Phytotoxin Produced by *Bipolaris euphorbiae* in-vitro is Effective Against the Weed *Euphorbia heterophylla*

Aneli M. Barbosa\(^1\); Cristina G. M. Souza\(^1,2\); Robert F. H. Dekker\(^3\); Rafael C. Fonseca\(^1\) and Dalva T. Ferreira\(^4\)

Departamentos de: \(^1\) Bioquímica; \(^3\) Tecnologia de Alimentos e Medicamentos; e \(^4\) Química; Universidade Estadual de Londrina; Londrina - PR - Brazil. \(^2\) Presently at: Departamento de Bioquímica; Universidade Estadual de Maringá; Maringá - PR - Brazil

**ABSTRACT**

Four virulent strain isolates of the fungus, *Bipolaris euphorbiae* (previously identified as a *Helminthosporium* sp.), isolated from host plants in four states within Brazil were screened for the production of phytotoxins that promoted wilting and defoliation of the Brazilian weed, *Euphorbia heterophylla*, commonly found growing among soyabean crops. Only one isolate, *B. euphorbiae* Strain I (EUPH petropar from Mato Grosso state), produced phytotoxin in-vitro when grown in stationary culture for 7 d at 28 °C on minimum salts medium supplemented with 1.5 % glucose as the sole carbon source. Phytotoxin was also produced when the fungal strain was grown on fructose, galactose, mannose, xylose and sucrose. The addition of nitrogen source (yeast extract, peptone or malt extract) to the culture medium did not influence phytotoxin production. The phytotoxin produced by Strain I was most active at pH 6.0, stable between pH 3-9, and was highly thermostable, remaining fully active when heated at 90 °C for 1 h.

**Key words:** *Bipolaris euphorbiae, Euphorbia heterophylla*, phytotoxin production, phytotoxin stability

**INTRODUCTION**

The milk weed, or wild poinsettia, *Euphorbia heterophylla* is widely distributed throughout Brazil and infests up to 200,000 ha of cultivated soyabean (*Glycine max* L) land in the southern soyabean-growing states (Yorinori and Gazziero, 1989). The weed is difficult to control and herbicides that are effective against *E. heterophylla* can be expensive, as more than one application is frequently required during a crop season. The need for better control measures has led to a search for alternative methods to limit this weed’s distribution. Biocontrol agents have potential and often synthesise phytotoxins that produce one or more specific disease symptoms in host plants when infected. Phytotoxins can elicit symptoms such as chlorosis, growth abnormalities, necrosis and wilting. The exact mechanisms by which this occurs is still not fully understood, but is thought to involve complex biochemical changes within the host plant (Harborne, 1993).

The phytopathogenic fungus, *Bipolaris euphorbiae* previously identified as a *Helminthosporium* species), is the causal agent for the major disease of *E. heterophylla* within Brazil (Barreto and Evans, 1998), and has been reported to be highly efficient and promising as a biological...
control agent for this weed; as efficient as the best post-emergence herbicides (Yorinori, 1985; Yorinori and Gazziero, 1989). This microorganism produces host-specific phytotoxin(s) that elicits its effect during germination, and affects the leaves of susceptible *E. heterophylla* plants causing defoliation, but does not affect soybeans. Many fungal species pathogenic to agricultural crops of economic importance are known to produce phytotoxins. For example, those produced by the genus *Helminthosporium* have been shown to belong to several classes of natural products, e.g., macrocyclic peptides, polyketides, quinones, anthraquinones, and the free and glycosidic form of terpenes, phenols and flavonoids (Santos, 1997). An isolated toxin from *Helminthosporium sacchari* when applied to host-specific sugarcane plants, caused disease symptoms similar to those produced by the fungus (Steiner and Byther, 1971). The phytotoxins produced by *Bipolaris* species generally belong to a group of sesterpenoids called ophiobolins (tricyclic 5-8-5 ring systems) and have recently been reviewed by Au et al., (2000). Another related species, *B. cynodontis*, however, produced phytotoxins that belonged to a different group called cochlitoquinones (Lim et al., 1998). The phytotoxin(s) from *B. euphorbiae* have not yet been characterised, but are present in culture filtrates of the fungus (Ferreira et al., 1987).

