Biological Control Potential of Colletotrichum gloeosporioides for Coffee Senna (Cassia occidentalis)

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ABSTRACT

A fungal pathogen, Colletotrichum gloeosporioides was isolated from a greenhouse-grown seedling of coffee senna (Cassia occidentalis) and evaluated as a mycoherbicide for that weed. Host range tests revealed that coffee senna, wild senna (C. marilandica), and sicklepod (C. obtusifolia) were also affected by this pathogen, but 35 other crop and weed species, representing 8 botanical families were not affected. The fungus sporulated prolifically on solid and liquid media with maximum spore germination and growth occurring at 20°C - 30°C. Optimal environmental conditions included at least 12 h of free moisture (dew) at 20°C - 30°C. Spray mixtures containing approximately 1.0 × 10⁵ or more conidia·ml⁻¹ gave maximum control when coffee senna seedlings were sprayed until runoff occurred. Coffee senna seedlings that were in the cotyledon to first-leaf growth stage were most susceptible to this pathogen. Weed control efficacy studies under field conditions demonstrated that control of coffee senna was directly proportional to the inoculum concentration applied. Results of these tests suggest that this fungus has potential as a mycoherbicide to control coffee senna, a serious weed in the southeastern U.S.

Keywords: Bioherbicide; Mycoherbicide; Coffee Senna; Cassia occidentalis; Colletotrichum gloeosporioides

1. Introduction

Coffee senna (Cassia occidentalis L.) is a non-nodulating legume that was originally introduced as a potential crop plant [1,2]. It has escaped cultivation and has become widely distributed in the south-central and southeastern regions of the United States [3]. It is a very troublesome weed in soybean [Glycine max (L.) Merr.], cotton (Gossypium hirsutum L.), and peanut (Arachis hypogaea L.) fields in much of the southeastern U.S. [4-8].

Coffee senna is economically important because it causes yield loss, seed quality degradation, and difficulty with harvest [9]. Higgins et al. [10] reported that one coffee senna plant 7.5 m⁻¹ of row can reduce seed cotton yields by 117 kg·ha⁻¹. Coffee senna seeds can germinate and emerge throughout the growing season and its germination and emergence have been recently assessed [11]. Its seeds have impermeable seed coats and scarification is required to allow imbibition of water, and the breaking of dormancy [3].

Control of this weed is difficult because of its tolerance to many commonly used herbicides, its prolific growth habit, and emergence throughout the growing season [4,9,12-14]. Alternative controls are needed to replace or supplement existing methods. The technical and biological feasibility of mycoherbicides for controlling various weeds has been established [15], and this method warrants consideration for controlling coffee senna.

An anthracnose disease was observed on greenhouse-grown seedlings of coffee senna and the fungus Colletotrichum gloeosporioides (Penz.) Penz. & Sacc, an anamorph of Glomerella cingulata (Stoneman) Spauld. and Schrenk, was isolated from infected leaf and stem tissue. Coffee senna seedlings sprayed with conidial suspensions of this fungus developed severe disease symptoms under controlled conditions (suggesting potential bioherbicidal activity) and a patent was issued for the use of this fungus as a biological control agent (16). Later studies indicated that this fungus was also useful in the biological control of sicklepod, a closely related weed of several crops [17,18]. More extensive studies on the potential of using this fungus as a mycoherbicide for coffee senna comprise the subject of this report. C. gloeosporioides was evaluated in laboratory, growth chamber, and field studies to determine the effects of temperature on germination and growth of the fungus in vitro, the effect of dew period, dew temperature, inoculum concentration, and plant
growth stage on biocontrol efficacy, and the host range of this pathogen. Determination of these parameters is essential for evaluating this fungus as a mycoherbicide.

