Epidemiology and Pathology of Strawberry Anthracnose: A North American Perspective

Barbara J. Smith

U.S. Department of Agriculture, Agricultural Research Service, Thad Cochran Southern Horticultural Laboratory, Small Fruit Research Unit, P.O. Box 287, Poplarville, MS 39470

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Abstract. Three Colletotrichum species—Colletotrichum acutatum J.H. Simmonds (teleomorph Glomerella acutata J.C. Guerber & J.C. Correll), Colletotrichum fragariae A.N. Brooks, and Colletotrichum gloeosporioides (Penz.) Penz. & Sacc. in Penz. [teleomorph Glomerella cingulata (Stoneman) Spauld. & H. Schrenk]—are major pathogens of strawberry (Fragaria xananassa). Strawberry anthracnose crown rot has been a destructive disease in commercial strawberry fields in the southeastern United States since the 1930s. The causal fungus, C. fragariae, may infect all aboveground plant parts; however, the disease is most severe when the fungus infects the crown, causing crown rot, wilt, and death. Colletotrichum gloeosporioides was responsible for an epidemic of anthracnose crown rot in strawberry nurseries in Arkansas and North Carolina in the late 1970s. The anthracnose fruit rot pathogen, C. acutatum, was first reported in 1986 on strawberry in the United States. Since the 1980s, increased losses due to anthracnose fruit and crown rots in the United States may be related to changes in cultivars and to widespread use of annual plasticulture production rather than the matted-row production system. Anthracnose investigations in the United States have concentrated on its epidemiology and differences among the three causal Colletotrichum spp. in their cultural, morphological, and molecular characteristics; their infection processes; and their pathogenicity. Results from these studies have resulted in a better understanding of the diseases and have led to better disease control. Strawberries grown in soils with high nitrogen levels are more susceptible to anthracnose than are those grown in soils with lower nitrogen levels or those amended with calcium nitrate. Anthracnose is spread more rapidly in fields that have overhead irrigation and plastic mulch than in fields where drip irrigation and straw mulch are used. Fungicide efficacy has been determined in in-vitro, greenhouse, and field studies, and pathogen resistance to some fungicides has been detected. Anthracnose-resistant cultivars are a major objective of most strawberry breeding programs in the southern United States.

Colletotrichum species cause serious diseases of many fruit and vegetable crops worldwide, and three species, C. fragariae, C. acutatum, and C. gloeosporioides, cause anthracnose diseases of strawberry (Smith and Black, 1990). Since the 1930s, anthracnose crown rot caused by C. fragariae and C. gloeosporioides has been a destructive disease in strawberry nurseries and fruit-production fields in the southeastern United States (Brooks, 1931). In 1986, the presence of the anthracnose fruit rot pathogen, C. acutatum, was first reported on strawberry in the United States (Smith and Black, 1986). Increased losses due to anthracnose fruit and crown rots in the United States since the 1980s may be related to the shift from matted-row culture to the annual plasticulture production system, as well as to changes in cultivars. Anthracnose diseases are increasing in importance and resulting in major economic losses to strawberry growers worldwide. The objective of this report is to summarize anthracnose related research from the United States (excluding research from California and Florida) so that we can better understand these diseases and their control.

