Costus spiralis (Costaceae) is a species native to the Amazon region and is used in traditional medicine. The endophytic fungi used in this study were obtained from leaves of this plant. 13 strains were selected to obtain hydroethanolic extracts and were submitted to hydroalcoholic extraction and evaluated for antioxidant activity by DPPH (2,2-difenil-1-picrilhidrazil) and FRAP (ferric reducing antioxidant power), and all of the fungi had positive results. The antimicrobial action of crude extracts had a good range of activities. All extracts had inhibitory activities against the yeasts of Candida albicans and C. parapsilosis, with 125 to 500 μg/mL MIC. Eight extracts had antimicrobial activities against Bacillus subtilis (MIC from 62.4 to 125 μg/mL), 5 against Pseudomonas aeruginosa (MIC from 125 to 500 μg/mL), 2 against Salmonella enterica (MIC from 125 to 62.5 μg/mL), and 2 against Enterococcus faecalis (MIC from 500 to 125 μg/mL). The presence of secondary metabolites, including coumarins, was observed during chemical evaluation by thin layer chromatography. Total phenol content was estimated, and a strong positive correlation to antioxidant activity was observed, according to its Pearson coefficient. This is the first report of the bioactive potential of endophytic fungi isolated from the Costaceae family in Brazilian ecosystems.

1. Introduction

Costus spiralis is a Brazilian Amazon plant that is recognized for its medical and ornamental values. This plant is commonly used in popular medicine to treat urinary infections and kidney stones and for diabetes management. The plant also acts as an antioxidant, antibacterial agent and diuretic and to promote wound healing [1–3]. Normally, plants from tropical regions have a higher diversity of endophytic microorganisms compared with those found in temperate regions [4]. Endophytic fungi are microorganisms that colonize the vegetal tissues, either inter- or intracellularly, without causing any damage to the vegetal host. There are few studies that focus on the diversity, ecology, and other factors involved in endophyte plant interaction [5–7]. These microorganisms are a good resource for identifying new bioactive products, with more than 20,000 substances described to date [8]. Among these substances, 51% had novel structures and 80% exhibited some biological activity making endophytes a promising resource for the identification of new bioactive molecules [9, 10]. A variety of pharmacological activities have already been described from endophytic fungi, including antibacterial [11–13], antifungal [14–16], antiparasitic [17], trypanocidal [18,19], leishmanicidal [19], antimalarial [20], anti-inflammatory [21], neuroprotective [22], antioxidant [7], immunosuppres- sion [23], antiviral [23, 24], anticolinesterasic [12], antineoplastic [25–29], and cytotoxic [12, 22, 30] properties.
Despite scientific advances, infectious diseases remain a major contributor to mortality and morbidity in public health. The main explanation for this is the ability of microorganisms to acquire resistance against antimicrobial substances. This characteristic results in a constant need to discover and develop new drugs [31].

Oxygen radicals and superoxide anions play important roles in biochemical/physiological reactions in the human body. However, when these species are produced in excess due to pathophysiological processes or environmental interference, they can promote tissue damage and result in disease [32]. Currently, few antioxidant substances can be used in clinical situations, thus underscoring the necessity to identify new, safe, and efficient molecules for this purpose [33]. Bioactive molecules from new sources, such as endophytic fungi, deserve attention because they may lead to drugs with different pathologies, food additives, or cosmetics.

2. Materials and Methods

2.1. Endophytic Fungi. Endophytic fungi were isolated from healthy leaves of C. spiralis. From each leaf three 0.5 cm² pieces were cut. The leaf sample fragments were surface sterilized by successive dipping in 2% Extran detergent (Sigma-Aldrich) for 2 min, 70% ethanol for 1 min, and 2% sodium hypochlorite for 3 min and then by washing with sterile distilled water for 2 min; they were then placed on potato dextrose agar (PDA; Difco) containing 100 mg/mL chloramphenicol. The sterile water wash was plated on PDA to confirm the infection process. Plates were incubated for up to 60 days at 25°C. The mycelia from the margins of fungal colonies growing from the leaf fragments were transferred to fresh PDA and further purified by transferring hyphal plugs to new PDA plates. Thirteen endophytic fungi were isolated from leaves of C. spiralis. These microorganisms were previously tested with phytopathogen controls (data not shown) and are preserved in Culture Collection from the Laboratory of General and Applied Microbiology at the Federal University of Tocantins, Brazil.

