CHARACTERIZATION OF FUNGAL ISOLATES FROM PYCNIDIA AND PSEUDOTHECIA FROM LESIONS OF PHAEOSPHAERIA LEAF SPOT IN MAIZE

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ABSTRACT - Phaeosphaeria leaf spot, a maize disease of occurrence in Brazil, can reduce the grains production of susceptible cultivars up to 60%. The causal agent of the disease was described on the basis of visual descriptions of sexual and asexual reproductive structures (pseudothecia and pycnidia) found in the lesions, and identified as being the fungus *Phaeosphaeria maydis*, imperfect form *Phyllosticta* sp. However, the isolation of this fungus is difficult and its inoculation in maize plants under controlled conditions, has not been successful in the reproduction of the symptoms. In this work, isolates obtained from pycnidia and pseudothecia present in foliar lesions were compared using isoenzymatic (α and β esterases) and RAPD patterns, in order to determine whether these two states belong to the same etiological agent. These techniques clearly distinguished between isolates from pycnidia and pseudothecia, indicating that these structures belong to distinct fungal species, and not to the anamorphic and teleomorphic states of the same agent, as previously suggested. Based on morphological characteristics, the isolate from pycnidia was classified as *Phoma tropica*, an opportunistic fungus that colonizes pre-established lesions, and not as a *Phyllosticta* sp. as described initially.

Key-words: *Zea mays*, *Phaeosphaeria maydis*, *Phyllosticta*, isozymes, RAPD analysis

CARACTERIZAÇÃO DE ISOLADOS FÚNGICOS DE PICNÍDIOS E PSEUDOTÉCIOS DE LESÕES DA MANCHA FOLIAR DE PHAEOSPHAERIA EM MILHO

RESUMO - A doença foliar descrita no Brasil como sendo mancha por phaeosphaeria em milho pode reduzir a produção de grãos em até 60%, quando se utilizam cultivares suscetíveis. O agente causal da doença foi descrito a partir de observações visuais de estruturas reprodutivas sexuadas e assexuadas (pseudotécios e pícnidios) encontradas no interior das lesões, e identificado como sendo o fungo *Phaeosphaeria maydis*, forma imperfeita *Phyllosticta* sp. No entanto, esse fungo é de difícil isolamento e sua inoculação em plantas de milho, sob condições controladas, não tem apresentado sucesso na reprodução dos sintomas. Isolados obtidos a partir de pícnidios e pseudotécios presentes nas lesões foram comparados através do padrão isoenzimático (α e β esterases) e RAPD, visando determinar se esses dois estados são realmente pertencentes ao mesmo agente etiológico. Os resultados obtidos permitiram a separação dos isolados oriundos de pícnidios daqueles oriundos de pseudotécios em grupos bem distintos, indicando que essas estruturas pertencem a espécies fúngicas distintas, e não ao estado anamórfico e teleomórfico do mesmo agente, como vem sendo atribuído. A partir da
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classificação baseada em caracteres morfológicos, o fungo isolado de picnídios foi identifica-
do como Phoma tropica, um fungo oportunista colonizador de lesões preestabelecidas, e não
Phyllosticta sp como descrito inicialmente.

Palavras-chave: Zea mays, Phaeosphaeria maydis, Phyllosticta, isoenzimas, RAPD

The foliar disease of maize commonly known
as phaeosphaeria leaf spot was first described in India
by Rane et al. (1965) and, by the mid-1980s, was
reported in Brazil (Fantim, 1994). Balmer & Pereira
(1987) and Casela (1998) noted that, in recent years,
there has been an increase in the significance of this
disease in Brazil, where plants are generally affected
at the end of the crop growing season, although
younger plants may also be affected causing
premature dryness in more severe attacks. The
symptoms begin with the appearance of slightly
chlorotic, water-soaked lesions on the leaves, which
later become necrotic and straw-colored. The
incidence of infection increases in conditions of high
relative humidity and moderate temperatures (Fantim,
1994), and may reduce grain yield up to 60%
(Fernandes & Oliveira, 1997).

