Presence and significance of Bacillus thuringiensis Cry proteins associated with the Andean weevil Premnotrypes vorax (Coleoptera: Curculionidae)

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Abstract: The Andean weevil *Premnotrypes vorax* represents an important cause of damage to Colombian potato crops. Due to the impact of this plague on the economy of the country, we searched for new alternatives for its biological control, based on the entomopathogenic bacteria *Bacillus thuringiensis*. A total of 300 *B. thuringiensis* strains obtained from potato plantations infested with *P. vorax* were analyzed through crystal morphology, SDS-PAGE, PCR and bioassays. We used site-directed mutagenesis to modify the Cry3Aa protein. Most of the *B. thuringiensis* isolates had a bipyramidal crystal morphology. SDS-PAGE analyses had seven strains groups with δ-endotoxins from 35 to 135 kDa. The genes *cry* 2 and *cry* 1 were significantly more frequent in the *P. vorax* habitat (PCR analyses). Three mutant toxins, 1 (D354E), 2 (R345A, ΔY350, ΔY351), and 3 (Q482A, S484A, R485A), were analyzed to assess their activity against *P. vorax* larvae. Toxicity was low, or absent, against *P. vorax* for isolates, wild type *cry* 3Aa and *cry* 3Aa mutants. The genetic characterization of the collection provides opportunities for the selection of strains to be tested in bioassays against other insect pests of agricultural importance, and for designing Cry proteins with improved insecticidal toxicity. Rev. Biol. Trop. 57 (4): 1235-1243. Epub 2009 December 01.

Key words: native strain, Andean weevil, Cry3Aa protein, potato crop, site-directed mutagenesis.

It has been estimated that 9000 species of insect pests affect commercial crops in the world. Sustainable control of insects in agriculture is very important since it was estimated that chemical control cost 7500 million dollars (Arrieta & Espinoza 2006). In addition, the use of synthetic insecticides is not recommended because of the long residual action and toxicity to a wide spectrum of organisms, including human (Song *et al.* 2008). Consequently, interest has developed in the use of alternative strategies for biological control, such as *Bacillus thuringiensis* (*Bravo et al.* 1998).

The entomopathogenic activity of Gram positive *B. thuringiensis* is mainly due to the

synthesis of large quantities of insecticidal crystal proteins (ICPs), or δ -endotoxins, during sporulation that can be distinguished as distinctively shaped crystals under electronic scanning microscopy (Kati *et al.* 2007). These inclusions are comprised of toxins known as Cry and Cyt proteins. Up to date, more than 300 toxins have been cloned, sequenced, and classified into 55 groups of Cry proteins and two groups of Cyt proteins based on the amino acid homology (Crickmore *et al.* 2008). Cry proteins have been used as biopesticide sprays on a significant scale for more than 50 years, and their safety has been demonstrated (Schnepf *et al.* 1998). Some of the ICP genes have

been applied for bioengineered crop protection, resulting in transgenic crop plants with excellent insect protection (Song et al. 2008). The active spectra of ICPs includes larvae of Lepidoptera, Coleoptera, Diptera, Hymenoptera, Homoptera, Mallophaga and Orthoptera (López-Pazos & Cerón 2007). B. thuringiensis strains are distributed worldwide in soil, stored products, insects, insect breeding environments, and the phylloplane (Mahalakshmi et al. 2005). It is still necessary to search for more strains and toxins, since a significant number of pests are not controlled with the available. It is also important to provide alternatives for insect resistance, especially with regard to the transgenic crops (Bravo et al. 1998).

The Andean potato weevil (or White Worm as named in Colombia) Premnotrypes vorax (Coleoptera: Curculionidae) is a major cause of damage to potato crops in the Andean regions of Colombia, affects 75% of cultivated area, and 10-90% of the potato harvest losses are due to this insect (Herrera 1997). The aim of this study was to analyze the effect of B. thuringiensis on the survival of P. vorax. Firstly, to characterize a *B. thuringiensis* native strain collection from P. vorax ecosystem, and secondly, to identify, through bioassays, the biological activity of B. thuringiensis native strains against Andean potato weevil. Also, the toxicity of the Cry 3A δ -endotoxin in P. vorax larvae was evaluated by engineering the specific residues D354, R345, Y350 and Y351 (loop I in the binding domain), and Q482, S484 and R485 (loop III) in domain II (Wu & Dean 1996, Wu et al. 2000).