The program on the biological control of *E. heterophylla* by *B. euphorbiae* in soybean cultivation commenced at EMBRAPA, Londrina, Paraná, Brazil, in 1980 by Dr. J.T. Yorinori of National Centre of Soybean Research. The work reported here is an extension of those studies and concerns the examination of phytotoxin production in-vitro by four virulent strains of *B. euphorbiae* isolated from *E. heterophylla* plants found growing within four states of Brazil. We furthermore report on the production of phytotoxin(s) in one of those isolates (*B. euphorbiae* Strain I: EUPH petropar) when cultured on simple nutrient medium, and comment on the effect of some cultivation parameters and properties of the phytotoxin itself.

**MATERIALS AND METHODS**

Virulent strains of the fungus, *Bipolaris euphorbiae*, were isolated from *Euphorbia heterophylla* leaves collected in four states of Brazil: Mato Grosso (Strain I: EUPH petropar), Mato Grosso do Sul (Strain II: EUPH fc bom), Paraná (Strain III: a sporulating mutant), and Rio Grande do Sul (Strain IV: HEM C95 A). All of the isolates of the above strains were kindly provided by Dr. J.T. Yorinori of EMBRAPA, Londrina, Paraná, Brazil. The fungi were maintained at 4 °C on potato-dextrose agar (PDA) slants, and subcultured at intervals of 1 - 4 months.

Inoculum was prepared by growing the fungal strains on PDA at 28 ± 0.2 °C for 7 days, and using plugs of 13 mm diameter colonized with mycelium. Three plugs were used as inoculum per flask containing 50 ml of culture medium. The *B. euphorbiae* strains were grown in 250 ml Erlenmeyer flasks containing 50 ml basal medium (Vogel minimum salts medium (Vogel, 1956) with glucose as carbon source). Glucose and the salts solutions were separately sterilized (121 °C, 20 min), and combined to bring the final glucose concentration in the basal medium to either 5, 15, 20 or 30 g 1⁻¹. After inoculation, the flasks were incubated at 28 ± 2 °C either in a rotary shaker (180 rev min⁻¹), or in a stationary incubator, for 7 days. *B. euphorbiae* Strain I was also grown on a large scale in 2 l Erlenmeyer flasks containing 400 ml basal medium (glucose 15 g 1⁻¹) under stationary conditions at 28 ± 2 °C for 7 days. All experiments were carried out in triplicate, except for those determining the growth profile of the fungus (followed over a period of 13 days), which was performed in replicates of four. The reported data represent average values ± SD. Flasks without inoculum were maintained in parallel with the growing cultures as controls. Experiments evaluating the influence of the air-to-medium ratio were conducted in flasks of 250 ml capacity containing 25 ml, 50 ml and 100 ml of basal medium (15 g 1⁻¹ glucose). In experiments examining the effects of exogenous-added nitrogen, either yeast extract (Difco), malt extract (Difco) or bacteriological peptone (BioBras, Brazil), were added to the basal medium (15 g 1⁻¹ glucose) at a concentration of 2 g 1⁻¹. Potato-dextrose medium (no added salts) was prepared from whole potato (200 g) boiled in water (1 l) for 1h, followed by mashing, and the contents strained through cotton gauze. Glucose was added to the filtrate at a final concentration of 15 g 1⁻¹.
Cell-free fluid (ECF) was recovered by centrifugation (480 x g for 10 min) and the supernatant filtered and stored at 4 °C. Total sugar was measured by the phenol-sulfuric acid method (Dubois et al., 1956). Reducing sugars were determined by the methods of Nelson (1944) and Somogyi (1945). Glucose was used as standard in both sugar determinations. Protein was determined by a modified Lowry method using bovine serum albumen as a standard (Hartree, 1972). Fungal biomass was determined gravimetrically after washing the recovered mycelium with isotonic saline, centrifuging and drying the mycelium to constant weight at 100 °C. Mycelial weights were corrected for the agar plugs used as inoculum. All assays were conducted in triplicate and represent mean values ± SD.

