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Characterization of a *Bacillus thuringiensis* strain collection isolated from diverse Costa Rican natural ecosystems

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Abstract: Costa Rican natural ecosystems are among the most diverse in the world. For this reason, we isolated strains of the entomopathogenic bacteria *Bacillus thuringiensis* (*Bt*) to determine their diversity, distribution and abundance. A total of 146 *Bt* strains were obtained from environmental samples collected from diverse natural ecosystems and life zones of Costa Rica. We recovered *Bt* strains from 71%, 63%, 61% and 54% of soil samples, fresh leaves, other substrates and leaf litter respectively. *Bt* was isolated in 65% of the samples collected in the humid tropical forest in national parks (Braulio Carrillo, Gandoca Manzanillo, Sierpe, Hitoy Cerere, and Cahuita), and in 59% of the samples collected in the dry tropical forest (Parque Nacional Marino las Baulas, Palo Verde and Santa Rosa). In the very humid tropical forest (Tortuguero) *Bt* was isolated in 75% of the samples and in the very humid tropical forest transition perhumid (Carara) it was found in 69% of the samples. The strains exhibit a diverse number, size and morphology of parasporal inclusion bodies: irregular (47%), oval (20%), bipyramidal (3%), bipyramidal and cubic (1%), bipyramidal, oval and irregular (5%) and bipyramidal, oval and cubic crystals (2%). Strains isolated from Braulio Carrillo, Tortuguero and Cahuita, presented predominantly irregular crystals. On the other hand, more than 60% of the isolates from Terraba-Sierpe and Hitoy-Cerere had medium oval crystals. Strains from Gandoca-Manzanillo, Palo Verde and Carara presented mainly combinations of oval and irregular crystals. Nevertheless, the greatest diversity in crystal morphology was observed in those from Santa Rosa, Llanos del Río Medio Queso and Parque Marino las Baulas. Protein analyses of the crystal-spore preparations showed δ -endotoxin with diverse electrophoretic patterns, with molecular weights in the range of 20 to 160 kDa. Fifty six percent of the strains amplified with the *cry2* primer, 54% with *vip3*, 20% with *cry1*, 9% with *cry3-cry7* and 8% with *cry8*. The *cry11* and *cyt* genes were found in 8% and 7% of the strains, respectively. When analyzed with specific primers for the *cry1* subfamily, 13 different genetic profiles were obtained. In addition, twenty-four strains did not amplify with any of the primers used, suggesting they contain novel *cry* genes. The diversity of *Bt* genes found in this collection indicates it could have great potential for the control of different species of insect pests. The toxicological characterization of the strains by bioassays against important insect pests will provide useful information about their potential use for the formulation of biological insecticides and their respective *cry* and *vip* genes for the transformation of crops to confer resistance to insects. Rev. Biol. Trop. 54(1): 13-27. Epub 2006 Mar 31.

Key words: *Bacillus thuringiensis*, crystals, *cry*, *vip* genes, δ -endotoxins, national parks, Costa Rica.

Costa Rica is one of the 20 richest countries in terms of biodiversity, harboring 4% of the total biodiversity of the world (Obando 2002). The geographic location of the country in the neotropics, its geology, two closely separated coasts, a complex mountain range

system and diverse microclimates and ecosystems are some of the determining factors for the biological diversity of the country. The neotropics provide a rich source for the discovery of new species and strains of microorganisms. Therefore prospecting for gene

diversity is envisaged as a promising investigative area of the 21st century, giving Costa Rica a unique opportunity to lead the process in the region. The genetic resources obtained from microorganisms play an important role in the production of new enzymes, antibiotics and bioinsecticides by the biotechnological industry (Bull *et al.* 1992, Samsonov *et al.* 1997).

It has been estimated that 67 000 species of plagues affect agriculture in the world, and approximately 9 000 species are insect pests (Ross and Lembi 1985). As a result, sustainable control of insects in agriculture is crucial since, for 2001, it was estimated that chemical worldwide control of insects cost 7 500 million dollars (James 2002). In addition, the use of synthetic insecticides is not recommended because of the undesirable effects on human health, ecological problems caused by their slow degradation and the lack of specificity in their insecticidal action. This situation has stimulated the search of new alternatives of insect control based on the entomopathogenic bacteria *Bacillus thuringiensis* (*Bt*).

Bt is a Gram positive bacteria of the Bacillaceae family that has been used as a bioinsecticide for the biological control of plagues of economic importance in agriculture over the last decades (Aronson *et al.* 1986). This bacteria synthesizes crystalline insecticidal proteins or δ -endotoxins. These δ -endotoxins form parasporal inclusion bodies, which are very toxic and highly specific to the target insect but are innocuous to animals and humans (Betz *et al.* 2000). *Bt*-based bioinsecticides have been formulated against lepidopteran and coleopteran larva of plant pests as well as against mosquitoes and black flies, which are vectors for a variety of human diseases (Bravo *et al.* 1998).

Most *Bt* strains produce δ -endotoxins encoded in the *cry* and *cyt* genes contained in megaplasmids of more than 30 Mda. These toxins were originally classified by Hofte and Whiteley (1989) into four classes (Cry1, Cry2, Cry3, and Cry4), according to their amino acid sequence similarity and their insecticidal toxicity. More recently, Crickmore *et al.* (1998) established a new classification system based

solely on the amino acid homology, resulting in 34 classes of Cry proteins. In addition, Estruch *et al.* (1996) discovered that some *Bt* strains also synthesize Vip insecticidal proteins during the vegetative growth phase. Both Vip and Cry proteins act in the midgut epithelium causing intestinal paralysis, cellular lysis and finally insect death.

The identification of *Bt cry* and *cyt* genes by PCR has proven to be a very useful method for strain characterization, offering several advantages in terms of rapidity and reproducibility (Ben-Dov *et al.* 1997, Porcar and Juárez-Pérez 2003). A single *Bt* strain can harbor up to eight different *cry* genes (Martínez 2002). In general, the type of *cry* and *cyt* genes present in a strain correlates to some extent with its insecticidal activity (Porcar and Juárez-Pérez 2003). Thus, the identification of the gene profile in a *Bt* collection can be a useful tool to predict its potential insecticidal activity.