MATERIALS AND METHODS

Samples collection: Different colombian potato crops (*Solanum tuberosum* spp. *andigena* var. Diacol Capiro R-12) (Andean region) were sampled randomly chosen. They were located in Cerro La Conejera ecological park (Geographic coordinates 4°47'14,3"-5°35'13" N, 74°4'3,4"-73°15'49,7" W) and Romeral municipal rural settlement of Sibate municipality (Geographic coordinates 4°26'3" N,

74°14'8" W). No commercial *B. thuringiensis* based product had been used in any of the sampled areas. Two hundred grams of samples (soil and stored products) was collected with a sterile spatula at 10 cm deep. Dead *P. vorax* insects were obtained from soil and tubers. Samples were stored in labeled sterile plastic bags at room temperature until processed.

Bacterial strains: *B. thuringiensis* subsp *kurstaki* (HD1), *B. thuringiensis* subsp *aizawai* (HD137), *B. thuringiensis* subsp. *tenebrionis* and *B. thuringiensis* subsp. *san diego* were supplied by *Bacillus* Genetic Stock Center (The Ohio State University, Columbus, OH). Acetate selection method developed by Travers *et al.* (1987) was used to isolate the native strains.

Phase contrast microscopy and SDS-PAGE: Parasporal inclusions of each isolate were classified through phase contrast microscopy in one of the following types: bipyramidal, rectangular, cuboidal, spherical, triangular, and amorphous. Spore-crystal suspensions were used for protein analyses in 10% SDS-polyacrylamide gels.

PCR cry gene analysis: Native B. thuringiensis strains were characterized in terms of cry gene presence. Those strains which reacted with the general cry primers were characterized subsequently through additional PCRs with specific primers, in order to identify cry 1Aa, cry 1Ab, cry 1Ac, cry 1Ba, cry 1Da, cry 2, and cry 3 and cry 7 genes presence. The Cry primers used were designed by Cerón et al. (1994, 1995) and Ben-Dov et al. (1997). DNA preparation and PCRs were made in a thermocycler (Perkins-Elmer model 480) using Cerón et al. (1995) methods.

Mutagenesis and protein isolation: Three Cry 3Aa-mutants have been described elsewhere (Wu *et al.* 2000, Wu & Dean 1996) and were constructed by introduction of the mutations D354E (loop 1 in domain II), R345A, ΔΥ350, ΔΥ351 (loop 1 in domain II), Q482A, S484A and R485A (loop 3 in domain

II) in expression plasmid pMH10 derived from pKK233-2 (Pharmacy LKB Biotechnology), using the Kit GeneTailor™ Site-Directed Mutagenesis System (Invitrogen). After mutagenesis, single-strand DNA sequencing was carried out (Macrogen, Korea).

Wild-type *cry* 3Aa and mutant genes were expressed in *Escherichia coli* XL-1-Blue. Cells were grown for 48 h in 500 ml of TB medium (Sambrook *et al.* 1989) containing 100 μg/ml ampicillin. Crystal extracts were prepared and solubilized in 50 mM sodium carbonate buffer (pH 10.0). The concentration of solubilized protoxin was determined by Lowry assay (Lowry *et al.* 1951). When necessary, activation of protoxin was performed by adding trypsin at a ratio of 1:10 (trypsin:protoxin, w/v) and incubating for 2 h at 37 °C. The activated toxin was used immediately. Protoxin and toxins were analyzed by SDS-polyacrylamide gel (8%) electrophoresis (SDS-PAGE).

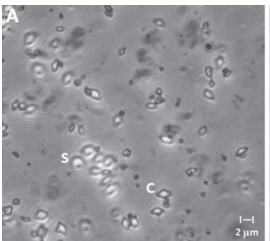
Bioassays: The biopesticide activity of different *B. thuringiensis* strains, Cry 3Aa and mutants, was evaluated on first instar larvae of *P. vorax* from a laboratory colony. Potato cubes were surface contaminated with a toxin concentration of 10000 (bacterial strains) and 2000 (Wild-type *cry* 3Aa and mutants) ng/cm²

diluted in 60 μ l deionized water. Cubes were distributed in plate wells and each one was infested with one *P. vorax* first instar larvae. Plates were sealed and incubated into a 18°C, $60 \pm 5\%$ relative humidity, and 12:12 h light:dark photoperiod room. In insect bioassay 72 neonate larvae per treatment were used and mortality was recorded after 7 days (Martínez & Cerón 2002).