Soybean plants, and E. heterophylla plants susceptible to virulent strains of B. euphorbiae, were cultivated in pots from germinating seeds in a greenhouse under lights with a 12 h on-off cycle. Biological assays for phytotoxin activity were conducted on the leaves of plants that were at least 25 days old. Bioassays were performed by applying ECF topically with the aid of a cotton swab to the leaves of susceptible E. heterophylla and soybean plants in the late afternoon. The plants were then covered with black plastic bags to facilitate the absorption of phytotoxin, and left overnight. Next morning the bags were removed, and observations recorded at 12, 24 and 96 h following application of the ECF. Appropriate dilutions of the ECF were made using distilled water to determine the potency of phytotoxin concentration on biological activity. In some experiments, freeze-dried ECF was used (ECF from B. euphorbiae Strain I was dialysed (cellophane membrane) against distilled water at room temperature for 24 h, the water changed every 2 h, and then lyophilised). A 2 % w/v solution of the lyophilised powder was used in the bioassays.

Biological activity was scored numerically in arbitrary units: (-) negative, (+) positive; (+1), when disease spots appeared on the leaves after 12 h and wilted after 24 h; (+2), when the leaves wilted after 12 h (highly positive) and thereafter resulted in defoliation.

Thermostability: A 2 % (w/v) solution of the lyophilized ECF in water was heated for 15, 30 and 60 min at 50 °C, and the solution cooled to room temperature prior to the bioassay as described above. In another experiment, the solutions containing phytotoxin were heated at 70, 80, 90 and 100 °C for 60 min, cooled and assayed for biological activity. Evidence of biological activity was scored numerically in arbitrary units: (0), no observable changes on the leaves; (1), spots dispersed on the leaves; (2), dark spots on all the leaves; (3), partial wilting of the leaves with the presence of dark spots; and (4), leaves completely wilted and defoliated.

pH dependence on phytotoxin activity: A 4 % (w/v) aqueous solution of the lyophilised ECF was diluted 1:1 in buffer (McIlvaine’s citrate-phosphate buffers of pH 3-7, and glycine-NaOH buffers of pH 8-12) to examine the effect of pH on phytotoxin activity. Freshly made buffered solutions containing the phytotoxin were assayed for biological activity, and scored numerically as outlined above. Appropriate solutions consisting of only buffer at each pH were diluted 1:1 with water and applied topically to E. heterophylla leaves as controls, and were assessed and scored in the same way as the ECF. Positive results from controls were deducted from the tests screening phytotoxin activity to correct for buffer interference. To determine the potency of the phytotoxin, the 2 % buffered ECF solution, at each pH where there was a biological response, was diluted using deionized water to 1.0, 0.5, and 0.25 % (w/v) final ECF concentration, the pH adjusted where necessary, and further evaluated for biological activity.

RESULTS AND DISCUSSION

The four virulent B. euphorbiae strains used in this work grew on basal medium, but did not produce phytotoxin activity (Table 1) when cultivated in-vitro under submerged conditions as judged by the negative results for the bioassays on the leaves of susceptible E. heterophylla plants. Of the four fungal strains tested, only one, Strain I (EUPH petropar), produced phytotoxin activity indicating pathogenicity when cultivated under stationary conditions on basal medium (Table 1).

In the field, all four strains of B. euphorbiae were host-specific disease-producing pathogens. The reason why the ECF of the other three fungal strains failed to produce biological responses
could be explained twofold; the phytotoxin could be produced at low levels on the growth medium used that did not permit their detection (Pringle and Scheffer, 1967), and secondly, these strains could require specific components to induce the synthesis of phytotoxin, such as exist in the complex biochemical make-up of the host plant that determined these fungal strains to be host-specific. For example, plants containing activators such as 2-amino-1,3-propanediol (serinol) produced in sugarcane leaves have been reported to induce phytotoxin production in attenuated strains of Helminthosporium sacchari (Pinkerton and Strobel, 1976). Additionally, we have observed that after successive transfers through long-term subculturing on PDA, B. euphorbiae Strain I lost the ability to produce phytotoxin in vitro.

**Table 1.** The influence of submerged and stationary culture conditions on phytotoxin production by four strains of Bipolaris euphorbiae grown on basal medium containing 20 g l\(^{-1}\) glucose.