2.2. Molecular Identification

2.2.1. Fungal DNA Extraction. Endophytic fungi were cultured on Sabouraud dextrose agar slants for 7 days at 25°C. The DNA extraction was performed according to the method described by de Hoog et al. [34]. Briefly, approximately 0.1 g of fungal mycelia was transferred to 1.5 mL tubes containing 400 μL of lysis buffer (50 mM Tris-HCl, pH 8.0, 5 mM EDTA pH 8.0, 100 mM NaCl, 1% SDS) and kept at −20°C for 10 min. The fungi cells were mixed with 1 mm glass beads and agitated using a Vortex for 30 sec followed by a 30 sec interval on ice, then repeated. Cells were incubated for 30 min at 60°C with 5 μL 20 mg/mL Proteinase K. The samples were then incubated for 10 min at 65°C with 162 μL of CTAB solution (200 mM Tris-HCl, pH 7.5, 200 mM Na-EDTA, 8.2% NaCl, 2% CTAB). Following incubation, 570 μL chloroform:isooamylic alcohol (24:1 v/v) was added, and the tubes were kept for 10 min on ice. The homogenate was centrifuged at 14,500 xg for 10 min at 4°C. Then, the supernatant was transferred to a new tube, and 10% of the residual volume of 3 M sodium acetate was added. The mixture was then incubated on ice for 30 min. Precipitates were removed by centrifugation, and the DNA was recovered by isopropanol precipitation, washed with 70% ethanol, allowed to air dry, and resuspended in 50 μL of TE buffer (10 mM Tris-HCl, pH 8.0, and 0.1 mM EDTA). The concentration was measured by monitoring the UV absorbance at 260 nm using Nanodrop 2000 (Thermo USA), and the DNA was diluted to 10 ng/μL with ultrapure water.

2.2.2. DNA Amplification and Sequencing. DNA sequencing was performed using a BigDye Direct Cycle Sequencing Kit (Life Technologies). For PCR amplification, 1 μL of diluted DNA (10 ng/μL) was mixed with 5.0 μL of BigDye Direct PCR Master Mix, 0.75 μL of each forward ITS1-M13 (5′-TGTAAAACGACGGCCAGT-3′) and reverse ITS4-M13 (5′-TCCGCTTATGATATGC-3′) primers at a 10 μM concentration, which include the M13 universal primer sequences, and 2.4 μL of ultrapure water. PCR cycling conditions were as follows: initial denaturation of 10 min at 94°C, followed by 35 cycles (30 s) of denaturation at 96°C, annealing for 15 sec at 62°C, and extension for 30 sec at 72°C. Following PCR, 3 μL of BigDye Direct Sequencing Master Mix and 1 μL of forward BigDye Direct M13 forward primer (provided) were added directly to the PCR product, and back into the PCR thermal cycler, and incubated as follows: 15 min at 37°C, 2 min at 80°C, and 1 min at 96°C, followed by 25 cycles of 10 sec at 96°C, 5 sec at 50°C, and 75 sec at 60°C.

At the completion of the sequencing reaction, the sequencing products were purified following the ethanol/EDTA protocol for BigDye Terminator version 3.1 Cycle Sequencing Kit. Electrophoresis was performed on the Applied Biosystems 3500XL Genetic Analyzer. Electropherograms were proofread with the software BioEdit Sequence Alignment Editor (1997–2013). The sequencing results for each sample were analyzed using BLASTn (Basic Local Alignment Search Tool, version 2.2.15 of BLAST 2.0) to verify the results and to identify the fungus for each sample.

2.3. Crude Extract Acquisition. The 13 endophytes were multiplicated in PDA at 25°C ± 2 for 5 days. Subsequently, 6 mm discs were placed in the center of 4 Petri dishes with PDA agar and incubated at 25°C ± 2 for 14 days. These cultures were extracted (1:20 p/v) by hydroethanolic maceration (1:3 v/v) for 48 h at 25°C, under agitation. The samples were centrifuged at 2,400 × g for 20 min at 8°C and then filtered. The solvent from each supernatant was removed on a rotary evaporator at 35°C, lyophilized, and stored in a desiccator without light. The same extraction process was performed for 4 sterile BDA Petri dishes that were used as negative control.