ANTHRACNOSE PATHOGENS

Colletotrichum fragariae, C. acutatum, and C. gloeosporioides (Howard et al., 1992; Smith, 1998a, 1988b, 1988c) cause anthracnose diseases of strawberries. C. fragariae was assumed to be the causal agent of strawberry anthracnose in the United States until 1986 when Smith and Black (1986) reported the presence of C. acutatum on strawberry in the United States. This fungus had previously been reported to cause anthracnose fruit rot of strawberry in Queensland, Australia (Simmonds, 1965); however, it had probably been present in the United States for some time under the name of Gloeosporium spp. (Maas, 1984). Colletotrichum gloeosporioides causes anthracnose crown rot, petiole lesions, and leaf spots indistinguishable from those caused by C. fragariae. In a series of comprehensive studies of these three major causal agents of strawberry anthracnose, Smith and Black (1990) examined the cultural characteristics, conidia, appressoria, and setae of 24 Colletotrichum isolates. Colletotrichum fragariae isolates developed beige to olive to dark gray colonies, did not form the ascosporogenous culture, and their conidia were typically cylindrical with one end sharply tapered and the other end rounded. Colletotrichum gloeosporioides isolates were very similar except they formed the Glomerella cingulata ascosporogenous culture in which their conidia were usually rounded on both ends. Isolates of both C. fragariae and C. gloeosporioides produced dark black setae, visible with a hand lens, in acervuli in culture and on petiole, stolon, and fruit lesions. Colletotrichum acutatum isolates produced fusiform conidia tapered on both ends; developed white, pink, orange, rose, or beige colonies; and did not form setae or the ascigerous state in culture. The growth rate of C. acutatum in culture was slower than the other two species at all temperatures tested with the greatest difference being at 32 °C. Colletotrichum fragariae, the “original” anthracnose fungus, was first identified in Florida in 1931 (Brooks, 1931). It spread throughout the southeastern United States and was responsible for crown rot and death of many plants in strawberry nurseries in the 1970s (Horn et al., 1972). It has a narrow host range, infecting only strawberry and a few weed hosts, and is rarely found outside the southeastern United States. Colletotrichum fragariae generally causes more severe petiole and crown symptoms than C. acutatum, and C. fragariae is considered by some to be a host-specific or con-specific form of C. gloeosporioides (Howard, 1983; Howard et al., 1992; Sutton, 1992).

In the late 1970s, C. gloeosporioides was identified as the causal agent on plants obtained from Arkansas and North Carolina nurseries that died from a crown rot identical to that caused by C. fragariae (Howard et al., 1992). It has a wide host and geographic range, causing diseases of many plant hosts worldwide.

The greatest economic losses due to anthracnose on strawberry are from fruit rot caused by C. acutatum, which also infects many other fruit and vegetable crops, including apples, tomatoes, peppers, peas, peaches, blueberries, blackberries, and grapes (Bernstein et al., 1995; Howard et al., 1992; Smith, 2002). The presence of the pathogen has been reported on strawberries in almost all areas of the world where they are grown. Crown infections of strawberry plants by C. acutatum often result in stunted plants rather than
plant death. Infected plants usually do not thrive after transplantation and produce few berries at harvest.

Historically, *C. acutatum* has been considered to be the anthracnose fruit-rotting pathogen, and *C. fragariae* and *C. gloeosporioides* have been associated with petiole and stolon lesions and crown rot; however, all three species may cause similar symptoms and may be found to occur on the same plant (Howard et al., 1992). Identification of these pathogens should be based on classical taxonomic characteristics or molecular techniques, not symptoms.

Results of several studies suggest that *C. acutatum* may have evolved into a subgroup that is highly virulent and host-specific to strawberries. In one of these studies (Denoyes-Rothan et al., 2002), 95 isolates of *Colletotrichum*, including 81 isolates of *C. acutatum* (62 from strawberry) and 14 isolates of *C. gloeosporioides* (13 from strawberry), were characterized by various molecular methods and pathogenicity tests. Results based on random amplified polymorphic DNA (RAPD) polymorphism and internal transcribed spacer 2 (ITS2) sequence data provided genetic evidence of two subgroups within *C. acutatum*. The first subgroup, characterized as CA-clonal, included only isolates from strawberry and exhibited identical RAPD patterns and nearly identical ITS2 sequence analyses. A larger genetic group, CA-variable, included isolates from various hosts and exhibited variable RAPD patterns and divergent ITS2 sequence analysis. On the bases of these molecular data, Denoyes-Rothan et al. (2002) proposed that the CA-clonal subgroup contained closely related, highly virulent *C. acutatum* isolates that may have developed host specialization to strawberry. Isolates of *Colletotrichum* spp. have been disseminated worldwide, probably through international plant exchanges, as their genetic polymorphism and geographical origins are not correlated (Denoyes-Rothan et al., 2002; Sreenivasaprasad and Tallinnas, 2005).