Since Costa Rican insects diversity is estimated to be 360 000 species (Obando 2002), and because a co-evolution of *Bt* strains and their susceptible insect hosts has been proposed (Apoyolo *et al.* 1995), prospecting for *Bt* strains in diverse natural ecosystems could result in the identification of Cry proteins with new specificities. The objective of this research was to isolate and characterize a collection of *Bt* strains from natural ecosystems, representing the diverse life zones of Costa Rica. The results obtained offered information about the prevalence of *Bt* in the country and the distribution, abundance and diversity of *Bt* strains. Those strains could be used in the near future in the formulation of insecticides for the biological control of insects of economic importance for the country. In addition, their *cry* genes could be used for the genetic transformation of plants.

MATERIALS AND METHODS

Sample collection: Two hundred and sixty five environmental samples (soil, leaf litter, fresh leaves and other substrates) were

collected from protected areas that include the diverse ecosystems and life zones of Costa Rica (Fig. 1). One hundred samples were collected from the humid tropical forest (Braulio Carrillo, Gandoca Manzanillo, Sierpe, Hitoy Cerere, and Cahuita), 80 from the dry tropical forest (National Park Marino las Baulas, Palo Verde, Santa Rosa), 24 of the very humid tropical forest (Tortuguero) and, finally, 61 samples from the humid tropical forest transition perhumid (Carara). The samples were dried at 50°C for 24 hours and stored at room temperature.

Isolation of *Bt*: Bacteria were isolated using the protocol described by *Travers et al.* (1987) using T3 as selective enrichment medium. Sporulated cultures showing the typical *Bt* morphology were preserved both on filter paper at room temperature and in 50% glycerol at -70°C.

Light microscopy: Cultures of approximately five days were analyzed by light microscopy (Nikon E-200 Eclipse) by staining with

Coomassie blue (0,25% (w/v) in 60% ethanol (v/v) and 7% of acetic acid (v/v), with the purpose of determining crystal morphology.

Polyacrylamide gel electrophoresis:

Crystal-spore preparations were analyzed for the presence of the δ -endotoxins in 10% SDS-PAGE gels. The molecular weight of the Cry proteins was determined by comparing with protein markers of known molecular weight (BIO-RAD 161-0304).

DNA extractions: The protocol for DNA extraction described by Chen and Kuo (1993) was used and DNA concentrations were estimated by fluorometry at 280 nm (Quantech fluorometer, model FM1 109535).

PCR analyses: The general primer for the *cry1*, *cry3-7*, *cry5*, *cry8*, *cry11*, *cry12*, *cry14*, and *cyt* genes and specific primer for *cry1Aa*, *cry1Ab*, *cry1Ac*, *cry1Ad*, *cry1B*, *cry1C*, *cry1D*, *cry1E*, *cry1F* (Bravo *et al.*, 1998), also *vip1*, *vip2*, *vip3* (A. Bravo, personal communication) and the general primer for *cry2* gene (Ben-Dov



Fig. 1. Costa Rica: *Bacillus thuringiensis* collection localities.

et al. 1997) were used. For the PCR the following conditions were used: 25 mM MgCl₂, 10x buffer, 10mM dNTPs, 20 µM each primer, 2.5 U *Taq* polymerase and 5 to 20 ng of DNA. The PCR program was: one denaturing cycle of two min. at 95°C, 30 cycles of one min. at 95°C, one min. at 48-54°C, one min. at 72°C, and a final extension cycle of 5 min. at 72°C. PCR products were analyzed by gel electrophoresis in 1% (w/v) agarose gels. The reference strains HD-137, HD-1, Btt, HD-916, were obtained from the Bacillus Genetic Center Stock; Department of Biochemistry (Ohio State University).

RESULTS

A total of 146 *Bt* strains were isolated from environmental samples from diverse natural ecosystems collected from 9 of the 12 life zones of Costa Rica. *Bt* strains were obtained from 60% of the samples. It was possible to recover *Bt* strains with an efficiency of 71%, 63%, 61% and 54% from soil samples, fresh leaves, other

substrates and leaf litter respectively. *Bt* was isolated in 65% of the samples collected in the humid tropical forest of the national parks (Braulio Carrillo, Gandoca Manzanillo, Sierpe, Hitoy Cerere, and Cahuita), in 59% of the samples collected in the dry tropical forest (Parque Nacional Marino las Baulas, Palo Verde and Santa Rosa). In the very humid tropical forest (Tortuguero) *Bt* was isolated in 75% of the samples and in the very humid tropical forest transition perhumid (Carara) was found in 69% of the samples (Fig. 1). Statistical analysis (χ^2) indicated no significant differences in relation to the efficiency of *Bt* strain recovery from different life zones.

The microscopic observation of the crystals showed high diversity in the morphology, number as well as in the size of the parasporal inclusions. Small, medium and large oval crystals were frequently found. Also, some strains presented small and large bipyramidal as well as irregular crystals (Fig. 2). This diversity was reflected also in the presence of different crystal inclusions, since some strains produce crystals