<table>
<thead>
<tr>
<th>Strains</th>
<th>Biomass (g l(^{-1}) ± SD)</th>
<th>Residual sugars** (g l(^{-1}) ± SD)</th>
<th>Biological assay(^{#})</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>SM*</td>
<td>ST*</td>
<td>SM</td>
</tr>
<tr>
<td>I</td>
<td>11.3 ± 0.84</td>
<td>10.7 ± 1.06</td>
<td>0.0 ± 0.84</td>
</tr>
<tr>
<td></td>
<td>± 0.16</td>
<td>± 0.11</td>
<td>(-)</td>
</tr>
<tr>
<td>II</td>
<td>8.0 ± 0.97</td>
<td>7.2 ± 1.91</td>
<td>1.3 ± 0.07</td>
</tr>
<tr>
<td></td>
<td>± 1.91</td>
<td>± 0.16</td>
<td>(-)</td>
</tr>
<tr>
<td>III</td>
<td>9.0 ± 1.01</td>
<td>6.5 ± 1.64</td>
<td>0.0 ± 0.14</td>
</tr>
<tr>
<td></td>
<td>± 1.64</td>
<td>± 0.14</td>
<td>(-)</td>
</tr>
<tr>
<td>IV</td>
<td>6.3 ± 0.98</td>
<td>3.4 ± 1.85</td>
<td>4.2 ± 0.16</td>
</tr>
<tr>
<td></td>
<td>± 1.85</td>
<td>± 0.16</td>
<td>(-)</td>
</tr>
</tbody>
</table>

* SM submerged, * ST stationary cultivation. ** as reducing sugars, # undiluted ECF

Phytotoxin activity, however, was restored when the strain was again grown on the leaves of susceptible E. heterophylla plants. This phenomenon has also been observed in H. sacchari (Pinkerton and Strobel, 1976). However, we didn’t believe that reversion was the prime reason why the other B. euphorbiae strains failed to produce phytotoxin, as these isolates were demonstrated to be virulent when examined on susceptible E. heterophylla plants. Physiological factors were more likely to be responsible for lack of production of phytotoxin. The ECF from Strain I when applied to the leaves of soybean plants failed to cause any signs of disease (spotting or wilting) over a four-day period, demonstrating that the fungal phytotoxin did not affect soybean plants. It has been reported that fungal phytotoxins can also interact with other plants besides the host-specific plant (Hudson, 1986), however, this was not the case with B. euphorbiae. The phytotoxin of B. euphorbiae, furthermore, did not affect germination of soybean seeds, nor did it show any disease symptoms with plant development during growth. These results demonstrated that the phytotoxin produced by B. euphorbiae is specific for E. heterophylla, and indicates its suitability as a biocontrol agent of this weed when growing among soyabean crops. Field-grown soybean plants have similarly been shown not to be affected when treated with spores and live cultures of B. euphorbiae (Yorinori and Gazziero, 1989).

The glucose concentration in the medium was found to influence the production of phytotoxin by B. euphorbiae Strain I (Table 2). This is probably due to the production of more biomass by the growing fungal cells at the higher glucose levels, hence the cells can produce more phytotoxin. When these filtrates were diluted and assayed biologically, strong wilting of the leaves could be detected at dilutions of 1:20 for filtrates from the fungus grown on 15 and 30 g l\(^{-1}\) glucose, but some spots on the leaves were still noticeable at 1:80 dilution. However, filtrates from cultures grown on basal medium containing 5 g l\(^{-1}\) glucose registered a positive response only when undiluted.

The growth profile for B. euphorbiae Strain I on basal medium (15 g l\(^{-1}\) glucose) over 13 days is shown in Figure 1. Maximum biomass was
reached by day 6-7 which coincided with a drop in the pH to 4.2. The maximum specific growth rate ($\mu_{\text{max}}$) over the first 5 days was 0.423 day$^{-1}$. The effect of inoculum concentration when doubled did not produce a lag phase nor more biomass. Nor did it result in higher phytotoxin activity being produced. However, when the inoculum was halved, a lag phase occurred, and proportionately less biomass and phytotoxin were produced. The bioassay results presented in (Figure 1) showed that phytotoxin production reached its maximum at 6-7 days growth coinciding with maximal mycelium production.

![Graph showing residual sugars, pH, and biomass over time](image)

**Figure 1** - Production of phytotoxin by Bipolaris euphorbiae Strain I (EUPH petropar) grown in stationary culture on basal medium containing 15 g l$^{-1}$ glucose.

