ABSTRACT - The present study combines the examination of toxins produced by *C. cassiicola* and the effects of the fungus colonization on *L. camara*. *C. cassiicola* was cultivated on solid media and the crude extracts CAE and CE were produced. Both extracts were submitted to a seed germination and growth assay utilizing *Physalis ixocarpa*, *Trifolium alexandrinum*, *Lolium multiflorum* and *Amaranthus hypochoridius*. The effect of the extracts on the ATP-synthesis in isolated spinach chloroplasts was also tested. Bioassay guided chromatographic fractionation identified the most active extract (CAE). From this extract ergosta-4,6,8(14),22-tetraen-3-one (C1) and fatty acids were isolated. The C1 compound reduce ATP synthesis in isolated spinach chloroplasts. The interference of fatty acids with ATP synthesis and also with weed growth provides one explanation of the phytogrowth-inhibitory properties of such fungal extracts. Histological observations involving fungus-plant interaction were made on *L. camara* plants inoculated with *C. cassiicola* conidia suspension. After inoculations, fragments of the leaf blades were prepared for observation by light and scanning electron microscopy. Fungal colonization of *Lantana camara* was typical of a necrotroph and penetration initiated a hypersensitive response. *L. camara* reacted to the pathogen penetration through thickening of the epidermis walls, cytoplasm granulation and a cicatrisation tissue.

Keywords: allelopathy, bioassays, photosynthesis inhibitor, phytotoxicity.

RESUMO - O presente estudo combina a investigação de toxinas produzidas por *C. cassiicola* e os efeitos da colonização do fungo sobre *L. camara*. *C. cassiicola* foi cultivado em meio sólido do qual se obtiveram os extratos brutos CAE e CE. Ambos os extratos foram submetidos aos testes de germinação e crescimento utilizando *Physalis ixocarpa*, *Trifolium alexandrinum*, *Lolium multiflorum* e *Amaranthus hypochoridius*. O efeito dos extratos na síntese de ATP em cloroplastos isolados de espinafre. Os bioensaios direcionaram o fracionamento cromatográfico permitindo a identificação do extrato mais ativo (CAE). Desse extrato isolou-se o composto ergosta-4,6,8(14),22-tetraen-3-on (C1) e ácidos graxos. O composto C1 reduz a síntese de ATP em cloroplasto isolados de espinafre. A interferência dos ácidos graxos sobre a síntese de ATP e crescimento das plantas daninhas fornece uma explicação para as propriedades fitoinibitórias dos extratos fúngicos. Realizaram-se observações histológicas envolvendo a interação fungo-planta em plantas de *Lantana camara* inoculadas com suspensão de conídios de *C. cassiicola*. Após as inoculações, fragmentos da lâmina foliar foram preparados para observações por microscopia de luz e eletrônica de varredura. A colonização fúngica de *L. camara* foi tipicamente necrotrófica e a penetração iniciou uma resposta hipersensível. *L. camara* reagiu à penetração do patógeno pelo espessamento da parede da epiderme, granulação do citoplasma e tecido de cicatrização.

Palavras-chave: alelopatia, bioensaio, inibidor da fotossíntese, fitotoxicidade.
INTRODUCTION

*Corynespora cassiicola* is an anamorphic fungus, which causes foliar spots in more than 70 species of plants worldwide (Silva et al., 1998). It has been reported to infect numerous economically important crops both in tropical and subtropical countries (Breton et al., 2000). Its economic importance as a pathogen of several crops has been described and is responsible for extensive damage to rubber plantations becoming an important limiting factor to cultivations in Asia. For instance, in Sri Lanka, the fungus has spread to all rubber plantations, becoming the most destructive foliar disease to affect the growth of this crop (Silva et al., 1998). In Northern Brazil, *C. cassiicola* is considered to be one of the worst pathogens of tomato (Kurozawa & Pavan, 2006). However, some studies have indicated that *C. cassiicola* is a complex species, including populations that are physiologically distinct and show host-specificity (Onesirosan et al., 1975, Silva et al., 1998).

*Lantana camara* (Verbenaceae) is regarded as one of the world’s worst weeds (Holm et al., 1977). It is native to tropical and subtropical America and has been dispersed throughout the world as a popular ornamental plant (Sanders, 1946) benefiting its spread as a pantropical weed. During the last two decades, surveys of the fungal pathogens associated with *L. camara* were performed in Brazil aimed at finding useful biological control agents (Barreto et al., 1995; Pereira & Barreto, 2001). Among the fungi that were collected during such surveys was *C. cassiicola* which later was demonstrated to be a *forma specialis* that was physiologically specialized to *L. camara* and named *C. cassiicola* f. sp. *lantanae*. This is a severe pathogen of this plant, capable of provoking defoliation and debilitation of the attacked plants (Pereira et al., 2003). Investigation of this fungus as a mycoherbicide suggested production of an undefined toxin (Pereira et al., 2003; Onesirosan et al., 1975). This toxin secreted by the fungus may translocate between cells upon application of conidia extracts upon host leaves causing foliar necrosis identical to that observed upon direct inoculation with conidia (Breton et al., 2000).

