Toxic Activity, Molecular Modeling and Docking Simulations of Bacillus thuringiensis Cry11 Toxin Variants Obtained via DNA Shuffling

Alvaro Mauricio Florez1*, Miguel Orlando Suarez-Barrera2,3, Gloria M. Morales2, Karen Viviana Rivera2, Sergio Orduz4, Rodrigo Ochoa5, Diego Guerra5 and Carlos Muskus6

1 RG Microbial Ecology: Metabolism, Genomics & Evolution, Microbiomas Foundation, Chía, Colombia, 2 Laboratorio de Biología Molecular y Biotecnología, Universidad de Santander, Bucaramanga, Colombia, 3 Escuela de Medicina, Facultad de Salud, Universidad Industrial de Santander, Bucaramanga, Colombia, 4 Grupo Biología Funcional, Laboratorio de Prospección y Diseño de Biomoléculas, Escuela de Ciencias, Universidad Nacional, Sede Medellín, Colombia, 5 Programa de Estudio y Control de Enfermedades Tropicales PECET, Unidad de Biología Molecular y Computacional-UBMC, Universidad de Antioquia, Medellín, Colombia

The Cry11 family belongs to a large group of δ-endotoxins that share three distinct structural domains. Among the dipteran-active toxins referred to as three-domain Cry11 toxins, the Cry11Aa protein from Bacillus thuringiensis subsp. israelensis (Bti) has been the most extensively studied. Despite the potential of Bti as an effective biological control agent, the understanding of Cry11 toxins remains incomplete. In this study, five Cry11 variants obtained via DNA shuffling displayed toxic activity against Aedes aegypti and Culex quinquefasciatus. Three of these Cry11 variants (8, 23, and 79) were characterized via 3D modeling and analysis of docking with ALP1. The relevant mutations in these variants, such as deletions, insertions and point mutations, are discussed in relation to their structural domains, toxic activities and toxin-receptor interactions. Importantly, deletion of the N-terminal segment in domain I was not associated with any change in toxic activity, and domain III exhibited higher sequence variability than domains I and II. Variant 8 exhibited up to 3.78- and 6.09-fold higher toxicity to A. aegypti than Cry11Bb and Cry11Aa, respectively. Importantly, variant 79 showed an α-helix conformation at the C-terminus and formed crystals retaining toxic activity. These findings indicate that five Cry11 variants were preferentially reassembled from the cry11Aa gene during DNA shuffling. The mutations described in loop 2 and loop 3 of domain II provide valuable information regarding the activity of Cry11 toxins against A. aegypti and C. quinquefasciatus larvae and reveal new insights into the application of directed evolution strategies to study the genetic variability of specific domains in cry11 family genes.

Keywords: Bacillus thuringiensis, Cry11, DNA shuffling, docking, Aedes aegypti, Culex quinquefasciatus
INTRODUCTION

*Bacillus thuringiensis* (Bt), a Gram-positive bacterium characterized by the production of Cry δ-endotoxins capable of killing insects, has been used since the late 1930s as a biological control agent (Melo et al., 2016). A total of 308 holotype toxins are clustered into 75 Cry proteins (Crickmore et al., 2014) (revised February, 2018). The tertiary structures of nine Cry toxins determined via X-ray crystallography to date contain three conserved domains with specific functions and implicated in the structural stability of the protein. The domain I is a bundle of 7-8 α helices involved in pore formation, domain II is a β-prism with exposed loops regions involved in receptor binding and, domain III is a β-sandwich and has influence on receptor binding, ion channel formation and insect specificity (Li et al., 1991; Grochulski et al., 1995; Derbyshire et al., 2001; Galitsky et al., 2001; Morse et al., 2001; Guo et al., 2009; Hui et al., 2012; Figure 1A).

The mechanisms by which Bt induces death in insects are controversial and have not been completely elucidated (Vachon et al., 2012). Currently, there are two mechanisms related to Bt-induced toxicity in insects that have been accepted; the sequential binding and signaling pathways (Zhang Q. et al., 2017). The sequential binding mechanism has been extensively studied and is based on the formation of pores in epithelial cells in the midgut of targeted insects, which results in toxin-receptor interactions, osmotic imbalance and cell death (Bravo et al., 2007; Pardo-Lopez et al., 2013). After crystal ingestion, Cry toxins become solubilized, and specific proteases present in the lumen of the midgut activate the toxins, which then bind to specific receptors located in the insect midgut. In some Cry toxins, this event induces the proteolytic removal of helix α1 (Aronson, 2000), triggering Cry toxin oligomerization, insertion of oligomeric structures altering membrane stability, receptor and production of channels or pores, ultimately leading to cell lysis and insect death (Bravo et al., 2004; Figure 1B). The signaling pathways is a recent proposed mechanism in which the activation of signaling cascades, leads to increased cyclic AMP and protein kinase activities, resulting in cell death (Zhang et al., 2006). Both of these mechanisms contain gaps. In the sequential binding mechanism, the presence of several types of resistance and the link between proteolysis and pore formation are not understood (Melo et al., 2016). In the signaling pathway mechanism, how Cry toxin-receptor interactions mediate toxic activity is unclear (Melo et al., 2016). In this context, it has been suggested that Cry toxins can cause death based on their ability to induce both pore formation and ion channel activation (Zhang Q. et al., 2017).

The pB toxin megaplasmid from Bt subsp. *israelensis* (Bti) contains four Cry proteins encoded by the cry4Aa, cry4Ba, cry10Aa, and cry11Aa genes and two Cyt proteins encoded by the cyt1Aa and cyt2Ba genes (Berry et al., 2002). The cry genes produce 134, 128, 78, and 72 kDa polypeptides, respectively, all of which possess larvicidal activity higher than of the Cyt’s proteins (Ben-Dov, 2014). However, the high toxic activity of Bti is the result of synergistic interactions between all of them (Ben-Dov, 2014). That is the case of Cry1Aa that despite the low toxicity, it is highly synergistic with Bti Cry toxins and aids to overcome resistance in mosquitoes to Cry toxins (Wirth et al., 1997). Due to the synergistic interactions of Bt subsp. *israelensis* toxins, this bacterium has been used worldwide to control mosquito larvae of the genera *Aedes*, *Culex* and *Anopheles*, which are involved in the transmission of diseases including malaria, hemorrhagic fever, dengue fever, lymphatic filariasis, yellow fever (Ben-Dov, 2014), Chikungunya and Zika (Chouin-Carneiro et al., 2016; Gardner et al., 2016; Tsetsarkin et al., 2016).