**ANTHRACNOCOE DISEASE SYMPTOMS**

Anthracnose crown rot (Smith, 1998a), caused by either *C. fragariae* or *C. gloeosporioides*, is first apparent by the wilting of the youngest leaves in the hottest part of the day. The young wilted leaves may appear to recover and become turgid in the evenings; however, most will wilt and die after a few days. Shortly after plants wilt, a red discoloration appears within the crown tissue, and the causal pathogen may be isolated from discolored tissue. After the plants have been dead for several days, the crown tissue will turn dark brown to black, and then *Colletotrichum* spp. is difficult to isolate. *Colletotrichum acutatum* also may cause crown death; however, typically a single side crown is infected rather than the entire crown, and infected plants are stunted but do not die. Each of the three *Colletotrichum* spp. may cause petiole and stolon lesions which are dark brown or black and sunken and often girdle the petiole or stolon. Pink masses of conidia are usually visible near the center of each lesion. All three species also cause leaf spots (Howard et al., 1992; Maas and Palm, 1997; Smith, 1998c). Black leaf spot, typically caused by *C. fragariae* and *C. gloeosporioides*, is characterized by gray or light black spots, usually not necrotic, peppered across the top surface of the strawberry leaflets. *C. acutatum* more typically causes irregular leaf spot, the primary symptom of which is the appearance of necrotic black lesions at the tip of the leaflets. All three *Colletotrichum* spp. also cause flower blights and fruit rots (Smith, 1998b). Fully open flowers are much more susceptible than closed buds (Smith, 1993). Infected green fruit are often hard and brown and mucromify rather than ripen. Anthracnose lesions on ripe fruit are firm, slightly sunken and covered with pink spore masses. *Colletotrichum acutatum* also causes root lesions.

**ANTHRACNOCOE INFECTION PROCESS AND PATHOGEN DISPERAL**

Curry et al. (2002) studied the infection process of strawberry petioles and stolons by *C. acutatum* and *C. fragariae* using light and electron microscopy. Both fungal species invaded the host tissue in a similar manner; however, *C. fragariae* invaded the plants more rapidly than *C. acutatum*. Both species penetrated the cuticle via an appressorium, and their hyphae grew within the cuticle and cell walls of epidermal, subepidermal, and subtending cells. They began invasion with a brief biotrophic phase, in which they invaded living cells, before entering an extended necrotrophic phase, in which they proliferated among dead cells. Acervuli formed once the cortical tissue had been moderately disrupted and developed as a stroma just beneath the outer periclinial epidermal walls. Acervuli erupted through the cuticle and released conidia. Invasion of the vascular tissue typically occurred after acervuli matured but remained minimal.

The time from infection of the strawberry by *Colletotrichum* spp. to first sporulation (the latent period) is an important factor in the speed at which anthracnose may spread within a field. The latent period depends on the temperature and ranges from 2–3 d at 25°C to 6–17 d at 5°C (King et al., 1997). At 5 and 10°C, the latent period was shorter for *C. acutatum* than for *C. gloeosporioides* and *C. fragariae*; however, at higher temperatures the latent period for all species was similar. Appressoria and secondary conidia produced by *C. acutatum* on symptomless foliage may be a significant source of inoculum for fruit infections (Leandro et al., 2001) and may also contribute to the availability of inoculum throughout the growing season (Leandro et al., 2003a). Conidial germination, appressorial production, and secondary conidiation are all favored by longer periods of wetness than the 4 h required for secondary conidia to form. *Colletotrichum acutatum* survived up to 8 weeks on leaves in greenhouse studies (Leandro et al., 2003a) and up to 5 weeks on fabric (Norman and Strandberg, 1997). More conidia formed on leaves when exposed to flower extracts than when exposed to leaf extracts or water (Leandro et al., 2003b), suggesting that *C. acutatum* inoculum levels on strawberry foliage may increase during flowering.

