Identification of *Fusarium cf. verticillioides* as The Causal Agent of Pokka Boheng Disease in Sugarcane in the Department of Antioquia, Colombia

Juliana Giraldo-Arias¹, Santiago Celis-Zapata², Nicolás D. Franco-Sierra³, Juan J. Arroyave-Toro⁴, Claudia Jaramillo-Mazo ⁵ and Javier Correa Alvarez⁶

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Abstract

The cultivation of sugarcane represents an important part in the Colombian economy due to the diverse applications in industries like liquor, food, paper and biofuels. Sugarcane worldwide production is affected by the presence of phytopathogenic agents, mainly filamentous fungi such as *Physalospora tucumanensis* (red rot disease) and *Fusarium* spp. To date in Colombia, Pokka boheng disease whose causal agent is the fungus
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*Fusarium verticillioides*, has not been reported, which is why it is necessary to identify appropriately this microorganism, responsible for losses in productivity and food contamination. In order to isolate and identify the infectious agent from symptomatic tissues, disinfection and culture in liquid and solid culture mediums were performed in malt extract (2%) and yeast extract (0.2%) both liquid and solid, for 7 days. After several replicates in agar plate, a purification was made along with a morphological characterization based on the shape and color of the mycelium, as well as the type of spores generated. Additionally, the genetic material was extracted and gene markers (ITS, Elongation Factor 1-α (EF) and β-Tubulin (Btub)) were amplified by PCR. Then, DNA sequencing was used to obtain the data to make a phylogenetic reconstruction by probabilistic methods (Maximum Likelihood and Bayesian Inference). The isolated strain, named as EA-FP0013 was located in the Fujikuroi complex group, with high probable identity to *Fusarium verticillioides*. Thus, early and species-specific identification of these fungal isolates by molecular methods may allow the timely diagnosis of emerging pathophysiological diseases of interest in the region, and thus propose the respective control strategies.

**Keywords:** Sugarcane pathogens; *Fusarium* sp.; Molecular phytopathology; phylogenetics.

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**Resumen**

El cultivo de la caña de azúcar representa una parte importante en la economía de Colombia debido a las diversas aplicaciones en industrias como licor, alimentos, papel y biocombustibles. Su producción en todo el mundo se ve afectada por la presencia de agentes fitopatógenos, principalmente hongos filamentosos como *Physalospora tucumanensis* (agente causal de la enfermedad “Pudrición roja”) y *Fusarium* spp. Hasta la fecha, en Colombia, no se ha reportado la enfermedad de Pokka boheng en este cultivo cuyo agente causal es el hongo *Fusarium verticillioides*, por lo que es necesario identificar adecuadamente este microorganismo, responsable de las pérdidas en productividad y contaminación de los alimentos. Con el fin de aislar e identificar el agente infeccioso de los tejidos sintomáticos, se realizó la desinfección y posterior siembra de éstos en medios de cultivo extracto de malta (2%) y extracto de levadura (0.2%) tanto líquido como sólido durante 7 días. A partir de varias placas petri, se realizó la respectiva purificación y una caracterización morfológica basada en la forma y el color del micelio, así como el tipo de esporas generadas. Adicionalmente,
el material genético se extrajo y se amplificaron los marcadores genéticos (ITS, factor de elongación 1-α (EF) y β-tubulina (Btub)) por PCR. Luego, se utilizó la secuenciación del ADN para obtener los datos para realizar una reconstrucción filogenética mediante métodos probabilísticos (máxima verosimilitud e inferencia bayesiana). La cepa aislada, nombrada como EA-FP0013 se localizó en el grupo del complejo Fujikuroi, con una alta identidad probable para *Fusarium verticillioides*. Por lo tanto, la identificación temprana y específica de especie de estos aislados fúngicos, utilizando métodos moleculares, puede permitir el diagnóstico oportuno de enfermedades fisiopatológicas emergentes de interés para la región, y así proponer las estrategias de control respectivas.