Initially, Phyllosticta sp was described as
the etiological agent of phaeosphaeria leaf spot, based
on observations of pycnidia inserted on plant tissue
in the center of the lesions. Depending on climatic
conditions and season, sexual reproductive
structures, described as perithecium, may also be
seen (Rane et al., 1965). More recent reports have
identified these sexual structures as pseudothecium
(Fantim, 1994). Fungi producing these pseudothecia
have been identified as Phaeosphaeria maydis, the
perfect form of Phyllosticta sp, an ascomycete of
the family Pleosporaceae. The anamorphic and
teleomorphic states of the causal agent were
described on the basis of visual descriptions of the
reproductive structures (pycnidium and pseudothecium) present in the lesions.

In this work we have used biochemical and
molecular techniques to compare the isolates from
pycnidia with those from pseudothecia, in order to
establish whether these two structures are really
different states of the same agent.

Material and Methods

Fungal isolates

Fungal isolates used in this study are
described in Table 1. All cultures were prepared from
monoconidial isolates. Isolate 6, obtained from
pycnidia was provided by the Plant Pathology
Laboratory of EMBRAPA Maize and Sorghum in
Sete Lagoas, Minas Gerais, Brazil, and isolate 89,
obtained from pseudothecium, was kindly provided
by Dra G. Fantim of the Biological Institute of Cam-
pinas, São Paulo, Brazil. All other isolates were
obtained from foliar lesions containing reproductive
structures. The isolates were cultivated in oat media
(60 g of oat flakes + 15 g of agar + 1000 mL of
distilled water) for 20 days at 22 ± 2 °C (12 h light /
12 h dark).

Polyacrylamide gel electrophoresis (Paccola-
Meirelles et al., 1988)

The esterase isoenzyme patterns of 30
isolates (15 from pycnidia and 15 from pseudothecia)
were evaluated. The isolates were cultivated in potato
dextrose broth, pH 6.9, at 22 ± 2 °C (12 h light/12 h
dark) for seven days. The resulting mycelia were
washed three times in distilled water, filtered,
macerated in liquid N2, and homogenized in 0.5 M
Tris-HCl, pH 6.8 (1 g of mycelia: 1 mL of buffer).
After centrifugation at 40,000xg for 20 min, the
samples were separated by electrophoresis
(Paccola-Meirelles et al., 1988) on 10%
polyacrylamide gel. The pattern of α and β esterases
was determined using the substrate 1% α- naphthyl acetate and 1% β- naphthyl acetate.

**DNA extraction**

For total DNA extraction, according to Maki et al. (2001), 400 mg of mycelia of each isolate were macerated in liquid N₂, and extracted in 4 mL of extraction buffer (200 mM Tris-HCl, pH 8.0 + 250 mM NaCl + 25 mM EDTA, pH 8.0 + 1% SDS + 1 µl β- mercaptoethanol per mL of buffer) and incubated at 65°C for 15 min. Two milliliters of phenol were then added to the samples and centrifuged for 15 min at 10.000xg. The supernatant was collected and transferred to centrifuge tubes, followed by addition of phenol and chloroform (1mL each) to the tubes. After centrifugation for 15 min at 10.000xg, 2 mL of chloroform-octanol (24:1v/v) was added to the supernatant and again centrifuged for 15 min. Ice-cold isopropanol (8 mL) was added to the supernatants. After precipitation, the DNA was transferred to microtubes containing 100 µl of TE (0.5 mL of 1M Tris + 0.1 mL of 0.5 M EDTA, pH 8.0). The DNA concentration was estimated by comparison with DNA standards on 0.8% agarose gel electrophoresis.

