayres et al., 2007).

2.7. Molecular characterization

The strains of *Bt* with larvicidal activity were investigated for the presence of the *cry* (14), *cyt* (6), and *chi* genes encoding toxins active against mosquito larvae using polymerase chain reaction (PCR). Eleven universal primers—*cry1*, *cry2*, *cry4*, *cry10*, *cry11*, *cry24*, *cry32*, *cry44*, *cyt1*, *cyt2* and *chi*—and 10 specific primers—*cry1Ab*, *cry4Aa*, *cry4Ba*, *cry10Aa*, *cry11Aa*, *cry11Ba*, *cyt1Aa*, *cyt1Ab*, *cyt2Aa* and *cyt2Ba*—were used for gene amplification (Table 1). The following strains were used as positive controls: *B. thuringiensis* var. *aizawai* (XenTari-WDG; *cry1*, *cry1Ab*), *B. thuringiensis* var. *kurstaki* (Dipel WP; *cry2*), *B. thuringiensis* *sotto* (T03 001; *cry24*), *B. thuringiensis* var. *yunnanensis* (T20 001; *cry32*),

Table 1
lists the studied *B. thuringiensis* genes with the respective primers used and the expected fragment sizes from PCR as well as the annealing temperature.

Gene	Sequence of primers (5'–3')	TF of pb	TM
<i>cry1</i> ¹	CTGGATTTACAGGTGGGGATAT (f) TGAGTCGCTTCGCATATTTGACT (r)	543–594	52
<i>cry1Ab</i> ²	AAGCAAGGGTTATTACATTACG (f) CCAATACTAAGATCAGAGGG (r)	~550	56
<i>cry2</i> ³	GTTATTCTTAATGCAGATGAATGGG (f) CGGATAAAAATAATCTGGGAAATAGT (r)	689–701	52
<i>cry4</i> ³	GCATATGATGTAGCGAAACAAGCC (f) GGGTGACATACCATTTCAGGTC (r)	439–459	52
<i>cry4Aa</i> ⁴	GAACCTGGGTATGGCACTCAAC (f) CTCACAACGATTAGACCCTTC (r)	777	50
<i>cry4Ba</i> ⁴	GCGAGGTTTCCCATGTCTAC (f) GTTGTAGGGTGGAAITGTTATC (r)	347	52
<i>cry10</i> ³	TCAATGCTCCATCCAATG (f) CTTGATAGGCCCTTCCTCCG (r)	348	51
<i>cry10Aa</i> ⁴	ATTGTTGGAGITAGTGCAAGG (f) AATACTTTGGATGTGTCTTGAG (r)	995	50
<i>cry11</i> ¹	TTAGAAGATACGCCAGATCAAGC (f) CATTGTACTTGAAGTTGTAATCCC (r)	305	51
<i>cry11Aa</i> ⁴	AGGATGGATAGGAAACGGGAAG (f) CCGTATCCAGCAGGTAAGC (r)	470	50
<i>cry11Ba</i> ⁴	TACAGGATGGATAGGGAATGG (f) TAATACTGCCATCTGTGCTTG (r)	608	52
<i>cry24</i> ³	TTATCAATGTTAAGGGATGC (f) ACTGGATCTGTGTATATTTTCTAG (r)	304	48
<i>cry32</i> ³	TGGTCGGGAGAGAATGGATGGA (f) ATGTTTGGCACCATTTC (r)	676–677	54
<i>cry44Aa</i> ⁵	CATTACACGGGGTGCCTTAT (f) CCGCACCTTACATGTGTCCA (r)	444	60
<i>cyt1</i> ³	CCTCAATCAACAGCAAGGGTATT (f) TGCAAAACAGGACATTGTATGTGTAATT (r)	477–480	52
<i>cyt1Aa</i> ⁴	AACTCAAACGAATAACCAAG (f) TGTTCTTTACTGCTGATAC (r)	300	53
<i>cyt1Ab</i> ⁴	AAGCAAGGGTTATTACATTACG (f) CCAATACTAAGATCAGAGGG (r)	698	54
<i>cyt2</i> ³	ATTACAAAATTGCAAATGGTATTCC (f) TTTCAACATCCACAGTAATTTCAAATGC (r)	355–356	52
<i>cyt2Aa</i> ⁴	GCATTAGGAAGACCATTTG (f) AAGGCTAAGAGITGATATCG (r)	361	53
<i>cyt2Ba</i> ⁶	CAGGAACCTCTTAATCAAAGTGAAT (f) CATCTACTGTAGGTTCTAAATTTGT (r)	177	50
<i>chi</i> ⁷	ATGGTCATGAGGTCTC (f) CTATTCGCTAATGACG (r)	2027	45

Legend (f) = forward; (r) = reverse; Pb = base pair. ¹ Bravo et al. (1998), ² Fatoretto et al. (2007); ³ Jouzani et al. (2008), ⁴ Costa et al. (2010), ⁵ Vidal-Quist and Castañera (2009), ⁶ Costa et al. (2014), ⁷ Lin and Xiong (2004). Legend: TF = Fragment size, pb = pairs of base and TM = melting temperature.

B. thuringiensis entomocidus (T06a 001; *cry44*) and *B. thuringiensis* var. *israelensis* (T04 001) for the remaining genes, provided by the Laboratory of Bacterial Genetics and Applied Biotechnology (LGBBA) of the School of Agrarian and Veterinary Sciences, São Paulo State University, Universidade Estadual Paulista – FCAV/UNESP Jaboticabal).

2.8. Total DNA extraction and amplification reaction of mosquito-specific genes

Total DNA extraction from the strains was performed using the InstaGene Matrix DNA extraction kit (Bio-Rad, USA), according to the manufacturer's recommendations. One colony of each isolate was grown for approximately 12 h in nutrient agar was transferred to a microtube containing 1 mL of sterile Milli-Q water; this sample was then centrifuged for 1 min at 12,000 rpm and 20 °C. The supernatant was discarded; then, 200 µL of InstaGene Matrix (Bio-Rad) was added, and the material was incubated in a water bath at 56 °C for 25 min, after vortexed thoroughly for 10 s and then incubated at 100 °C for 8 min. The sample was again vortexed at moderate speed for 10 s and centrifuged at 20 °C for 2.5 min. The DNA which were stored in a freezer at –20 °C until the time of use.