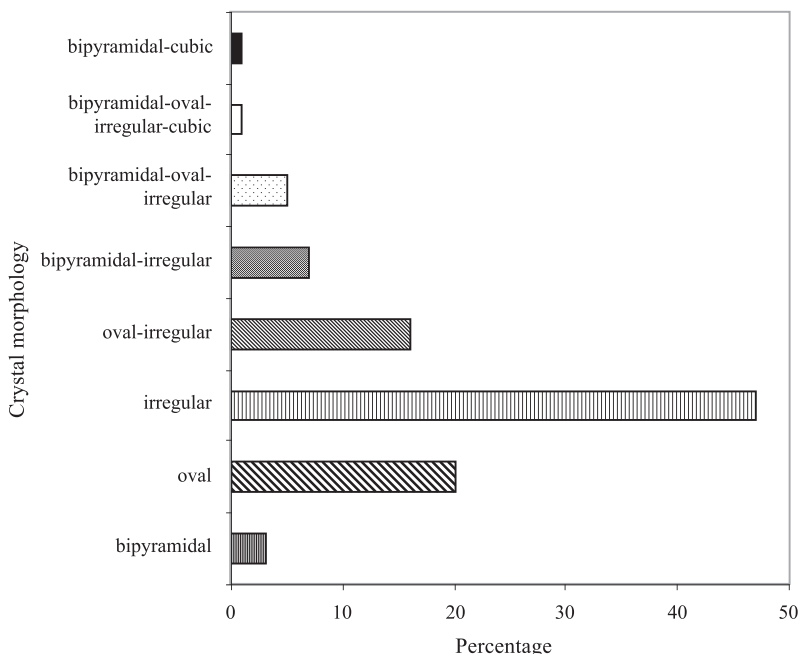


Fig. 2. Percentage distribution of crystal morphologies of the Costa Rican *Bacillus thuringiensis* strain collection, isolated from natural ecosystems.

with several morphologies, whereas others show only one type of crystal. Most strains presented more than one crystal morphology, for example up to six different crystals in the same strain. Figure 2 shows the different crystal morphologies, the irregular being the most common (47%), followed by the oval (20%). Twenty-four strains (16%) showed a combination of both. The combination of bipyramidal and cubical crystals was rare.

No correlation was found when crystal morphology and isolation sites were compared. However, strains isolated from Braulio Carrillo, Tortuguero and Cahuita, presented abundant irregular crystals. One of the few strains that presented large cubic, oval and bipyramidal crystals was isolated from Tortuguero. On the other hand, more than 60% of the isolated strains of Térraba-Sierpe, and Hitoy-Cerere presented medium oval crystals. Several strains isolated from Gandoca-Manzanillo, Palo Verde and Carara presented mainly combinations of oval and irregular crystals. The greatest diversity in morphologies was observed in those from Santa Rosa, Llanos del Río Medio Queso and Parque Marino las Baulas.

The collection was also characterized by SDS-PAGE to determine the number and molecular weight (MW) of the Cry proteins. The analyses showed diverse electrophoretic patterns, with MWs in the range of 20 to 160 kDa (Appendix 1). Strains isolated from Parque Marino las Baulas presented very different δ -endotoxin profiles. Strain CIBCM-134 expressed proteins of 130 and 60 kDa, while the CIBCM-142, produced proteins of 20 or 40 kDa. These strains presented small and large bipyramidal crystals respectively. There was no correlation between the morphology of the inclusion body and the MW of the δ -endotoxins (Appendix 1). The irregular and the oval crystals presented the greatest diversity in the MW of their Cry proteins, while the bipyramidal crystals often showed high MW δ -endotoxins (140 or 120 kDa). For example, the CIBCM-251 strain with oval crystals presented Cry proteins of 90 or 100 kDa. In addition, there

was no correlation between crystal morphology and the number of Cry proteins detected. For example, one strain from Braulio Carrillo with pleomorphic crystals showed proteins of 60, 70 and 100 kDa, while another with the same crystal morphology isolated from Santa Rosa contained a single polypeptide of 70 kDa. It should be pointed out that strains that showed four δ -endotoxins in the gel also presented several crystals, while strains with single crystal morphology presented three Cry proteins (for example CIBCM-251).

The characterization of the collection by PCR showed that *cry2* and *vip3* genes were found in 87 and 85 strains respectively. Thirty-one strains amplified with the general primer *cry1*, 14 with *cry3-7*, 12 with *cry8*, 11 harbored the *cry11* gene and finally 13 amplified with the *cyt* general primer. Amplification with the general primers for the *cry5*, *cry12*, *cry14* and *vip1* genes was not detected. Also it is important to mention that 24 strains did not amplify with any of the primers used. Figure 3 shows the distribution of the genes of the collection according to forest type. All types of forest contained *Bt* strains with same type of genes, but their frequencies differed. Nevertheless, the dry tropical forest showed the greatest number of strains with *cry2*, *cry3-7*, *cry8* and *cyt* genes. Several strains presented great diversity of *cry* genes, for example CIBCM-165 amplified with *cry1*, *cry2*, *cry11*, *cyt* and *vip3* genes. Also CIBCM-154 amplify with the primers for *cry1*, *cry2*, *cry3-7* and *cry8* genes (Appendix 1).

Strains that amplified with the *cry1* general primer were further analyzed with specific primers for different subfamilies. Twenty-two strains presented at least one gene of the *cryIA* subfamily, 11 strains showed the *cryID* and four the *cryIB* gene (Appendix 1). The *cryIE* gene was scarce (only two strains). None of the *Bt* strains of this collection contained the *cryIC* or *cryIF* genes. On the other hand, eight strains did not amplify with any of specific primers tested for *cry1* subfamilies. A total of 13 different genetic profiles were found for the *cryI* family (Appendix 1).

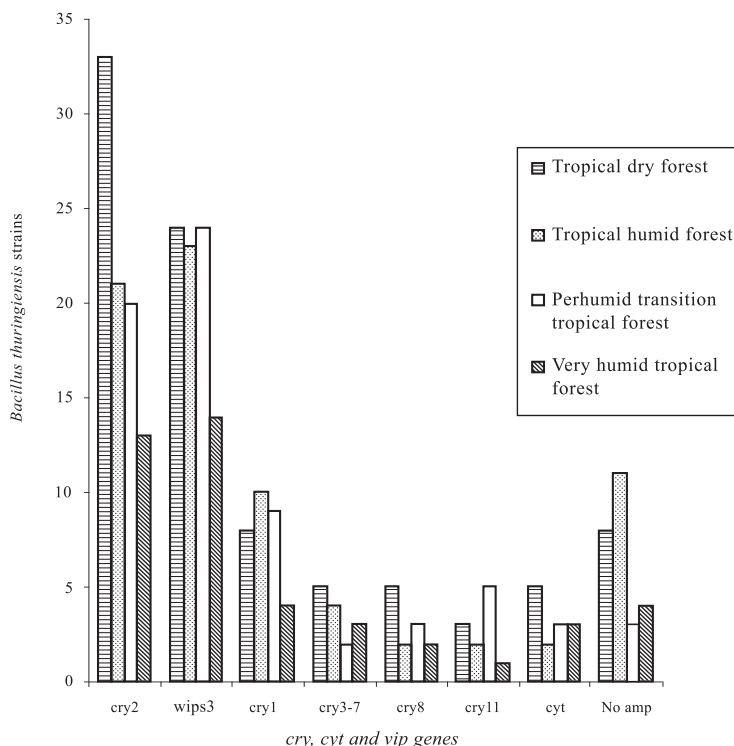


Fig. 3. Distribution of cry, cyt and vip genes of the Costa Rican *Bacillus thuringiensis* strain collection, according to forest type.