Rain splash is the primary means by which *Colletotrichum* spp. conidia are spread from plant to plant in the field. Madden and Boudreau (1997) found that anthracnose fruit rot incidence generally declined as plant density increased and concluded that plant density reduced the amount of rain that penetrated the plant canopy, thus reducing the amount of splash. Most fruit infection occurred in a 25-cm radius of the source of the inoculum, an infected fruit (Madden and Wilson, 1997). Nathimpera et al. (1999) studied splash dispersal of the conidia of the three *Colletotrichum* spp. and found that conidia of *C. fragariae* dispersed over the shortest distance and those of *C. acutatum* dispersed over the longest distance. This was probably due to the greater amount of spores produced on infected fruit by *C. acutatum*. *Colletotrichum acutatum* conidia may survive in soil and plant debris under dry conditions for up to 12 months, but conidia and sclerotia die rapidly under moist conditions, i.e., soil moisture ≥12% (Norman and Strandberg, 1997).

**ANTHRACNOCOE CULTURAL CONTROL MEASURES**

Because the primary source of infection in most fruiting fields appeared to be infected transplants, strawberry growers in the southeastern United States were advised after the anthracnose crown root epidemics in the 1980s to obtain their transplants from nurseries in the northern United States, Canada, or California that were believed to be outside the range of *C. fragariae*. More recently, *C. acutatum* has been found in some of the nurseries in these areas. McNelis et al. (1992a) demonstrated that anthracnose-free transplants can be produced in the southeastern United States by locating nurseries in areas where strawberries are not grown commercially. Disease-free transplants remain the primary control of anthracnose crown rot and fruit rot.

Because *Colletotrichum* spp. may infect many other hosts, primary infection in strawberry fields sometimes is assumed to come from these other hosts growing near the strawberry field. To test the hypothesis that *Colletotrichum* spp. may move from other fruit or vegetable hosts to strawberry, 37 *Colletotrichum* isolates, representing nine species collected from 12 hosts, were wound-inoculated onto the leaves and stems of strawberry, blueberry, blackberry, muscadine grape, tomato, and pepper (Smith, 2002). *Colletotrichum fragariae* isolates...
were the most aggressive and caused lesions at an average of 38% of inoculation sites on all hosts except pepper. Percentages of infection for the other species were 25% *Colletotrichum capsici*, 18% *Gloeosporiella*, 15% *C. acutatum*, 11% *C. destructivum*, 9% *C. truncatum*, 8% *C. cucodes*, 6% *C. higginsianum*, and 5% *C. orbiculare*. Strawberry was the most susceptible host with 58% of petiole and 14% of leaf inoculations of all isolates resulting in lesion development. Pepper was the most resistant host with no symptom development on leaves or stems following inoculation with any isolate. These results suggest that primary anthracnose infections in strawberry fields are most often from infected strawberry transplants and only rarely from other diseased fruit or vegetable hosts.

Anthracnose spreads within a field by splashing water, and living mulches (such as wheat, rye, or rye grass) in row middles have been shown to reduce disease spread within a field. Sublethal doses of grass-specific herbicides such as sethoxydim, may be used to prevent excessive growth of rye grass (Gupton, 2000). Organic mulches, such as wheat straw or pine needles, will also reduce splash and result in lower incidence of anthracnose compared with rows mulched with plastic (Madden, 1992; Smith and Spiers, 1986). Anthracnose is less severe when water is supplied to plants using drip irrigation rather than overhead irrigation (Madden, 1992; Smith and Spiers, 1986).

Anthracnose crown rot was observed to be less severe in commercial fields when strawberries were grown on soils with low nitrogen fertility (Howard, et al., 1992). In greenhouse studies, Smith (1987, 1989) determined that strawberries grown in soils with high levels of nitrogen, especially from ammonium sources, are more susceptible to anthracnose than plants grown in soils with lower nitrogen levels or those with high levels of calcium nitrate. Anthracnose fruit rot caused by *C. acutatum* was less severe on fruit from greenhouse-grown plants receiving drench or foliar applications of calcium sulfate than on fruit from plants receiving water, calcium chloride, or calcium nitrate treatments. Fruit from plants receiving foliar applications of CaCl₂ developed less fruit rot than that from plants receiving soil applications of CaCl₂ (Smith and Gupnon, 1993).