PCR reactions were performed at a final volume of 25 μ L with: 1X GoTaq[®] Flexi DNA Polymerase buffer, 2.0 mM MgCl₂, 0.2 mM dNTPs (Promega), 0.2 pmol/ μ L of each primer (Invitrogen), 1U of GoTaq[®] DNA Polymerase enzyme (Promega), and 50 ng of DNA. The PCR was performed for all genes mentioned above; the reaction was performed in a Gencycler-G96G thermocycler (Biosystems). The general amplification conditions were programmed according to the following specifications: 94 °C for 5 min; followed by 30 cycles at 94 °C for 30 s, with the annealing temperature optimized according to each primer (Table 1), and final extension at 72 °C for 5 min. For amplification of the *chi* (chitinase) gene, the program was as follows: 5 min at 94 °C; 30 cycles of 1 min at 94 °C, 1 min at 45 °C, and 1.5 min at 72 °C; and 10 min at 72 °C for a final extension (Costa et al., 2010).

Amplification products were analyzed by electrophoresis in a 1,5% agarose gel with current applied at 90 V in 1X TBE (Tris/Borate/EDTA) buffer with alkaline pH and photographed in an L-PIX device (Loccus Biotechnology).

3. Results

3.1. Isolation of *Bacillus thuringiensis*

Four-hundred strains of *Bt* were obtained, 244 from insects, 85 from the Amazon soil, and 71 from the Caatinga soil. The Amazon soil was the substrate with the highest isolation rates, with 72.7% of the samples presenting *Bt*, followed by the Caatinga soil with 53.3% and then dead insects with 43.2%.

However, the highest mean occurrence of *Bt* per sample was observed for dead insects, with 5.5 strains, whereas for soil samples, the rates were 4.7 for the Caatinga and 3.9 for the Amazon.

In the insects, Hymenoptera was the group with the highest number of *Bt* strains, with 193 strains and a mean of 12.9 for sample, a value three times higher than that found for the other orders, Coleoptera and Hemiptera, which had 100 and 21 strains and means of 4.5 and 3, respectively.

3.2. Selection bioassays of *Bacillus thuringiensis* strains pathogenic for mosquitoes

From the total of 553 strains of *Bt* selectively tested against *Ae. aegypti*, 37 (6.7%) showed pathogenicity, 12 of which killed 100% of the larvae in 24 h (Table 2). The remaining 25 strains showing pathogenicity reached 100% mortality after 48 h.

Among the 37 pathogenic strains, approximately one-half (47.7%) were obtained from the Cerrado soil. The rate of strains from this biome that are active against mosquitoes was 11.11% of the total number of strains tested, three times higher than that obtained for the Caatinga biome, for which the lowest rate (3.5%) was found.

3.3. Molecular characterization

In all 37 strains that showed pathogenic activity, positive amplification occurred for the genes tested using the universal and specific primers (Table 3).

The PCR reactions showed positive amplification of all genes studied; however, variation was observed in the number of *cry* and *cyt* genes per strain. Twelve different gene combinations were observed, with three strains (BtMA-37, BtMA-626, and BtMA-215) containing a single gene, whereas five strains (BtMA-676, BtMA-684, BtMA-685, BtMA-688, and BtMA-690) amplified fragments with the expected size for 15 genes, thus confirming the presence of six families of genes active against Diptera (*cry4*, *cry10*, *cry11*, *cyt1*, *cyt2*, and *chi*) and the presence of nine genes specific to mosquitos. This profile was similar to the one obtained with the *Bti*-T4001 standard strain, for which positive amplification was also observed for the nine genes encoding toxins active against mosquitoes and for the *chi* gene. The BtMA-679 and 687

strains, in addition to the nine *Bti* genes, showed amplification for the *cry32* gene (Table 3).

The *cry4Aa* and *cry4Ba* genes had a higher occurrence rate in the strains (59.4%), followed by the *cyt1Aa* and *cyt2Aa* genes, which were present in 56.7% and 48% of the strains, respectively. Next, in decreasing order of frequency, were *cry10Aa* (45%), *cyt1Ab* (43.24%), *cry11Aa* and *cyt2Ba* (37.8%), *chi* (35.3%), *cry11Ba* (32.4%), *cry32* (27%), *cry1* (8.1%), and *cry44Aa* (2.7%). No positive amplification was found for the *cry1Ab*, *cry2*, and *cry24* genes.

3.4. Bioassays for estimating the lethal concentration (LC₅₀) and (LC₉₀)

The toxicity bioassays were performed with the 12 strains of *Bt* that achieved 100% mortality after 24 h. Based on the obtained values of LC₅₀ and LC₉₀ and the respective confidence intervals, four toxicity groups were formed. The most toxic strains were BtMA-690, BtMA-1114, and the *Bti* T4001 standard strain, followed by the group composed of BtMA-679, BtMA-687, and BtMA-688, and then the strains BtMA-37, BtMA-681, BtMA-684, BtMA-685, BtMA-689, and 703. The lowest toxicity was observed for the BtMA-691 strain, which showed the highest values for LC₅₀ and LC₉₀ (Table 4).

The quantitative bioassays showed that two strains (BtMA-690 and BtMA-1114) had similar performance to the *Bti* standard strain, for which no significant difference between the LC₅₀ values was observed in the three evaluation periods (ANOVA: F = 16, p = 0.06). In the 24-h evaluation, BtMA-690 and *Bti* obtained the same LC₅₀ value of 0.003 mg/L; however, the LC₉₀ was 0.009 mg/L and 0.014 mg/L, respectively. For BtMA-1114, the LC₅₀ was 0.004 mg/L, and the LC₉₀ was 0.008 mg/L (Table 4).

In the period (48 h), BtMA-690, BtMA-1114, and the *Bti* T4001 obtained the same value of LC₅₀ (0.003 mg/L), but the LC₉₀ was 0.009 mg/L for the two isolates and 0.011 mg/L for the standard strain. At the evaluation after 72 h, the LC₅₀ value was 0.001 mg/L for *Bti* and 0.002 and 0.003 mg/L for BtMA-690 and BtMA-1114, respectively, whereas the LC₉₀ was 0.005 mg/L for BtMA-690 and 0.006 mg/L for the other two strains (Table 4).

Based on the LC₅₀ values obtained at the three evaluation periods, the less toxic isolates (BtMA-37 and BtMA-691) showed LC₅₀ values approximately 20 times higher than the BtMA-690, BtMA-1114, and *Bti* standard strains, whereas the groups BtMA-679, BtMA-687, and BtMA-688 were five times less toxic than *Bti*.