DISCUSSION

A collection of 146 strains of *Bacillus thuringiensis* was obtained from environmental samples derived from diverse natural ecosystems of Costa Rica. This research corroborated the broad distribution of *Bt* in different microhabitats. Isolation efficiency was of 60%, a very similar figure to that reported by other authors (Martin and Travers 1989). It is important to indicate that the dry tropical forest seems to present the greatest diversity, not only in crystal morphologies but also in the insecticidal protein genes.

This collection showed great diversity in morphology, size and number of parasporal inclusions. It was not possible to establish a correlation between crystal morphology and the origin of the strain. Although it was possible to isolate strains from different areas with

similar crystal morphologies, it is interesting to emphasize that some strains from specific sites presented rare morphologies, for example the CIBCM-142 from the Parque Marino las Baulas showed long bipyramidal crystals and CIBCM-296 from Tortuguero had large cubic crystals.

Diversity in crystal protein patterns was also revealed by electrophoresis. Our results differed from those obtained in strains isolated from agricultural ecosystems (Viquez 2000). It was reported previously that the characterization by SDS-PAGE of these strains showed homogeneous δ -endotoxin profiles that presented mainly two bands, one of 130 and another of 60 kDa. The strains isolated from natural ecosystems showed, in addition, other proteins of 20, 40, 70, 80 and 90 kDa. Similar results were reported by Bravo *et al.* (1998) who found that the strains with novel Cry proteins were isolated from Mexican humid tropical forests.

Our results and those obtained by Bravo *et al.* (1998) showed that natural ecosystems with more insect species present a greater diversity of *Bt* strains. Since a co-evolution of *Bt* strains and their susceptible insect hosts has been proposed (Apyolo *et al.* 1995), prospection of *Bt* strains in diverse natural ecosystems should be considered as an efficient strategy for the identification of new Cry proteins with novel specificities.

The *Bt* crystals of some strains are made up of a single protein, for example *Bt var kurstaki* HD-73 that contained only the Cry1Ac protein. Some of the characterized strains of this collection, showed a single protein in the SDS-PAGE, suggesting that their crystals are comprised of a single protein, or by two or more proteins with the same molecular weight. Nevertheless, one type of crystal could be constituted by different proteins as in the strain *Bt var morrisoni*, where proteins Cry4, Cry1A and Cyt form several inclusions covered by a common membrane (Ibarra *et al.* 1986). In other strains, like the HD-1, several proteins (Cry1Aa, Cry1Ab, and Cry1Ac) form a single crystal (Hofte *et al.* 1998). Abundant oval crystals were observed in the strain CIBCM-251, but when analyzed by electrophoresis, three δ -endotoxin of 60, 90 and 100 kDa were detected. This complexity was observed in some of the strains that synthesize single bipyramidal crystals, for example CIBCM-281 that presented three δ -endotoxins with different MWs.

It was also noticed that strains with bipyramidal crystals frequently presented two proteins, one of high MW of 120 or 140 kDa and another of 65 or 60 kDa. These proteins could be the protoxin and the active toxin respectively. As *Bt* regulates the synthesis of proteases that can process protoxin to toxins (Rukmini *et al.* 2000), the proteolysis could imply reduction of activity, since the δ -endotoxins are less stable than protoxin or, on the other hand, would permit the bacteria to respond faster because the insect is ingesting the activated toxin.

Some of the δ -endotoxins of low MW (30-40 kDa) detected in the gels could be either Cyt proteins or binary toxins. In addition to the Cry proteins, it is probable that some of strains

produce Vip proteins, since a high percentage of the strains amplified with Vip3 primers. Unfortunately it was not possible to determine if those proteins were expressed during the vegetative phase and secreted to the medium.

The most common *cry* genes found in nature are those within the *cryI* family (Porcar and Juárez-Pérez 2003). In our collection strains containing *cryI* genes were not so abundant (20%). The *cry2* gene was the most frequently found, like in other collections (Ibarra, personal communication). It has been reported that these proteins have MWs of 70 to 75 kDa. Table 1 shows that several strains of the collection expressed a protein of 70 kDa that could be the Cry2 protein, a protein toxic for insects of the Lepidoptera and Diptera orders. The high frequency of these proteins in several *Bt* strains could have permitted them to extend the host range.

The second most frequent gene of this collection was *vip3*, (58 %). Estruch *et al.* (1996) found this gene in only 15% of the strains of their collection. Arrieta *et al.* (2004) also reported a widespread distribution of this gene in strains isolated from Costa Rican agricultural ecosystems. In contrast, the *vip1* gene was not detected in this collection and only one strain presented the *vip2* gene. Different results were obtained in a Costa Rican collection isolated from coffee plantations where the frequency of the *vip2* gene was higher (Arrieta *et al.* 2004).

The occurrence of *cryI* gene varies greatly among different *Bt* collections. For example *cryIA* genes were frequently present in more than 50% of the strains, whereas other genes of the subfamily, such as *cryIE* and *cryIF*, were less frequent. However there are some exceptions, such as the high frequency of *cryIE* in a Chinese collection (Porcar and Juárez-Pérez 2003).

Several reports showed a high frequency of certain combinations of *cryI* genes, for example the linkage of the *cryIC* and *cryID* genes (Bravo *et al.* 1998, Ferrandis *et al.* 1999, Hongyu *et al.* 2000). This *cryIC* and *cryID* linkage may be explained by their location on the same replicon (Sanchis *et al.* 1988).