In the course of our continuing efforts to discover new natural herbicides we describe in this study the examination of lipophilic chemical compounds produced by *C. cassiicola* and the tissue and cell changes in *L. camara* upon fungus colonization.

MATERIALS AND METHODS

**General procedures:** Column chromatography was performed using Crosfield Sorbil C60 (32-63 µm) silica gel. Infrared spectra were recorded on a Perkin Elmer FTIR PARAGON 1000 spectrometer, using potassium bromide disk. ¹H and ¹³C NMR spectra were recorded on a Varian Mercury 300 instrument (300 MHz and 75 MHz respectively), using deuterated chloroform as solvent and tetramethylsilane (TMS) as internal reference (δ = 0). Mass spectra were recorded under electron impact (70 eV) in a SHIMADZU GCMS-QP5050A instrument.

**Fungus cultivation and extraction of toxins:** *Corynespora cassiicola* (RWB 01 isolate) was aseptically cultivated on autoclaved commercial polished rice plastic bags (each containing 350 g of rice and 200 cm³ of water) and incubated at 25 ºC for 15 days. After incubation, the colonized substrate was submitted to a sequential extraction of increasing polarity i.e. ethyl acetate (named CAE extract) and then ethanol (named CE extract) using a Soxhlet apparatus for 6 hours (Carvalho et al., 2001). The solvents were removed under reduced pressure in a rotary evaporator at 40 ºC.

Prior to chromatographic fractioning, crude extracts CAE and CE were submitted to several biological assays. Initially, the effect of these extracts, from 0 µg g⁻¹ to 100 µg g⁻¹, was tested on the ATP-synthesis in isolated spinach chloroplasts (*Spinacea oleracea*) using a methodology previously reported (Barbosa et al., 2006).

The effect of the same extracts (CAE and CE) on seed germination and growth of dicotyledonous (*Physalis ixocarpa* and *Trifolium alexandrinum*) and the monocotyledonous (*Lolium multiflorum* and *Amaranthus hypochondriacus*) plants was examined. Bioassays were performed by germinating 40 seeds of each species for five days in 9 cm
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Petri dishes containing two sheets of Whatman n° 1 paper and 3 cm³ of test or control solution. Seeds were incubated in the dark at 25 °C in a controlled chamber (Kato-Noguchi & Tanaka, 2003/4). A solution was prepared using DMSO in concentration of 50 µM. Control experiments were also conducted with the same DMSO concentration. The seed germination data is presented as percent differences from control after three days of incubation. After incubation, the root and shoot lengths were measured to the nearest millimeter. All treatments were replicated four times using a completely randomized design. The percentages of root and shoot growth inhibitions were calculated in relation to the control. The data were analyzed using Tukey’s test at 0.05 probability level.

Chromatographic fractionation of CAE extract: The crude ethyl acetate extract (202 g) was fractionated on a silica gel column chromatography using a series of solvent gradients. Initial elution with hexane (fraction F1) was followed by dichloromethane (fraction F2), then hexane:ethyl acetate (1:1v/v) (fraction F3), ethyl acetate (fraction F4), ethyl acetate:methanol (2:1v/v) (fraction F5) and finally methanol (2:1v/v) (fraction F6). Each fraction was concentrated using a rotary evaporator and submitted to the ATP-synthesis screening bioassays as previously described (Barbosa et al., 2006).

The most active fractions (F1, F2 and F3) on the ATP-synthesis bioassays were further analyzed for fatty acids and triacylglycerols content by gas chromatography as described previously (Barbosa et al., 1999). Further chromatographic purification of fraction F2 (14 g) with hexane:ethyl acetate (3:1 v/v) resulted in the isolation of a white solid identified as ergosta-4,6,8(14),22-tetraen-3-one (named C1). The structure of this compound was elucidated by infrared and RMN (¹H and ¹³C) spectroscopy and mass spectrometry.