2. Materials and Methods

2.1. Isolation and Culture of Colletotrichum gloeosporioides

The fungus was isolated from diseased coffee senna tissue by surface sterilizing sections of diseased tissue in 0.05% NaOCl for 1 min, then placing the sections on potato-dextrose agar (PDA) amended with the antibiotics chloramphenicol (0.75 mg·ml⁻¹) and streptomycin sulfate (1.25 mg·ml⁻¹). The plates were incubated for 48 h at 25°C. Advancing edges of fungal colonies were transferred to PDA and incubated for 5 days at 25°C under alternating 12-h light/12-h dark regimens, provided by cool white fluorescent lights. The fungus was sub-cultured on PDA without antibiotics, and preserved under refrigeration in sterilized sandy loam soil (25% water holding capacity), and on sterile silica gel containing skim milk [19]. Conidia of C. gloeosporioides were produced in a 30-L fermenter (B. Braun, Model C-30, Bethlehem, PA, USA) at a 20-L working volume containing 20-L of modified Richards’ medium [20] plus V-8 vegetable juice (15%; v/v) (Campbell’s Soup Co., Camden, NJ, USA), with an aeration rate of 10 L air min⁻¹ and an agitation of 250 rpm using 2 Rushton-style impellers. A silicon-based antifoam (HODAG FD-62, Lambent Technologies, Gurnee, IL, USA) was used to control foaming, as required. Inoculum for the fermentations consisted of 250 ml of conidia and mycelium grown in identical medium that was produced in 500 ml Erlenmeyer shake-flasks, incubated at 25°C and 250 rpm for 5 days. Fermenter batch fermentations were maintained at 25°C for 5 days. Spores were separated from the spent medium by filtering through double-layered cheesecloth. The spore suspension was centrifuged at 2500 x g for 15 min at 15°C. The medium had an initial pH of 5.5, and pH was not controlled or monitored during culture growth. Conidial concentrations used in the tests were determined with hemacytometers.

2.2. Effect of Temperature on Germination and Radial Growth Rate

Conidial germination was measured by spreading 0.1 ml of a suspension (1.0 × 10⁷ conidia·ml⁻¹) on PDA plates, and incubating them at 10°C, 15°C, 20°C, 25°C, 30°C, or 35°C on open-mesh wire shelves of an incubator (Precision Scientific Inc., Chicago, IL, USA). Twelve-hour photoperiods were provided by two-20 W, cool-white fluorescent lamps positioned in the incubator door. Light intensity at plate level was 200 μE·m⁻²·s⁻¹ photosynthetically active radiation (PAR) as measured with a light meter (LI-COR Inc., Lincoln, NE, USA). Germinated conidia (500 plate⁻¹) were counted after 16 h.

For radial growth studies, 5-mm plugs were taken from advancing margins of a 7-day old colony of the fungus and placed in the centers of PDA plates. The plates were incubated at temperatures of 10°C, 15°C, 20°C, 25°C, 30°C, or 35°C as described above for the conidial germination studies.

Colony diameters were measured after 7 days of incubation. In each experiment, five replicate plates for each temperature were utilized. Both experiments were conducted twice, and the results of each experiment were pooled following testing for homogeneity.

2.3. Plant Production

In all experiments coffee senna plants were grown from seed in a commercial potting mix contained in peat strips. Each strip contained 12 plants. The potting mix was supplemented with a slow-release fertilizer (14:14:14, NPK). The plants were placed in sub-irrigated trays on benches in the greenhouse. Greenhouse temperatures ranged from 25°C to 30°C with 40% to 60% relative humidity (RH). The photoperiod was 12 h with 1650 μE·m⁻²·s⁻¹ (PAR) measured at midday.

2.4. Effect of Air and Dew Temperature

Five to seven-day old seedlings (cotyledonary to early first-leaf stage) of coffee senna were sprayed until runoff with 1.0 × 10⁷ conidia·ml⁻¹ in water. Control plants were sprayed with distilled water only. The plants were placed in darkened dew chambers at 100% RH at temperatures of 15°C, 20°C, 25°C, 30°C, or 35°C. The plants were then transferred to environmental growth chambers (Conviron, Model E-7, Pembina, ND) with day/night air temperatures of either 20°C/10°C, 25°C/15°C, 30°C/15°C, 35°C/25°C, or 40°C/30°C. Photoperiods were 14 h with 820 to 840 μE·m⁻²·s⁻¹ PAR and a RH of 65%.