**Primers and RAPD (Random Amplified Polymorphic DNA) analysis**

Nineteen previously selected primers were used in the analysis: OPA3 (5’ AGT CAG CCA C 3’), OPA4 (5’ AAT CGG GCT G 3’), OPA8 (5’ GTG ACG TAG G 3’), OPA9 (5’ GGG TAA AGC C 3’), OPA10 (5’ GTG ATC GCA G 3’), OPA13 (5’ CAG CAC CCA C 3’), OPA14 (5’ TCT GTG CTG G 3’), OPW3 (5’ GTC CGG AGT G 3’), OPW4 (5’ CAG AAG CGG G 3’), OPW5 (5’ GGC GGA TAA G 3’), OPW6 (5’ AGG CCC GAT G 3’), OPW8 (5’ GAC TGC TCT C 3’), OPW9 (5’ GTG ACC GAG T 3’), OPW10 (5’ TCG CAT CCC T 3’), OPW11 (5’ CTG ATG CGT G 3’), OPW12 (5’ TGG GCA GAA G 3’), OPW13 (5’ CAC AGC GAC A 3’), OPW16 (5’ CAG CCT ACC A 3’), OPW20 (5’ TGT GGC AGC A 3’). Each amplification reaction contained 9.5 µl of sterile, ultra-pure water, 2.5 µl of PCR buffer 10X (50 mM KCl + 100 mM Tris-HCl, pH 8.6 + 0.1% Gelatin + 2 mM MgCl₂), 1µl of dNTP solution (containing 100 µM each of dATP, dTTP, dGTP and dTTP), 1 µl of primer (4 µM), 1 µl of DNA-Taq polymerase 1 U/mL and 10µL of DNA solution (5 ng/µl). Polymerase chain reaction (PCR) was done using a 100 PTC (programmable thermal controller) thermocycler (Perkin Elmer – Gene Amp ). The program used consisted of 1 min at 95°C for initial denaturation followed by 35 cycles of amplification (10 seconds at 94°C for denaturation, 1 min at 36°C for annealing and 2 min at 72°C for extension by Taq-polymerase and incorporation of the nucleotides) and a final extension for 7 min at 72°C after which the tubes were maintained at 4°C (Maki et al. 2001). The amplification products were separated on 1% agarose gels, stained with ethidium bromide and photographed under UV light with Polaroid film. The data obtained by RAPD were analyzed by the presence (1) and absence (0) of a specific band on the gel for each one of the 19 primers tested. These analyses were carried out considering a total number of 163 polymorphic bands and the program used was Statistica 5.0 version (StatSoft, Inc.)

**Pathogenicity tests**

Pathogenicity tests were performed on 30-day-old plants, of the susceptible maize cultivar HS200. Conidia and ascospores of isolates 6 and 89, were produced in oat culture medium at 22 ºC, and a 10⁶ mL⁻¹ spore suspension was sprayed onto the leaves. Control plants were sprayed with sterilized water. Three pots with two plants per pot were used in each treatment. After inoculation plants were kept for 60 h in a dew chamber with 100% relative humidity.
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**Results and Discussion**

The frequency of pycnidia and pseudothecia in phaeosphaeria leaf spot lesions of 30 maize hybrids were evaluated. Pycnidia were seen in only 8.8% of the lesions and pseudothecia in 7.3%; about 79.3% of the lesions showed no reproductive structures and 2% had both structures. The isolation frequency of the fungus described as the patogenic agent was low, about 4.4%. Several fungal species other than the ones described as being the causal agent of phaeosphaeria leaf spot were also isolated. Isolates obtained from picnidia and pseudothecia are listed in Table 1.

**TABLE 1.** Isolates obtained from lesions containing reproductive structures and their respective origins.

<table>
<thead>
<tr>
<th>Isolates</th>
<th>Locality</th>
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<tbody>
<tr>
<td>1</td>
<td>Chapadão do Céu (GO)</td>
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<td>5</td>
<td>Goianésia (GO)</td>
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<td>6</td>
<td>Sete Lagoas (MG)</td>
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<td>9</td>
<td>Sete Lagoas (MG)</td>
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<td>28</td>
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<td>29</td>
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<td>30</td>
<td>Passo Fundo (MG)</td>
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<td>41</td>
<td>Araçatuba (SP)</td>
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<td>Batatais (SP)</td>
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<td>43</td>
<td>Jardimópolis (SP)</td>
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<td>89</td>
<td>Anhembi (SP)</td>
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<td>90</td>
<td>Ponta Grossa (PR)</td>
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<td>93</td>
<td>Cascavel (PR)</td>
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*1–51: isolates from pycnidia; 52–93: isolates from pseudothecia.

The electrophoretic profiles of $\alpha$ and $\beta$ esterases allowed the separation of isolates obtained from pseudothecia and pycnidia, suggesting that these isolates belong to different species. Esterases profiles of isolates obtained from pseudothecia were very similar, with only one polymorphic band seen in isolate 68 (Figure 1). The electrophoretic pattern of this isolate indicates that it belongs to a different species. By contrast, there was an extensive variation among the esterase patterns of isolates obtained from pycnidia (Figure 2).