Cry11Aa from Bti is a 72 kDa protein toxin that is activated by gut enzymes via the proteolytic removal of 28 residues from its N-terminus and proteolytic cleavage into two fragments of 38 and 30 kDa that remain associated and retain toxicity (Dai and Gill, 1993; Revina et al., 2004; de Barros Moreira Beltrao and Silva-Filha, 2007). The Cry11Aa toxin has higher activity against *Aedes* and *Culex* than against *Anopheles* (Revina et al., 2004; Otieno-Ayayo et al., 2008). In *A. aegypti*, this toxin interacts with two midgut brush border membrane receptors; a GPI anchored and alkaline phosphatase (ALP1) (Fernandez et al., 2006) and also binds to Cyt1Aa as a kind of membrane-bound receptor of Cry11Aa increasing the toxic activity (Perez et al., 2005). Other midgut proteins different to the receptor alkaline phosphatase (ALP1) such as ATP binding protein, increases the toxicity of Cry11Aa against *C. quinquefasciatus* (Zhang L. et al., 2017). Other two toxins, Cry11Bb (94 kDa) and Cry11Ba (81 kDa), share a similar insect specificity and are phylogenetically related to Cry11Aa. Cry11Bb and Cry11Ba are produced by Bt subsp. *medellin* and Bt subsp. *jegathesan*, respectively (Delecluse et al., 1995; Orduz et al., 1998).

Although the tertiary structure of Cry11 toxins have not been determined by X-ray crystallography, Cry11Aa have been the most studied among this group using protein engineering tools. Several mutations haven been introduced into different domains that are implicated in its toxicity. Therefore, studies have focused on domain I developing N-terminally truncated forms of Cry11Aa (Pang et al., 1992) or modifications in domains II and III altering the interactions with its receptor in the midgut, confirming the importance of these domains for Cry11Aa-mediated toxicity (Fernandez et al., 2005, 2009). Based on phage display and site-directed mutagenesis, the exposed regions of loop α8, β4 and loop 3 in domain II of Cry11Aa have been shown to be involved in the interaction of Cry11Aa with *A. aegypti* brush border membrane vesicles (BBMVs). Specifically, two mutations in loop α8, V262E and E266A, reduced the toxic activity of Cry11Aa against *A. aegypti* (Fernandez et al., 2005). There are also mutations in loop α-8 that are involved in Cry11Aa-ALP1 receptor interaction, that affect the Cyt1Aa and Cry11Aa interaction reducing the synergism between these proteins and decreasing their toxic activity (Perez et al., 2005). Other binding sites have been described for the interaction of Cry11Aa with the receptor ALP1 as an important secondary receptor for Cry11Aa and Cry11Ba (Chen et al., 2017). The involved regions are located in loop 2 of domain II and β18-β19 of domain III of Cry11Aa, which interact with ALP1 regions R258-G102 and N287-S288, respectively (Fernandez et al., 2009).
Since the three Cry11 toxins are phylogenetically related and exhibit similar specificity to insect species, it is possible to infer similarities at structural level that can be analyzed after mixing their genes in order to create novel proteins with improved properties. Therefore, considering the lack of studies focused on this approach, we designed a DNA shuffling strategy to obtain variants with increased toxicity to A. aegypti and C. quinquefasciatus. DNA shuffling has been used alone or in combination with phage display and the staggered extension process via homologous recombination to increase the activity of Cry toxins against specific insect pests (Lucena et al., 2014). This technique is a powerful approach based on recombination between parental genes in a single DNA shuffling reaction following random fragmentation (Stemmer, 1994).

Here, we report five variants that were reassembled from the cry11Aa gene that exhibit from moderate to high toxic activities against A. aegypti and C. quinquefasciatus mosquito larvae. Variant 8 was the most toxic to the mosquito larvae, and variants 23 and 79 displayed important differences in 3D structure, toxin-ALP1 interactions and toxicity in which are implicated domains II and III preferably.

**MATERIALS AND METHODS**

**Microbial Strains, Clone Selection and Gene Constructs**

For DNA manipulation, *Escherichia coli* DH5α, JM109 (Promega) and DH5αTOP10® (Life Technologies) cells were grown at 37°C in Luria Bertani (LB) culture medium supplemented with ampicillin (50 µg/ml) and X-gal (50 mg/ml). A Bt 4Q2-81 strain carrying the pBTM3 plasmid and expressing the cry11Bb gene from Bt subsp. medellin (Restrepo et al., 1997) and a second Bt 4Q2-81 strain carrying the plJEG90.1 plasmid and expressing the cry11Ba gene from Bt subsp. jegathesan (Delecluse et al., 1995) were cultured as previously described (Restrepo et al., 1997) in M1 medium supplemented with 30 µg/ml tetracycline and 20 µg/ml erythromycin. Crystal production was evaluated via phase contrast microscopy. *E. coli* DH5α cells harboring the pSV2 plasmid, which carried the p19 gene upstream of cry11Aa from
**Isolation of cry11 Genes via PCR**

The cry11 genes were amplified via PCR using specific primers and plasmid DNA from constructs pBTM3, pSV2 and pIEG90.1 as templates. Briefly, reactions were conducted in a final volume of 50 µl that contained 20 ng of plasmid DNA, 0.5 µM primers, 1× Taq polymerase buffer, 0.4 mM dNTPs, 1.5 mM MgCl2 and 0.5 U GoTaq polymerase (Promega). The amplification conditions were denaturation at 94°C for 4 min followed by 35 cycles of 45 s at 94°C, 45 s at 55°C, and 4 min 30 s at 72°C, and a final extension at 72°C for 10 min. The PCR products were separated via electrophoresis and purified using the PCR Clean-Up System (Promega). The cry11Ba gene was obtained via two independent PCRs, producing a 1.8-kb product (denoted Ba1) corresponding to cry11Ba with a deletion of 246-bp downstream of the ATG start site and a 0.9-kb fragment (denoted as Ba-2) that contained a 0.75-kb fragment of cry11Ba including the stop codon and a 249-bp segment homologous to the multiple cloning site (MCS) of the pHT315 shuttle vector.

**cry11 Gene Cloning, Insert Validation, and Sequencing**

The cry11 genes obtained via PCR were cloned using the TOPO TA system. Recombinant variants for each product were selected, and plasmid DNA was extracted using the Wizard Plus Miniprep kit (Promega). Verification of each insert was performed via digestion of 50 ng of DNA with EcoRI, followed by separation via agarose gel electrophoresis. The released inserts were also used as templates for PCR to confirm the presence of the cry11 genes in the inserts. Each reaction was conducted in a final volume of 25 µl that contained 0.4 µM each of the forward primer 5′-TTCAGAAGATAGCGCCAGATCAAGC-3′ and the reverse primer 5′-TTCAGAAGATAGCGCCAGATCAAGC-3′ (Bravo, 1997; Bravo et al., 1998) in 1× Taq polymerase buffer, 0.4 mM dNTPs, 2.5 mM MgCl2 and 0.12 U GoTaq polymerase. The amplification conditions were 5 min at 94°C followed by 35 cycles of 45 s at 94°C, 45 s at 51°C, and 1 min at 72°C and a final extension step of 6 min at 72°C. pTOAa containing a 2.5-kb insert, pTOBa-1 containing a 1.6-kb insert, pTOBa-2 containing a 0.78-kb insert and pTOBb containing a 3.5-kb insert were sequenced by Macrogen, Inc. (Seoul, South Korea) using M13/T7 primers and the primer pair pCR4F (5′-GATAACATTTCACACAGGA-3′) and pCR4R (5′-TTGTAAACACGCGCCAGTG-3′).