2.5. Effect of Dew Period Duration

Coffee senna seedlings in the cotyledonary to early first leaf stage of growth were sprayed until runoff occurred with a spray mix containing 1.0 × 10⁷ conidia·ml⁻¹. Control plants were sprayed with distilled water. The inoculated plants were then placed in darkened dew chambers at 25°C and 100% RH for periods of 4, 8, 12, 16, 20, or 24 h. Following the dew period inoculation, the plants were placed on sub-irrigated trays in the greenhouse and monitored for 14 days for disease development and mortality. Greenhouse temperatures were 28°C to 32°C, with 40% to 60% RH at day lengths of 12 h with an average of 1650 mol·m⁻²·s⁻¹ photosynthetically active radiation at
mid-day. Mortality and dry weight reductions were recorded after 14 days after treatment. The experiment was conducted twice with 3 sets of 12 plants for each experiment.

2.6. Effect of Inoculum Concentration and Plant Growth Stage

Coffee senna plants in the cotyledonary, 1 to 2 true-leaf, 3 to 4 true-leaf and 5 to 7 true-leaf stages of growth were sprayed with conidial suspensions of $1.0 \times 10^3$ to $1.0 \times 10^7$ conidia·ml$^{-1}$ and held in a dew chamber for 16 h at 25°C. Control plants were sprayed with distilled water only. Plants were moved to the greenhouse, and mortality and dry weight reductions were recorded after 14 days after treatment. Experiments were conducted twice with 3 sets of 12 plants for each experiment.

2.7. Host Range

2.7.1. Crop Plants

A variety of crop plants were inoculated with the fungus in order to evaluate its host range: pumpkin (Cucurbita pepo L.) cv. Jack-o-Lantern; squash [Cucurbita pepo var. melopepo (L.) Alef.] cv. Golden Summer Crookneck; watermelon (Citrullus vulgaris Schr.) Charleston Grey; sweet corn (Zea mays L.) cv. Truckers Favorite and Honey Butter; rice (Oryza sativa L.) cvs. Labelle and Starbonnett; grain sorghum [Sorghum bicolor (L.) Moench] cv. Texas C-424; alfalfa (Medicago sativa L.) cv. Delta; soybean cvs. Bedford, Bragg, Davis, Dare, Centennial, Forrest, Hill, Hood, and Tracey; peanut (Arachis hypogaea L.) cv. Tennessee Reds; garden bean (Phaseolus vulgaris L.) cvs. Kentucky Wonder, Romano Pole, Ohio Pole, Jackson Wonder, and Henderson Bush; blackeyed pea [Vigna unguiculata (L.) Wolf.] cvs. Lady Cowpea, and White Cowder Pea; cotton cvs. Stoneville 506 and Deltapine 61; tomato (Lycopersicon esculentum Mill.) cvs. Beefsteak and Marion. Seeds of these plant species were planted and grown under greenhouse conditions to various growth stages when they were subjected to the experimental procedures related to the objectives of these studies.

