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Strawberry Anthracnose: Histopathology of *Colletotrichum acutatum* and *C. fragariae*

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ABSTRACT


Ontogeny of the invasion process by *Colletotrichum acutatum* and *C. fragariae* was studied on petioles and stolons of the strawberry cultivar Chandler using light and electron microscopy. The invasion of host tissue by each fungal species was similar; however, each invasion event occurred more rapidly with *C. fragariae* than with *C. acutatum*. Following cuticular penetration via an appressorium, subsequent steps of invasion involved hyphal growth within the cuticle and within the cell walls of epidermal, subepidermal, and subtending cells. Both species of fungi began invasion with a brief biotrophic phase before entering an extended necrotrophic phase. Acervuli formed once the cortical tissue had been moderately disrupted and began with the development of a stroma just beneath the outer periclinal epidermal walls. Acervuli erupted through the cuticle and released conidia. Invasion of the vascular tissue typically occurred after acervulus maturation and remained minimal. Chitin distribution in walls of *C. fragariae* was visualized with gold-labeled wheat germ agglutinin. The outer layer of bilayered walls of conidia, germ tubes, and appressoria contained less chitin than unilayered hyphae in planta.

Additional keywords: colloidal gold, hemibiotrophy, host–pathogen interactions, lectin, ultrastructure.

MATERIALS AND METHODS

Propagation of strawberry plants. Strawberry plants (cv. Chandler) were planted in 10- × 10-cm plastic pots containing a 1:1 (vol/vol) mixture of Jiffy-Mix (JPA, West Chicago, IL) and pasteurized sand and grown for at least 6 weeks before inoculation in a greenhouse maintained at 30°C day/18°C night ±6°C with a 16-h photoperiod. Older leaves, runners, and flowers were removed 1 to 7 days before inoculation, and three or four young leaves remained on each plant at inoculation.

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Inoculation of plants. Isolates of *C. acutatum* Goff and *C. fragariae* CF-63 and CF-75 (26) were maintained on silica gel at USDA-ARS in Poplarville, MS. These isolates were grown on potato dextrose agar (PDA) for 14 to 21 days at 25°C under fluorescent light at 14/10 h of light/dark intervals. The cultures were located approximately 18 cm beneath four fluorescent tubes (Sylvania Super Saver Lite White, 34 W, 16.7 µmol-m⁻²-s⁻¹; OSRAM Sylvania, Danvers, MA). Conidial suspensions used for inoculations were prepared by flooding each PDA culture plate with sterile deionized water and gently scraping the agar surface with a glass rod to remove conidia. The resulting conidial suspension was filtered through cheesecloth and a final conidial suspension of 1.5 × 10⁶ conidia per ml was used to inoculate strawberry plants.

Attached petioles and stolons were inoculated by misting to the point of runoff or by spot inoculation. Plants were incubated in a

### Table 1. Time from inoculation of strawberry (*Fragaria × ananassa*) cv. Chandler with two *Colletotrichum* spp. to first observation of infection events

<table>
<thead>
<tr>
<th>Event</th>
<th><em>Colletotrichum acutatum</em></th>
<th><em>Colletotrichum fragariae</em></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Petiole&lt;sup&gt;a&lt;/sup&gt;</td>
<td>Stolon&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>Visible lesion</td>
<td>4 days</td>
<td>3 days</td>
</tr>
<tr>
<td>Girdling lesion</td>
<td>5 days</td>
<td>5 days</td>
</tr>
<tr>
<td>Acervulus formation</td>
<td>5 days</td>
<td>5 days</td>
</tr>
<tr>
<td>Hyphal visible in cortex</td>
<td>4 days</td>
<td>3 days</td>
</tr>
<tr>
<td>Cortical cell collapse</td>
<td>5 days</td>
<td>4 days</td>
</tr>
<tr>
<td>Epidermal cell collapse</td>
<td>Did not occur</td>
<td>Did not occur</td>
</tr>
</tbody>
</table>

<sup>a</sup> Based on 25 or more observations.

<sup>b</sup> Based on 10 or more observations.