2.4. Antimicrobial Activity

2.4.1. Microorganisms. All microorganisms were obtained from the American Type Culture Collection (ATCC, Rockville,
Bacillus subtilis (G+ve) bacteria were treated culture was spread on plates with antibiotic-free medium at a density adjusted to a 0.5 McFarland turbidity standard (10⁶ CFU/mL) and diluted 1:10 for the broth microdilution procedure. Microtiter plates were incubated at 37°C, and the MICs were recorded after 24 h of incubation. The lyophilized crude extracts of endophytes were diluted in dimethylsulfoxide (Sigma, D8418), filtered in sterile membranes, and tested at 1000, 500, 250, 125, 62.5, 31, 25, 7.8, 3.9, and 1.9 µg/mL concentrations. The MIC was defined as the lowest concentration of crude extracts that inhibit the target microorganisms’ growth.

The minimal bactericidal concentrations (MBCs) were determined from the results obtained in MIC. For this, an aliquot of 10 µL from the wells that had inhibition was spread in Petri dishes with Mueller-Hinton agar (Fluka, 70191) and incubated at 37°C for 24 h. The MBCs were defined as the lowest concentration of crude extract that resulted in no growth when the treated culture was spread on antibiotic-free medium plates after the incubation. All tests were repeated and confirmed.

2.4.2. Antibacterial Susceptibility Test. The minimal inhibitory concentrations (MICs) of all extracts and the reference antibiotics tetracycline (Sigma, T3258) and penicillin (Sigma, P3032) were determined using microdilution techniques in Mueller-Hinton broth (Merck) following the protocol established for bacteria [35]. Inoculates were prepared in the same medium at a density adjusted to a 0.5 McFarland turbidity standard (10⁸ CFU/mL) and diluted 1:10 for the broth microdilution procedure. Microtiter plates were incubated at 37°C, and the MICs were recorded after 24 h of incubation. The lyophilized crude extracts of endophytes were diluted in dimethylsulfoxide (Sigma, D8418), filtered in sterile membranes, and tested at 1000, 500, 250, 125, 62.5, 31, 25, 7.8, 3.9, and 1.9 µg/mL concentrations. The MIC was defined as the lowest concentration of crude extracts that inhibit the target microorganisms’ growth.

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2.4.3. Antifungal Susceptibility Test. The MICs of crude extracts were determined against yeasts using broth microdilution techniques, according to the method described by CLSI [36]. MICs were determined in RPMI 1640 medium (Sigma, R6504), pH 7.0. The starting inoculum was 1.0 × 10⁸ CFU/mL. Microtiter plates were incubated at 37°C in a dark humid chamber, and MICs were recorded after 48 h. The lyophilized hydroethanolic extracts from endophytes were diluted in dimethylsulfoxide (Sigma, D8418), filtered in sterile membranes, and tested at 1000, 500, 250, 125, 62.5, 31, 25, 7.8, 3.9, and 1.9 µg/mL concentrations. The MIC was defined as the lowest concentration of compounds without microorganism growth. Nystatin (Sigma, N6261) was used as the drug control. The minimal fungicidal concentrations (MFCs) were obtained from the MIC results. For this, an aliquot of 10 µL from the wells that showed inhibition was spread in Petri dishes with Sabouraud agar (Merck, 105438) and incubated at 37°C for 24 h. The MFCs were defined as the lowest concentration of crude extract that resulted in no growth when the treated culture was spread on plates with antibiotic-free medium after the incubation. All tests were repeated and confirmed.

2.5. Antioxidant Assay

2.5.1. Thin Layer Chromatography (TLC). The antioxidant activity from the hydroethanolic extracts of the 13 fungi strains was evaluated by thin layer chromatography (TLC). The extracts were dissolved in a hydroethanolic solution (3:1 v/v) to achieve a final concentration of 20 mg/mL. The solution was loaded onto a TLC plate (20 × 20 cm, Merck) and eluted with ethyl acetate, formic acid, acetic acid, and water (100:11:27 v/v). A methanolic solution of 2,2-difenil-1-picrilhidrazil (DPPH) (Aldrich, D9132) at 0.2% was used to visualize the products. Quercetin (Sigma, Q0125) and ascorbic acid (Sigma, A0278) were used as standards. Positive samples showed yellow bands on a purple background in chromatograms.