However, eleven *Bt* strains of the Costa Rican collection contained only the *cryID*. Ferrandis *et al.* (1999) suggested that the absence of the *cryIC* gene might be explained by a deletion or negative selection of the *cryIC* gene from an ancestral *cryIC-cryID* linkage.

In summary, the diversity of *Bt* toxins indicate that the collection analyzed has great potential for the control of different species of insect pests of economic importance. Strains with a diversity of genes for the control of lepidopterans, dipterans and coleopterans were found. It is interesting to mention that this *Bt* collection obtained from natural ecosystems presented greater diversity of *cry* genes in comparison to other collections isolated from Costa Rican agricultural ecosystems (Arrieta *et al.* 2004, Mora and Espinoza 2005). The characterization of this collection will offer very useful information for the selection of *Bt* strains with particular Cry protein profiles to be evaluated for their toxicity against specific insect pests of important food crops.

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RESUMEN

Como los ecosistemas naturales de Costa Rica figuran entre los más diversos del mundo, se propuso aislar la

bacteria entomopatógena *Bacillus thuringiensis* (*Bt*) con el fin de conformar una colección de cepas y caracterizarlas molecularmente. Se obtuvieron 146 cepas a partir de muestras ambientales de diversas áreas protegidas, que incluían 9 de las 12 zonas de vida de Costa Rica. Se recuperaron cepas del 71%, 63%, 61% y 54% de las muestras de suelo, hojas frescas, otros sustratos y hojarasca respectivamente. Se aisló *Bt* del 65% de las muestras del bosque tropical húmedo, un 59% de las muestras del bosque tropical seco. Del bosque tropical muy húmedo se aisló *Bt* del 75% de las muestras y finalmente del bosque tropical muy húmedo transición perhúmedo se encontró en el 69% de las muestras. Las cepas se caracterizaron según la morfología de los cuerpos paraesporales de inclusión, el peso molecular de las δ -endotoxinas y de genes *cry*, *cyt* y *vip* que contenían. Las cepas exhibieron cristales de diferente morfología, tamaño y número: irregulares, ovales, bipiramidales, cúbicicos o mezclas de uno u otro. No se encontró correlación al comparar la forma de los cristales con el sitio de origen de la cepa. El análisis proteico de las mezclas de esporas y cristales mostró que las cepas contenían δ -endotoxinas de 20 a 160 kDa. El 66 por ciento de las cepas amplificaron con los imprimadores específicos para el gen *cry2*, 54% con *vip3*, 20% con el *cry1*, 9% con el *cry3-cry7* y 8% con el gen *cry8*. Los genes *cry11* y *cyt* se encontraron en el 8% y 7% de las cepas respectivamente. Veinticuatro cepas no amplificaron con los imprimadores utilizados por lo que podrían contener genes novedosos. Las cepas que contenían el gen *cry1* se amplificaron posteriormente con imprimadores específicos para la subfamilia de dicho gen, obteniéndose 13 perfiles diferentes. En síntesis, la diversidad genética de las cepas sugiere que la colección tiene un gran potencial para el control de diferentes especies de insectos de importancia económica en la agricultura y en salud pública. El análisis toxicológico de las cepas mediante bioensayos con insectos plaga proveerá información muy útil acerca del uso potencial de estas cepas para la formulación de bioinsecticida. Asimismo, los genes *cry* y *vip* podrían utilizarse para, mediante ingeniería genética, conferir resistencia a los cultivos.

Palabras clave: *Bacillus thuringiensis*, cristales, genes *cry*, *vip*, δ -endotoxinas, parques nacionales, Costa Rica.

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APPENDIX 1

Summary of the characterization of the Costa Rican Bacillus thuringiensis strain collection isolated from natural ecosystems, according to crystal morphology, molecular weight of the δ -endotoxin and insecticidal genes

<i>Bacillus thuringiensis</i> strains	Protected area	Enviromental sample	Crystal morphology	Molecular weight of Cry & Cyt proteins (kDa)	<i>Cry, cyt and vip</i> genes
CIBCM-1	Braulio Carrillo	Soil	Ir	65, 50, 45	<i>cry2, vip3</i>
CIBCM-4	Braulio Carrillo	Soil	Ir	110, 100 60	<i>N amp</i>
CIBCM-5	Braulio Carrillo	Soil	Ir	125, 70, 50	<i>cry1, cry2, vip3</i>
CIBCM-7	Braulio Carrillo	Soil	Ir	70, 30	<i>cry2, vip3</i>
CIBCM-8	Braulio Carrillo	Soil	Ov	60, 50	<i>N amp</i>
CIBCM-9	Braulio Carrillo	Soil	Ov	60, 45, 30	<i>vip3</i>
CIBCM-11	Braulio Carrillo	Leaf litter	Ir	65, 50, 40	<i>cry2, vip3</i>
CIBCM-12	Braulio Carrillo	Soil	Ov	65	<i>vip3</i>
CIBCM-13	Braulio Carrillo	Other substrate	Ov	65, 44	<i>N amp</i>
CIBCM-15	Braulio Carrillo	Fresh leaves	Ir	110, 90, 70	<i>N amp</i>
CIBCM-16	Braulio Carrillo	Leaf litter	Ov, Ir	130, 75, 60	<i>cry1, cry2, vip3</i>
CIBCM-17	Braulio Carrillo	Soil	Ir	75, 60	<i>cry2, vip3</i>
CIBCM-18	Braulio Carrillo	Other substrate	Ir	100, 70, 60	<i>cry1, cry2, vip3</i>
CIBCM-19	Braulio Carrillo	Other substrate	Bi, Ir	85, 65	<i>cry2, vip3, cry11</i>
CIBCM-63	Carara	Fresh leaves	Bi, Ir	140, 110, 70	<i>cry1, cry2, vip3, cry8</i>
CIBCM-64	Carara	Other substrate	Bi, Ir	90, 72, 45	<i>cry2, cry8, vip3</i>
CIBCM-65	Carara	Other substrate	Bi	140, 85, 65	<i>cry1, vip3</i>
CIBCM-66	Carara	Other substrate	Ir	85, 65, 56	<i>cry2, vip3</i>
CIBCM-67	Carara	Other substrate	Ir	70, 60, 52	<i>cry2, cry11, vip3</i>
CIBCM-68	Carara	Other substrate	Bi, Ir	140, 75, 68, 45	<i>cry1, cry2, vip3</i>
CIBCM-69	Carara	Soil	Bi, Ir	81, 60	<i>cry1, cry2</i>
CIBCM-70	Carara	Other substrate	Bi, Ov, Ir	110, 75, 45	<i>cry1, cry2, vip3</i>
CIBCM-74	Carara	Soil	Ov	45, 20	<i>cry2</i>
CIBCM-75	Carara	Soil	Bi	85, 66	<i>cry2, vip3</i>
CIBCM-76	Carara	Leaf litter	Ov, Ir	85, 66, 56	<i>cry2, cry11, vip3</i>
CIBCM-80	Carara	Soil	Ov	85, 65	<i>cry2,vip3</i>
CIBCM-81	Carara	Soil	Ov	82, 55, 40	<i>cyt, vip3</i>