In the RAPD analysis, of the 19 primers evaluated, 11 (OPA3, OPA4, OPA8, OPA9, OPA10, OPA13, OPW4, OPW5, OPW6, OPW12...
the first, contained the isolates of *Phyllosticta* sp (with the exception of isolate 68); the second contained isolates of *Phaeosphaeria maydis* and, the third included only isolate 90. The RAPD analysis confirmed the results obtained by the isoenzymatic analysis allowing the separation of the isolates of *Phyllosticta* sp and *Phaeosphaeria maydis* into distinct groups. These results suggested that pycnidia and pseudeotheca present in *phaeosphaeria* leaf spot lesions belong to two different species rather than being the anamorphic and teleomorphic states of the same fungus.

The genetic variation among isolates from the same geographic origin such as isolates 6, 9, 16, 17 and 23 collected in Sete Lagos, (MG) was sometimes greater than that among isolates from different regions. This observation may reflect the action of mutations and, possibly, genetic exchange between strains, since anastomosis of hyphae has been reported in this species (Cervelatti *et al.*, 1998). Similar observations were made by Machado *et al.* (1997), in isolates of *Cercospora sojina* collected in Minas Gerais.

Isolate 6 (from pycnidia) and isolate 89 (from pseudeotheca) were classified according to their morphological characteristics. Isolate 6 originally described as *Phyllosticta* sp was classified as *Phoma tropica* and isolate 89 described as *Phaeosphaeria maydis* was classified as belonging to the *Phaeosphaeria* genus.

The species *Phoma tropica* has been described in association with leaf spots and stem lesions in more than 20 species of ornamental plants in green houses in Holland and Germany (Schneider & Boerema, 1975).

Errors in the classification of *Phoma* and *Phyllosticta* species are relatively common. As an example, *Phoma glomerata* classified by Simay (1994) was originally deposited in culture banks as *Phyllosticta maculiformis*, *Phyllosticta sojoecola* and *Phoma pomorum*. Many fungi are erroneously
classified as *Phyllosticta* because of this species' ability to establish itself in lesions caused by other agents and, in many cases, the classification of *Phoma* and *Phyllosticta* is based on the type of host, and not on characteristics observed during fungus cultivation.

In the pathogenicity tests, conidia and ascospores from isolates 6 and 89, did not reproduce symptoms of phaeosphaeria leaf spot under controlled conditions, and Paccola-Meirelles *et al.* (2001) has suggested that these fungi are not the primary agents of the infection. Although phaeosphaeria leaf spot has been described as a fungal disease, these authors suggest the possibility of an association with bacteria, the *Pantoea ananas*. According to their hypothesis, the bacteria initiates infection, and the lesion, later on, is invaded by a co-inhabiting fungus. This explains the high incidence of lesions without fungal structures and the low frequency of *Phaeosphaeria* sp. isolated from reproductive structures.

Our results indicate that the pycnidia observed in the center of the lesions of the disease described as phaeosphaeria leaf spot in maize do not belong to the fungus *Phyllosticta* sp as it has been attributed, but rather belong to the saprophytic species, *Phoma tropica*, that develops in lesions produced by other pathogenic agents.
FIGURE 4. Amplification products generated from 17 isolates obtained from pycnidia (A) and 17 isolates obtained from pseudothecia (B) from phaeosphaeria leaf spot lesions in maize, using primer OPA13. C: negative control (sample containing only PCR buffer, primer dNTPs, Taq polymerase and ultra pure water). M: Molecular weight markers; 100 pb DNA Ladder (GIBCO BRL). Fragment sizes in Kilobases are indicated on the left.

FIGURE 5. Dendrogram of the relationships among 34 isolates obtained from pycnidia and pseudothecia present in the lesions of phaeosphaeria leaf spot in maize, based on random amplified polymorphic DNA (RAPD) analysis.
Conclusion

The pycnidia and pseudothecia present in the lesions of phaeosphaeria leaf spot belong to distinct fungal species (Phoma tropica and Phaeosphaeria sp., respectively) and not to the anamorphic and teleomorphic states of the same agent.

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