2.5.2. Determination of Antioxidant Activities by DPPH Method. Antioxidant activity was determined in accordance with the published methods [37, 38]. Ascorbic acid was used as a standard (2–10 µg/mL, Sigma, A0278). Fungi extracts and negative controls were diluted to 25, 50, 100, 150, and 200 µg/mL. A methanolic solution of DPPH (0.1 mM, 2 mL) was added to 1 mL of each dilution. For each experiment, solutions with 1 mL sample and 2 mL ethanol were used to establish a baseline. Controls were performed with 1 mL of ethanol and 2 mL of methanolic solution of DPPH (0.1 mM). These solutions were homogenized and kept in the dark for 30 min at 25°C, and the absorbance was obtained at 517 nm with a spectrophotometer (Biochrom, Model BioWave II). All tests were conducted in triplicate.

Antioxidant activity of each extract concentration was determined using the following equation:

\[ AA\% = 100 \left( 1 - \frac{\text{Abs}_{\text{sample}} - \text{Abs}_{\text{blank}}}{\text{Abs}_{\text{blank}}} \right) \times \frac{100}{\text{Abs}_{\text{contr}}} \]  

where AA is the total antioxidant activity, Abs_{sample} is the absorbance of samples or standard, Abs_{blank} is the baseline absorbance, and Abs_{contr} is the absorbance of the control solution.

Linear regression curves were obtained from the AA% results and their respective results and predictive equations were expressed as CE₅₀.

2.5.3. Determining the Antioxidant Activity by Ferric Reducing Antioxidant Power (FRAP). FRAP analyses were performed in accordance with the published protocols [39]. An acetate buffer solution (0.3 M, pH 3.6), a 10 mM solution of 2,4,6-tris (2piridil)-s-triazine (TPTZ) in 40 mM HCl, and an aqueous solution of ferric chloride (20 mM) were prepared. The FRAP reagent was prepared few minutes before mixing the solutions for analysis (10:1:1 v/v).

The total antioxidant activity (TAA) for each of the crude extracts was determined from three methanolic dilutions (0.5, 0.25, and 0.125 mg/mL) and 6 aqueous dilutions using ferrous sulfate (Sigma-Aldrich, 61230) as a standard in 500 and 2000 µM concentrations. For each sample, 0.1 mL of each dilution was added to 2.9 mL of FRAP and kept in the dark at 37°C for 30 min. The absorbance of each solution was
determined at 593 nm using a spectrophotometer (Biochrom, Model Biowave II) and FRAP as an internal reference. From these results, calibration curves were calculated for each sample. The result of TAA from each sample was expressed in millimole equivalents of iron per mg of lyophilized fungi extract (mM Fe\(^{2+}\)/mg Ext).

2.6. Chemical Evaluation of the Extracts by TLC. The secondary metabolites from the 13 hydroethanolic extracts and from the negative control were evaluated by TLC using different mobile phases and visualizing agents in aluminum sheets (20 × 20 cm) preactivated, covered by silicagel GF\(_{254}\) (Merck). The samples were dissolved in hydroethanolic solution (1:3 v/v). The sheets were placed in a chromatography chamber, eluted along a 16 cm path in one direction, and observed by luminescence (UV365 nm). Using the previously described techniques and parameters for identification [40] and identical standards, we were able to evaluate the following classes of secondary metabolites: alkaloids, with quinine standard (Aldrich, 145904) and caffeine (Calbiochem, 205548); monoterpenoids, sesquiterpenoids, and diterpenoids using a thymol standard (Sigma, T0501); triterpenoids and steroids, with a stigmasterol standard (Sigma, S2424); phenolic compounds, with a tannic acid standard (Sigma-Aldrich, S2424); condensed proanthocyanidins and leucoanthocyanidins, with a catechin standard (Fluka, 43412).

2.7. Quantification of Total Phenolic Compounds. The quantification of phenols in hydroethanolic extracts was realized using the Folin-Ciocalteu method [41], using tannic acid (Sigma-Aldrich, 403040) as a standard. An aliquot (150 μL) from each sample (1.0 mg/mL) was added to 250 μL of Folin-Ciocalteu reagent. 500 μL of sodium carbonate solution (7.5%, w/v) was then added to the mixture. The sample was diluted to 5.0 mL with water and left to stand in the dark for 30 min at 25°C. Each sample had its absorbance verified at 760 nm in spectrophotometer (Biochrom, Model Biowave II) using water to establish a baseline.