4. Discussion

The bacterium *Bt* is found in all environments, but soil has been the most used source of isolation (Polanczyk et al., 2004; Gobatto et al., 2010; El-Kersh et al., 2016; Reyaz et al., 2017). In the present work, in which soil samples and insects were analyzed, the insects were the substrates with the highest number of *B. thuringiensis* per sample, and the presence of the bacterium was detected in 56% of the insect samples used.

The presence of *Bt* in insects is generally high (Gobatto et al., 2010; Pinto et al., 2003; Assaeedi et al., 2011). These bacteria develop with these organisms, thus making them a natural source of the pathogen and making it possible to find new strains of this bacterium both in dead and live insects (Bernhard et al., 1997; Abulreesh et al., 2012 Abulreesh et al., 2012).

In the present work, detection of the bacterium was possible in three orders of insects, with Hymenoptera being the most promising, for which was found in 66% of the samples, corresponding to more than twice the percentage occurrence for the orders Coleoptera with 31.8% and Hemiptera with 28.5%.

Other studies have shown different results for *Bt* occurrence rates in insects, such as the order Coleoptera with 60% (Hernandez et al., 2005), whereas for Hymenoptera, 40% of the bacterial colonies obtained from two species of this order of insects were identified as *Bt*

Table 2

Bacillus thuringiensis isolates with 100% larvicidal activity for *Aedes aegypti* larvae at 24 and 48 h in the laboratory conditions according to the origin of the isolation substrate in municipality the isolated.

Biome	substrate	Larvicidal activity		Municipality	Latitude (S) Longitude (W)
		24 h	48 h		
Amazônia					
BtMA- 37	solo	100	–	Viana	S 03°13'12.3" W 045°08'88.7"
BtMA-179	solo		100	Santa Luzia	S 04°38'20.5" W 046°23'30.1"
BtMA-215	solo		100	Bela Vista	S 03°75'60.3"W 045°22'62.9"
BtMA-229	solo		100	Santa Inês	S 03°85'73.3"W 045°53'49.2"
BtMA-233	solo		100	Santa Inês	S 03°85'73.3"W 045°53'49.2"
BtMA-237	solo		100	Santa Inês	S 03°85'73.3"W 045°53'49.2"
BtMA-241	solo		100	Santa Inês	S 03°85'73.3"W 045°53'49.2"
Cerrado					
BtMA-459	solo		100	São J. dos Patos	S 06°50'37.5" W 043°68'65.8"
BtMA-527	solo		100	Benedito Leite	S07°22'55.5" W 044°55'97.2"
BtMA-559	solo		100	Balsas	S 07°53'53.3"W 046°03'91.1"
BtMA-626	solo		100	Coelho Neto	S 04°25'31.0" W 043°01'38.9"
BtMA-676	solo		100	Duque Barcelar	S 04°13'72.5" W 042°94'91.8"
BtMA-679	solo	100	–	Duque Barcelar	S 04°13'72.5" W 042°94'91.8"
BtMA-681	solo	100	–	Duque Barcelar	S 04°13'72.5"W 042°94'91.8"
BtMA-682	solo		100	Duque Barcelar	S 04°13'72.5" W 042°94'91.8"
BtMA-684	solo	100	–	Duque Barcelar	S 04°13'72.5"W 042°94'91.8"
BtMA-685	solo	100	–	Duque Barcelar	S 04°13'72.5"W 042°94'91.8"
BtMA-686	solo		100	Duque Barcelar	S 04°13'72.5"W 042°94'91.8"
BtMA-687	solo	100	–	Duque Barcelar	S 04°13'72.5" W 042°94'91.8"
BtMA-688	solo	100	–	Duque Barcelar	S 04°13'72.5" W 042°94'91.8"
BtMA-689	solo	100	–	Duque Barcelar	S 04°13'72.5" W 042°94'91.8"
BtMA-690	solo	100	–	Duque Barcelar	S 04°13'72.5" W 042°94'91.8"
BtMA-691	solo	100	–	Duque Barcelar	S 04°13'72.5" W 042°94'91.8"
BtMA-694	solo		100	Duque Barcelar	S 04°13'72.5" W 042°94'91.8"
Insetos					
BtMA-1054	Hymenoptera		100	Mirador	N° da Amostra 1
BtMA-1061	Hymenoptera		100	Mirador	1
BtMA-1107	Coleoptera		100	Mirador	2
BtMA-1108	Coleoptera		100	Mirador	2
BtMA-1109	Coleoptera		100	Mirador	2
BtMA-1114	Coleoptera	100	–	Mirador	3
BtMA-1115	Coleoptera		100	Mirador	3
BtMA-1116	Coleoptera		100	Mirador	3
BtMA-1119	Coleoptera		100	Mirador	3
BtMA-1120	Coleoptera		100	Mirador	3
BtMA-1134	Hymenoptera		100	Mirador	4
BtMA-1147	Hymenoptera		100	Mirador	5
Caatinga					
BtMA –703	solo	100		Santa Quitéria	S 03°48'34.6"W 042°55'59.8"

(Pinto et al., 2003).

The Caatinga biome showed a higher mean number of strains per samples compared to the Amazon soil when was isolated from the soil of the two biomes. The Caatinga, located in the northeastern region of Brazil, had already been reported as the region with the greatest abundance of *Bt* in the country (Silva et al., 2012), for which the presence of the bacteria was detected in 16.9% of the samples (Silva et al., 2002).

The number of strains of *Bt* per sample in the Amazon biome found in this study corroborates previous findings, in which even lower mean values 0.48 and 2.28 strains per sample were observed (Pereira et al., 2013; Soares-da-Silva et al., 2015). In contrast, *Bt* occurred in approximately 70% of the soil samples of this biome used for isolation, which demonstrates the wide distribution of this bacterium in this environment.

The persistence of *Bt* spores in the soil involves different factors, with the soil chemical constituents being suggested as one of the main factors affecting this persistence (Polanczyk et al., 2004). The number *Bt* of strains obtained from the different biomes varied considerably, the presence of this bacterium in all types of environments indicates that *Bt* must undergo intense selective pressure and, to survive, has developed different ways of resisting natural enemies (Habib and Andrade, 1998).

This is confirmed by the pathogenicity of *Bt* to different groups of insects (Frankenhuyzen, 2009; 2013).

About the pathogenicity of *Bt* strains active against mosquitoes, mortality, in general, is low compared to other groups of insects (Armengol et al., 2006; Gobatto et al., 2010). In the present study, 6.7% of the tested strains showed pathogenic activity against larvae of the three species, confirming the low occurrence of the mosquito-specific strains.