**Palabras clave:** Patógenos de la caña de azúcar; *Fusarium* sp.; Fitopatología molecular; filogenia.

### 1 Introduction

Sugarcane is an important product for Colombian and world economy, representing 70% of the sugar total production. This plant is focus of study and research because of its different utilities in the food industry like the production of biofuels, liquors, fertilizers and fibers like paper. Sugarcane is susceptible to multiple microorganism’s attacks, including filamentous fungi such as *Ceratocystis paradoxa*, *Colletotrichum falcatum*, *Fusarium verticillioides*, *Puccinia melanocephala*, among others [1], several of this pathogens exist in Colombia invading the sugarcane crops, among these are the species from the genus Fusarium, most of them are highly pathogenic and toxic [2].

The particular fungus *Fusarium verticillioides*, acts rotting host plants infecting from the stem to the leaves during the first 4 to 6 months of life [1]; those plants who manage to survive this period are severely affected morphologically as well as physiologically. These symptoms reduce the productivity of crops, strongly and directly impacting the commercial production of sugarcane [3]. Also, reports of poisoning and development pathologies due to intake of products infected by *F. verticillioides* have been found [4] this related to the high production and release of mycotoxins generated by the Fusarium genus fungi, these mycotoxins are known as fumonisins [4]. The species *Fusarium verticillioides* is highlighted within the species of its genus for being a large producer of fumonisins, with the highest productivity. Fumonisins will not break down or denature in the...
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The presence of chemical agents neither at high temperatures, so it is almost certain that if these mycotoxins are present in sugarcane crops used for human consumption products, they will reach up to processed products and finally they would be consumed, representing a serious risk to health. Due to the wide range of products that derives from sugarcane, the exposure of people to these toxins would be significantly high [5].

*Fusarium verticillioides* belongs to the *Gibberella fujikuroi* complex, this is a monophyletic taxon that includes at least 50 species from the genus *Fusarium*, with high similarity and overlapping morphological traits that makes their differentiation difficult and confusing [6]. It’s here where the importance of molecular identification lies, as relying only on morphological characterization may probably lead to errors on determining *Fusarium* species. This complex is also recognized because of the remarkable ability of its species to produce a wide range of mycotoxins and secondary metabolites that produce severe diseases in humans and animals [7], but the amount of toxins produced varies between species [8], therefore, knowing which taxonomic group was isolated in this work, it allows to establish the size of the threat that represents for food consumption security, crops safety and productivity.

To date in Colombia, *Fusarium verticillioides* has only been reported in the South of the country by [2], but there was not a formal report on the presence of the fungus *Fusarium verticillioides* as causal agent of Pokkah boeng disease and as a phytopathogenic agent for the case of sugarcane in Antioquia. For this reason, the principal aim of this project is to make a report of the presence of a sugarcane pathogenic fungus unknown in the region and contributing to molecular systematics and plant pathology studies.

2 Materials and methods

2.1 Isolation and cultivation of the fungus.

Sugarcane samples, which showed symptoms of red coloration and rotting characteristics typical of Pokkah Boeng disease, were collected in the municipality of Medellín (Antioquia, Colombia). Fungus-infested plant mate-
rial was collected under the frame collection permit granted by the Agen-
cia Nacional de Licencias Ambientales (ANLA) to CIBIOP research group
(Universidad EAFIT, Resolution No. 1566 of 2014). The infested cane
was acquired around the Poblado metro-station in Medellín, Colombia in a
local sale of “guarapo” (sugarcane juice). The cane used in those beverages
regularly comes from the municipalities of Barbosa and Copacabana where
it is cultivated. The tissue samples from infected stems were immersed in
1.5% sodium hypochlorite for 1 minute, then in 75% ethanol for 30 sec-
onds and finally washed in distilled water to eliminate the excess of these
two compounds; all this with the aim of eliminating superficial pathogens
as our microorganism of interest is a vascular fungus. Finally, disinfected
tissue was cultured in acidified potato dextrose agar medium (PDA) (0,2%
lactic acid) to prevent the growth of contaminant bacteria. A monoconoi-
dial culture was not obtained, nevertheless, what was observed through
microscopy demonstrates the purity of the culture.