RESULTS

A total of 300 (42.85%, out of 700) strains, of *P. vorax* environment samples examined showed colonial and microscopic morphology identical to *B. thuringiensis*. The value of Bt index, according to Iriarte *et al.* (1998), calculated as a number of *B. thuringiensis* isolates, divided by the total number of *Bacillus* strains, was assessed to be 0.43. Phase contrast microscopy demonstrated that all strains presented diverse crystal morphologies: bipyramidal, cuboidal, point and pleomorphic. Isolates with bipyramidal and cuboidal-shape crystals predominated on the soil and dead insects (63% and 25% respectively) (Fig. 1).

All *B. thuringiensis* isolates of their crystal protein products were further characterized by SDS-PAGE. The results revealed that the



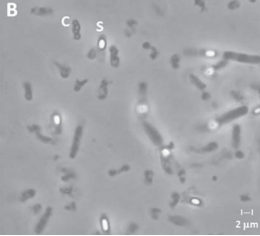


Fig. 1. Phase contrast micrograph of *B. thuringiensis* isolates showing the bipyramidal and cuboidal crystals. (A) Isolate IBUN27p, (B) Isolate IBUN72.1. (S) Spore, (C) Crystal.

colombian *P. vorax* environment strains synthesize a protein, or group of proteins, with a molecular mass between 35 and 135 kDa (Fig. 2). The highest percentage of isolates contained two protein types, 130 and 60 kDa; and, in addition to these proteins, an intermediate molecular weight protein of approximately 100 kDa. However, insects isolates possessing others toxins (125, 40 and 35 kDa) combinations were represented by less than 2%.

The profiles of all PCR products were compared with those of standard strains. An isolate was considered to contain a determined gene only when the amplification product was of the expected size. Analysis of the B. thuringiensis isolates indicated that most contained several cry 2 genes, as well as cry 1 (Table 1). Within the 300 strains, 212 isolates (70.6%) harbored cry 1, cry 2, cry 3 or cry 7 gene. Several strains carried the cry 1-type genes (107 or 35.6%) or cry 2 gene (134 or 44.6%) whereas only 51 strains (17%) contained the cry 3- cry 7 genes (Table 1). Eighty-eight of the 300 isolates (29.3%) did not harbor any of the cry 1, cry 2 or cry 3- cry 7 genes. All of the different Cry 1 subfamilies were observed with different frequencies. Among them, cry 1A in 95 strains (31.6%) and *cry* 1B in 50 strains (16.6%) were considered abundant, while cry 1D in 39

strains (13%) and cry 1C in 21 strains (7%) were less prevalent (Table 1). The study of cry gene combinations in cry gene-containing isolates showed that more than 49% (147) harbored more than one type of cry gene. About 74.7% of the *cry* 1 gene-containing isolates also harbored a cry 2 gene while 59.7% of the cry 2 gene-containing isolates also harbored a cry 1-type gene and about 5.5% of the cry 3- cry 7 gene-containing isolates also harbored cry 1 gene. However, only 2.8% of the cry 1 gene-containing isolates also harbored a cry 3cry 7 gene. These results indicated that cry 1 and cry 2 genes have a high tendency to occur together and the cry 3- cry 7 gene frequently occurred alone.

The expression of the *cry* 3Aa gene in XL-1-Blue was analyzed by sequencing (data not shown) and SDS-PAGE (Fig. 3). The predominant band of expressed protein visualized on SDS-PAGE was 73 kDa. These crystal proteins could be further reduced to 67 and 55 kDa if treated with trypsin (Fig. 3). All mutants were highly expressed as inclusion bodies in *E. coli*, similar to the wild type Cry 3Aa.