**Test Primers for Reassembly via PCR**

The pTOAa, pTOBa-1 and pTOBa-2 constructs were used as templates in 50-µl PCRs that contained 0.32 µM PCR4 primers as described in **Supplementary Table 1**, 1× Pfx polymerase buffer, 0.4 mM dNTPs, 2.5 mM MgCl2 and 0.5 U Pfx50 polymerase (Life Technologies) in 1× reaction buffer that contained 1 mM MgSO4. The amplification conditions were denaturation at 94°C for 4 min followed by 35 cycles of 45 s at 94°C, 45 s at 59–67°C, and 4 min 30 s at 68°C and a final extension step of 10 min at 68°C. The PCR products were separated via electrophoresis and purified using the Wizard purification system (Promega).

To generate the pGE7 construct, 50 ng of pGEBb-1 plasmid DNA was used as a template. The PCR was performed in a final volume of 50 µl that contained 0.32 µM of each primer as described in **Supplementary Table 1**, 1× Pfx polymerase buffer, 0.3 mM dNTPs, 1 mM MgSO4 and 0.5 U Pfx50 polymerase. The PCR conditions were denaturation for 4 min at 94°C followed by 35 cycles of 45°C for 45 s, 68°C for 45 s, and 68°C for 4 min 30 s and a final extension step at 68°C for 10 min. The PCR products were separated via agarose gel electrophoresis and purified using the Wizard® SV Gel and PCR Clean-Up System (Promega).

**DNA Shuffling**

Three micrograms of each PCR product obtained from the TA cloning constructs and from pGEBb with lengths of 2.5 kb (cry11Aa), 1.6 kb (cry11Ba-1), 0.78 kb (cry11Ba-2), and 3.5 kb (cry11Bb) were mixed in 25 µl of 50 mM Tris–HCl, pH 7.4, and 10 mM of MnCl2. In the same tube, 0.0006 U DNase I (Life Technologies) was added to a final volume of 50 µl. The reaction was incubated between 5 and 20 min at room temperature to optimize production of fragments ranging between 25 and 250 bp. The reaction was stopped by adding 25 µl of 25 mM EDTA. The DNase I digestion products were separated via electrophoresis and purified with the QIAEX II Gel Extraction Kit (QIAGEN). Forty microliters of the pooled of purified fragments were used as a template for PCR without primers in 1× Pfx buffer, 0.3 mM dNTPs, and 2.5 U Pfx50 polymerase in a final volume of 50 µl under the following conditions: 94°C for 3 min, 45 cycles of 94°C for 30 s, 48°C for 3 min and 68°C for 1 min (with a 12-s increase in extension time per cycle), and a final extension step at 68°C for 7 min. The products for reassembly were validated via agarose gel electrophoresis and purified using the Wizard PCR Clean-Up system.

The full-length sequences were amplified using two sets of primers, PCR4F/R and PGE7F/R, and the combination of these primers, PCR4F/PGE7/R, along with a template of 1 µl of the products of the primerless PCR, to a final volume of 50 µl. The PCR was conducted in 1× Pfx50 buffer, 0.3 µM dNTPs, 0.3 µM primers, and 5 U Pfx50 under the following conditions: 4 min at 94°C, 25 cycles of 94°C for 45 s, 55°C for 1 min, and 68°C for
4 min (with a 20-s increase in extension time per cycle), and a final extension step at 68°C for 10 min. The second PCR product, corresponding to the DNA shuffling product, was analyzed via agarose gel electrophoresis and purified using the Wizard PCR Prep DNA Purification System.

**Cloning, DNA Sequencing and Homology Analysis**

The shuffled PCR products were cloned using the TOPO Zero Blunt PCR cloning kit (Life Technologies) according to the manufacturer’s instructions and chemically competent E. coli DH5αTOP10 cells (Life Technologies). Sequencing data were used to select clones according to their open reading frame (ORF) and DNA identity to the parental genes. Plasmid DNA from each clone was isolated using a Wizard Minipreps kit, and the DNA was sent to Macrogen, Inc., South Korea, for sequencing. The forward and reverse primers used for sequencing were 

\[ 
\text{M13 forward (−17): 5'}-\text{GTAAACGACGGCCAG-3'}; 
\text{M13 reverse, 5'}-\text{CAGGAAAACAGCTATGAC-3'}.
\]

Sequence alignments were performed using ClustalW (Thompson et al., 1994), available on the web.

**Transformation of Bacillus thuringiensis**

The PCR products were cloned using TOPO Zero Blunt PCR cloning kit and then subcloned into the pSV2 expression vector using *HindIII* and *SacI*. The resulting vectors were transformed into BMB171 cells via electroporation using a Bio-Rad Micropulser™. BMB171 cells were grown in LB-glycine 0.12% up to an OD600 of 0.15, corresponding to the early exponential phase, and then transformed with 500 ng of the corresponding to the DNA shuffling product, was analyzed via scanning electron microscopy (SEM).

**Cultures, Solubilization, Cry Protein Quantification, and SDS–PAGE**

Cultures expressing variants 1, 8, 23, 79, and 81 and recombinant Cry1Aa and Cry1Bb, including the *Bt* acrystaliferous BMB171 strain, were grown in 10 ml of LB supplemented with chloramphenicol (6 μg/ml) for 7 days at 30°C and 300 rpm. After 48 h of incubation, the culture purity was confirmed via microscopic observation of spores, crystals, and lysed cells. The final culture was collected via centrifugation at 11,200 × g for 15 min at 4°C. The supernatant was discarded, and 1 ml of 1 M NaCl was added to the pellet, which was shaken for 1 h at 30°C and 50 rpm to neutralize protease activity. Then, the suspension was washed twice with 1 ml of 1× PBS and centrifuged at 11,200 × g for 5 min at 4°C. The number of spores was determined via heat shock using 100 μl of each culture and incubated at 72°C for 20 min. Afterward, the samples were incubated at 4°C for 10 min and were diluted by 10⁻¹ to 10⁻⁵ in a final volume of 100 μl. The dilutions were plated on LB agar supplemented with chloramphenicol (6 μg/ml) and incubated at 30°C for 24 h.

To quantify Cry protein production, 200 μl of the final sporulated culture of each variant was solubilized by adding 800 μl of solubilization buffer (50 mM NaOH, 10 mM EDTA, pH 11.7), incubating the culture at 4°C overnight and centrifuging the culture at 25,200 × g for 1 h at 4°C. The supernatant was collected, and the volume was adjusted to 1 ml with Tris-base (0.1 M, pH 7.4). The protein concentration was determined using the Bradford protein assay (Bradford, 1976) and was confirmed via SDS–PAGE using bovine serum albumin (BSA) as a standard. The protein samples were electrophoresed on 10% SDS–PAGE gels at 80 V for 90 min using a Bio-Rad mini protein system (Bio-Rad Laboratories). A total of 5 μg of protein was loaded per lane, and the protein bands were visualized by staining with Coomassie Brilliant Blue R-250 solution for 30 min.