2.7.2. Weeds

Various grass and dicotyledonous weeds were inoculated to determine this pathogen’s host range as follows: common cocklebur (Xanthium strumarium L.); tall morning-glory (Ipomoea purpurea (L.) Roth. and pitted morning-glory (Ipomoea lacunosa L.); johnsongrass [Sorghum halepense (L.) Pers.]; coffee senna (Cassia occidentalis L.); sicklepod (C. obtusifolia L.), partridge pea (C. fasciculata Michx.), wild senna (C. marilandica L.), hairy sensitive pea (C. pilosa L.), and round-leaf cassia (C. rotundifolia L.); Florida beggarweed (Desmodium tortuosum L.); showy crotonaria (Crotalaria spectabilis Roth); hemp sesbania (Sesbania exaltata Rydb. ex. A.W. Hill; rattlebox (S. drummondii L.), northern jointvetch [Aeschynomene virginica (L.) B.S.P.], prickly sida (Sida spinosa L.); velvetleaf (Abutilon theophrasti Medik); and jimsonweed (Datura stramonium L.). Seedlings ranging from the cotyledonary to the third true-leaf stage were sprayed until runoff with conidial suspension of $1.0 \times 10^7$ ml$^{-1}$ and incubated in dew chambers for 24 h at 25°C. Non-inoculated strips containing 12 plants of each species were included as a control. Coffee senna seedlings were included in all experiments as susceptible controls. Plants were considered either resistant (no visible reaction) or susceptible (necrosis, flecking) by visual observation after 10 days of incubation in the greenhouse. Seeds of these weeds were grown under greenhouse conditions and tested as described above for the crop species.

2.8. Field Experiments

Two field experiments were conducted on a Dundee very fine sandy loam (Aeric Ochraqualf) soil at the USDA-ARS, Southern Weed Science Experimental Farm at Stoneville, MS. Plots consisted of four rows of soybeans (cv. “Centennial”), 12.2 m long and 1 m apart, with the two center rows receiving treatment. All rows were planted with scarified coffee senna seed at a density of about 100 seeds·m$^{-1}$ of row. Treatments consisted of: 1) $1.0 \times 10^6$ conidia·ml$^{-1}$ in distilled water and 0.2% Silwet L-77 surfactant; 2) $1.0 \times 10^6$ conidia·ml$^{-1}$ in distilled water and 0.2% Silwet L-77 surfactant; 3) $1.0 \times 10^5$ conidia·ml$^{-1}$ in distilled water and 0.2% Silwet L-77 surfactant; and 5) distilled water only. The conidia·ha$^{-1}$ in those plots that received fungal treatments. Applications were made at midday with a hand-held pressurized sprayer. For the first experiment, environmental conditions at the time of inoculation and for 24 h following treatment were: temperature at inoculation, 29°C with a RH of 69%. The high temperature for the 24 h period was 31°C and the low temperature was 25°C. Maximum RH was 94%, with a short dew period of 3 h. For the second experiment, environmental conditions at the time of inoculation and for 24 h following treatment were: temperature at inoculation, 34°C with a RH of 58%. The high temperature for the 24 h period was 34°C, and the low temperature was 24°C. Maximum RH was 92%, with a short dew period of 4 h. The plants were monitored for disease development at 7 day intervals for 21 days. A randomized complete block design was utilized, and the treatments were replicated four times. Data over the 2 years from the two experiments were pooled fol-
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2.9. Statistical Procedures

All experiments utilized a randomized complete block experimental design, with a minimum of three replications of 12 plants for each experiment. All experiments were repeated. Data were subjected to the analysis of variance and values are presented as the means of replicated experiments. Duncan’s multiple range test at the 0.05 level of probability was used to separate means in the dew and air temperature experiment. Confidence limits (95% level) were used to analyze the data in remaining experiments.

3. Results

The fungus was identified as *Colletotrichum gloeosporioides* based on microscopic examination of diseased tissue and cultural characteristics. The morphology of the isolate is identical to that reported previously for this species [21]. The acervuli are glabrous (non-setose) and are slightly sunken in the host tissue. Under moist conditions, slimy masses of conidia accumulate on the upper surface of the acervulus and cause breaking of the epidermal layer and cuticle. Identification was confirmed by B. C. Sutton at the International Mycological Institute (IMI) in Kew, England (the fungus is an anamorph of the ascomycetous fungus *Glomerella cingulata*). Cultures of this pathogen have been deposited in the International Mycological Institute as No. IMI 325028 and with the Agricultural Research Culture Collection (NRRL) as accession No. NRRL 21046.