The calibration curve was calculated using tannic acid at 0.1–0.5 μg/mL as a standard. The total phenol content was expressed in milligram equivalents of tannic acid per gram of extract (mg EAT/g) using linear equations. The assays were conducted in triplicate, and the experimental design was completely analyzed using the Tukey test (ANOVA) with the ASSISTAT 7.6 beta software.

Pearson correlation analysis was performed using the BioEstat 5.3 software, and graphics were prepared with OriginPro 8.6. We used the following ranges of the Pearson coefficient (r) to represent an absent or very weak correlation (0.00 to 0.19), a weak correlation (0.20 to 0.39), a moderate correlation (0.40 to 0.59), a strong correlation (0.60 to 0.79), and a very strong correlation (0.80 to 1) [42].

### 3. Results and Discussions

3.1. Molecular Identification. Molecular identification was achieved by sequencing the Internal Transcribed Spacer (ITS) region. Sequences with 99% or more similarity were considered to be from the same species. Those with similarities between 93% and 98% were considered to be from the same genus. Sequences below 93% similarity were considered to be a previously unidentified strain [43]. Among the thirteen endophyte fungi analyzed, 11 were identified as being from the Phomopsis/Diaporthe, Cochliobolus, or Sordariomycetes genus (Table 1).

The unidentified fungi (4426 and 4400) may represent new species, and further studies are needed to determine their phylogenetic classification.

3.2. Antibacterial and Antifungal Activities. The endophytic fungi extracts were tested against G+ve and G–ve bacteria.
Table 2: Minimum inhibitory concentration (MIC, in $\mu$g/mL) of crude extracts obtained from endophytic fungi isolated from Costus spiralis against yeasts and bacteria.

| Extract code | Endophytic species | Yeasts | Bacteria |
|--------------|--------------------|--------|----------|
|              |                    | Ca     | Cp       | Ec       | Pa       | Kp       | Se       | Bs       | Sa       | Ef       |
| CEBP1        | Phomopsis sp.      | 125    | 125 a   | 125 a    | a        | a        | 125 a    | a        | a        |
| CEDp2        | Diaporthe phaseolorum | 500    | 500 a   | 500 a    | a        | a        | a        | a        | a        |
| CED3         | Diaporthe sp.      | 500    | 500 a   | 500 a    | a        | a        | a        | a        | a        |
| CED4         | Diaporthe sp.      | 500    | 500 a   | 500 a    | a        | a        | a        | a        | a        |
| CEP4         | Phomopsis sp.      | 125    | 125 a   | 125 a    | a        | a        | 62.5 a   | a        | a        |
| CE6          | Not identified     | 500    | 500 a   | 500 a    | a        | a        | a        | a        | a        |
| CED7         | Diaporthe sp.      | 250    | 250 a   | 250 a    | a        | a        | 125 a    | a        | a        |
| CES8         | Sordariomycetes sp.| 250    | 250 a   | 250 a    | a        | a        | 125 a    | a        | a        |
| CE9          | Not identified     | 500    | 500 a   | 125 a    | a        | a        | 125 a    | a        | a        |
| CEP10        | Phomopsis sp.      | 500    | 500 a   | 500 a    | a        | a        | a        | a        | a        |
| CEDp11       | Diaporthe phaseolorum | 125    | 125 a   | 250 a    | a        | 125 a    | 62.5 a   | 500 a    |
| CEC12        | Cochliobolus ssp.  | 125    | 125 a   | 125 a    | 62.5 a   | 125 a    | 125 a    |
| CES12        | Sordariomycetes sp.| 250    | 250 a   | 250 a    | a        | a        | 125 a    | a        | a        |
| CECn         | Not identified     | a      | a       | a        | a        | a        | a        | a        | a        |
| Nystatin     | 1.9                | 1.9    | —       | —        | —        | —        | —        | —        | —        |
| Penicillin   | —                  | —      | —       | —        | —        | —        | 1.9      | 3.9      | 3.9      |
| Tetracycline | —                  | —      | 1.9     | 3.9      | 15.6     | —        | —        | —        | —        |

Note: Ca: Candida albicans, Cp: C. parapsilosis, Ec: Escherichia coli, Pa: Pseudomonas aeruginosa, Kp: Klebsiella pneumoniae, Se: Salmonella enterica subsp. enterica serovar Typhi, Bs: Bacillus subtilis, Sa: Staphylococcus aureus, and Ef: Enterococcus faecalis.

a: MIC/MFC/MBC above 1,000 $\mu$g/mL; —: not available; CECn: crude extract control negative.

and yeasts. These microorganisms were selected according to their medical importance. The minimum inhibitory concentration (MIC) of extracts was determined using the microdilution method (Table 2) [44].