APPENDIX 1 (continued...)
Summary of the characterization of the Costa Rican Bacillus thuringiensis strain collection isolated from natural ecosystems, according to crystal morphology, molecular weight of the δ -endotoxin and insecticidal genes

<i>Bacillus thuringiensis</i> strains	Protected area	Enviromental sample	Crystal morphology	Molecular weight of Cry & Cyt proteins (kDa)	Cry, cyt and vip genes
CIBCM-85	Carara	Leaf litter	Ir	55, 40	<i>N amp</i>
CIBCM-90	Carara	Other substrate	Bi, Ir	80, 72, 50, 35	<i>cry1, cry2, vip3</i>
CIBCM-92	Carara	Leaf litter	Ir	80, 67, 55, 40	<i>cry11, cyt</i>
CIBCM-94	Carara	Leaf litter	Ov, Ir	95, 65, 40	<i>cry2, cry11, vip3</i>
CIBCM-97	Carara	Soil	Ov, Ir	70, 60, 45	<i>cry2, cry11, vip2, vip3</i>
CIBCM-101	Carara	Soil	Ov	80, 50	<i>vip3</i>
CIBCM-103	Carara	Soil	Ov, Ir	80, 55, 45	<i>cry2, vip3</i>
CIBCM-105	Carara	Soil	Ir	90, 55, 50, 40	<i>vip3</i>
CIBCM-107(A)	Carara	Soil	Ov, Ir	100, 70, 60	<i>cry2, vip3</i>
CIBCM-112	Carara	Leaf litter	Ov, Ir	120, 80, 60	<i>cry2, vip3</i>
CIBCM-122	Carara	Leaf litter	Ir	65, 55, 40	<i>N amp</i>
CIBCM-127	Marino las Baulas	Soil	Ir	85, 70, 60	<i>cry8</i>
CIBCM-128	Marino las Baulas	Leaf litter	Ir	120, 80, 40	<i>cry2</i>
CIBCM-129	Marino las Baulas	Soil	Bi, Ir	100, 90, 70, 45	<i>cry2, cry8</i>
CIBCM-131	Marino las Baulas	Soil	Ir	120, 80, 40	<i>cry2</i>
CIBCM-134	Marino las Baulas	Leaf litter	Bi, Ov, Ir, Cu	130, 60	<i>cry1, cry2, vip3</i>
CIBCM-135	Marino las Baulas	Soil	Ov	120, 70	<i>vip3</i>
CIBCM-137	Marino las Baulas	Leaf litter	Ov	80, 70, 45	<i>cry2</i>
CIBCM-142	Marino las Baulas	Soil	Bi, Ov	140, 40, 20	<i>cry1</i>
CIBCM-147	Gandoca-Manzanillo	Soil	Ov, Ir	70, 62, 52	<i>vip3</i>
CIBCM-150	Gandoca-Manzanillo	Fresh leaves	Ov	70, 60, 40	<i>N amp</i>
CIBCM-151	Gandoca-Manzanillo	Soil	Ov, Ir	60, 50, 35	<i>vip3</i>
CIBCM-153	Gandoca-Manzanillo	Fresh leaves	Ov, Ir	65, 55, 45	<i>N amp</i>
CIBCM-154	Gandoca-Manzanillo	Soil	Ov, Ir	80, 75, 65, 50	<i>cry1, cry2, cry3-7, cry8</i>
CIBCM-155	Gandoca-Manzanillo	Soil	Ir	80, 64	<i>cry1, cry2, cry8</i>
CIBCM-156	Gandoca-Manzanillo	Leaf litter	Ir	130, 90, 70, 60	<i>cry1, cry2, vip3</i>
CIBCM-161	Medio-Queso	Soil	Ov	80, 65	<i>cry2</i>

APPENDIX 1 (continued...)
Summary of the characterization of the Costa Rican Bacillus thuringiensis strain collection isolated from natural ecosystems, according to crystal morphology, molecular weight of the δ -endotoxin and insecticidal genes