**Fig. 1.** Light micrographs of stolon inoculated with *Colletotrichum* spp. **A and B,** *C. acutatum* 24 h after inoculation. **C and D,** Cross sections showing *C. fragariae* 48 and 72 h after inoculation. Bar = 25 µm. **A,** Septate (arrow) conidium (co) with a typical melanized appressorium (a) as a distinct swollen area located at the end of a germ tube (g) and an immature, nonmelanized appressorium (n). The pore (arrowhead) from which the penetration peg develops is visible. Bar = 5 µm. **B,** Cross section showing hypha (h) entering a stolon via an open stoma (arrow). Invading hyphae (arrowheads) in host apoplast. Bar = 25 µm. **C,** Colonization of cortical cells and periplastic spaces by fungal hyphae (h). Cortical cells show wall undulation (arrowhead) and plasmalemmal disruption (arrow). **D,** Host walls showing increasing distortion (arrowhead) accompanied by disruption of the host plasmalemma (hp) and invasion of dead cells by fungal hyphae (h).
sealed 10-gallon tank at 100% relative humidity for 24 h and then at reduced relative humidity by removing half of the plastic tank cover for the remainder of the incubation time (25,26). Plants inoculated with \textit{C. acutatum} were incubated at 28°C; those inoculated with \textit{C. fragariae} were incubated at 32°C. Following inoculation, petioles and stolons were examined microscopically at 8- to 10-h intervals for 96 h to document the morphology and time required for germ tube development, appressorial formation, penetration of the epidermis, subsequent invasion of tissue, development of disease symptoms, and acervulus production.

**Tissue processing.** Symptomatic tissue was fixed for 2 h in 0.1 M sodium cacodylate buffer (pH 7.0), 2.0% formaldehyde, and 2.5% glutaraldehyde. Tissue was rinsed with distilled water and postfixed 45 min in cacodylate buffered (pH 7.0) 1% osmium tetroxide. Tissue was dehydrated with an ethanol series (50, 70, 85, 95, and 100%) followed by propylene oxide. Tissue was then infiltrated in 50/50% by volume ERL 4206 resin (Spurr’s)/propylene oxide for 4 h on a rotator, followed by 75/25% by volume resin/propylene oxide change overnight on a rotator. Tissue was infiltrated with two changes of 100% resin during an 8-h period and cured at 70°C for 24 to 36 h. ERL 4206 resin contained nonenyl succinic anhydride (26 g), vinyl cyclohexene dioxide (10 g), diglycidyl ether (6 g), and dimethylaminoethanol (0.2 g).

Slides with semithin sections (≈1 µm) were placed on a hotplate at 55°C, stained for 1 to 2 min with 0.1% aqueous toluidine blue-O, and examined with a light microscope. Ultrathin sections (≈100 nm) were taken with a Porter-Blum MT-2B microtome (Ivan Sorvall, Inc., Newton, CT) and a diamond knife (Diatome-U.S., Fort Washington, PA) and collected on 200-mesh copper grids. Grids were stained with lead citrate (19) and 2% aqueous uranyl acetate. Sections were viewed using a Zeiss EM 10-C transmission electron microscope (Carl Zeiss, Inc., Thornwood, NY).

We developed a procedure to aid in the location of appressoria on fixed, embedded material. Slivers of epidermal tissue on which appressoria had been seen with a compound microscope were flat embedded in 2 to 3 mm of resin and cured at 70°C for 20 to 24 h. Appressoria on the host tissue were identified through the thin plastic with a compound microscope, and hand sections containing appressoria were obtained. Those hand sections were then re-mounted with Duco cement (Devcon, Danvers, MA) on prefaced stubs (created by curing Spurr’s plastic in polyethylene Beem

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**Fig. 2.** Light micrographs 72 h after inoculation with \textit{Colletotrichum} spp. **A,** Petiole inoculated with \textit{C. acutatum}. Bar = 50 µm. **B, C, and D,** Stolon inoculated with \textit{C. fragariae}. Bar = 25 µm. **A,** Advanced stage of invasion showing disrupted cortex and mature acervulus on intact epidermal and subepidermal layers. Vascular tissue (phloem and xylem) remains intact at this point. **B,** Hyphae invading parenchyma cells (arrowhead) and vessel members (arrow). **C,** Expansion of the host cuticle as the acervulus enlarges. **D,** The mature acervulus and stroma usually occupy the epidermal (e) and subepidermal (se) host layers while the cortex experiences massive destruction. ac = acervulus; s = stroma; c = cuticle; cx = cortex; hw = host wall; p = phloem; x = xylem.
Fig. 3. Electron micrograph of a germinated conidium (co) of *Colletotrichum fragariae* showing the gold probe labeling chitin on the inner wall (arrows) and the septum (s). The probe is absent from the outer wall and the matrix (m). c = cuticle; hw = host wall; iw = inner wall; ow = outer wall. Bar = 50 nm.