3.1. Isolation and Culture

The fungus was readily isolated from diseased tissue and it sporulated abundantly on PDA. When reinoculated onto healthy seedlings, the fungus was highly virulent and killed all inoculated plants within 5 days, while the controls remained healthy (data not shown). The organism produced typical anthracnose lesions on leaves and stems, with acervuli scattered throughout the lesions. The acervuli were glabrous (non-setose) and were slightly sunken in the host tissue. Conidia were elliptical, with rounded ends, and range in size from 15 to 19 µm by 4 to 6 µm and averaged 16 by 6 µm.

3.2. Effect of Temperature on Germination and Radial Growth

Maximal germination of conidia on PDA occurred at 25 (86%) and 30°C (79%), with significant decreases in germination at the other temperatures (Figure 1(A)). The fungus grew well at 20°C to 30°C, but growth was significantly reduced when incubated at 10°C or 15°C and slow growth occurred at incubation temperatures of 35°C or higher (Figure 1(B)). The fungus also sporulated prolifically under submerged culture in modified Richard’s medium.
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Medium. Conidial yields averaged $3.0 \times 10^8$ conidia·ml$^{-1}$ after 7 days at 25°C and 250 rpm (data not shown). This growth is comparable to conidial yields produced by other *C. gloeosporioides* formae speciales that have been evaluated as bioherbicides against other weeds [22,23].

3.3. Effect of Dew and Air Temperatures

Optimal day, night, and dew temperatures for maximum weed control (100%) were 30°C/25°C/20°C (Figure 2). Both weed mortality and biomass reduction were significantly reduced when the day/dew/night temperatures were lowered to 20°C/15°C/10°C. No pathogenesis or mortality occurred on coffee senna seedlings under a day/dew/night temperature regime of 40°C/35°C/30°C (Figure 2).

3.4. Effect of Dew Period Duration

The fungus killed coffee senna over a broad range of dew period durations conducted at 25°C (Figure 3). Although some infection occurred at all dew periods tested, a dew period of at least 12 h was required to kill about 80% of fungus-inoculated plants. Complete mortality was not achieved at any dew period, but many plants were severely stunted, which resulted in greatly reduced biomass.

3.5. Host Range

In greenhouse tests the fungus had no effect on other weed and crop plants tested except for sicklepod, which was slightly affected by the pathogen, and wild senna, which was nearly as susceptible as coffee senna (data not shown). Plants were grown to the cotyledonary to third-leaf growth stage and tested as described in the material and methods. Severe disease developed on coffee senna and wild senna within 2 to 3 days of inoculation, and seedlings were killed within 4 to 6 days.

3.6. Effect of Plant Growth Stage and Inoculum Concentration

Weed control (mortality) under greenhouse conditions was significantly increased at all growth stages by increasing the fungal inoculum concentration (Figure 4). Coffee senna seedlings in the 5 to 7 leaf stage were more resistant to infection than younger plants. Weed control was significantly less than that achieved with plants at

![Figure 2](image)

Figure 2. Effect of temperatures of day/night/dew periods on weed control of coffee senna under growth chamber conditions. Black bars = mortality; gray bars = reduction in biomass. Values of bars denoted by the same letter are not significantly different at the 95% confidence level.

![Figure 3](image)

Figure 3. Effect of dew period duration on weed control of coffee senna inoculated with *C. gloeosporioides* at 1.0 × 10$^7$ under greenhouse conditions. Black bars = mortality; gray bars = reduction in biomass. Values of bars denoted by the same letter are not significantly different at the 95% confidence level.

![Figure 4](image)

Figure 4. Effect of plant growth stage on weed control (mortality) of coffee senna inoculated with *C. gloeosporioides* at various inoculums concentrations under greenhouse conditions. Growth stages: cotyledon-to-1 leaf stage (○), 2 to 4 leaves (▲); 5 to 7 leaves (▽); 8 to 10 leaves (□); 11 to 13 leaves (●). Error bars are ± 1SEM.
earlier growth stages at the inoculum concentrations tested. Similar results were found on the dry weight reductions of plants at these growth stages and conidia concentrations (data not shown).