These studies carried out in Brazil with native strains of *Bt* active against mosquito larvae also showed low occurrence of these strains, rate below 2% of *Bt* active against mosquitoes (Dias et al., 2002; Praça et al., 2004; Ootani et al., 2011). In Saudi Arabia, 33.8% of the native strains isolated from distinct parts in that country showing pathogenicity for *Anopheles gambiae*, an important vector of African malaria (El-kerh et al., 2016).

Studies performed on Amazon soil showed similar rates to that found in the present study, 8.7%and 2% of strains were active against *Ae. aegypti*, Pereira et al. (2013) and Soares-da-Silva et al. (2015), respectively.

The variation in the number of native strains of *Bt* with larvicidal activity against mosquitoes observed for the different biomes can be explained by the different profiles of the insecticidal toxin-producing

Table 3
Gene profile of the 37 isolates of *Bacillus thuringiensis* active against *Aedes aegypti* larvae obtained from the three soil e insects Brazilian biomes.

Genes	Isolates																					
	cy 1	cry 1Ab	cry 2	cry 4	cy4Aa	cry4Ba	cry 10	cry10Aa	cry 11	cry11Aa	cry11Ba	cry1	cry1Aa	cry1Ab	cry2	cy2Aa	cy2Ba	chi	cry 24	cry 32	cry 44Aa	
<i>Bt israelensis</i>																						
<i>Bt azawai</i>	+	+																				
<i>Bt kurstaki</i>		+																				
<i>Bt sotto</i>																				+		
<i>Bt. entomocidius</i>																						
<i>Bt. yuannanensis</i>																						
Amazônia																						
BtMA-37					+																	
BtMA-179																						
BtMA-215																						
BtMA-229																						
BtMA-233																						
BtMA-237																						
Cerrado																						
BtMA-241																						
BtMA-459																						
BtMA-527																						
BtMA 626																						
BtMA-559																						
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BtMA-691																						
BtMA-694																						
Insetos																						
BtMA-1054																						
BtMA-1061																						
BtMA-1107																						
BtMA-1108																						
BtMA-1109																						
BtMA-1114																						
BtMA-1115																						
BtMA-1116																						
BtMA-1119																						
BtMA-1120																						
BtMA-1134																						
BtMA-1147																						
Caatinga																						
BtMA-703																						

Legend: + = presence and - = absence.

Table 4
Lethal Concentrations LC₅₀ and LC₉₀ in mg/L at the 24-, 48-, and 72-hour evaluations for *Bacillus thuringiensis* isolates pathogenic to mosquitos.

Isolados	LC ₅₀ (IC 95%)	LC ₉₀ (IC 95%)	Slope ± SE	χ ² (GL = 3)
24 h				
BtMA-37	0.063 (0.056–0.073)	0.158 (0.114–0.380)	3.184 (0.323)	6.3
BtMA-679	0.015 (0.012–0.019)	0.042 (0.029–0.082)	2.904(0.143)	14.5
BtMA-681	–	–	–	–
BtMA-684	0.078 (0.065–0.120)	0.161 (0.110–0.415)	4.047 (0.881)	2.4
BtMA-685	0.032 (0.021–0.038)	0.049 (0.040–0.103)	6.608 (0.428)	32.2
BtMA-687	0.017 (0.014–0.019)	0.052 (0.040–0.076)	2.603 (0.138)	5.1
BtMA-688	0.012 (0.011–0.013)	0.030 (0.027–0.034)	3.205 (0.148)	1.2
BtMA-689	0.049 (0.030–0.166)	0.319 (0.113–6.440)	1.579 (0.153)	7.7
BtMA-690	0.003 (0.002–0.003)	0.009 (0.007–0.011)	2.763 (0.137)	5.2
BtMA-691	0.437 (0.252–1.290)	3.997 (1.338–35.925)	1.334 (0.230)	0.5
BtMA-703	0.038 (0.029–0.044)	0.057 (0.047–0.130)	7.208 (0.549)	24.1
BtMA –1114	0.004 (0.003–0.005)	0.008 (0.006–0.018)	4.827 (0.271)	30.6
Bti t4001	0.003 (0.002–0.003)	0.014 (0.011–0.017)	2.010 (0.075)	12.5
48 h				
BtMA-37*	0.0474 (0.038–0.053)	0.1 27 (0.098–0.240)	2.994 (0.319)	5.0
BtMA-679	0.011 (0.011–0.012)	0.027 (0.025–0.030)	3.408 (0.154)	1.1
BtMA-681*	0.044(0.039–0.050)	0.072 (0.059–0.117)	5.968 (0.596)	6.5
BtMA-684	0.053 (0.049–0.058)	0.114 (0.093–0.157)	3.828 (0.466)	2.3
BtMA-685	0.015 (0.009–0.025)	0.057 (0.031–0.204)	2.163 (0.111)	19.5
BtMA-687*	0.015 (0.008–0.029)	0.046 (0.025–0.709)	2.593 (0.140)	48.8
BtMA-688*	0.009 (0.008–0.010)	0.022 (0.020–0.025)	3.276 (0.168)	1.8
BtMA-689*	0.040 (0.029–0.109)	0.131 (0.066–3.681)	2.480 (0.371)	5.9
BtMA-690	0.003(0.002–0.004)	0.009 (0.006–0.016)	2.650 (0.122)	20.0
BtMA-691*	–	–	–	–
BtMA-703*	0.036 (0.033–0.038)	0.051 (0.047–0.059)	8.287 (0.633)	5.1
BtMA-1114	0.003(0.0026–0.0034)	0.006 (0.005–0.008)	4.006 (0.206)	6.7
Bti t4001	0.003 (0.0021–0.0033)	0.011 (0.009–0.015)	2.055 (0.093)	3.8
72 h				
BtMA-37	0.023(0.006–0.033)	0.104 (0.085–0.196)	1.945 (0.542)	1.2
BtMA-679	0.008 (0.007–0.008)	0.018(0.016–0.019)	3.622 (0.173)	2.3
BtMA-681	0.034 (0.030–0.037)	0.072(0.061–0.096)	3.880 (0.351)	3.01
BtMA-684	0.044 (0.035–0.049)	0.064 (0.056–0.101)	7.963 (0.693)	37.2
BtMA-685	0.011 (0.009–0.013)	0.051 (0.036–0.081)	1.901 (0.091)	4.05
BtMA-687	0.011 (0.008–0.014)	0.031 (0.021–0.069)	2.799 (0.138)	21.7
BtMA-688	0.008 (0.007–0.009)	0.018 (0.016–0.023)	3.587 (0.211)	3.3
BtMA-689	0.033(0.023–0.069)	0.207 (0.089–1.638)	1.605 (0.138)	7.4
BtMA-690	0.002 (0.001–0.002)	0.005 (0.004–0.006)	3.009 (0.150)	0.6
BtMA-691	1.487 (0.440–39.74)	548.28 (25.29- NE)	0.499 (0.119)	0.4
BtMA-703	0.034 (0.027–0.038)	0.049 (0.043–0.068)	8.282 (0.695)	12.5
BtMA-1114	0.003 (0.002–0.004)	0.006 (0.005–0.001)	4.048 (0.206)	28.9
Bti T4001	0.001 (0.001–0.002)	0.006 (0.005–0.008)	1.924 (0.090)	2.7