**SEM**

The final cultures were centrifuged at 11,200 × g at 4°C for 10 min, and the precipitate was washed twice with 1× PBS. The pellet was resuspended in 1/10 of the original volume, and 100 μl of the samples were placed on glass slides and dried overnight at room temperature. The samples were coated with a thin layer of gold on a Denton Desk Vacuum IV and analyzed using a JEOL JSM 5010 LV scanning electron microscope.

**Half Lethal Concentration (LC₅₀) in A. aegypti and C. quinquefasciatus Larvae**

Each bioassay consisted of two replicates, each with 30 first instar larvae in 1 ml at 24°C for each variant at 7 different concentrations under the same environmental conditions. A total of 420 larvae were used for each variant. Larval mortality was determined by counting the number of live larvae after 24 and 48 h, and 50% lethal concentrations were determined statistically via Probit analysis which employ a transformation from sigmoid dose-response curve to a straight linear and then analyzed by a regression on the relationship. The calculation of the average lethal concentration (LC₅₀) was made using the R-Project Software.

**3D Structure Prediction and Validation and Secondary Structure Analysis of Non-conserved Regions**

The amino acid sequences of parental Cry11Aa and variants 8, 23, and 79 were modeled via threading methodology using the free local server I-TASSER. From the five models obtained by the

1. http://blast.ncbi.nlm.nih.gov/Blast.cgi
2. http://www.expasy.org
3. https://www.r-project.org/
4. http://zhanglab.ccmb.med.umich.edu/I-TASSER/
program, the first model of each structure was selected according to the best C and TM scores. These structures were geometrically and energetically validated to assess the quality of the generated 3D model using different servers, such as the Ramachandran SWISS-MODEL\(^5\), the Z-score and energy graph in the ProSA-web server\(^6\), ERRAT\(^7\), and Verify3D\(^8\). The structures obtained for variants 8, 23, and 79 were aligned with the structure of Cry11Aa. For variant 79, complementary analysis based on the predicted secondary structure of non-conserved regions was performed using JPred\(^9\). Subsequently, the ab initio method was used to predict the 3D structure of variant 79 using Robetta server\(^10\).

### Molecular Docking of Cry11 Domains With ALP1

Parental Cry11Aa and variants 8, 23, and 79 were analyzed to identify their interactions with the receptor ALP1 from *A. aegypti* (UniProtKB ID: Q16WV8). For this protein, two regions that interact with Cry11Aa were identified by epitope mapping. These regions are located within R59-G102 and N257-1296 in ALP1, which interact with residues in loop 8 of Cry11Aa domain II and residues R561-N570 in Cry11Aa domain III, respectively (Fernandez et al., 2009). The structure of ALP1 was modeled using the I-TASSER server taking into account folding recognition by threading. The model was evaluated according to Z-scores obtained using ProSA-web (see text footnote\(^5\)), and a Ramachandran plot was generated using Swiss-MODEL (see text footnote\(^5\)). For docking analysis, the interactions between Cry11Aa and ALP1 regions were analyzed using the Cry11Aa-interacting domains as peptides with rigid conformations based on the predicted 3D structures obtained in the previous step. The ALP1 structure and the peptides obtained from Cry11Aa and each variant were parameterized using AutoDockTools\(^11\) via the addition of polar hydrogens to each residue's side chain to facilitate the formation of hydrogen bonds. The structures were also treated with Gasteiger partial charges to facilitate electrostatic interactions among other molecular entities. Docking analysis was performed using AutoDock Vina\(^12\) considering an exhaustiveness set to 80, which is proportional to the length of the ligand. For the simulated interactions with both regions of ALP1, 3D grid cubic boxes with sides of 32 Å in length and a grid space of 1.0Å were located on the defined active site center, covering all the residues of interest and allowing the entrance of the full peptide structures into the protein cavities. Subsequently, the different docking conformations for each variant were illustrated and analyzed using LIGPLOT\(^13\).

---

\(^{5}\)http://swissmodel.expasy.org/workspace/?func=tools_structureassessment1

\(^{6}\)https://prosa.services.came.sbg.ac.at/prosa.php

\(^{7}\)http://services.mbi.ucla.edu/ERRAT/

\(^{8}\)http://services.mbi.ucla.edu/Verify_3D/

\(^{9}\)http://www.compbio.dundee.ac.uk/jpred/

\(^{10}\)http://robetta.bakerlab.org/

\(^{11}\)http://autodock.scripps.edu/resources/adt

\(^{12}\)http://vina.scripps.edu/

\(^{13}\)http://www.ebi.ac.uk/thornton-srv/software/LIGPLOT/

### RESULTS

#### Parental cry11 Genes

*cry11Aa, cry11Ba,* and *cry11Bb* were used as parental genes to be fragmented in DNA shuffling based on their closely phylogenetical relationship, similarities at structural protein level and toxic specificity to similar insect species. This approach has been used for in vitro recombination of families of homologous genes in order to create novel proteins with improved properties and is useful for those in which the three-dimensional structure is unknown. In contrast to other random mutagenesis protocols, this technique introduces mutations by random DNA fragmentation and PCR reassembly in a cyclic process that alternates gene diversification, screening and selection of functional variants (Stemmer, 1994).

Four PCR products of 3.5, 2.7, 1.8, and 0.9 kb corresponding to *cry11* genes were obtained (Supplementary Figure 1A), as confirmed via DNA sequencing. The 3.5-kb PCR product contained a 2.2-kb fragment encoding the Cry11Bb protein. Three segments were also identified downstream of the last stop codon. The first segment consisted of 234-bp and showed 93% identity to *cry11Bb2* (accession number HM68615.1). The second segment was a 129-bp fragment that showed 93% identity to the complementary strand of the *IS2140* insertion element (accession number M23740.1) and was used to distinguish the reassembled products from the *cry11Bb* gene. The third segment, a 173-bp fragment, showed 83% identity to the complementary strand of the *cry30Aa* gene (accession number AJ251978.1).

The 2.7-kb PCR product contained a 1.9-kb fragment encoding the Cry11Aa protein as well as two additional segments of 580- and 90-bp that were identified upstream of the first ATG and downstream from the stop codon, respectively. The 580-bp fragment was homologous to the p19 accessory protein gene (GenBank: CAD30080.1) and was used to distinguish the reassembled products from *cry11Aa* gene. The downstream 90-bp segment was homologous to the MCS of pSV2. Sequence analysis showed that the 1.8-kb and 0.9-kb PCR products from *cry11Ba*, denoted as Ba1 and Ba2, respectively, shared a 407-bp segment. A deletion of 246-bp downstream of ATG start site was used to recognize the reassembled products from *cry11Ba* gene. No mutations were detected in the DNA sequences of the PCR products of any of the parental genes.