Fraction F3 (39 g) was purified on a silica gel column chromatography, eluted with a mixture of hexane:ethyl acetate (6:1 to 1:1 v/v), resulting in the isolation of 0.07 g of compound C1. The remaining of this fraction (38.81 g) consisted of a combination of fatty acids and triacylglycerols, as characterized by gas chromatography analysis (Barbosa et al., 1999).

Histopathological observations: Observations of fungus-plant interaction were made on L. camara plants inoculated with a suspension of conidia of C. cassiicola (RWB 01 isolate). Inoculations were performed following the procedure described by Pereira et al. (2003). Seven leaves of L. camara (mainly the third to fifth leaves counting from the apex of a branch) were treated with C. cassiicola. The treatment consisted of brush-inoculating the adaxial and abaxial sides of leaves with a conidial suspension at concentrations of 1 x 10⁶ conidia cm⁻³ then incubating the plants in a dew chamber for 48 h in the dark. The plants were then transferred to a thermostatted greenhouse at 25 °C. Selected leaves were collected regularly at 24 hour intervals, for seven days. Fragments of the basal, median and apical portions of the leaf blades were fixed in FAA and prepared for observation under a light microscope (Olympus AX 70, fitted with a camera Olympus U-Photo). The samples were embedded in paraffin and transversally sectioned in a rotating microtome (Spencer, 820). Sections were stained with safranin and astra blue (Kraus & Arduim, 1997) and mounted in Permount.

Observations of C. cassiicola structures within the plant tissue were made by means of a clearing and staining method described by Keane et al. (1988).

Leaf segments from selected material were prepared for SEM analysis by immersion in 2.5% glutaraldehyde and post-fixed in osmium tetroxide (1% m/v). Samples were dehydrated in an ethanol series and then dehydrated in a critical point dryer (model CPD 030, BAL-TEC, Liechtenstein) using CO₂ as transition fluid. The samples were subsequently coated with gold (20 nm thickness) in a sputter coater (SCA 010, Balzers, Liechtenstein) using the procedure described by Bozzola & Russell (1992) and examined using an SEM Zeiss model LEO 1430VP (Cambridge, England). Each experiment was repeated three times.

RESULTS AND DISCUSSIONS

The effect of CAE and CE extracts on the germination and growth (root and shoot development) of T. alexandrinum, L. multiflorum, P. ixocarpa and A. hypochondriacus at 50 µg g⁻¹
concentration was evaluated (Table 1). The CAE extract caused 37, 42 and 13% inhibition on germination and the radicle and shoot growth of *P. ixocarpa*, respectively. The CE extract caused 122% growth induction on the roots of *P. ixocarpa* and had no significant effect on the aerial parts or upon germination. As for the development of *T. alexandrinum*, the CAE extract caused 34% and 21% inhibition on radicle and shoot growth, respectively. The CE extract had no significant effect on the radicle and shoots development, but reduced germination by 14%. Ethyl acetate extract (CAE) had no effect on germination but caused 44% and 27% inhibition on the roots and shoots development of *L. multiflorum*. The CE extract was less active on this species causing 21% and 6% inhibition of the roots and aerials parts, respectively. In contrast to the CAE extract, it caused a 58% inhibition on the germination of *L. multiflorum*. The CAE extract had no significant effect on the development of the root and shoot, but caused 17% inhibition on the germination of *A. hypochondriacus*. The CE extract, however, caused 25% growth induction upon radicle development for this monocotyledonous species but development of germination and aerial parts of the plant remained unaffected. The phytotoxicity results of extracts CAE and CE on the germination and growth (root and shoot development) inhibition of selected dicotyledonous and the monocotyledonous species indicated that the CAE and CE extract have unspecific behavior since they inhibited both monocotyledonous and dicotyledonous plants.

Having confirmed the phytotoxicity of both extracts, especially the CAE, and considering that the fungus causes chlorosis upon infected plant tissue, we suspected that the toxin or toxins produced by this pathogen could affect the photosynthesis (Mills et al., 1980; Barbosa et al., 2006; Demuner et al., 2006; King-Díaz et al., 2006). Accordingly, the CAE and CE extract were able to inhibit the ATP synthesis in isolated spinach chloroplasts, being the CAE the more effective one, presenting an IC$_{50}$ of 20 µg g$^{-1}$ (Figure 1). Fractions F1 and F2 caused significant inhibitory effect on the ATP synthesis (IC$_{50}$ < 25 µg g$^{-1}$). Fraction F3 also affected the ATP synthesis (Figure 2).