In the literature, there are no representative criteria for the MIC of endophytic fungi extracts. Therefore, we used the criteria cited in Table 3 for comparison [45].

The MIC values obtained for C. albicans and C. parapsilosis show moderate activities for all of the extracts tested. The CEBP1, CEP4, CEDp11, and CEC12 had a minimum fungicidal concentration (MFC) equal to 125 $\mu$g/mL for both yeasts, showing a fungicidal effect in the Phomopsis sp., D. phaseolorum, and Cochliobolus sp. extracts. All other extracts showed no fungicidal activity.

For the antibacterial evaluation, the tested extracts were active against G+ve and G−ve bacteria. Eight extracts were able to inhibit the growth of B. subtilis, with MIC between 62.5 and 125 $\mu$g/mL. B. subtilis was the microorganism most inhibited by the tested extracts, followed by P. aeruginosa, which was inhibited by five extracts, and S. enterica and E. faecalis, which were both inhibited by two extracts. The lower MICs (62.5 $\mu$g/mL), which are considered to be good activities, were obtained from CEC12 against S. enterica and from CEP5 and CEDp11 against B. subtilis, the last of which showed an MBC of 62.5 $\mu$g/mL. The other antibacterial assays did not report MBC. Strains CEDp11 and CEC12 inhibited more pathogens (C. albicans, C. parapsilosis, P. aeruginosa, S. enterica, B. subtilis, and E. faecalis).

Antimicrobial activities from endophytic fungi of C. spiralis have not previously been reported in the literature.

Table 3: Criteria of selection of positive antimicrobial activities of crude extracts.

| MIC of crude extract | Result |
|----------------------|--------|
| Below 100 $\mu$g/mL  | Good antimicrobial activity |
| Between 100 and 500 $\mu$g/mL | Moderate antimicrobial activity |
| Between 500 and 1000 $\mu$g/mL | Weak antimicrobial activity |
| Above 1000 $\mu$g/mL | Inactive |

MIC = minimum inhibitory concentration.

Our results thus demonstrate the biotechnological potential of these strains.

3.3. Chemical Evaluation of Extracts. Different classes of secondary metabolites were found by evaluating the 13 fungi extracts. We observed variations between the detected compounds and the fungi species (Table 4). Particularly noteworthy are the presence of triterpenes in five extracts and the presence of steroids and coumarins and the absence of alkaloids and condensable tannins in all analyzed extracts.

We conducted TLC using different mobile phases for the 13 analyzed extracts. The best resolution among the compounds was observed with toluene:ethyl-acetate (75:25 v/v), observed by UV at 365 nm. The fingerprints of all fungi extracts showed compounds with common characteristics of coumarins by emission of blue or green fluorescence, and these results were confirmed by NEU visualization at UV365 nm (Figure 1).
These results confirmed the importance of using secondary metabolites in the extracts, similar to the coumarins identified in previous assays.

Further, the organic fractions show higher resolution among the compounds, being visualized in higher number and others. This class of compounds was related to several biological activities, including protease inhibition, acetylcholinesterase, K vitamin antagonism, antimicrobial, growth regulation, antiallergenic, antimalaric, antiviral, immunosuppression, hypolipidemic, hypotensor, antispasmodic, and antioxidant activities [30, 47–50].

To obtain more information about the chemical compounds observed in the fingerprints, we conducted the fractionation of the crude extracts by liquid-liquid partition, using dichloromethane as the nonpolar organic phase and water as the polar phase. Fractions were analyzed by TLC and compared with the crude extracts. The results from CEDpII and CEC12 show the nonpolar nature of most of the compounds in the extracts, similar to the coumarins identified in previous assays.

The fingerprints show the presence of one complex matrix of compounds in the extracts. With respect to the molecular composition, we observed differences and a single profile for CEP6, CEDpII, CEC12, and CES13, and we observed similarities between CEP1 and CEDp2, the samples CED3, CED4, and CEP5, and the samples CED7, CES8, and CED9. It was also possible to observe variations in this matrix of compounds between different strains from D. phaseolorum species (CEDp2 and CEDpII).

Experiments that showed differences in the production of secondary metabolites were observed with Streptomyces sp. [46]. These results confirmed the importance of using fingerprint acquisition and chemical screening as a tool for the simple selection of chemical characteristics in bioprospection of fungi strains.