#### Assembly of Full-Length cry11 Genes

The primers used for DNA shuffling were tested via conventional PCR and random fragmentation, and the results indicated that the parental gene amplifications were successful (Supplementary Figure 1A). The purified PCR products (Supplementary Figure 1B) were mixed and treated with DNase I for 7, 8, or 9 min. However, only the products treated for 8 min (Supplementary Figure 1C) generated fragments between 25 and 200-bp. These products were reassembled, resulting fragments between 1 and 10-kb (Supplementary Figure 1D). After the final assembly using the PCR4F and pGE7R primers, we observed fragments between 0.25 and 2-kb (Supplementary Figure 1E). The assembly reaction products were cloned as described in the
Materials and Methods. A total of 94 variants were obtained, and 10 of these variants did not contain an insert. For the remaining variants, 34 were <1.0-kb, 14 were between 1.1 and 2-kb, 22 were >2.1-kb and, and 14 did not show homology.

Characteristics and Sequence Homology of cry11 Variants

According to the sequence analysis of the Cry11 variants, 14 of them did not show homology to any known endogenous Cry11 toxin. Among those 14 variants, six variants were >2.1-kb and eight were between 1 and 2-kb. The 22 variants that displayed sequence homology and a similar size to the full-length parental genes (>2.1-kb) were clustered into three groups. The first group consisted of variants 1, 8, 23, 28, 54, 79, and 81; these variants contained the p19 gene located upstream from the ATG start site and showed homology to cry11Aa. A second group of ten variants, including 16, 36, 51, 57, 61, 68, 71, 75, 77, and 85, showed DNA homology to cry11Aa but lacked the p19 gene. The third group of variants, 14, 17, 67, 76, and 86, showed homology to the cry11B genes.

Among the 22 variants that were between 1 and 2-kb, 14 showed homology to the cry11B genes, and eight did not have homology to any of the cry11 genes used. All 34 variants that were <1-kb showed homology to the cry11B genes, and their reassembled products contained only domain III.

Among all variants obtained via DNA shuffling, variants 1, 8, 23, 28, 54, 79, and 81 were selected for characterization. Homology analysis of the deduced amino acid sequences of Cry11Aa and variants 1, 8, 23 and 81 showed a high degree of conservation with few amino acid changes, which were preferentially located in domain III (Supplementary Figure 2A). Variants 28 and 54 exhibited 100% identity to Cry11Aa. Therefore, these mutants were excluded from further analysis. Comparative analysis of Cry11Aa, variant 23 (aa 1-643), variant 8 (aa 1-568), and variant 79 (aa 1-551) showed that the two first variants are highly conserved in the extension of the sequences. However, many variations relative to Cry11Aa were present in variant 79 at the end of the sequence, particularly beginning from aa 286. In addition to polymorphisms, several insertions/deletions were present at the end of this sequence (Supplementary Figures 2B,C).

Finally, secondary structure analysis of the deduced amino acid sequence from variants 8, 23, and 79 predicted the presence of α helix, β-sheets and loops (Supplementary Figure 2C). The accession numbers and particular characteristics of the variants 1, 8, 23, 79, and 81 genes are described in Table 1.

Protein Expression and Crystal Formation

SDS–PAGE revealed that all variants contained a similar pattern of solubilized proteins, and degradation was not observed (Supplementary Figure 3A). According to SEM analysis, variants 1, 8, 23, 79, and 81 as well as the parental proteins Cry11Aa and Cry11Bb form crystals (Supplementary Figure 3B). Strain BMB171, which was used as the plasmid recipient for protein expression, did not show crystal formation (Supplementary Figure 3B).

The Selected Variants Exhibited Moderate to High Toxic Activity Against A. aegypti and C. quinquefasciatus.

In accordance with sequence identity, the variants 8, 23, and 79 were reassembled products from the cry11Aa parental gene. The toxic activity against A. aegypti larvae was up to 3.78- and 6.09-fold higher for variant 8 than for Cry11Bb and Cry11Aa, respectively. No significant differences were observed against C. quinquefasciatus. Variants 23 and 79 showed lower and higher mutation rates than Cry11Aa, respectively, although both variants retained toxic activity against C. quinquefasciatus. Additionally, variants 23 and 79 exhibited moderate and high toxicity to A. aegypti, respectively. Surprisingly, variant 79, despite of high variations still retaining toxic activity. Recombinant Cry11Bb exhibited high toxic activity to both A. aegypti and C. quinquefasciatus larvae, while variants 23 and 81 exhibited lower toxic activity than control against A. aegypti. The results of toxicity assays of five variants for the two types of larvae are shown in Figure 2. Based on their mutations and bioassay results, variants 8, 23, and 79 were chosen for analysis of their 3D structure and interaction with ALP1.

Variants 8 and 23 Are Similar to Their Parental Protein, Whereas Variant 79 Shows Structural Differences

The Z-score, ERRAT and Verify3D results obtained for Cry11Aa and variants 8, 23, and 79 are shown in Supplementary Table 2. Multiple alignment including structural alignment of Cry11Aa and its variants showed that variants 8 and 23 are similar to their parental protein due to high sequence conservation (Supplementary Figures 2A,B). The similarity of these variants with respect to Cry11Aa was 87.4 and 98.9%, respectively. These variants also showed similar structural conformation to Cry11Aa (Figures 3A–D). However, variant 79 exhibited structural differences in the non-conserved region compared with its parental protein (Figures 3E,F). The similarity of variant 79 with respect to Cry11Aa was 55.7%. The predicted secondary structure in the non-conserved region of variant 79 predominantly contains α helices instead of β-sheets, which are found in the corresponding region of Cry11A. This result was also found based on ab initio analysis, thus confirming the high prevalence of α helices in this region (Figure 3G).

The Interactions of Cry11Aa With ALP1 Are Conserved in Variants 8 and 23

The Ramachandran analysis of ALP1 showed that 76% of its residues were in favorable zones according to phi and psi angle positions and were involved in interactions with the parental Cry11Aa protein or its variants. None of the residues of ALP1 positioned in unfavorable zones (Supplementary Figure 4A) were involved in interactions with the parental Cry11Aa protein or its variants. The Z-score (~7.27) and the energy obtained from the ProSA-web server were below 0. Both
TABLE 1 | Molecular characteristics of Cry11 variants obtained via DNA shuffling.

| Variants   | GenBank accession number | Identity (%) cry11Aa | Mutation rate (%) | Del. (nt) | Subs. (nt) | Ins. (nt) | Subs. (nt) | Del. (nt) |
|------------|--------------------------|----------------------|-------------------|-----------|------------|-----------|------------|-----------|
| Variant 1  | MH068786                 | 84.6                 | 15                | 219       | 0          | 73        | 6          | 1         |
| Variant 8  | MH068787                 | 87.7                 | 13                | 219       | 6          | 0         | 13         | 0         |
| Variant 23 | MH068788                 | 98.9                 | 1                 | 9         | 6          | 5         | 2          | 0         |
| Variant 79 | MH068789                 | 80.1                 | 20                | 326       | 7          | 42        | 21         | 0         |
| Variant 81 | MH068790                 | 90.7                 | 8                 | 153       | 0          | 8         | 2          | 0         |
| TOTAL      |                          |                      |                   | 926       | 19         | 128       | 44         | 1         |

D, domain; Del, deletions; Subs., substitutions; Ins., insertions.