Legend: LC₅₀ and LC₉₀ = Lethal Concentration; CI, Confidence interval. NE = NE, not estimated.
* Strains that showed a difference in toxicity for T Student. The control showed mortality ≤ 5%.

cry and *cyt* genes (Bravo et al., 1998; Armengol et al., 2006; Abulreesh et al., 2012; Reyaz et al., 2017).

In the isolates used in the present study, the variation of the gene profile of naturally occurring strains active against mosquitos showed different combinations of the genes encoding insecticidal proteins (Bravo et al., 1998; Ibarra et al., 2003; Jouzani et al., 2008; González et al., 2011; El-kersh et al., 2016). Gene profile variation was also observed in *Bt* isolates native to Colombia in a study of selection of strains in *C. quinquefasciatus* and Lepidopteran larvae (Armengol et al., 2006).

The Cerrado biome showed the largest number of strains with combinations of mosquito-specific toxin-encoding genes, including *cry4*, *cry11*, *cry10*, *cyt1*, and *cyt2*, similar to the standard strain, and the only Caatinga isolate with larvicidal activity also has the same *Bti* genes. On the other hand, this gene profile was not observed for the Amazon strains.

The high frequency of mosquito-specific *cry* genes presents in 97.2% of isolates with larvicidal activity demonstrates the importance of this class of genes in the pathogenicity of *Bt* for this insect group. Among the mosquito-active *cry* genes described in the literature, the genes of the *cry4* and *cry11* stand out due to their larvicidal potential. In the present study, it was found that the *cry4Aa* and *cry4Ba* genes were the most frequent, presents in more than half (59.4%) of the strains. Higher frequency of *cry4Ba* genes in isolates with larvicidal activity against *Ae.*

aegypti (Costa et al., 2010; Campanini et al., 2012).

In the present study, the frequency of *cyt* genes was lower than that of *cry*, the *cyt1Aa* gene was the most frequent. The presence of this gene is often detected in strains active against mosquito larvae (Costa et al., 2010; El-kersh et al., 2016).

This gene plays a key role in the activity of *Bt* against mosquitoes, as the Cyt1Aa proteins act directly in the insertion of the Cry toxins into the intestinal epithelium of the larvae, which can increase the toxicity of the strains where they are found (Pérez et al., 2005; Elleuch et al., 2015b). The toxin *cyt1Aa* may hinder selection by mosquito populations for resistance (Pérez et al., 2005).

The synergism between Cry and Cyt proteins presents greater toxicity than the use of Cry proteins alone or the combination between two or more Cry (Crickmore et al., 1995; Xu et al., 2014; Elleuch et al., 2015b). In the present study, the presence of the combination of *cry/cyt* genes was observed in 78% of isolates with pathogenic activity, and the results showed that the most toxic strains obtained in this study contain different combinations of *cry* and *cyt* genes.

The bioassays with *Ae. aegypti* help relate the toxicity information of each strain to the molecular identification of the genes that may be directly involved in the larvicidal activity in mosquitoes, this is the culicid species with the highest number of toxins with larvicidal activity already described (Frankenhuyzen, 2009; 2013).

In the present study, two strains (BtMA-690 and BtMA-1114) were found to have similar toxicity to the standard strain *Bti*-T4001, with equal LC₅₀ values in at least one of the three mortality evaluations.

Notably, the two isolates with the highest toxicity were obtained from different substrates BtMA-690 from a soil sample and BtMA-1114 from Coleoptera insects—both collected in a Cerrado area. These data are important for obtaining the diversity of toxic strains.

Considering the gene profile of the most toxic strains, all genes present in *Bti* were detected in BtMA-690, whereas BtMA-1114 presented positive amplification for the genes *cry4*, *cry11*, *cry10*, *cyt1*, and *cyt2*, but did not show amplification for the *cry11Ba* and *cyt1Ab* genes; in addition, the *cry32* gene was also identified in this strain.

Cry32 insecticidal proteins have three different genes *cry32Ba*, *cry32Ca*, and *cry32Da* encoding toxins active against *Ae. aegypti* (Frankenhuyzen, 2009). The presence of the *cry32* gene is an indication that for the BtMA-1114 strain, indicate the diversity of the gene profile of the pathogenic strains for mosquitoes obtained in this study.

Several studies show the variation of the gene profile and toxicity of *Bt González et al.* (2011), while studying strains native to Cuba, observed that for three of these, the LC₅₀ values were better than that of the standard strain, and two strains contained the same gene profile as *Bti*, whereas the other strain showed different plasmid and protein profiles. This was also observed in two strains—BLB355 from Portugal and BLB196 from Saudi Arabia—that presented larvicidal activity against *Ae. aegypti*, but the presence of *Bti* genes was not detected in these strains (Elleuch et al., 2015a).

On the other hand, other studies have reported that strains with a *Bti*-like gene profile are the most effective (Costa et al., 2010; Santos et al., 2012). In the present study, the BtMA-690 isolate presented a gene profile and a toxicity similar to those of the standard strain; the same profile was detected in seven other strains, but for those, the insecticidal activity was lower than that of the standard.

However, despite the BtMA-679 and BtMA-687 strains not showing the same degree of toxicity as *Bti*, the genes present in the standard strain, as well as the *cry32* gene, were found in these strains. Thus, as with BtMA-1114, these strains are promising for preventing the emergence of resistance to the combination of *Bti* Cry/Cyt toxins, as the use of new strains with gene profiles different from those of the strains already being used is a way to avoid selection of resistant mosquito populations (Peralta and Palma, 2017).

In addition to the combination of the *cry/cyt* genes found in the strains of this study, the presence of the chitinase gene was also observed in 35% of the strains. The presence of the insecticidal proteins Cry/Cyt combined with chitinase contributes to the overall toxicity of the strains because chitinases have the potential to destroy the peritrophic matrix of larvae, thus facilitating the contact between δ -endotoxins and their receptors in the intestinal epithelium (Sampson and Gooday, 1998; Juárez-Hernández et al., 2015). The presence of these toxins in strains active against mosquitoes was also described in other Brazilian strains (Costa et al., 2010).