Fig. 4. Electron micrograph of an appressorium (a) of *Colletotrichum fragariae* separated from an adjacent fungal cell (f) by a matrix (m). The probe labeled the inner walls (arrows) and not the outer walls or matrix. The basal pore (arrowhead) is located on the basal portion of the appressorium and delimited by a membrane. c = cuticle; hw = host wall. Bar = 50 nm.
capsules size “00” and trimmed with a 1 × 1 mm face) for ultra-thin sectioning.

Application of wheat germ agglutinin lectin probe for chitin. The application of the wheat germ agglutinin probe (WGA; L-9640, Sigma, St. Louis) was modified from Benhamou (2). The ultrathin sections on 200-mesh nickel grids were floated on a drop of 0.01 M phosphate buffered saline (PBS; 40 ml of 0.02 M potassium hydroxide, 50 ml of 0.02 M potassium dihydrogen phosphate, and 10 ml of distilled water), pH 7.4, in a moist chamber for 1 h. Grids were subsequently incubated for 2 h in a moist chamber in 15 µl of lectin-gold complex at the concentration supplied by the manufacturer of 1.4 × 10^{13} particles per ml. The probe was lectin from Triticum vulgaris conjugated to 10 nm colloidal gold particles. Each grid was then washed by dipping repeatedly and gently for 10 s into PBS. Grids were air dried and then stained with lead citrate and uranyl acetate as described above.

Controls. Four-day-old cultures of C. fragariae grown on PDA as described above were overlaid by a thin layer of 2% agar to hold the mycelium together. Pieces of the culture were cut into plugs 3 × 3 mm, fixed and processed for ultrathin sectioning, and probed with gold-labeled lectin as previously described.

The specificity of the WGA lectin binding was determined by the following tests used by Benhamou (2). These controls were run simultaneously with the application of the normal probe on symptomatic tissue. (1) The gold-labeled lectin was first pre-adsorbed with poly-(1-4)-β-N-acetyl-D-glucosamine purified from crab shells (C-9752, Sigma, St. Louis) at a concentration of 1 mg per 2 ml of probe for 1 h and then applied as described above to sections containing fungi from a strawberry petiole lesion 1 mm or larger. (2) The ultrathin sections were floated on a drop of PBS for 1 h and probed with wheat germ agglutinin without a gold label for 1 h. The remainder of the probe application was as described above.

RESULTS

Light microscopy. The ontogeny of the infection processes of C. acutatum and C. fragariae on inoculated plants were similar (Table 1) at the light or electron microscopic levels. Lesions most commonly developed on stolon tips and the upper portion of the youngest petiole with fully expanded leaflets. Lesions were first observed 2 days after inoculation as lenticular brown spots that developed into the dark, sunken, and leathery brown spots that developed into the dark, sunken, and leathery brown spots that developed into the dark, sunken, and leathery.

Fig. 5. Electron micrographs of strawberry tissue inoculated with Colletotrichum spp. A and B, Early stages of invasion of petiole 116 h after inoculation with C. acutatum. C and D, Fungal cells of C. fragariae penetrating living host cells of stolon. A, Fungal cell under the host cuticle. B, Fungal cell in anticlinal host wall. C, 116 h after inoculation. D, 48 h after inoculation. c = cuticle; f = fungal cell; h = hypha; hw = host wall; m = matrix. Bar = 2 µm.
lesions typical of anthracnose that eventually girdled the stolon or petiole.

Spore germination began as early as 16 h after inoculation, with the germ tubes originating from either end of a conidium and occasionally from both ends. Germ tubes generally spanned one plant cell length or less, but occasionally grew across several cells. Appressorial formation began as early as 20 h after inoculation. Development of an appressorium began as a swelling of the germ tube tip. Appressorial development could occur on such a short germ tube as to appear sessile. Appressoria were initially unmelanized, but by 24 h they were fully pigmented (Fig. 1A). Small circular areas on the basal sides of appressoria, seen by focusing through an appressorium with the light microscope, were interpreted to be pores from which the penetration pegs developed (Fig. 1A). The typical mature appressorium for either species was globose and approximately 3 to 5 µm in diameter. Appressoria frequently developed over anticlinal host walls, but occasionally, were found over periclinal walls or near stomata. Appressoria and their basal pores were found on the surface of both healthy and diseased plant tissue.