2.2 Sugarcane infection and control

The same infection methodology described in [9] was used. Three sugarcane
plantlets (4 weeks old) were infected with the fungus strain EA-FP0013 and
three others were used as control in greenhouse. Two horizontal cuts were
made perpendicular from two vertical cuts separated by 5 mm each one
of them at the upper part of the stem. Then a piece of a similar size
was taken from the medium where the fungus was growing and put on the
wound generated to favor the contact between the fungus and the plant.
The controls were subjected to the same procedure but they were treated
only with agar without any microorganism. This same test was also carried
out using 8 banana plantlets (Musa acuminata cultivar Williams, 4 weeks
old) in the same way, 4 as control and 4 infected, this to verify the possible
host range of the EA-FP0013 strain. After 5 weeks of treatment tissue
samples were taken and the same isolation process was made to determine
if the same microorganism used for inoculation was obtained.
2.3 Molecular identification

The isolate was grown in Sabouraud dextrose broth for one week at 28 °C and constant agitation (200 rpm). Total DNA was extracted from the mycelium of the fungus following a CTAB protocol modified for the isolation of DNA from filamentous fungi, as described in [10]. The polymerase chain reaction (PCR) method was used to amplify marker genes ITS, Elongation Factor 1-α (EF) and β-Tubulin (Btub). The sequences of the primers used to amplify the ITS regions were taken from [11] and from [12], [13] for Btub and EF, see supplemental Table 1. The amplification programs were used with the following parameters: for ITS; 5 min at 95°C, 35 cycles x (1 min 94 °C, 1 min 55 °C, 2 min 72 °C), 5 min at 72 °C. For B-tubulin 5 min at 95 °C, 35 cycles x (35 sec 94 °C, 55 sec 55.4 °C, 2 min 72 °C), 5 min at 72 °C. For EF-1 it was 5 min at 95 °C, 35 cycles x (1 min 94 °C, 55 sec 50 °C, 2 min 72 °C), 5 min at 72 °C. All reactions were performed in a C1000 Thermal Cycler (BioRad Technologies).

The amplification products were visualized on 1% agarose gel and quantified in Nanodrop (ThermoFisher Nano2000) for subsequent shipment to the Sanger sequencing service at the facility of the Universidad de los Andes, Colombia. The ab1 files obtained from the facility were analyzed (trimming bases below P20 and assembled to obtain consensus) using scripts in Python and tools from Biopython libraries [14]. The final consensuses from each gene were deposited in GenBank database of NCBI (National Center for Biotechnology Information - USA) under the following accession numbers: ITS (MH050788), Btub (MH899102), EF (MH899103).

2.4 Phylogenetic analysis

Two phylogenetic analysis were performed: 1) a single locus analysis based on ITS sequences and 2) a MultiLocus Sequence Analysis (MLSA) using three loci; the first corresponds to internal transcribed spacer region of ribosomal RNA locus (ITS-1, 5.8S and ITS-2). The other two correspond to conserved functional genes: β-Tubulin (B-tub) and Elongation Factor 1-α (EF). Sequences for comparison were retrieved using NCBI-BLAST against nucleotide database, using Biopython 1.67 modules [14]. The taxa
used for each analysis an its accession numbers are listed in Supplemental File 1 presented as annexes.

The PhyPipe automated pipeline [15] (available at: https://gitlab.com/cibiop/phypipe/) was used for phylogenetic reconstruction by Maximum Likelihood (ML) and Bayesian Inference (BI) methods. The pipeline of PhyPipe comprises alignment of DNA sequences with MAFFT 7.222, then partition analysis with PartitionFinder and phylogenetic reconstruction with RAxML 8.2.8, MrBayes 3.2.6 or Garli 2.01. For BI analysis, MrBayes was executed under the following parameters: two independent MCMC runs, four chains, 1,000,000 generations, 35% of the relative frequency of burning and sampling of 100. For ML analysis, Garli was executed by first performing a search ML (5 independent searches), 1000 bootstrap pseudoreplicates were made and mapped to the best ML topology using SumTrees from the DendroPy 4.1.0 package.