Amino acids F<sup>316</sup> and W<sup>319</sup> of variant 8 formed three hydrogen bonds with amino acids E<sup>498</sup> and Q<sup>400</sup> of ALP1 (Figure 4C) in contrast to the two hydrogen bonds found in the interaction between F<sup>389</sup> and Q<sup>391</sup> of the parental Cry11Aa protein with Y<sup>478</sup> and E<sup>105</sup> of ALP1 (Figure 4D). For variant 23, a single hydrogen bond between G<sup>388</sup> and Q<sup>486</sup> of ALP1 was found (Figure 4E), whereas variant 79 formed five hydrophobic interactions with nine amino acids of ALP1 (Figure 4F).

According to the data for Cry11Aa, the peptide 564<sup>RVQSQNSGNN</sup>573, located in the β<sub>18</sub>β<sub>19</sub> region of domain III, was found in all variants with exception of variant 79 (Supplementary Figure 2C). The LIGPLOT analysis of Cry11Aa showed three hydrophobic interactions of Cry11Aa with surface-exposed amino acids of ALP1 (Figure 5A). However, in variant 23, only one stable interaction through a hydrogen bond between R<sup>561</sup> of the variant protein and G<sup>261</sup> of ALP1 was observed (Figure 5B). In variant 8, two hydrogen bonds between R<sup>491</sup> of the variant protein and N<sup>229</sup> of ALP1 as well as three hydrophobic interactions of R<sup>491</sup>, V<sup>492</sup>, and Q<sup>493</sup> of the variant protein with V<sup>258</sup>, G<sup>257</sup>, and G<sup>261</sup> of ALP1 were formed (Figure 5C).
FIGURE 3 | Prediction of the 3D Structures of Cry11Aa and Variants 8, 23, and 79. (A) Conserved region of variant 8 in light blue and Cry11Aa in beige, RMSD: 1,084 with 247 aa. (B) Non-conserved region of variant 8 in light blue and Cry11Aa in beige. (C) Conserved region of variant 23 in light blue and Cry11Aa in beige, RMSD: 1,132 with 488 aa. (D) Non-conserved region of variant 23 in light blue and Cry11Aa in beige. (E) Conserved region of variant 79 in light blue and Cry11Aa in beige, RMSD: 1,084 with 247 aa. (F) Non-conserved region of variant 79 in light blue and Cry11Aa in beige. (G) Ribbon representation of the non-conserved region of variant 79 generated using the Robetta server.

DISCUSSION

Directed evolution approaches such as phage display, DNA shuffling and staggered extension process shuffling combined with Red/En homologous recombination have been proposed to increase the activity of Bt Cry toxins (Lucena et al., 2014). Other approaches based on phage-assisted continuous evolution (PACE) (Badran et al., 2016), in vitro template-change PCR (Shu et al., 2016), site-directed mutagenesis, and error-prone PCR have also been used successfully to identify novel
receptors expressed on the surface of insect midgut cells and to understand the effects of different cry gene mutations on the mechanism of action of Cry toxins (Lucena et al., 2014). In this study, we report five Cry toxin variants produced via reassembly during DNA shuffling of the cry11Aa gene that showed toxic activity against A. aegypti and C. quinquefasciatus larvae.

DNA shuffling was designed using internal sequences in the parental constructs, including the identification of specific sites for priming during reassembly. We used the upstream p19 gene to identify the genes that were reassembled from cry11Aa. A deletion of 246 bp downstream of the ATG start site was used to recognize those genes reassembled from cry11Ba, and an internal sequence corresponding to the specific primers was used to reassemble the variants from the construct containing the cry11Bb gene. The p19 gene was present upstream of the first ATG codon in variants 1, 8, 23, 28, 54, 79, and 81. According to sequence analysis, all variants displayed some degree of identity to cry11Aa; this observation indicated that all variants were preferentially reassembled from this parental gene during DNA shuffling (Table 1).

Variants 1 and 79 retained toxic activity against A. aegypti and C. quinquefasciatus despite lacking 8.0 and 11.9 kDa N-terminal regions, respectively. These variants exhibited the
highest mutation rates, 15 and 20%, respectively. The mutations found in variant 1 were not in regions involved in pore formation or toxin-receptor interactions, explaining the toxicity of variant 1 to *A. aegypti* based on bioassays. The insertion of 22 amino acids at the end of the C-terminus with seven substitutions did not affect the toxic activity of variant 1. The results of the bioassays with *C. quinquefasciatus* larvae showed that variant 1 has similar toxic activity to Cry11Aa but 7.63-fold lower toxic activity than Cry11Bb. In the other hand, toxic activity of variant 79 was unexpected, despite its numerous mutations, its toxic activity against *A. aegypti* and *C. quinquefasciatus* was high ([Figure 2](#fig2)). The deduced amino acid sequence of variant 79 did not show mutations in helices α-4 and α-5, which are implicated in pore formation, or in regions involved in toxin-receptor interactions, such as loop α8 and strand β4 ([Supplementary Figure 2C](#sup1)). However, among the 260 amino acids at the C-terminus that were modified in variant 79, we found five amino acids located in loop α-2 (F<sup>281</sup>, T<sup>282</sup>, Q<sup>283</sup>, W<sup>284</sup>, and F<sup>285</sup>) that generate nine hydrophobic interactions with ALP1 ([Figure 4F](#fig4)). These five amino acids in Cry11Aa and in variants 8 and 23 also form more stable interactions with ALP1 than hydrogen bonds ([Figures 4C–E](#fig4)). Additionally, according to 3D and secondary structural analyses, the non-conserved C-terminal region of variant 79 has an unusual

![Figure 5](#fig5) Interactions visualized in LigPlot of ALP1 against peptides of Cry11Aa and its variants 8 and 23. (A) Cry11Aa, (B) Variant 23, and (C) Variant 8.
and strand $\alpha$. Importantly, this variant did not show substitutions in regions significant differences against $\beta$. We suggest that the changes found in loop 2 of domain II and strands $\beta_{18}-\beta_{19}$ in domain III suggest a role in the stability of the interaction with ALP1. The formation of two hydrogen bonds in loop 2 of domain II produced by W$_{319}$ and F$_{320}$ with two amino acids of ALP1 including one hydrophobic interaction (Q$_{321}$) in the same cavity (Figure 4A), as well as two hydrogen bonds formed in domain III by R$_{491}$ with N$_{259}$ of ALP1 and three hydrophobic interactions, could explain the toxicity toward $A. aegypti$ mediated by ALP1 (Figure 4C).