The coumarins are secondary metabolites found in different organisms such as vegetables, bacteria, fungi, lichens,

**Table 4: TLC results of secondary metabolites in the fungal extracts.**

| Crude extract code | Endophytic species       | UV 365 nm | NEU | Chloridric vanillin | KOH 10% | KOH 10% | Dragendorf | Sulfuric vanillin | Sulfuric Burchard | Liebermann Burchard |
|-------------------|--------------------------|-----------|-----|---------------------|---------|---------|------------|------------------|-------------------|---------------------|
| CEP1              | *Phomopsis* sp.          | (+)       | (+) | (-)                 | (+)     | (-)     | (-)        | (+)              | (+)               | (+)                 |
| CEDp2             | *Diaporthe phaseolorum* | (+)       | (+) | (-)                 | (+)     | (-)     | (-)        | (+)              | (+)               | (+)                 |
| CED3              | *Diaporthe* sp.          | (+)       | (+) | (-)                 | (+)     | (-)     | (-)        | (+)              | (+)               | (+)                 |
| CED4              | *Diaporthe* sp.          | (+)       | (+) | (-)                 | (+)     | (-)     | (-)        | (+)              | (+)               | (+)                 |
| CEP5              | *Phomopsis* sp.          | (+)       | (+) | (-)                 | (+)     | (-)     | (-)        | (+)              | (+)               | (+)                 |
| CEP9              | Not identified           | (+)       | (+) | (-)                 | (+)     | (-)     | (-)        | (+)              | (+)               | (+)                 |
| CES8              | *Sordariomyces* sp.      | (+)       | (+) | (-)                 | (+)     | (-)     | (-)        | (+)              | (+)               | (+)                 |
| CES9              | Not identified           | (+)       | (+) | (-)                 | (+)     | (-)     | (-)        | (+)              | (+)               | (+)                 |
| CEDp10            | *Phomopsis* sp.          | (+)       | (+) | (-)                 | (+)     | (-)     | (-)        | (+)              | (+)               | (+)                 |
| CEDp11            | *Diaporthe phaseolorum* | (+)       | (+) | (-)                 | (+)     | (-)     | (-)        | (+)              | (+)               | (+)                 |
| CEC12             | Cochliobolus ssp.        | (+)       | (+) | (-)                 | (+)     | (-)     | (-)        | (+)              | (+)               | (+)                 |
| CES13             | *Sordariomyces* sp.      | (+)       | (+) | (-)                 | (+)     | (-)     | (-)        | (+)              | (+)               | (+)                 |
| CECn              | Not identified           | (-)       | (-) | (-)                 | (-)     | (-)     | (-)        | (-)              | (-)               | (-)                 |

(+): positive; (-): negative; CECn: crude extract control negative.

1. Coumarins; 2. Phenolic compounds; 3. Proanthocyanins, leucoanthocyanins; 4. Coumarins (mobile phase to coumarins); 5. Anthraquinones, naphthoquinone, and anthocyanins (mobile phase to anthraquinones); 6. Alkaloids; 7. Phenols, sterols, terpenes, higher alcohols, polyketides, and saponins (mobile phase to saponins); 8. Phenols, sterols, terpenes, higher alcohols, polyketides, and saponins (mobile phase to mono-, sesqui-, and diterpenes); 9. Triterpenes (pink-reddish band); 10. Sterols (gray band).

**Figure 1:** Fingerprints of 13 fungi extracts obtained by TLC observed by UV at 365 nm. Mobile phase: toluene : ethyl acetate (75 : 25 v/v). 1: CEP1, 2: CEDp2, 3: CED3, 4: CED4, 5: CEP5, 6: CEP6, 7: CED7, 8: CES8, 9: CES9, 10: CEP10, 11: CEDp11, 12: CEC12, 13: CES13, C: negative control.
when compared with the crude extracts. These data show
the complex matrix of compounds present in the extracts
obtained from these fungi species. This complex matrix
can increase the interest in the possible biotechnological
applications of these extracts and encourage the elucidation
of the chemical structures of these molecules (Figure 2).

The CEC12 fingerprint (ethyl acetate: toluene, 25:75 v/v)
demonstrated 10 different fluorescent bands in the organic
fraction. Bands 2, 3, and 6 (R_f 0.34, 0.42, and 0.62) represent
compounds that are exclusive to this extract (Table 5).