To test the toxicity of *B. thuringiensis* native strains, Cry 3Aa and mutants, first instar larvae of *P. vorax* were exposed to media whose surface was spread with 10 000 ng/cm² protoxins

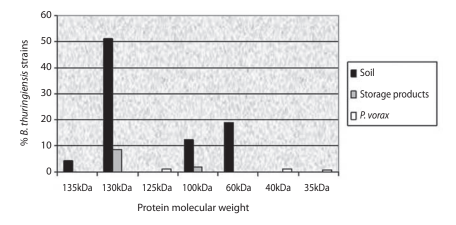


Fig. 2. Protein band patterns in *B. thuringiensis* strains from *P. vorax* ecosystem. (\blacksquare) *B. thuringiensis* strains isolated from soil; (\blacksquare) *B. thuringiensis* strains isolated from storage products; (\square) *B. thuringiensis* strains isolated from dead insects. Polypeptides were visualized by Coomassie blue staining.

TABLE 1 The cry gene combination profiles present in the B. thuringiensis strain collection from P. vorax ecosystem (n = 300)

No.	cry gene profiles	No. of isolates	Freq. (%)
1	cry 3, cry 7	51	17
2	cry 1Aa, cry 1Ab, cry 1Ba, cry 1Ca, cry 1Da	5	1.6
3	cry 1Aa, cry 1Ab, cry 1Ba, cry 1Ca, cry 1Da, cry 2	11	3.6
4	cry 1Aa, cry 1Ab, cry 1Ba, cry 1Ca, cry 1Da, cry 3, cry 7	1	0.3
5	cry 1Aa, cry 1Ba, cry 1Ca, cry 1Da	1	0.3
6	cry 1Aa, cry 1Ba, cry 1Ca, cry 1Da, cry 3, cry 7	2	0.6
7	cry 1Ba, cry 1Ca, cry 1Da	1	0.3
8	cry 1Aa, cry 1Ab, cry 1Ac, cry 2	5	1.6
9	cry 1Aa, cry 1Ab, cry 1Ac, cry 1Ba, cry 1Da	1	0.3
10	cry 1Aa, cry 1Ab, cry 1Ac, cry 1Ba, cry 1Da, cry 2	17	5.6
11	cry 1Aa, cry 1Ab, cry 1Ac	4	1.3
12	cry 1Aa, cry 1Ac, cry 2	8	2.6
13	cry 1Aa, cry 2	2	0.6
14	cry 1Ab, cry 1Ac, cry 2	8	2.6
15	cry 1Aa, cry 1Ab	1	0.3
16	cry 1Aa, cry 1Ab, cry 2	2	0.6
17	cry 1Ab	2	0.6
18	cry 1Ab, cry 2	25	8.3
19	cry 1Ba	9	3
20	cry 1Ba, cry 2	2	0.6
21	cry 2	54	18
22	NA	88	29.3

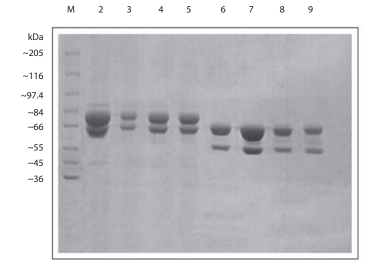


Fig. 3. Coomassie blue stained 12.5% SDS-PAGE comparing the expression of Cry3Aa proteins and trypsin digested toxins. Lanes: (1) Molecular weight marker (M); (2) wild-type Cry3Aa purified from *E. coli*; (3) loop 3 triple mutant (Q482A, S484A, R485A); (4) loop 1 triple mutant (R345A, Δ Y350, Δ Y351); (5) loop 1 mutant (D354E); (6) loop 3 triple mutant treated with trypsin; (7) wild-type Cry3Aa treated with trypsin; (8) loop 1 mutant treated with trypsin; (9) loop 1 triple mutant treated with trypsin. Each lane contained 12 μ g of protein.