The N-terminal deletions found in variants 1, 8, 23, 79, and 81 were between 3 and 108 amino acids in length (0.33 and 11.8 kDa). However, these deletions are not implicated in the differences in toxic activity against $A. aegypti$ and $C. quinquefasciatus$ larvae between variants. Only one study has reported an N-terminal deletion in the Cry11Aa toxin where a truncated protein lacking 9.6 kDa was non-toxic to $A. aegypti$ (Pang et al., 1992). In other Cry toxins such as Cry2a, deletion of 42 amino acids at the N-terminus increased the toxic activity against Spodoptera littoralis, Helicoverpa armigera, and Agrotis ipsilon (Mandal et al., 2007). Furthermore, in Cry1Ac, a deletion of 56 amino acids at the N-terminus, which included helix $\alpha$-1, increase the toxic activity against Pectinophora gossypii by 107-fold (Mandal et al., 2007) and against Plutella xylostella and Ostrinia nubilalis by 350-fold (Tabashnik et al., 2011). However, the toxic activity of Cry4Ba against $A. aegypti$ was abolished when more than 38 amino acids were removed from the N-terminus (Pao-intara et al., 1988) and, in the case of chimeric proteins formed by a fusion of N-terminus of Cry4Ba and the C-terminus of Cry1Ac, an increase of toxicity against $C. pipiens$ larvae was observed (Zghal et al., 2017). Although these findings confirm the importance of the N-terminal region in the toxicity of Cry proteins, the N-terminal deletions found in our variants did not affect their toxic activity against $A. aegypti$ or $C. quinquefasciatus$ larvae.

The difference in toxicity against $A. aegypti$ and $C. quinquefasciatus$ (Figure 2) could be explained by the presence of compounds either in midgut juice or membranes-bond proteases. Although the roles of these compounds and proteases have not been tested in variants obtained by DNA shuffling, there is evidence that the capacity to processing the protoxin depends on specific proteases located in the larval midgut and favored by alkaline conditions (Ben-Dov, 2014). In Cry11Aa toxin, the treatment with proteases generates fragments with different molecular weight and toxic activity against $A. aegypti$ (Revina et al., 2004; de Barros Moreira Beltrao and Silva-Filha, 2007), whereas in $C. quinquefasciatus$, the processing pattern differ from those that are active to $A. aegypti$ (Dai and Gill, 1993). Therefore, we suggest that the toxicity differences found in $A. aegypti$ and $C. quinquefasciatus$ could be explained by mechanism that dependent on the host.
Overall, the data presented in this report indicate that the N-terminal deletions observed in all variants did not affect their toxicity to *A. aegypti* or *C. quinquemaculatus*. Variant 8, which contained several substitutions in domains II and III, was the most interesting variant produced in this study due to its high toxicity to the two mosquito species. These findings confirm the importance of these domains in Cry toxin-receptor interactions and in Cry protein toxicity. The substitutions found in variants 8 and 23 provide new information about the role of loops 2 and 3 of domain II in Cry toxin-ALP1 interactions. Importantly, the α helix conformation of the C-terminus of variant 79 based on secondary structure analysis corresponds to a new protein structure with toxic activity. We believe that DNA reassembly via DNA shuffling following random fragmentation could be a good strategy to generate random mutations in specific cry genes to design new and more potent toxins.

**AUTHOR CONTRIBUTIONS**

AF conceived the study, was in charge of overall direction and planning and wrote the manuscript with input from all authors. MS-B carried out the experiments and worked out almost all of the technical details. GM performed directed evolution techniques and sequencing analysis. KR assisted with MS-B measurements and bioassays. SO contributed to electronic microscopy analysis and devised the project. RO carried out the molecular docking analysis. DG contributed to the analysis of 3D structure prediction and validation and secondary structure analysis of non-conserved regions. CM contributed to the interpretation of the 3D structure prediction and docking results.

**REFERENCES**

Abdullah, M. A., Alkate, O., Mohammad, M., McNall, R. J., Adang, M. J., and Dean, D. H. (2003). Introduction of CoxI toxicity into Bacillus thuringiensis Cry4Ba by protein engineering. *Appl. Environ. Microbiol.* 69, 5343–5353. doi: 10.1128/AEM.69.9.5343-5353.2003

Aronson, A. (2000). Incorporation of protease K into larval insect membrane vesicles does not result in disruption of integrity or function of the pore-forming *Bacillus thuringiensis* delta-endotoxin. *Appl. Environ. Microbiol.* 66, 4568–4570. doi: 10.1128/AEM.66.10.4568-4570.2000

Badran, A. H., Guzov, V. M., Huai, Q., Kemp, M. M., Vishwanath, P., Kain, W., et al. (2016). Continuous evolution of *Bacillus thuringiensis* toxins overcomes insect resistance. *Nature* 533, 58–63. doi: 10.1038/nature17938

Ben-Dov, E. (2014). Bacillus thuringiensis subsp. israelensis and its Dipteran-Specific Toxins. *Toxins* 6, 1222–1243. doi: 10.3390/toxins6041222

Berry, C., O’Neill, S., Ben-Dov, E., Jones, A. F., Murphy, L., Quail, M. A., et al. (2002). Complete sequence and organization of pBtox, the toxin-coding plasmid of *Bacillus thuringiensis* subsp. *israelensis*. *Appl. Environ. Microbiol.* 68, 5082–5095. doi: 10.1128/AEM.68.10.5082-5095.2002

Bradford, M. M. (1976). A rapid and sensitive method for the quantitation of microgram quantities of protein utilizing the principle of protein-dye binding. *Anal. Biochem.* 72, 248–254. doi: 10.1016/0003-2697(76)90527-3

Bravo, A. (1997). Phylogenetic relationships of *Bacillus thuringiensis* delta-endotoxin family proteins and their functional domains. *J. Bacteriol.* 179, 2793–2801. doi: 10.1128/JB.179.9.2793-2801.1997

Bravo, A., Gill, S. S., and Soberon, M. (2007). Mode of action of *Bacillus thuringiensis* Cry and Cyt toxins and their potential for insect control. *Toxicon* 49, 423–435. doi: 10.1016/j.toxicon.2006.11.022

Bravo, A., Gomez, I., Conde, J., Munoz-Garay, C., Sanchez, J., Miranda, R., et al. (2004). Oligomerization triggers binding of a *Bacillus thuringiensis* Cry1Ab pore-forming toxin to aminopeptidase N receptor leading to insertion into membrane microdomains. *Biochim. Biophys. Acta* 1667, 38–46. doi: 10.1016/j.bbamem.2004.08.013

**FUNDING**

This work was supported by grants from the Colombian Department of Science, Technology and Innovation, COLCIENCIAS, 1299–12-16813 and 5201-545-31565.

**ACKNOWLEDGMENTS**

We thank Andres Castañeda, Laura Natalia Parra, Luz Dary Rodriguez, Liliana Torcoroma, and Juan David Estupiñan for technical assistance.

**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb.2018.02461/full#supplementary-material
Florez et al.

Cry11 Variants Obtained via DNA Shuffling

Bacillus thuringiensis

Pardo-Lopez, L., Soberon, M., and Bravo, A. (2013).