The first signs of host cell distress appeared when the cell wall contour became distorted, and the plasmalemma pulled away from the cell wall (Fig. 1B to D). Hyphae during early stages of lesion formation were less abundant in epidermal and subepidermal cells than in cortical cells. A gradual increase of fungal hyphae in the host cuticle, epidermis, and subepidermis was observed with hyphae restricted to the apoplast in early stages of invasion and then invading dead cells (Fig. 1B to D). As the infection progressed, cell walls became increasingly distorted (Fig. 1D). In contrast to disrupted cortical cell walls, epidermal and subepidermal walls remained intact even though hyphae penetrated their walls (Fig. 2A). Invasion of the vascular tissue and acervulus

Fig. 6. Electron micrographs showing later stages of invasion. A, Stolon 72 h after inoculation with Colletotrichum fragariae. B, Petiole 116 h after inoculation with C. acutatum. A and B, Penetration of dead host cells. Some host cells were characterized by electron dense osmiophilic material associated with the tonoplast (A), while others were not (B). C and D, Early development of a stroma in stolon 116 h after inoculation with C. fragariae. C, Hypha (h) growing from the subepidermis to the epidermal layer. D, Initial stromatic fungal cells located between the cuticle and the host wall of the epidermal layer. Note the increasing degree of vacuolation (v) of fungal cells, the disruption of the tonoplast, and the presence of abundant osmiophilic material. c = cuticle; e = epidermal layer; f = fungal cell; hw = host wall; o = osmiophilic material; arrow = tonoplast. Bar = 2 µm.
formation occurred concurrently (Fig. 2A and B). The fungus was seen in vessel elements (Fig. 2B) during this last phase of the invasion process.

Acervulus formation began with the development of a stroma within the cuticle of epidermal cells. The cuticle expanded considerably to accommodate the growing acervulus (Fig. 2C). The maturing acervulus broke through the cuticle and eventually produced conidia (Fig. 2D). The acervulus expanded radially with the stroma also extending from the epidermis to the subepidermis. Acervuli were produced over a disrupted cortex, or in extreme cases, over a completely disintegrated cortex (Fig. 2D). In either case, the epidermal and some of the subepidermal tissue remained intact.

The invasion of host tissue by *C. fragariae* and *C. acutatum* was similar. However, each invasion event occurred more rapidly with *C. fragariae* than with *C. acutatum* (Table 1). Lesions were visible on petioles 3 days after inoculation with *C. fragariae*, but were not visible until 4 days after inoculation with *C. acutatum*. A similar delay in invasion events was observed between petiole and stolons following inoculation of either species. Most events were observed a day sooner on the stolons than on the petioles within species.

**Electron microscopy.** The conidial and germ tube walls comprised two layers, with the outer layer being thinner and more electron dense than the inner layer. Only the inner conidial layer was labeled by the WGA probe (Fig. 3). Upon germination, a septum was laid down bisecting the conidium (Fig. 3), and one also developed between the conidium and the germ tube. The WGA probe was observed on the inner wall layers of the germ tube and its septa. A few vesicles were present in the germ tube after maturation of the appressorium, although no distinct organelles were seen. The probe was not found in the cytoplasm of germ tubes or appressoria.

Appressoria were sometimes covered by an extracellular matrix that was somewhat amorphous with dispersed fibrillar material. The appressorial matrix was not labeled by the WGA probe (Fig. 4). The upper wall of the appressorium appeared to have two distinct layers, with the probe being restricted to the inner layer (Fig. 4). The two layers became gradually indistinct as they approached the basal side of the appressorium. The appressorial pore was defined by the appressorial wall tapering toward a central area on the basal side. The pore was usually occluded by a membrane (Fig. 4). No evidence of labeling was found on either the host cytoplasm or walls (Fig. 4).

Some hyphae invaded the cuticle (Fig. 5A), outer epidermal wall, and anticlinal walls of the epidermis (Fig. 5B). Some hyphae initially penetrated what appeared to be living host cells (Fig. 5C and D). The host plasmalemma invaginated to accommodate hyphal penetration. The interface of the host membrane and hyphal wall was frequently associated with a particular arrangement of electron dense material (Fig. 5C and D). Several degrees of disturbance and relaxation of the microfibrillar components of cell walls were observed during penetration of host cells including early penetration events (Fig. 5A). Death of fungal cells was a rare occurrence.