TABLE 2 Classification of the B. thuringiensis isolates in groups according to their toxicity levels against P. vorax (10000 ng/cm²)

Sample habitat	Number of isolates causing a mortality of 0–10% (three assays)	Number of isolates causing a mortality of 10–20% (three assays)	Total isolates examined
Soil	224	17	241
Storage products	36	11	47
Dead insects	12	0	12
Total	272	28	300

TABLE 3

Toxicity against P. vorax of Cry3Aa wild type and mutants proteins (2000 ng/cm²)

Proteins	Mortality (%) with protoxin (three assays)		Mortality (%) with toxin (three assays)		Protoxin variation coefficient	Toxin variation coefficient		
Cry3Aa wild type	12.9	15.62	15.62	9.37	9.37	9.37	0.10	0
R345A, ΔΥ350, ΔΥ351	6.25	9.37	9.37	15.62	21.87	21.87	0.21	0.18
Q482A, S484A, R485A	9.37	6.25	9.37	9.37	12.5	9.37	0.21	0.17
D354E	6.25	6.25	9.37	15.62	15.62	12.5	0.2	0.12

and activated toxins, and 2 000 ng/cm² recombinant protoxins and toxins. Larvae exposed to toxin treated media exhibited a mortality percentage lower than 22% (Tables 2 and 3).

DISCUSSION

Since potatoes are the most important crop in the Andean region, all the samples collected were related to this crop, and 100% of them had an origin associated with the important pest in the area: the Andean weevil P. vorax. Here, we determined the presence of different cry genes within the P. vorax ecosystem collection. The selection of B. thuringiensis isolates was based principally on phase-contrast microscopy. The Bt index was high (0.43). Probably, humidity and excrements produced by P. vorax larvae in the galleries of the infested potatoes make a favorable environment for B. thuringiensis colonization. Similar to the lepidopteran-active reference strains (especially B. thuringiensis kurstaki), most of the isolates exhibited two major polypeptides

with a relative molecular mass in the range of 130 to 140 kDa and approximately 70 kDa, respectively. The SDS-PAGE analysis showed other proteins (125, 100, 40 and 35 kDa from soil, store products and dead *P. vorax* samples) in *B. thuringiensis* strains, thus, these results predicted that our insect isolates contain possible novel Coleopteran-specific proteins. Based on the varied parasporal polypeptide patterns of the isolates, we conclude that this *B. thuringiensis* collection has quite an attractive diversity of Cry proteins.

The important biological activity of *B. thuringiensis* isolates appears to be derived from the combined attributes of different Cry toxins formed into a proteinaceous body. Determination of their *Cry* gene contents is useful for the prediction of their toxicity. *B. thuringiensis* strains recovered from *P. vorax* environment were diverse in terms of their *cry* gene contents. According to PCR analysis, the *cry* 1, *cry* 2 and *cry* 3- *cry* 7 genes were observed in our *B. thuringiensis* collection but in different frequency. The *cry* 2 genes were the most

frequently found in the native isolates collection (Table 1). A high frequency of cry 1 genes seems to be common to all B. thuringiensis strain collections analyzed so far (Bravo et al. 1998, Hernández et al. 2005, Cinar et al. 2008). It is possible that *cry* 1 gene-containing strains may be more abundant. We cannot exclude the probability that the high percentage of cry 2 genes in B. thuringiensis isolates may have been biased because of the procedure used for selection (Bravo et al. 1998). The second most abundant genes in the bacterial strain collection were the cry 1 genes and then the cry 3- cry 7 genes (Table 1). In our study, the cry 2 gene (44.6%) was detected more frequently than in other reports. For instant Ben-Dov et al. (1997) found that strains containing cry 1 genes were the most abundant; Bravo et al. (1998) detected 49.5% of cry 1 gene; Uribe et al. (2003) reported 73% of cry 1 gene; Wang et al. (2003) detected 76.5% of cry 1 gene and 70% of cry 2 gene; Thammasittirong & Attathom (2008) reported strains containing cry 1-type genes (81.3%) at the same frequency as strains harboring cry 2 gene (80.6%). There is low presence of cry 3- cry 7 gene in potato crops infested with P. vorax. It is important to mention that many of the isolates harbored more than one cry gene, suggesting that B. thuringiensis strains have a high frequency of genetic information exchange. Twenty-one distinct cry gene profiles were identified in our collection (Table 1), which indicates the high diversity in the *cry* gene contents of the *B*. thuringiensis strains. The analysis of the genes showed that the cry 1 and cry 2 genes were most often present together and most of the bacterial strains contained more than one type of lepidopteran-active cry gene. Some strains containing combinations of cry genes that were less frequently observed, such as lepidopteranactive cry 1 genes and coleopteran-active cry 3- cry 7 genes, were identified (Table 1), these isolates are good possibilities in the search for biological control agents with a wider spectrum of activity. Other groups have reported the presence of cry 1 genes and cry 3- cry 7 genes in the same B. thuringiensis strain (Bravo et al.