For the phylogenetic tree in Figure 4 all the other species belong to the Gibberella fujikuroi complex to have a balanced matrix. In addition the species used were only those that had the available sequences of the 3 genes (ITS, EF and B-tub) in the databases.

3 Results and discussion

3.1 Identification and morphological characterization of isolated fungus

A filamentous fungus strain EA-FP0013 was successfully isolated in PDA medium from infected sugarcane tissue, whose characteristics coincide with the most representative characters of the strains of Fusarium verticillioides reported in the literature by [16,17,18,19,20]. The isolated fungus presents mycelial growth with colorations between pink, purple and red in the center with white edges (Figure 1A), radial growth, oval microconidia with flattened base and grouped in chains (Figure 1B and 1D). Also long, thin and septate macroconidia, with a curved apical cell (Figure 1C). The strain did not form chlamydospores at in vitro level in the evaluated medium.
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*F. verticillioides* is one of the species of the *Gibberella fujikuroi* complex, so that, a differentiation of its morphological structures is needed in order to discard other fungus identities as Fusarium subglutinans, Fusarium sacchari and Fusarium pseudonygamai. According to previous articles, *F. pseudonygamai* forms large microconidia aseptate [17]; *F. sacchari* forms septate macroconidia thin, curved and with the apical cell curved in tip and basal cell in foot shape [20] and *F. subglutinans* forms curved, thin macroconidia and almost cylindrical microconidia [21]. None of these characteristics were observed under the microscope in our isolated strain. Another important distinctive character observed in EA-FP0013 was the absence of microconidia grouped in chains in *F. subglutinans*, *F. pseudonygamai* and *F. sacchari*. All this evidence suggests that EA-FP0013 strain belongs to the species *verticillioides*, but as it is a complex of species, it important to corroborate using molecular identification and phylogenetic analysis.

![Figure 1](image1.png)

**Figure 1:** A. Fungus isolated from the sampled sugar cane, with purple and white mycelium. B. Stained tissue of the fungus visualized under the microscope (40X) septate macroconidia with curved apical cell in red and oval microconidia in yellow. C. Stained tissue of the fungus visualized under the microscope (40X) microconidia in chains.

### 3.2 Infection experiments

After obtaining a sample of infected cane and having isolated a fungus with the characteristics of a *Fusarium* strain, it was necessary to confirm if indeed the isolated fungus was the causative agent of the visible infection in the sugarcane sample. For this, Koch’s postulates were tested to
determine its pathogenic activity. The results showed that isolated fungus EA-FP0013 was indeed the infectious agent generating the disease, causing the same symptoms of Pokka Boheng in the sugarcane used for this experiment. As can be seen in Figure 2 after the infection in the greenhouse, stems of healthy sugar cane developed the same red rot (Figure 2D and 2E) visible in Figure 2A and culturing of samples from those plants resulted in the isolation of a fungus (Figure 2B) with the same phenotypic characteristics as the one in Figure 1A. One control is visible in figure 2C, with no traces of the red rot. By looking tissue of the infected sugarcane under the stereoscope, it can be observed how the rot invades the stem fibers (Supplemental Figure 1A and 1B), and when observing the tissue under the microscope one can see how the fungus invades the phloem’s cells, this (Supplemental Figure, 1C and 1D) is the reason why the amount of juice that can be extracted from the sugarcane decreases when it is infected with Pokka boheng [21]. In all of the infected plants, the infection was extended through the vascular system to the leaves but it did not reach them and was neither found in the roots. This was due to the fact that the model of infection in the greenhouse does not allow verification in the foliar and radicular areas as this symptoms are usually visible in very advanced stages of the disease.