Thereafter both phytotoxin activity and mycelium concentration decreased, probably as a consequence of endogenous metabolism, as the sugar concentration in the medium at this stage was low (0.8 g l$^{-1}$). Furthermore, cell lysis was not observed during this period. Maximal production of phytotoxin towards the end of the growth phase has similarly been reported for *H. victoriae* (Luke and Wheeler, 1955), *H. maydis* (Smedegard and Nelson, 1969), and *H. nodulosum* (Vidhyasekaran, 1977). The air-to-nutrient medium ratio also influenced the production of phytotoxin by *B. euphorbiae* when grown under stationary conditions, and was maximal at 10:1, demonstrating that as the volume of air in the flasks decreased, this was accompanied by a corresponding decrease in glucose consumption, biomass and phytotoxin production. In this case, phytotoxin production appears to be related to the amount of mycelium produced, and this observation was further corroborated by the results correlating biomass levels with phytotoxin activity in Figure 1 and Table 2.

Glucose is commonly used as a carbon source by most fungi, including those producing phytotoxins. Fructose, mannose, galactose, xylose and sucrose can also serve as carbon sources, but require an adaptation phase as they must be phosphorylated prior to interconversion during glycolysis (Gadd, 1988). Xylose is metabolised via the pentose phosphate pathway, while sucrose must first be hydrolyzed to its constituent sugars. *B. euphorbiae* grew equally well on each of the different sugar substrates studied, and produced phytotoxin (Table 3) in each case. Sugar consumption was similar (98 %) for all of the substrates, with the exception of galactose which was lower (81 %). Vidhyasekaran (1977) has also reported the production of phytotoxin in *H. nodulosum* when cultivated on similar sugars and starch. Fructose and sucrose, along with glucose, are common plant sugars produced by photosynthesis (Raven, Evert and Eichdorn, 1992), and can support growth of microorganisms
including *B. euphorbiae*. The influence of various nitrogen sources added to the basal medium had no effect on the production of phytotoxin by *B. euphorbiae* Strain 1. When the ECF was diluted, the yeast extract-grown cultures lost phytotoxin activity, but the ECF from the other grown cultures, nonetheless, maintained their biological potency, even at dilutions of 1:20. The C:N ratio in the nitrogen-supplemented cultures was low (1.9 - 3.3) compared to the basal medium (8.6). A low C:N ratio has similarly been found to support phytotoxin production in *H. nodulosum* (Vidhyasekaran, 1977). Potato-dextrose medium could also serve as a suitable medium for the production of phytotoxin by *B. euphorbiae* Strain 1.

There did not appear to be any correlation between phytotoxin concentration and phytotoxin activity when a solution of lyophilised ECF was applied topically to the leaves of *E. heterophylla*. The reason for this may be governed by how well the applied ECF was absorbed and diffused into the surrounding tissues in the leaves to manifest a pathogenic response. A 2 % solution of the freeze-dried ECF was therefore selected in studies examining the influence of temperature and pH on biological activity.

**Table 2** - The effect of glucose concentration on phytotoxin activity and biomass production by Bipolaris euphorbiae Strain I after seven days under stationary culture conditions

| Glucose (g l\(^{-1}\)) | Biomass (g l\(^{-1}\) ± SD) | Residual sugars** (g l\(^{-1}\) ± SD) | Final pH* | Biological assay
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>5</td>
<td>5.0 ± 1.56</td>
<td>0</td>
<td>6.6</td>
<td>+1*</td>
</tr>
<tr>
<td>15</td>
<td>9.6 ± 1.28</td>
<td>1.7 ± 0.05</td>
<td>4.8</td>
<td>+2*</td>
</tr>
<tr>
<td>30</td>
<td>12.6 ± 0.72</td>
<td>10.1 ± 0.11</td>
<td>4.6</td>
<td>+2*</td>
</tr>
</tbody>
</table>

*initial pH 6.3, * undiluted ECF, **as reducing sugars, * ECF diluted 1:20

**Table 3** - The influence of carbon source on phytotoxin activity and biomass production by Bipolaris euphorbiae Strain I after seven days growth under stationary conditions.