Several dying host cells exhibited an amorphous, osmiophilic material around the inside portion of the tonoplast (Fig. 6A, C, and D); others did not (Fig. 6B). Some cortical fungal hyphae appeared to be growing back to the epidermis in the stage prior to acervulus formation (Fig. 6C). Acervulus formation involved the development of a subcuticular stroma within and beneath the outer host epidermal cell walls (Fig. 6D). A gradual vacuolation de-

![Fig. 7.](image)
veloped in stromatic cells (Fig. 6D). The stroma continued to develop into the epidermal and adjacent subepidermal cells, finally rupturing the cuticle (Fig. 7). Late development of the acervulus included the extension of the host cuticle to accommodate the expanding stroma, the rupture of the cuticle, and early development of conidia (Fig. 7). C. acutatum and C. fragariae had a pulvinate type of acervulus (27).

Controls. There were no apparent variations in the labeling of walls of fungi grown on PDA when compared to both young and older hyphae from lesions in plants. The WGA probe was evenly distributed on the unilayered walls. Septa were labeled with the same intensity as the outer walls.

Labeling was diminished as expected when the probe was preadsorbed with chitin (control 1). Labeling was diminished as expected when the sections were incubated with lectin before applying the lectin-gold complex (control 2).

DISCUSSION

C. acutatum and C. fragariae underwent a very brief biotrophic phase (less than 12 h) before entering their extended necrotrophic phase. Representatives of both species of fungi were observed entering living host cells and developing a matrix comparable to that described in reported biotrophic systems (4,11,13,15), but they killed the host cells within a few hours. Previous descriptions for other Colletotrichum species cite biotrophy at a minimum of “less than 24 h” (14,29). But a situation in which a living cell is entered and quickly killed might be considered a modification of necrotrrophy, rather than biotrophy or hemibiotrophy.

Both intracellular hemibiotrophic and subcuticular intramural Colletotrichum species avoid triggering resistance responses during the symptomless phase (17). Species with an intracellular hemibiotrophic strategy require an intimate cytoplasmic interaction between the plant and the pathogen, while species with a subcuticular intramural strategy do not require such intimacy. Since the former species have a restricted host range, they are considered specialists, while the latter species are considered generalists (1). Regardless of the length of the biotrophic phase, all species eventually revert to necrotrophic development. Since C. acutatum is a generalist invader, only necrotrophic development was predicted, while C. fragariae is a specialist invader (22) for which an extended biotrophic phase was predicted to precede the necrotrophic phase. The brevity of the biotrophic phase in C. acutatum and C. fragariae leaves some uncertainty about calling these fungi hemibiotrophs and comparing their initial infection strategies in terms of previously described species.

Penetration pegs were observed in our studies at the light microscope level but not the ultrastructural level. Appressoria of C. fragariae observed at the ultrastructural level may have been in the process of forming a penetration peg or may have been arrested just prior to formation of a penetration peg. Latent or quiescent stages have been observed in fungal infections of fruits (30) where the mature appressorium often will not produce a peg until the fruit is at a particular stage of ripeness. This concept has not been sufficiently studied in strawberry for the fungal penetration of stolon, petioles, leaves, or roots. Strawberry petioles are often sufficiently studied in strawberry for the fungal penetration of stolon, petioles, leaves, or roots. Strawberry petioles are often sufficiently studied in strawberry for the fungal penetration of stolon, petioles, leaves, or roots. Strawberry petioles are often sufficiently studied in strawberry for the fungal penetration of stolon, petioles, leaves, or roots.

The distribution of chitin visualized by WGA gold probe in C. lindemuthianum has been reported by O’Connell and Ride (15). We found several similarities as well as some differences between their findings and our results using the same probe to study C. fragariae. The outer appressorial wall layer of C. lindemuthianum was labeled more heavily by the probe than the inner layer. By contrast, the outer appressorial wall layer of C. fragariae was labeled less heavily by the probe than the inner layer. The matrices of both C. lindemuthianum and C. fragariae were unlabeled. The young intracellular hyphae of C. lindemuthianum were unlabeled, but the invasive hyphal walls of C. fragariae and walls of the comparable secondary hyphae of C. lindemuthianum were labeled. The actively growing hyphal tips of either species labeled with less intensity than the mature walls. The outer wall layers of preinvasive structures of C. fragariae did not show chitin-gold complex, but the single wall layers of invasive hyphae from the earliest stages observed in planta contain chitin distribution comparable to young hyphae observed in vitro. C. fragariae did not appear to modify its chitin distribution during its very brief biotrophic phase as does C. lindemuthianum. This supports O’Connell and Ride’s (15) suggestion that chitin masking is necessary during initial contact to establish a biotrophic relationship.

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LITERATURE CITED