In addition to carrying out the infection of sugarcane plants in the greenhouse, banana plants were also infected under the same procedure, in order to get more information about the fungus’s host range. The result showed that banana plants did not develop the disease but presented a rapid hypersensitivity response after several weeks of being infected (Figure 2F), where a small part of the cellular tissue surrounding the site of the infection passed through cell death in order to restrict the growth of the pathogen. This also helped us discard the possibility of the strain being F. oxysporum as this species of Fusarium is widely spread and recognized as a banana pathogen.
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**Figure 2:** A. Sample of sugar cane from which the fungus was isolated.  B. Fungus isolated from the sample in picture E.  C. Infection control plant from greenhouse experiment.  D and E. Symptoms developed by plants weeks after infection in greenhouse.  F. Banana plant weeks after infection in the greenhouse, it can be observed that the plant did not develop the disease.

### 3.3 Gene sequence analysis and molecular phylogenetics

An initial phylogenetic analysis was carried out with the ITS sequence from our fungal isolate and several other ITS sequences retrieved from GenBank, phylogenetic reconstructions were obtained from BI and ML analysis, obtaining in both cases the same topology (Figure 3). This topology shows a polytomy among the *Fusarium* genus belonging to the *Gibberella fujikuroi* complex, but it does show phylogenetic difference with *Fusarium proliferatum*, this being different from the polytomic sequences (with high support: BV>75 PP>0.95). Sequences from the *Gibberella fujikuroi* complex also differentiate from *F. oxysporum* and *F. subglutinans*, which also form a well supported clade. The lack of resolution may be due to the state of conservation of the ITS genes within the genus, that is to say that in this locus sufficient mutations have not accumulated enough so that the species can be differentiated based sole on this gene marker [22].
Figure 3: Phylogenetic tree derived from single analysis of related species of the genus *Fusarium* spp. using the nucleotide sequences of the ITS gene.
Looking for sequences of *F. verticillioides* in the databases, we only had ITS sequences in the GenBank, therefore *F. moniliformis* was used instead, but as the result shows, it does not group with the isolated sequence. It can be concluded that it is not *F. fujikuroi*, *F. moniliforme*, *F. proliferatum*, or *F. oxysporum*. In this point, there is no 100% certainty that the sequence isolated in this study is *F. verticillioides*, but there is sufficient evidence to say that it belongs to the *Gibberella fujikuroi* complex. Then, in order to give more resolution to this complex and differentiate EA-FP0013 from the other species, two other molecular markers were selected, EF and Btub. These markers have been used for many researchers around the world with successful results [23], [24], [25]. A phylogenetic analysis was carried out with concatenated genes and, through the ML and BI methodologies, a single phylogenetic tree was obtained (Figure 4). With this resolution it was possible to differentiate the species of the *Gibberella fujikuroi* complex with good support levels.

Subsequently, the Btub and EF genes were used as molecular markers with the aim of solving the polytomy. As a result, the new topology shows us that the sequences belong to different species and our sequence of interest (*Fusarium* sp. EA-FP0013) is the sister group of the clade containing *Fusarium moniliformis*, which is the old denomination of *Fusarium verticillioides* [22], [26] (Figure 4), all of them belonging to the *Gibberella fujikuroi* species complex. It can also be observed that all the species mentioned above differentiate from *Fusarium oxysporum*, including our strain in particular, discarding the possibility that EA-FP0013 is *F. oxysporum* [27] (Figure 4), this allows us to define with greater certainty that the isolated sequence of the sugarcane belongs to the phytopathogenic *Gibberella fujikuroi* species complex, possibly to *F. verticillioides* species according the morphologic similarities between this species and our sugarcane isolate [26]. Despite this, further studies are required in order to define the taxonomic status at species-level of this isolate. Finally, it was observed that *F. oxysporum* was not part of the complex, being this the reason why in the analysis that takes into account EF and B-tub genes this species was used as an outgroup. *F. oxysporum* is well reported as a pathogen of banana and other monocotyledons. Today this species counts with special attention thanks to owning races that devastate the cultivation of bananas. From this work we show that the isolated *Fusarium* strain does not belong
to the *F. oxysporum* species, since both molecularly and pathologically it does not present the characteristics of this lineage.