<table>
<thead>
<tr>
<th>Carbon source**</th>
<th>Biomass (g l(^{-1}) ± SD)</th>
<th>Sugar Consumption (%)</th>
<th>Final pH*</th>
<th>Biological assay*</th>
<th>**</th>
</tr>
</thead>
</table>
| Glucose         | 10.0 ± 0.84                 | 98                    | 6.5       | +2                | 15 g l\(^{-1}\) concentration, * as reducing sugars, * initial pH 5.8, ** undiluted ECF
| Xylose          | 8.4 ± 1.78                  | 98                    | 6.3       | +2                |
| Galactose       | 9.7 ± 1.56                  | 81                    | 6.1       | +2                |
| Mannose         | 9.8 ± 1.21                  | 99                    | 5.9       | +2                |
| Fructose        | 9.0 ± 1.46                  | 97                    | 6.8       | +2                |
| Sucrose         | 9.0 ± 1.65                  | 98                    | 6.9       | +2                |

The phytotoxin was stable when heated at 50 °C for up to 60 min losing no biological activity during this period. When the ECF was heated at temperatures up to 90 °C for 60 min, there was no apparent loss of biological activity under the assay conditions. Heating at 100 °C for 1 h, however, did result in a loss of about 25% of the initial activity of the phytotoxin, indicating that the phytotoxin produced by *B. euphorbiae* Strain I was highly thermostable. The toxin produced by *Helminthosporium nodulosum* has also been shown to be stable up to 90 °C, but lost total activity at 100 °C (Vidhyasekaran, 1977). The effect of pH on phytotoxin activity was studied using 2 buffer systems over a pH range of 3-12, and at different concentrations of ECF. It appeared that pH 6.0 was most effective in eliciting biological activity of the phytotoxin for all
Phytotoxin Produced by *Bipolaris euphorbiae*

Brazilian Archives of Biology and Technology 239

concentrations studied. This was within the range of the final pH of the culture medium following growth. The phytotoxin of *B. euphorbiae* was stable over a broad pH range (3-9). Comstock and Scheffer (1972) found that the phytotoxin produced by *H. maydis* was relatively stable at pH 3.5 when autoclaved (121 °C for 15 min) and progressively lost activity when autoclaved at higher pH values. Similarly, the toxin from *H. nodulosum* was stable at pH 3-7, but unstable within the alkaline range (Vidyasekaran, 1977), whereas the toxin from *H. sativum* was more stable at the alkaline range (Das and Srivastava, 1969).

Although the chemical structures of the phytotoxins produced by several species of *Bipolaris* are known (Au et al., 2000), that produced by *B. euphorbiae* Strain I has not yet been determined. Our studies, however, conclude that it’s phytotoxin is unlikely to be protein by nature, as the extreme temperatures it was exposed to would lead to it’s denaturation and hence, loss of biological activity.

ACKNOWLEDGMENTS

This work was supported by FINEP (Brazil) and PPG-UEL. C. Souza and R. Fonseca gratefully acknowledge CAPES and CNPq-PIBIC (Brazil) for providing scholarships at UEL. The authors are grateful to Dr. J.T. Yorinori of EMPRABA, Londrina, PR, Brazil, for providing the strain isolates of *Bipolaris euphorbiae*, and *Euphorbia heterophylla* seeds and plants. Erico Damineli is thanked for experimental assistance.

RESUMO

Quatro linhagens virulentas do fungo *Bipolaris euphorbiae* (previamente identificado como *Helminthosporium* sp.), isoladas de *Euphorbia heterophylla* de quatro estados do Brasil, foram estudadas quanto a produção de fitotoxina, que promove murcha e desfolha da referida erva daninha brasileira, popularmente denominada leiteiro e comumente encontrada nas culturas de soja. O isolado *Bipolaris euphorbiae* linhagem I (EUPH petropar do estado de Mato Grosso), produziu fitotoxina in-vitro quando cultivada sob condição estática, durante 7 dias, a 28 ºC em meio mínimo suplementado com glucose 1,5 %, como fonte única de carbono. A fitotoxina foi também produzida quando o isolado foi cultivado em frutose, galactose, manose, xilose e sacarose. A adição de fonte de nitrogênio (extrato de levureda, peptona e extrato de malte) ao meio de cultivo não influenciou a produção da mesma. A fitotoxina produzida pela linhagem I apresentou maior atividade em pH 6,0 , permanecendo estável na faixa de pH entre 3 e 9 e demonstrou elevada termoestabilidade, mantendo-se ativa após ter sido aquecida durante uma hora a 90 ºC.

REFERENCES


Hudson, H. J. (1986), Fungi as parasitic symbionts of...
Steiner, G. W. and Byther, R. S. (1971), Partial0 characterization and use of a host-specific toxin from Helminthosporium sacchari on sugarcane. Phytopathology, 61, 691-695.


Received: October 25, 2000; Revised: January 29, 2001; Accepted: June 25, 2001.