3.7. Field Tests

In field experiments, coffee senna plants treated with the highest inoculum concentration ($1.0 \times 10^7$ conidia·ml$^{-1}$) were controlled over 80% at 21 days after treatment, while only 40 and 20% weed control was achieved with $1.0 \times 10^6$ conidia·ml$^{-1}$ and $1.0 \times 10^5$ conidia·ml$^{-1}$, respectively, following this time period (Figure 5). The fungus sporulated profusely on infected tissue, and killed greenhouse-grown coffee senna seedlings when re-inoculated onto them, thus fulfilling Koch’s Postulates. No disease or mortality occurred to untreated control plants, or to plants treated with surfactant only (data not shown).

4. Discussions

These studies show that this isolate of *C. gloeosporioides* satisfies the requirements of a successful mycoherbicide as established by Daniel *et al.* [22]. It produces abundant inoculum in culture, is highly specific and virulent, and is pathogenic over a reasonably wide temperature range. The optimum temperatures for growth, germination, and pathogenic over a reasonably wide temperature range. Inoculum in culture, is highly specific and virulent, and is pathogenic under the conditions tested here, it is possible that through pathogen selection and formulation improvements, such as the use of crop oils, invert emulsions and/or surfactants, this weed could also be controlled by this isolate of *C. gloeosporioides*. Additional studies under field conditions will be required to further evaluate the potential for this pathogen as a mycoherbicide for coffee senna control.

![Figure 5. Effects of *C. gloeosporioides* conidia at several concentrations on control of coffee senna under field conditions.](image)

**Figure 5.** Effects of *C. gloeosporioides* conidia at several concentrations on control of coffee senna under field conditions. Concentrations are: ($\bullet$) $= 10^5$, ($\square$) $= 10^6$ and ($\mathrm{●}$) $= 10^7$ conidia·ml$^{-1}$. Error bars are ± 1SEM.

REFERENCES


[6] T. M. Webster and G. E. MacDonald, “A Survey of *C. gloeosporioides* spp. evaluated as mycoherbicides [24-28]. These temperatures are typical of those that occur in much of the southeastern U.S., where coffee senna is a serious weed problem [4]. This isolate of *C. gloeosporioides* did not affect the soybean cultivars or any other leguminous crop species tested. However, in order to further define the host range, more research should be conducted with this pathogen to include other soybean cultivars and other leguminous crop species that encompass a more diverse genetic background [29].

A major constraint of most fungi that have been evaluated as mycoherbicides, is the requirement for a lengthy period of free moisture (dew) following inoculation. However our finding that generally, better control was achieved under field versus greenhouse conditions was somewhat contrary to our expectations. This may have been caused by a more rapid drying of the dew and inoculum on plants soon after placement in the greenhouse compared to conditions in the field. This could be attributed to lower RH and the drying effects of air movement by fans. The temperature in the greenhouse tests were also generally lower than those in the field studies. The higher RH in the field could have prolonged favorable conditions for infectivity even though there the dew period in the field was shorter than for the greenhouse studies.

Proper timing of application to weeds in the most susceptible growth stages (cotyledonary to first-leaf stage) can optimize the chances for successful control. Water-in-oil formulations have shown promise in reducing the dew period requirements of some mycoherbicial fungi [18,24,30] that control various weeds. More specifically with coffee senna, granulated formulations [31, 32] and an emulsified (water-in oil) formulation [16] exhibited greater bioherbicidal efficacy and stabilization of this pathogen.

Although sicklepod was only slightly affected by this pathogen under the conditions tested here, it is possible that through pathogen selection and formulation improvements, such as the use of crop oils, invert emulsions and/or surfactants, this weed could also be controlled by this isolate of *C. gloeosporioides*. Additional studies under field conditions will be required to further evaluate the potential for this pathogen as a mycoherbicide for coffee senna control.