**Figure 4:** Phylogenetic tree derived from multilocus analysis of related species of the genus *Fusarium* spp. using the nucleotide sequences of the genes ITS, beta tubulin (ITS, B-tub) and elongation factor (EF).
4 Conclusions

From a sample of sugar cane showing the symptoms of Pokkah boeng, a strain of phytopathogenic fungus belonging to the genus *Fusarium* and to the *Gibberella fujikuroi* complex, probably of the species *Fusarium verticillioides*, was isolated through morphological and molecular techniques. We validated the Koch’s postulates infesting cane plantlets and determined its pathogenic activity, concluding also that the EA-FP0013 strain was not pathogenic for banana plants. The presence of this pathogenic species was unknown, thus, this report opens the doors to different perspectives such as research into the biology of this pathogen and about the disease it produces, structuring mitigation and control strategies to avoid economic losses and food contamination problems, alert health authorities and Studies of the toxins produced by the fungus and its impact on the health of consumers, all of them of highly importance for the industry and public health.

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**Supplementary data**

**Table 1:** Primers and annealing temperatures used for amplification and sequencing.

| Locus                  | Annealing temp (C) | Cycles | Primer | Direction | Primer sequence | Ref. |
|------------------------|--------------------|--------|--------|-----------|-----------------|------|
| Internal Transcribed Spacer (ITS) | 55                | 35     | ITS1   | Forward   | TCCGT AGGTG AACCT GCGG | [11] |
|                         |                    |        | ITS4   | Reverse   | TCCTC CGCTT ATTGA TATGC | [11] |
| Beta Tubulin           | 55.4               | 35     | T1     | Forward   | AACAT GCGTG AGATT GTAAG T | [12] |
| Beta Tubulin           | 55.4               | 35     | T22    | Reverse   | TCTGG ATGTG GTTGG GAATTC | [12] |
| Elongation Factor      | 50                 | 35     | EF-1   | Forward   | ATGGG TAAGG A(A/G)G ACAAG AC | [13] |
| Elongation Factor      | 50                 | 35     | EF-2   | Reverse   | GGA(G/A) GTACC AGT(G/C) ATCAT GTT | [13] |
**Table 2:** Taxa and sequences used for single locus phylogenetic analysis based on ITS marker

| Organism                  | Accession number          |
|---------------------------|---------------------------|
| *Fusarium* sp. EA-FP0013  | MH050788.1 (this study)   |
| *Fusarium verticillioides*| HQ248201.1                |
| *Fusarium pseudonygamai*  | DQ297561.1                |
| *Fusarium subglutinans*   | AY898263.1                |
| *Fusarium fujikuroi*      | AY898249.1                |
| *Fusarium solani*         | DQ094497.1                |
| *Fusarium falciforme*     | DQ094439.1                |
| *Fusarium napiforme*      | DQ297555.1                |
| *Fusarium proliferatum*   | GU074009.1                |
| *Fusarium oxysporum*      | GU361925.1                |
| *Colletotrichum falcatum* | NR_144793.1               |
| *Trichoderma viride*      | AM498442.1                |
| *Trichoderma harzianum*   | FJ442671.1                |
| *Pestalotiopsis disseminata* | EF055196.1           |
| *Pestalotiopsis versicolor* | DQ812940.1           |
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**Table 3:** Taxa and sequences used for single locus phylogenetic analysis based on ITS marker

| Organism                  | ITS          | B-Tub        | EF            |
|---------------------------|--------------|--------------|---------------|
| *Fusarium proliferatum*   | EU821492     | KT218534.1   | KT218533.1    |
| *Fusarium oxysporum*      | FJ466709.1   | KX253989.1   | KF918544.1    |
| *Fusarium moliniformis*   | JF499676.1   | FN545357.1   | JQ639211.1    |
| *Fusarium fujikuroi*      | KJ000433.1   | KF466438.1   | HF679028.1    |
| *Fusarium sp.* EA-FP0013 (this study) | MH050788.1 | MH899102.1  | MH899103.1    |
Figure 5: A and B. infected plant tissue seen in the stereoscope where you can see the characteristic red color product of stem rot, C and D. infected plant tissue seen in a microscope where the invasion of the cells of the stem can be observed phloem from the fungus.
Figure 6: Agarose gel electrophoresis showing DNA amplifications products of the three amplified loci in this study