1998), suggesting that strains with dual activity are also present in other regions. The great variability and distribution of cry gene content in B. thuringiensis strains even from the same country is well recognized, and the variations were likely associated with differences in the biological, geographical and ecological properties of the collected areas. The above evidences show the importance of establishing collections of this microorganism in different countries for developing strategies of biological insect control. Finally, 88 isolates did not react with any PCR primer. These isolates are candidates for harboring putative different cry genes. The identification of different B. thuringiensis isolates could be the first phase in the project for finding major toxicities, since different proteins may be toxic for new insects. The characterization of different cry genes should be encouraged once the susceptible insect is identified and more evidence on the potential of different proteins as pest control agents is available.

Under the conditions tested both native strain and recombinant and mutant cry 3Aa proteins presented in bioassays a mortality below 22% with the evaluated dose (Tables 2 and 3). The original objective of this project was to select strains, or recombinant proteins, with activity against P. vorax. Unfortunately, none of the isolates, or wild type Cry 3Aa and mutants, was toxic against this weevil. In spite of the fact that cry3 proteins bind to P. vorax brush border membrane vesicles (Martinez & Cerón 2004) it is still difficult to find active proteins to this pest. This result is not at all unexpected, as most screening studies of susceptibility to B. thuringiensis proteins have shown a frequent insecticidal toxicity of this organism to lepidopteran species, while only very few isolates and toxins have been found active against Coleoptera (Hernandez et al. 2005). Also, we tested the theory that surface loops in domain II of Cry 3Aa are involved in receptor binding of P. vorax larvae. Previous results revealed that substitutions within surface exposed loops of domain II on the cry 3Aa can affect reversible and irreversible receptor binding in coleopteran (Wu & Dean 1996, Wu

et al. 2000). However, none of the mutants was toxic to P. vorax larvae. The implications of our studies are important for new design of δ -endotoxins that overcome insect resistance to these toxins and for altered or improved insecticidal activity, as has been achieved in other Cry proteins.

RESUMEN

El gorgojo andino Premnotrypes vorax es una causa importante de daño en los cultivos colombianos de este tubérculo. Debido al impacto que esta plaga tiene sobre la economía del país, nos interesamos en buscar alternativas nuevas para el control biológico de P. vorax, basadas en la bacteria entomopatógena Bacillus thuringiensis. Se recolectaron un total de 300 cepas de B. thuringiensis a partir de plantaciones de papa infestadas con *P. vorax*, las cuales fueron analizadas por medio de la morfología del cristal, SDS-PAGE, PCR y ensayos biológicos. La mayoría de los aislamientos de B. thuringiensis presentaron cristales bipiramidales. Los análisis de SDS-PAGE indicaron la presencia de siete grupos de cepas con δ-endotoxinas que variaban entre 35 a 135 kDa. Las pruebas con PCR demostraron que los genes cry 2 y cry 1 fueron significativamente más frecuentes en el medioambiente de P. vorax. Además, se utilizó la mutagénesis sitio-dirigida para modificar la proteína Cry3Aa. Se analizaron tres toxinas mutantes, 1 (D354E), 2 (R345A, \(\Delta \text{Y350}, \(\Delta \text{Y351} \), y 3 (Q482A, S484A, R485A), para determinar su actividad contra larvas de P. vorax. Los ensayos de toxicidad señalaron escasa, o nula, actividad hacia P. vorax tanto para las cepas, la toxina Cry3Aa de referencia y las proteínas Cry3Aa mutantes. La caracterización genética de la colección puede proveer oportunidades para la selección de cepas que pueden evaluarse por medio de bioensayos contra otros insectos-plaga de importancia agrícola, y para el diseño de proteínas Cry con actividad toxica mejorada.

Palabras clave: cepa nativa, gorgojo andino, proteína Cry3Aa, cultivo de papa, mutagénesis sitio-dirigida.

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