Li, J. D., Carroll, J., and Ellar, D. J. (1991). Crystal structure of insecticidal delta-endotoxin from Bacillus thuringiensis. *Acta Crystallogr. D. Biol. Crystallogr.* 57, 1101–1109. doi: 10.1107/S090744010001816

Gardner, L. M., Chen, N., and Sarkar, S. (2016). Global risk of Zika virus depends critically on vector status of *Aedes albopictus*. *Lancet Infect. Dis.* 16, 522–523. doi: 10.1016/S1473-3099(16)00176-6

Morse, R. J., Yamamoto, T., and Stroud, R. M. (2001). Structure of Cry2Aa suggests a critical role for the ATP-binding protein from *Bacillus thuringiensis* in crystal toxins for mosquito larvae. *Proc. Natl. Acad. Sci. U.S.A.* 98, 154–159. doi: 10.1073/pnas.98.13.154

Pang, Y., Frutos, R., and Federici, B. A. (1992). Synthesis and toxicity of full-length and truncated bacterial Cry1D2 mosquitocidal proteins expressed in lepidopteran cells using a baculovirus vector. *J. Gen. Virol.* 73(Pt 1), 89–101. doi: 10.1099/0022-1317-73-1-89

Pao-intara, M., Angkunhasombat, C., and Panyim, S. (1988). The mosquito larvicidal activity of 130 kDa delta-endotoxin of *Bacillus thuringiensis* var. *israelensis* resides in the 72 kDa amino-terminal fragment. *Biochem. Biophys. Res. Commun.* 153, 294–300. doi: 10.1016/S0006-291X(88)81221-X

Pardo-Lopez, L., Soberon, M., and Bravo, A. (2013). *Bacillus thuringiensis* insecticidal three-domain Cry toxins: mode of action, insect resistance and consequences for crop protection. *FEBS Microbiol. Rev.* 37, 3–22. doi: 10.1111/j.1574-6976.2012.00341.x

Perez, C., Fernandez, I. E., Sun, J., Felch, J. L., Gill, S. S., Soberon, M., et al. (2005). *Bacillus thuringiensis* subsp. *israelensis* Cry11Aa synergizes Cry11Aa toxin by functioning as a membrane-bound receptor. *Proc. Natl. Acad. Sci. U.S.A.* 102, 18303–18308. doi: 10.1073/pnas.0505491102

Restrepo, N., Gutierrez, D., Patino, M. M., Thiery, L., Delecourse, A., and Orduz, S. (1997). Cloning, expression and toxicity of a mosquitoicidal toxin gene of *Bacillus thuringiensis* subsp. medellin. *Mem. Inst. Oswaldo Cruz* 92, 257–262. doi: 10.1590/S0074-02461997000200021

Reina, L. P., Kostina, L. I., Ganushkin, L. A., Mikhailova, L. A., Zulunin, I. A., and Chestukhina, G. G. (2004). Reconstruction of *Bacillus thuringiensis* sp. *israelensis* Cry11Aa endotoxin from fragments corresponding to its N- and C-moieties restores its original biological activity. *Biochemistry* 69, 181–187.

Shu, C., Zhou, J., Crickmore, N., Li, X., Song, F., Liang, G., et al. (2016). In vitro template-change PCR to create single crossover libraries: a case study with *B. thuringiensis* Cry2A toxins. *Sci. Rep.* 6:23356. doi: 10.1038/srep23356

Stemmer, W. P. (1994). DNA shuffling by random fragmentation and reassembly: in vitro recombination for molecular evolution. *Proc. Natl. Acad. Sci. U.S.A.* 91, 10747–10751. doi: 10.1073/pnas.91.22.10747

Tabashnik, B. E., Huang, F., Ghimire, M. N., Leonard, B. R., Siegfried, B. D., Rasangamay, M., et al. (2011). Efficacy of genetically modified Bt toxins against insects with different genetic mechanisms of resistance. *Nat. Biotechnol.* 29, 1128–1131. doi: 10.1038/nbt.1988

Thompson, J. D., Higgins, D. G., and Gibson, T. J. (1994). CLUSTAL W: improving the sensitivity of progressive multiple sequence alignment through sequence weighting, position-specific gap penalties and weight matrix choice. *Nucl. Acids Res.* 22, 4673–4680. doi: 10.1093/nar/22.22.4673

Tietzarkin, K. A., Chen, R., and Weaver, S. C. (2016). Interspecies transmission and chikungunya virus emergence. *Curr. Opin. Virol.* 16, 143–150. doi: 10.1016/j.coviro.2016.02.007

Vachon, V., Laprade, R., and Schwartz, J. L. (2012). Current models of the mode of action of *Bacillus thuringiensis* insecticidal crystal proteins: a critical review. *J. Invertebr. Pathol.* 111, 1–12. doi: 10.1016/j.jip.2012.05.001

van Frankenhuyzen, K. (2009). Insecticidal activity of *Bacillus thuringiensis* crystal proteins. *J. Invertebr. Pathol.* 101, 1–16. doi: 10.1016/j.jip.2009.02.009

Wirth, M. C., Georgihoiu, G. P., and Federici, B. A. (1997). Cry1A enables CryIV endotoxins of *Bacillus thuringiensis* to overcome high levels of CryIV resistance in the mosquito, *Culex quinquefasciatus*. *Proc. Natl. Acad. Sci. U.S.A.* 94, 10536–10540. doi: 10.1073/pnas.94.20.10536

Zghal, R. Z., Elleuch, J., Ben Ali, M., Darriet, F., Rebai, A., Chandre, F., et al. (2017). Towards novel Cry toxins with enhanced toxicity/broader: a new chimeric Cry4Ba/Cry1Ac toxin. *Appl. Microbiol. Biotechnol.* 101, 113–122. doi: 10.1007/s00253-016-7766-3

Zhang, L., Zhao, G., Hu, X., Liu, J., Li, M., Batool, K., et al. (2017). Cry11Aa interacts with the ATP-binding protein from *Culex quinquefasciatus* to improve the toxicity. *J. Agric. Food Chem.* 65, 10884–10890. doi: 10.1021/acs.jafc.7b04427

Zhang, Q., Hua, G., and Adang, M. J. (2017). Effects and mechanisms of Bacillus *thuringiensis* crystal toxins for mosquito larvae. *Insect Sci.* 24, 714–729. doi: 10.1111/j.1744-7917.12401

Zhang, X., Candás, M., Griko, N. B., Taussig, R., and Bulla, L. A. Jr. (2006). A mechanism of cell death involving an adenylyl cyclase/PKA signaling pathway is induced by the Cry1Ab toxin of *Bacillus thuringiensis*. *Proc. Natl. Acad. Sci. U.S.A.* 103, 9897–9902. doi: 10.1073/pnas.0604017103

Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Copyright © 2018 Florez, Suarez-Barrera, Morales, Rivera, Orduz, Ochoa, Guerra and Muskus. This is an open-access article distributed under the terms of the Creative Commons Attribution License (CC BY). The use, distribution or reproduction in other forums is permitted, provided the original author(s) and the copyright owner(s) are credited and that the original publication in this journal is cited, in accordance with accepted academic practice. No use, distribution or reproduction is permitted which does not comply with these terms.