When the mobile phase was used to separate anthraquinone
compounds, different from the other endophytic species,
the sample CEDp11 from D. phaseolorum showed peculiar
compounds. The CEDp11 fingerprint from the organic
fraction had 5 unique compounds, 3 of which were colored by
visible light in TLC (Table 5). The presence of orange and
yellow bands (R_f 0.45, 0.55, and 0.91) is characteristic of
anthraquinones. Typical fluorescence bands for coumarins
were visualized in this sample (Table 5).

These results show the complexity of the analyzed sam-
ples, particularly the organic fractions of CEDp11 and CEC12.

3.4. Quantification of Phenolic Compounds and Verification
of Antioxidant Activity. The quantification of phenolic com-
 pounds from the 13 crude extracts and negative controls
was conducted. Significant differences were found between
the samples (P < 0.01) and the negative control (Table 6).
Phenolic compounds have been very well known for their
antioxidant properties, owing to their unique ability to act as
free radical scavengers which, in turn, is an outstanding
attribute of their unique biochemical structure [51].

TLC screening revealed the presence of antioxidant sub-
stances in all samples, by the presence of yellow bands in
purple background resulting from the reduction of the DPPH
radical. As a result of these analyses, the samples were sub-
mited for characterization of their antioxidant activity using
the DPPH and FRAP methods. The results were significant,
particularly for CEC12, CE6, CES13, CEDp11, and CEPI
(Table 6).

The average of values of antioxidant activities obtained
from the FRAP and DPPH methods had a very strong
Pearson correlation (r = 0.901, P < 0.05), showing that these
methods have a positive correlation to the evaluated samples.
This correlation suggests that these compounds act as
hydrogen sources for DPPH radical and block the electron
donation, resulting in iron complex reduction in the FRAP
method. There was also a strong positive correlation between
the phenolic compound contents in samples, according to the
FRAP and DPPH results (r = 0.72, P < 0.05; r = 0.70,
P < 0.05, resp.), possibly attributed to the antioxidant activity
of phenolic substances, including coumarins.

Extracts that had higher antioxidant activities were sub-
mited to liquid-liquid fractionation with dichloromethane.
In the resulting organic fractions, the antioxidant activities
increased (Table 6), showing the nonpolar characteristic
of the antioxidant compounds present, as expected for
coumarins. This is the first report of antioxidant activities
from endophytic fungi from the Costaceae plants family.
Coumarins that contain dihydroxyl groups in the ortho posi-
tion, such as fraxetin (7,8-di-hydroxy-6-methoxy coumarin),
esculetin (6,7-di-hydroxy-coumarin), and 4-methyl esculetin
(6,7-di-hydroxy-4-methylcoumarin), are considered powerful
lipid peroxidation inhibitors and can eliminate the super-
oxide anion radical to promote iron chelation. These propri-
eties make these substances very interesting as antioxidants,
with possible applications in radial-free disease prevention
[50].

The extracts from endophytic fungi from C. spiralis
had antioxidant activities, and the important and complex
chemicals can be explored for biotechnological purposes.

Table 5: R_f values of bands from organic subfractions of CEDp11
(D. phaseolorum) and CEC12 (Cochliobolus ssp.) in TLC.

| Band | R_f | Characteristic                      |
|------|-----|------------------------------------|
| 1    | 0.19| Blue fluorescence (UV 365 nm)      |
| 2    | 0.45| Orange (visible), red fluorescence (UV 365 nm) |
| 3    | 0.52| Green fluorescence (UV 365 nm)     |
| 4    | 0.55| Orange (visible)                   |
| 5    | 0.91| Yellow (visible), fluorescence (UV 365 nm) |

CEDp11

| Band | R_f | Characteristic                      |
|------|-----|------------------------------------|
| 6    | 0.62| Blue fluorescence (UV 365 nm) — weak |
| 7    | 0.68| Blue fluorescence (UV 365 nm) — weak |
| 8    | 0.76| Blue fluorescence (UV 365 nm) — strong |
| 9    | 0.92| Blue fluorescence (UV 365 nm) — weak |
| 10   | 0.93| Blue fluorescence (UV 365 nm) — weak |

* Mobile phase: ethyl acetate : methanol : water (100: 13, 5: 10 (v/v)).
** Mobile phase: ethyl acetate : toluene (25: 75 (v/v)).
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