The present study provides evidence of the diversity the *Bt* with activity against larvae of *Ae. aegypti*, *Cx. quinquefasciatus*, and *An. darlingi* isolated from soils and insects different biomes Brazilians. The Cerrado biome showed more promise for obtaining strains with higher toxicity for mosquitoes (BtMA-1114 and BtMA-690). The data are promising for control mosquitoes of medical importance, since *Bt* is an effective component in the control of these insects, which are currently considered a public health problem worldwide.

Conflict of interest

The authors declare that there is no conflict of interest.

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References

- Abulreesh, H.H., Osman, G.L.E.H., Assaedi, A.S.A., 2012. Characterization of insecticidal genes of *Bacillus thuringiensis* strains isolated from arid environments. *Indian J. Microbiol.* 52, 500–503.
- Alves, S.B., Moraes, S.A., 1998. Quantificação de inoculo de patógenos de insetos. In: In: Alves, S.B. (Ed.), Controle microbio de insetos 765 FEALQ, Piracicaba p. 765 778.
- Armengol, G., Escobar, M.C., Maldonado, M.E., Orduz, S., 2006. Diversity of Colombian strains of *Bacillus thuringiensis* with insecticidal activity against dipteran and lepidopteran insects. *J. Appl. Microbiol.* 102, 77–88.
- Assaedi, A.S.A., Osman, G.E.H., Abulreesh, H.H., 2011. The occurrence and insecticidal activity of *Bacillus thuringiensis* in the arid environments. *Aust. J. Crop. Sci.* 5, 1185–1190.
- Ayres, M., Ayres Jr., M., Ayres, D.L., Santos, A.S., 2007. BioEstat versão 5.3: Aplicações estatísticas nas áreas das ciências Biológicas e médicas. Belém, Sociedade Civil Mamirauá, Brasília: MCT/CNPQ, Belém, Pará, Brasil.
- Ben-Dov, E., 2014. *Bacillus thuringiensis* subsp. israelensis and Its Dipteran-Specific Toxins. *Toxins* 6, 1222–1243.
- Bernhard, K., Jarrett, P., Meadows, M., Butt, J., Ellia, D.J., Roberts, G.M., Pauli, S., Rodgers, P., Burges, H.D., 1997. Natural isolates of *Bacillus thuringiensis*: worldwide distribution, characterization, and activity against insects pests. *J. Inverteb. Pathol.* 70, 59–68.
- Brasil. Ministério da Saúde (MS), 2016. Situação Epidemiológica da filariose linfática no Brasil. [http://portalquiquis.saude.gov.br/images/pdf/2016/marco/11/2015-039—Filariose-vers-ofinal.pdf](http://portalquiquis.saude.gov.br/images/pdf/2016/marco/11/2015-039-Filariose-vers-ofinal.pdf).
- Bravo, A., Sarabia, S., Lopez, L., Ontiveros, H., Abarca, C., Ortiz, A., Ortiz, M., Lina, L., Villalobos, F.J., Pena, G., Nuñez-Valdez, M.E., Soberón, M., Quintero, R., 1998. Characterization of *cry* genes in a Mexican *Bacillus thuringiensis* strain collection. *Appl. Environ. Microbiol.* 64, 4965–4972.
- Bravo, A., Gill, S.S., Soberón, M., 2007. Mode of action of *Bacillus thuringiensis* Cry e Cyt toxins and their potential for insect control. *Toxicol.* 49, 423–435.
- Bravo, A., Likitvivanavong, S., Gill, S.S., Soberón, M., 2011. *Bacillus thuringiensis Bacillus thuringiensis*: A story of a successful bioinsecticide. *Insect Biochem. Mol. Biol.* 41, 423–431.
- Cánthon, P.E., Cancino-Rodezno, A., Gill, S.S., Soberón, M., Bravo, A., 2015. Transcriptional cellular responses in midgut tissue of *Aedes aegypti* larvae following intoxication with Cry11Aa toxin from *Bacillus thuringiensis*. *BMC Genomics* 16, 1042.
- Campanini, E.B., Davolos, C.C., Alves, E.C.C., Lemos, M.V.F., 2012. Isolation of *Bacillus thuringiensis* strains that contain dipteran-specific cry genes from ilha bela (São paulo, Brazil) soil samples. *Braz. J. Biol.* 72, 243–247.
- Costa, J.R.V., Rossi, J.R., Marucci, S.C., Alves, E.C.C., Volpe, H.X.L., Ferraudo, A.S., Lemos, V.F.M., Desidério, J.A., 2010. Atividade tóxica de isolados de *Bacillus thuringiensis* a larvas de *Aedes aegypti* (L.) (Diptera: culicidae). *Neotrop. Entomol.* 39, 757–766.
- Crickmore, N., Bone, E.J., Williams, J.A., Ellar, D.J., 1995. Contribution of the individual components of the δ -endotoxin crystal to the mosquitocidal activity of *Bacillus thuringiensis* subsp. *israelensis*. *FEMS Micro. Lett.* 131, 249–254.
- Crickmore, N., 2017. Full List of Delta-endotoxins. <http://www.btnomenclature.info/>.
- Dias, D.G.S., Silva, S.F., Martins, E.S., Soares, C.M.S., Falcão, M., Gomes, A.C.M.M., Praça, L.B., Dias, J.M.C.S., Monerrat, R.G., 2002. Prospecção de estirpes de *Bacillus thuringiensis* efetivas contra mosquitos. *Bol. Pesqui. Des* 1340–1676.
- Djenane, Z., Nateche, F., Amziane, M., Gomis-Cebolla, J., El-Aichar, F., Khorf, H., Ferré, J., 2017. Assessment of the antimicrobial activity and the entomocidal potential of *Bacillus thuringiensis* isolates from Algeria. *Toxins* 9, 139–158.
- Ebi, K.L., Nealon, J., 2016. Dengue in a changing climate. *Environ. Res.* 151, 115–123.
- El-kersh, T.A., Ahmed, A.M., Al-sheikh, Y.A., Tripet, F., Ibrahim, M.S., Metwalli, A.A.M., 2016. Isolation and characterization of native *Bacillus thuringiensis* strains from Saudi Arabia with enhanced larvicidal toxicity against the mosquito vector *Anopheles gambiae* (s.l.). *Parasits Vectors* 9, 647.
- Elleuch, J., Tounsi, S., Hassen, N.B.B., Lacoix, M.N., Chandre, F., Jaoua, S., Zghal, R.Z., 2015a. Characterization of novel *Bacillus thuringiensis* isolates against *Aedes aegypti* (Diptera: Culicidae) and *Ceratitis capitata* (Diptera: Tephritidae). *J. Invert. Pathol.* 124, 90–97.
- Elleuch, J., Jaoua, S., Darriet, F., Chandre, F., Tounsi, S., Zghal, R.Z., 2015b. Cry4Ba and Cyt1Aa proteins from *Bacillus thuringiensis israelensis*: interactions and toxicity mechanism against *Aedes aegypti*. *Toxicol.* 104, 83–90.
- Fatoreto, J.C., Sena, J.A.D., Barreto, M.R., Lemos, M.L.V.F., Boiça Junior, A.L., 2007. Associação de bioensaios e caracterização molecular para seleção de novos isolados de *Bacillus thuringiensis* efetivos contra *Spodoptera frugiperda* (J. E. Smith) (Lepidoptera: Noctuidae). *Neotrop. Entomol.* 36, 737–745.
- Ferreira-de-Brito, A., Ribeiro, I.P., Miranda, R.M., Fernandes, R.S., Campos, S.S., Silva, K.A.B., Castro, M.G., Bonaldo, M.C., Brasil, P., Lourenço-de-Oliveira, C., 2016. First detection of natural infection of *Aedes aegypti* with Zika virus in Brazil and throughout South America. *Mem. Inst. Oswaldo. Cruz.* 111, 655–658.
- Finney, D.J., 1971. *Probit Analysis*. Cambridge University, London.
- Frankenhuyzen, K.V., 2009. Insecticidal activity of *Bacillus thuringiensis* crystal proteins. *J. Invert. Pathol.* 101, 1–16.
- Frankenhuyzen, K.V., 2013. Cross-order and cross-phylum activity of *Bacillus thuringiensis* pesticidal proteins. *J. Invert. Pathol.* 114, 76–85.