<i>Bacillus thuringiensis</i> strains	Protected area	Enviromental sample	Crystal morphology	Molecular weight of Cry & Cyt proteins (kDa)	Cry, cyt and vip genes
CIBCM-163	Medio-Queso	Soil	Ir	67, 55, 40	<i>N amp</i>
CIBCM-165	Medio-Queso	Other substrate	Bi, Ir	110, 85, 70, 40	<i>cry1, cry2, cry11, cyt, vip3</i>
CIBCM-166	Medio-Queso	Other substrate	Bi, Ir, Ov, Cu	140, 65	<i>cry1, cry2, vip3</i>
CIBCM-167	Medio-Queso	Soil	Bi, Ov, Ir	140, 110, 90, 73	<i>cry1, vip3</i>
CIBCM-168	Medio-Queso	Leaf litter	Ov, Ir	130, 95, 65	<i>N amp</i>
CIBCM-180	Medio-Queso	Soil	Ov, Ir	125, 100, 30	<i>cry3-7, cyt</i>
CIBCM-187	Palo Verde	Fresh leaves	Ov	140, 70, 50	<i>cry2, vip3</i>
CIBCM-188	Palo Verde	Soil	Ov, Ir	100, 70, 40	<i>cry2, cry11, cyt, vip3</i>
CIBCM-189	Palo Verde	Soil	Ov, Ir	125, 105, 40	<i>cry11, cyt, vip3</i>
CIBCM-191	Palo Verde	Soil	Ov	100, 90, 50, 30	<i>cry11, vip3</i>
CIBCM-193	Palo Verde	Fresh leaves	Ov	85, 45, 30	<i>cry2, cry3-7</i>
CIBCM-234	Palo Verde	Soil	Ir	80, 50	<i>N amp</i>
CIBCM-246	Palo Verde	Soil	Ov, Ir	70, 50, 40	<i>cry2</i>
CIBCM-250	Palo Verde	Leaf litter	Ov	100, 70, 60, 45	<i>vip3</i>
CIBCM-251	Palo Verde	Fresh leaves	Ov	100, 90	<i>cry2</i>
CIBCM-252	Palo Verde	Soil	Ov, Ir	70, 60, 50	<i>cry2</i>
CIBCM-254	Palo Verde	Fresh leaves	Ov	120, 90, 50	<i>cry2, cry3-7</i>
CIBCM-259	Santa Rosa	Soil	Ir	100, 90, 50	<i>cry2, vip3</i>
CIBCM-260	Santa Rosa	Leaf litter	Ir	105, 90, 60, 40	<i>cyt, vip3</i>
CIBCM-261	Santa Rosa	Fresh leaves	Ov, Ir	110, 95, 70	<i>cry2, vip3</i>
CIBCM-263	Santa Rosa	Soil	Ir	95, 70, 40	<i>cry2</i>
CIBCM-264	Santa Rosa	Fresh leaves	Ov, Ir	80, 72 40, 30	<i>N amp</i>
CIBCM-265	Santa Rosa	Soil	Ov	90, 62, 55	<i>N amp</i>
CIBCM-267	Santa Rosa	Fresh leaves	Bi, Ov, Ir	125, 90, 55	<i>cry1, cry2, cry8, vip3</i>
CIBCM-269	Santa Rosa	Soil	Ov	65, 55, 30	<i>N amp</i>
CIBCM-270	Santa Rosa	Fresh leaves	Ir	100, 90, 45	<i>cry2, vip3</i>

APPENDIX 1 (continued...)
Summary of the characterization of the Costa Rican Bacillus thuringiensis strain collection isolated from natural ecosystems, according to crystal morphology, molecular weight of the δ -endotoxin and insecticidal genes

<i>Bacillus thuringiensis</i> strains	Protected area	Enviromental sample	Crystal morphology	Molecular weight of Cry & Cyt proteins (kDa)	<i>Cry, cyt and vip genes</i>
CIBCM-271	Santa Rosa	Leaf litter	Ov	100, 90, 40	<i>cry2, vip3</i>
CIBCM-273	Santa Rosa	Soil	Ov, Ir	120, 85, 60	<i>cry1, cry2, vip3</i>
CIBCM-274	Santa Rosa	Soil	Ir	70	<i>cry2, cry3-7</i>
CIBCM-275	Santa Rosa	Fresh leaves	Ir	90, 65	<i>cry2, cry3-7, cyt</i>
CIBCM-276	Santa Rosa	Soil	Ir	100, 90, 50	<i>N amp</i>
CIBCM-277	Santa Rosa	Leaf litter	Ir	100, 45	<i>cry2, vip3</i>
CIBCM-279	Santa Rosa	Soil	Ov	50, 40, 20	<i>cry2, vip3</i>
CIBCM-281	Santa Rosa	Fresh leaves	Bi	130, 65, 40	<i>cry1</i>
CIBCM-285	Hitoy-Cerere	Soil	Ir	90, 80, 75, 60	<i>N amp</i>
CIBCM-290	Hitoy-Cerere	Leaf litter	Ir	90, 40	<i>cry3-7, vip3</i>
CIBCM-292	Hitoy-Cerere	Leaf litter	Ov	74, 60	<i>cry2, cry3-7</i>
CIBCM-293	Tortuguero	Soil	Ir	90, 65, 60, 45	<i>cry2, cyt, vip3</i>
CIBCM-294	Tortuguero	Leaf litter	Ir	100, 90, 52	<i>vips3</i>
CIBCM-296	Tortuguero	Soil	Bi, Cu	130, 65	<i>cry1, vips3</i>
CIBCM-300	Tortuguero	Leaf litter	Ir	100, 90, 50	<i>cry2</i>
CIBCM-302	Tortuguero	Soil	Ir	110, 85, 65	<i>N amp</i>
CIBCM-303	Tortuguero	Leaf litter	Ir	65, 60, 40, 35	<i>cry2</i>
CIBCM-304	Tortuguero	Fresh leaves	Bi, Ir	105, 65	<i>cry1, cry2, cry3-7, vip3</i>
CIBCM-305	Tortuguero	Soil	Ir	60, 40	<i>cry2</i>
CIBCM-306	Tortuguero	Leaf litter	Ir	40	<i>vip3</i>
CIBCM-307	Tortuguero	Fresh leaves	Ir	60, 40	<i>vip3</i>
CIBCM-308	Tortuguero	Soil	Ov	100, 90, 52, 40	<i>vip3</i>
CIBCM-310	Tortuguero	Fresh leaves	Ir	65, 50	<i>cry1, cry8, vip3</i>
CIBCM-311	Tortuguero	Soil	Ov	70, 65, 40	<i>N amp</i>
CIBCM-314	Tortuguero	Soil	Ir	95, 60	<i>cry2, cry3-7</i>
CIBCM-317	Santa Rosa	Soil	Ir	96, 76, 40	<i>N amp</i>
CIBCM-320	Santa Rosa	Soil	Ov-Ir	130, 105, 80, 60	<i>cry2, vip3</i>