The infrared spectra of fractions F1, F2 and F3 revealed that they were composed mainly of fatty acids and triacylglycerols, as suggested by absorptions at 1710 and 1742 cm$^{-1}$, respectively. Further gas chromatographic analysis of these fractions resulted in the characterization of the following acids in the approximate concentrations: myristic (C$_{14}$:0; 1%); palmitic (C$_{16}$:0; 24%); palmitoleic (C$_{16}$:1; 2%); stearic (C$_{18}$:0; 11%); linoleic (C$_{18}$:2; 20%); linolenic (C$_{18}$:3; 35%); arachidic acid (C$_{20}$:0; 4%). The presence of these lipophilic compounds might be of relevance in the context of the present study, since an investigation carried out by Tso (1964) have demonstrated that alkyl fatty esters derived from C6 to C36 saturated and unsaturated fatty acids cause plant-growth inhibition. Calvo et al. (1999) demonstrated the sporogenic effect of polyunsaturated fatty acids on the development of *Aspergillus* spp. as well as members of other fungal genera. The results presented in Figure 2 suggest that fatty acids and esters are responsible in part for the observed inhibition on the ATP synthesis.

<table>
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<tr>
<th>Table 1 - Results of germination and growth development bioassays at 50 µg g$^{-1}$ concentration</th>
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<td><em>P. ixocarpa</em></td>
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Zero: control; positive values: stimulation; negative values: inhibition.
The thin-layer chromatography analysis of all six fractions (F1 to F6) showed the presence of a fluorescent compound in fractions F2 and F3. This fluorescent compound (C1) was isolated by column chromatography fractionation as a white crystalline solid. The EI mass spectrum of C1 flagged a peak at \( m/z \) 392, corresponding to the molecular formula \( C_{28}H_{40}O \). This compound possessed twenty-eight unique resonance signals within the \( 13C \) NMR spectrum. DEPT 135 and DEPT 90 experiments indicated the multiplicity associated with each peak (6 C, 10 CH, 6 CH\(_2\), 6 CH\(_3\)). The signal at \( \delta \) 199.8 indicating the presence of a carbonyl group, was confirmed by the strong infrared spectrum absorption at 1669 cm\(^{-1}\). Signals at \( \delta \) 123.2, 124.6, 124.7, 132.7, 134.3, 135.2, 156.4 and 164.7, indicated the existence of eight different sp\(^2\) carbons, corresponding to four double bonds. A detailed analysis of the \( \text{ }^1\text{H} \) NMR and \( 13C \) NMR spectra led us to propose the structure of ergosta-4,6,8(14),22-tetraen-3-one for compound C1. Compound C1 inhibited the synthesis of ATP (27% inhibition at 250 \( \mu \)g g\(^{-1}\), Figure 3) in chloroplasts isolated from spinach. Since, this compound has been previously isolated from *Alternaria alternata* (Seitz & Paukstelis, 1977), *Pleurotus ostreatus* (Chobot et al., 1997) and *Tuber indicum* (Jinming et al., 2001) our spectroscopic data (not presented) were compared with those reported in the literature, confirming the identification. Although this steroid is a common constituent of fungus bio-membranes (Jinming et al., 2001), to the best of our knowledge, this is the first report describing its phytotoxicity. The results of the ATP bioassay showed that the phytotoxicity observed for this fungus may be explained in part for the presence of this steroid. Studies have shown that fungus and plants can be involved in allelophatic interactions (Bitencourt et al., 2007; Borges et al., 2007; Souza Filho, 2007; Rizzardi et al., 2008).

The histopathological observations showed that 24 h after healthy *L. camara* (Figure 4A)
Plants were inoculated with *C. cassiicola*, lesions were already visible upon leaves. Defoliation began 48 h after inoculation. Microscopic examinations have demonstrated that between 24-48 h after inoculation, conidial germination and tissue penetration had already occurred. At last, the histopathological observations differ from the results described by Pereira et al. (2003) that reported necrosis in *C. cassiicola* inoculated *L. camara* only 72 hours after inoculation. Our results are consistent with that of Breton et al. (1997) who have observed that *C. cassiicola* produced lesions on *Hevea brasiliensis* tissue leaves 24 hours after infection. Lesions were initiated by maceration of leaf tissue which resulted in necrosis (Figures 4B and 4C). Clarification of tissues allowed the observation of changes in necrotic tissues at the epidermis such as cell wall thickening and color changes (cells often becoming shiny golden brown) accompanying the fungal colonization (Figures 5A and 5B).