- Gobatto, V., Giani, S.G., Camassola, M., Dillon, A.J.P., Specht, A., Barros, N.M., 2010. *Bacillus thuringiensis* isolates entomopathogenic for *Culex quinquefasciatus* (Diptera: 83 Culicidae) and *Anticarsia gemmatalis* (Lepidoptera: Noctuidae). *Braz. J. Biol.* 70, 1039–1046.
- González, A., Díaz, R., Díaz, M., Borrero, Y., Bruzón, R.Y., Carreras, B., Gato, R., 2011. Characterization of *Bacillus thuringiensis* soil isolates from Cuba, with insecticidal activity against mosquitoes. *Rev. Biol. Trop.* 59, 1007–1016.
- Habib, M.E.M., Andrade, C.F.S., 1998. Bactérias entomopatogênicas. In: In: Alves, S.B. (Ed.), *Controle Microbiano de Insetos* 12. FEALQ, São Paulo, Piracicaba, pp. 383–446.
- Hernandez, C.S., Andrew, R., Bel, Y., Ferré, J., 2005. Isolation and toxicity of *Bacillus thuringiensis* from potato growing areas in Bolivia. *J. Invert. Pathol.* 88, 8–16.
- Honório, N.A., Câmara, D.C.P., Calvet, G.A., Brasil, P., 2015. Chikungunya: Uma arbovirose em estabelecimento e expansão no Brasil. *Cad. Saúde Pública* 31, 906–908.
- IBGE – Instituto Brasileiro de Geografia e Estatística 2017. <http://cidades.ibge.gov.br/xtras/uf.php?coduf=21>.
- Ibarra, J.E., Del Rincón, C., Ordúz, S., Noiega, D., Benintende, G., Monnerat, R., Regis, L., Oliveira, C.M.F., Lanz, H., Rodriguez, M.H., Sánchez, J., Pena, G., Bravo, A., 2003. Diversity of *Bacillus thuringiensis* strains from Latin America with insecticidal activity against different mosquito species. *Appl. Environ. Microbiol.* 69, 5269–5274.
- Jouzani, G.S., Abad, A.P., Seifinejad, A., Marzban, R., Kariman, K., Maleki, B., 2008. Distribution and diversity of dipteran-specific cry and cyt genes in native *Bacillus thuringiensis* strains obtained from different ecosystems of Iran. *J. Ind. Microbiol. Biotechnol.* 35, 83–94.
- Juárez-Hernández, E.O., Casados-Vázquez, L.E., Rincón-Castro, M.C., Salcedo-Hernández, R., Bideshi, D.K., Barboza-Corona, J.E., 2015. *Bacillus thuringiensis* subsp. *israelensis* producing endochitinase ChiA74(sp inclusions and its improved activity against *Aedes aegypti*. *J. Appl. Microbiol.* 119, 1692–1699.
- Jung, Y.C., Kim, S.U., Côte, J.-C., Lecadet, M.-M., Chung, Y.S., Bok, S.H., 1998. Characterization of a new *Bacillus thuringiensis* subsp. *higo* strain isolated from rice bran in Korea. *J. Invert. Pathol.* 71, 95–96.
- Lacey, L.A., Grzywacz, D., Shapiro-Ilan, D.I., Frutos, R., Brownbridge, M., Goettel, M.S., 2015. Insect pathogens as biological control agents: back to the future. *J. Invert. Pathol.* 132, 1–41.
- LeOra Software Company, 2003. *PoloPlus: Probit and Logit Analysis. User's Guide, Version 2.0*. Petaluma, CA, LeOra Software Company 39 pp.
- Lin, Y., Xiong, G., 2004. Molecular cloning and sequence analysis of the chitinase gene from *Bacillus thuringiensis* serovar *alesti*. *Biotechnol. Lett.* 26, 635–639.
- Moyes, C.L., Vontas, J., Martins, A.J., Ng, L.C., Koo, S.Y., Dusfour, I., Raghavendra, K., Pinto, J., Cordel, V., David, P., Weetman, D., 2017. Contemporary status of insecticide resistance in the major *Aedes* vectors of arboviruses infecting humans. *PLoS Neg. Trop. Dis.* 11, 1–20.
- Ootani, M.A., Ramos, A.C.C., Azevedo, E.B., Garcia, B.O., Santos, S.F., Aguiar, R.W.S., 2011. Avaliação da toxicidade de estirpes de *Bacillus thuringiensis* para *Aedes aegypti* Linnaeus, (Diptera: Culicidae). *J. Biotec. Biodivers* 2, 37–43.
- Pérez, C., Fernandez, L.E., Sun, J., Folch, J.L., Gill, S.S., Soberón, M., Bravo, A., 2005. *Bacillus thuringiensis* subsp. *israelensis* Cyt1Aa synergizes Cry11Aa toxin by functioning as a membrane-bound receptor. *Pnas* 102, 18303–18308.
- Peralta, C., Palma, L., 2017. Is the insect world overcoming the efficacy of *Bacillus thuringiensis*? *Toxins* 9, 1–5.
- Pereira, E., Teles, B., Martins, E., Praça, L., Santos, A., Ramos, F., Berry, C., Monnerat, R., 2013. Comparative toxicity of *Bacillus thuringiensis* Berliner strains to larvae of Simuliidae (Insecta: Diptera). *Bt Research.* 4, 8–13.
- Pinto, L.M.N., Azambuja, A.O., Diehl, E., Fiuza, L.M., 2003. Pathogenicity of *Bacillus thuringiensis* isolated from two species of Acromyrmex (Hymenoptera, Formicidae). *Braz. J. Biol.* 63, 301–306.