APPENDIX 1 (continued...)
Summary of the characterization of the Costa Rican Bacillus thuringiensis strain collection isolated from natural ecosystems, according to crystal morphology, molecular weight of the δ -endotoxin and insecticidal genes

<i>Bacillus thuringiensis</i> strains	Protected area	Enviromental sample	Crystal morphology	Molecular weight of Cry & Cyt proteins (kDa)	<i>Cry, cyt and vip genes</i>
CIBCM-321	Santa Rosa	Leaf litter	Ov	60, 44, 30	<i>cry2, vip3</i>
CIBCM-322	Santa Rosa	Fresh leaves	Bi, Ir	140, 80, 65	<i>cry1, cry2, cry8, vip3</i>
CIBCM-324	Santa Rosa	Leaf litter	Ov, Ir	90, 65, 40	<i>vip3</i>
CIBCM-325	Santa Rosa	Fresh leaves	Bi, Ov, Ir, Cu	150, 110, 80, 60	<i>cry1, cry2, vip3</i>
CIBCM-327	Cahuita	Soil	Ov, Ir	50, 30	<i>N amp</i>
CIBCM-328	Cahuita	Leaf litter	Ov	100, 80, 65	<i>cry3-7</i>
CIBCM-329	Cahuita	Fresh leaves	Ir	160, 90	<i>cry2, vip3</i>
CIBCM-330	Cahuita	Soil	Ir	86, 63, 44	<i>cry2, vip3</i>
CIBCM-331	Cahuita	Leaf litter	Ir	130, 110, 75	<i>cry2, vip3</i>
CIBCM-332	Cahuita	Fresh leaves	Ir	40, 35	<i>cry2, vip3</i>
CIBCM-334	Cahuita	Leaf litter	Bi	150, 130, 90, 80	<i>cry2</i>
CIBCM-335	Cahuita	Fresh leaves	Bi	160, 130, 80	<i>N amp</i>
CIBCM-338	Cahuita	Fresh leaves	Bi, Ov, Ir, Cu	135, 65	<i>cry1, cry2, vip3</i>
CIBCM-355	Tortuguero	Soil	Ir	160, 95, 65, 33	<i>cry11, cyt, vip3</i>
CIBCM-356	Tortuguero	Soil	Ir	100, 90, 40	<i>cry2, vip3</i>
CIBCM-357	Tortuguero	Soil	Ir	100, 90, 50	<i>cry2, vip3</i>
CIBCM-360	Tortuguero	Soil	Ir	65, 50	<i>cry8</i>
CIBCM-365	Tortuguero	Soil	Ir	65, 55	<i>cry2, cry3-7, vip3</i>
CIBCM-74(1)	Carara	Soil	Bi, Ov, Ir	65, 57, 50, 30	<i>cry1, cry2, cry3-7, cry8, vip3</i>
CIBCM-74(2)	Carara	Soil	Bi , Ov, Ir	65, 55	<i>cry1</i>
CIBCM-263(1)	Santa Rosa	Soil	Ir	100, 90, 40	<i>cry2</i>
CIBCM-294(1)	Tortuguero	Leaf litter	Bi -Ov-Ir	130, 100, 80, 60	<i>cry1, cry2, vips3</i>
CIBCM-88(98)	Carara	Soil	Ir	90, 65, 43	<i>cry2, vips3</i>
CIBCM-89(98)	Carara	Leaf litter	Ir	53, 40	<i>N amp</i>
CIBCM-90(97)	Carara	Other substrate	Ir	80, 65, 40	<i>cry3-7</i>
CIBCM-96(57)	Santa Rosa	Other substrate	Ir	140, 95, 70	<i>cry2</i>
CIBCM-96(63)	Santa Rosa	Other substrate	Ov, Ir	130, 70, 60	<i>cry1, cry2, vips3</i>

APPENDIX 1 (continued...)

Summary of the characterization of the Costa Rican Bacillus thuringiensis strain collection isolated from natural ecosystems, according to crystal morphology, molecular weight of the δ -endotoxin and insecticidal genes

<i>Bacillus thuringiensis</i> strains	Protected area	Enviromental sample	Crystal morphology	Molecular weight of Cry & Cyt proteins (kDa)	Cry, cyt and vip genes
CIBCM-96(72)	Santa Rosa	Other substrate	Ir	88, 77, 40	<i>N amp</i>
CIBCM-281(87)	Santa Rosa	Fresh leaves	Ir	60, 45	<i>cry2, cry8, vips3</i>
CIBCM-131(100)	Marino las Baulas	Soil	Ir	140, 80, 70	<i>cry2</i>
CIBCM-105(A)	Carara	Soil	Ir	90, 55, 50	<i>vips3</i>
CIBCM-107(A)	Carara	Soil	Ir	60, 53, 46	<i>cry1, cry2, vips3</i>
CIBCM-151(A)	Gandoca-Manzanillo	Soil	Ir	96, 80	<i>vip3</i>
CIBCM-1 (A)	Braulio Carrillo	Soil	Ir	70, 60, 45	<i>vip3</i>
CIBCM-251(A)	Palo Verde	Fresh leaves	Ov	110, 95, 85, 41	<i>cry2</i>
CIBCM-359(A)	Tortuguero	Soil	Ir	100, 63, 51	<i>N amp</i>
CIBCM-172(83)	Gandoca-Manzanillo	Soil	Ir, Ov	120, 110, 60	<i>cry2, vip3</i>
CIBCM-172(123)	Gandoca-Manzanillo	Soil	Ir, Ov	65, 53, 41	<i>cry2, vip3</i>
CIBCM-185(83)	Palo Verde	Soil	Ir, Ov	140, 100, 80	<i>N amp</i>
CIBCM-281(87)	Santa Rosa	Fresh leaves	Ir	70, 85	<i>cry2, cry8</i>

Bi= bipyramidal, Ir= irregular, Ov= oval, Cu= cubic, N amp = did not amplify with any of the primers used.