Penetration of *L. camara* by *C. cassiicola* occurred preferentially through the intercellular spaces, often next to the stomata between the subsidiary cells and the guard-cells (Figs. 5C, 5D). Purwantara (1987) observed that when *C. cassiicola* attacks *Hevea*, it penetrates preferentially through the intercellular spaces. Occasionally, penetration occurred through the stomata but no tropism towards stomata was apparent, as hyphae were mainly concentrated around the leaf hairs. The presence of such hyphal concentration surrounding the non-glandular hairs corresponded to the destruction of those structures (Figures 6A and 6B). The germinated conidia and fungal hyphae were much more abundant abaxially, which was clearly the first surface to show symptoms resulting from fungal infection. Furthermore, preference for colonization through the abaxial surface may be typical of each particular host-pathogen association. Duarte et al. (1983) observed morphological, physiological and

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**Figure 4** - *L. camara*. (A) Healthy flowering branch; (B) Inoculated leaf with necrotic leaf spot, 96 hours after inoculation with *C. cassiicola* (adaxial); (C) Ibid (abaxial).

**Figure 5** - Interaction between *C. cassiicola* and *L. camara* observed on clarified leaves. (A) Mycelial growth within plant tissue, 24 hours after inoculation (note necrotic area and reaction on walls of epidermal cells – arrowed); (B) Ibid 48 hours after inoculation (note granular reaction to fungal infection within epidermal cells - arrowed); (C) and (D) Hyphae penetrating between subsidiary cell and guard cell (arrowed). Ec = epidermal cell, Gc = germinating conidium, H = hyphae, S = stomata. (A) Bar = 20 μm. (B) Bar = 10 μm. (C) and (D) Bar = 15 μm.
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pathogenic differences among isolates of C. cassiicola from papaya and cocoa. Many hyphae developed following the depressions at contact areas of epidermal cells (Figure 6C), where exudates and other substances may accumulate aiding adherence of fungal structures. The cuticle could have represented a more effective barrier for an adaxial fungal penetration, can be interpreted as an adaptive strategy (Machado et al., 2008) although some penetration was also observed on that side (Figure 6D). Hyphae penetrated preferentially through the anticlinal walls of epiderm cells and extended from the lacunose parenchyma to the palisade parenchyma. Destruction of palisade tissue was less common since the fungus colonizes more aggressively upon the abaxial surface. In some areas that became necrotic after infection it was possible to observe cicatrized tissues. In such regions the cells usually stained intensely with safranin, indicating the presence of lignin. This was also observed at leaf vein areas where vascular bundles were disorganized when hyphae were present. Purwantara (1987) also noted that vascular bundles and associated tissues (epidermis, parenchyma and sclerenchyma) collapsed and stained red when exposed to safranin as observed for C. cassiicola. According to Purwantara (1987), the protoplast becomes granular after C. cassiicola infection because of chloroplast disintegration. Generally, plant response to hyphal infection varies from a slight darkening of the protoplasm of cells adjacent to the hyphae to the complete necrosis of the epidermal cells below the hyphae, which was similar to our findings (Dankyn & Milholland, 1984). This typical phenotypic browning response is a common event upon cell death, and normally it corresponds to accumulation of phenolic substances within the dead cells (Heath, 1998). Phenolic compounds are known to inhibit the fungal cell wall extension causing the swelling and subsequent rupture of the infecting hyphae (Mauseth, 1995). Plant cell degradation, or death, are known host defense strategies against invading pathogens. The hypersensitive response also was verified for C. cassiicola. This temporary response halts proliferation of pathogen until the host plant initiates phytoalexin production and lyses-inducing proteins with fungicidal activity (Brown et al., 1998). Since such barriers serve to immobilize the invading microorganism, allowing them exposure to a cocktail of antimicrobial products which include phytoalexins and enzymes involved in the production of active oxygen species (Breton et al., 1997, Brown et al., 1998). Furthermore, host resistance to C. cassiicola attack depends on a combination of structural barriers as well as the chemical substances produced by the host and fungus. Recent studies have suggested that the toxin cassicolin, produced by C. cassiicola, is fundamental to warrant its pathogenicity and may be a determinant factor in its pathogenicity (Breton et al., 2000; Lamotte et al., 2007). Application of this toxin reproduce the necrotic disease symptoms seen in fungal invasion of C. cassiicola on H. brasiliensis. To the best of our knowledge, the chemical structure of this interesting toxin remains undetermined.

In conclusion, the results from the present investigation indicated that C. cassiicola contains lipophilic phytogrowth inhibitors that could be involved in the allelopathic interactions with L. camara. The interference
of fatty acids and ergosta-4,6,8(14),22-tetraen-3-one isolated from the CAE extract (fractions F2 and F3) with ATP formation, weed growth, and L. camara infections might explain, in part, its phytogrowth inhibitory properties and its putative allelopathic effects.

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LITERATURE CITED