- Polanczyk, R.A., Rogério, F.P., Fiuza, L.M., 2004. Isolamento de *Bacillus thuringiensis* berliner a partir de amostras de solos e sua patogenicidade para *Spodoptera frugiperda* (J. E. Smith) (Lepidoptera: Noctuidae). *Rev. Bras. Agrociência* 10, 209–214.
- Praça, L.B., Batista, A.C., Martins, E.S., Siqueira, C.B., Dias, D.G.S., Gomes, A.C.M.M., Falcão, R., Monnerat, R.G., 2004. Estirpes de *Bacillus thuringiensis* efetivas contra insetos das ordens Lepidoptera, Coleoptera e Diptera. *Pesq. Agropec. Brasileira* 39, 11–16.
- Rebollo, M.P., Bockarie, M.J., 2017. Can Lymphatic Filariasis Be Eliminated by 2020? *Trends Parasitol.* 33, 83–92.
- Reyaz, A.L., Gunapriya, L., Arulselvi, P.I., 2017. Molecular characterization of indigenous *Bacillus thuringiensis* strains isolated from Kashmir valley. *Biotech* 7, 143–153.
- Sampson, M.N., Gooday, G.W., 1998. Involvement of chitinases of *Bacillus thuringiensis* during pathogenesis in insects. *Microbiology* 144, 2189–2194.
- Santos, F.P., Lopes, J., Vilas-Bóas, G.T., Zequi, J.A.C., 2012. Characterization of *Bacillus thuringiensis* isolates with potential for control of *Aedes aegypti* (Linnaeus, 1762) (Diptera: Culicidae). *Acta Trop.* 122, 64–70.
- Seixas, G., Grigoraki, L., Weetman, D., Vicente, J.L., Silva, A.C., Pinto, J., Vontas, J., Sousa, C.A., 2017. Insecticide resistance is mediated by multiple mechanisms in recently introduced *Aedes aegypti* from Madeira Island (Portugal). *PLoS Neg. Trop. Dis.* <http://dx.doi.org/10.1371/journal.pntd.0005799>.
- Silva, S.F., Dias, J.M.C., Monnerat, R.G., 2002. Isolamento, Identificação e Caracterização Entomopatogênica de Bacilos de Diferentes Regiões do Brasil 70. pp. 1–4.
- Silva, M.C., Siqueira, H.A., Marques, E.J., Silva, L.M., Barros, R., Lima Filho, J.V., Silva, S.M., 2012. *Bacillus thuringiensis* *Bacillus thuringiensis* isolates from northeastern Brazil and their activities against *Plutella xylostella* (Lepidoptera: Plutellidae) and *Spodoptera frugiperda* (Lepidoptera: Noctuidae). *Biocontrol Sci. Techn.* 22, 583–599.
- Siqueira, A.M., Quique Bassat, Q., Rodovalho, S., Lacerda, M.V.G., 2017. Raising the red flag for malaria elimination and integrated fever surveillance in the Brazilian amazon. *Comment* 5.
- Soares-da-Silva, J., Pinheiro, V.C.S., Litaiff-Abreu, E., Polanczyk, R.A., Tadei, W.P., 2015. Isolation of *Bacillus thuringiensis* from the state of Amazonas, in Brazil, and screening against *Aedes aegypti* (Diptera, Culicidae). *Rev. Bras. de Entomol.* 59.
- Stalinski, R., Laporte, F., Tetreau, G., Després, L., 2016. Receptors are affected by selection with each *Bacillus thuringiensis israelensis* Cry toxin but not with the full *Bti* mixture in *Aedes aegypti*. *Infection. Genet. Evol.* 44, 218–227.
- Tadei, W.P., Rodrigues, I.B., Rafael, M.S., Sampaio, R.T.M., Mesquita, H.G., Pinheiro, V.C.S., Zequi, J.A.C., Roque, R.A., Santos, J.M.M., 2017. Adaptive processes, control measures, genetic background, and resilience of malaria vectors and environmental changes in the Amazon region. *Hydrobiologia* 789, 179–196.
- Vidal-Quist, J.C., Castañera, J., 2009. Diversity of *Bacillus thuringiensis* strains isolated from citrus orchards in Spain and evaluation of their insecticidal activity against *Ceratitis capitata*. *J. Microb. Biotech.* 19, 749–759.
- WHO – World Health Organization, 1985. *Informal Consultation on the Development of Bacillus Sphaericus as a Microbial Larvicide*. UNDP/World Bank/WHO Special Programme For Research and Training in Tropical Diseases, Geneva.
- WHO – World Health Organization, 2005. *Guidelines for Laboratory and Field Testing of Mosquito Larvicides*. WHO/CDS/WHOPES/GCDPP/2005.13.
- WHO – World Health Organization, 2017. *Draft Global Vector Control Response 2017–2030*.
- Xu, C., Wang, B.C., Yu, Z., Sun, M., 2014. Structural insights into *Bacillus thuringiensis* Cry, Cyt and parasporin toxins. *Toxins* 6, 2732–2770.
- Yousten, A.A., 1984. *Bacillus sphaericus* *Bacillus sphaericus* microbiological factors related to its potential as a mosquito larvicide, New York. *Adv. Biotechnol. Proc.* 3, 315–343.
- Zhang, Q., Hua, G., Adang, M.J., 2016. Effects and mechanisms of *Bacillus thuringiensis* crystal toxins for mosquito larvae. *Insect Sci.* <http://dx.doi.org/10.1111/1744-7917.12401>.