The primary means to reduce the buildup of anthracnose fruit rot in the field is to harvest fruit frequently and remove all rotten fruit from the field. Following severe infections early in the fruiting season, all infected fruit should be stripped from the plants and removed from the field. Infected areas of a field should be harvested last in the day, or workers should wash up and change to clean clothes when they must enter un-infected areas of the field after they have harvested areas where fruit rot is present.

**IN-VITRO FUNGICIDE STUDIES**

Until highly resistant cultivars are available, growers must rely on chemical applica-

tions and cultural practices to reduce losses due to anthracnose. Failure of fungicides to control anthracnose epidemics may be due to the development of fungicide resistance in the *Colletotrichum spp.* population. For example, benzomyl was shown to effectively reduce the incidence of anthracnose crown rot (Horn et al., 1972; Howard, 1971) and was used intensively by strawberry growers for years to control anthracnose and other diseases. However, the anthracnose pathogens, *C. acutatum* and *C. fragariae*, developed resistance to it and other benzimidazole fungicides (LaMondia, 1995; McInnes et al., 1992a; Smith and Black, 1992, 1993), and benomyl was no longer effective for anthracnose control in strawberry fields. In-vitro trials have been used to screen fungicides for their ability to control anthracnose (LaMondia, 1993; McInnes et al., 1992b; Smith and Black, 1992). Smith and Black (1993) reported that all 16 *C. acutatum*, 14 out of 18 *C. fragariae*, and both *Gloeosporiella* isolates tested in vitro were resistant to benomyl and that all benomyl-resistant isolates were also resistant to carbendazim, which is in the same class of fungicides as benomyl. In a greenhouse study, plants treated with propiconazole had lower disease severity ratings than did plants treated with captan or benomyl (Smith and Black, 1991). However, the propiconazole-treated plants were shorter with dark green leaves that appeared thicker than the leaves of untreated plants.

**FIELD FUNGICIDE TRIALS**

Sixteen different fungicide treatments were evaluated in five fungicide studies conducted at Hammond, LA, and Poplarville, MS, during the 2002, 2003, and 2005 fruiting seasons (Wedge et al., 2007). Treatments were applied at 7- to 10-d intervals to three seasons (Wedge et al., 2007). Treatments were applied at 7- to 10-d intervals to three strawberry cultivars with different susceptibilities to anthracnose. The most frequent fruit rots at harvest were anthracnose fruit rot (caused by *Colletotrichum spp.*), stem end rot (caused by *Gnomonia comari* P. Karst.,), and Botrytis gray mold (caused by *Botrytis cinerea* Pers.:Fr.). Compared with the untreated control treatment, less anthracnose fruit rot occurred on berries from the pyraclostrobin + boscalid, cyprodinil + fludioxonil, azoxystrobin, pyraclostrobin, captan + fenhexamid, and captan treatments.

**USDA-ARS ANTHRACNOSE BREEDING PROGRAM**

In the early 1980s, when anthracnose became a major disease of strawberries in the southeastern United States, a breeding program to develop anthracnose-resistant cultivars adapted to the strawberry-growing areas in the southeastern United States was initiated by the USDA-ARS at the Small Fruit Research Station, Poplarville, MS, and the Fruit Laboratory, Beltsville, MD, with the collaboration of state experiment stations in Florida, Louisiana, and North Carolina (Galletta et al., 1997; Smith and Spiers, 1982). From 1976 to 1995, over 160,000 progeny from 448 crosses made at Beltsville, MD, primarily by G.J. Galletta, were screened for anthracnose resistance in the greenhouse at Poplarville, MS. Initially, parent lines were eastern cultivars and advanced selections from the breeding program at Beltsville. As the program progressed, resistant selections from the anthracnose program were crossed with commercial cultivars to improve the horticultural characteristics of the progeny. Seed from the crosses made at Beltsville were germinated at Poplarville, and resultant seedlings were inoculated with a conidial suspension of *C. fragariae*, incubated in a dew chamber for 48 h, moved to a warm greenhouse, and rated for anthracnose severity 30 d after inoculation. Resistant seedlings were evaluated in the field in Florida, Louisiana, Mississippi, Maryland, and North Carolina; selections were made on the bases of yield, fruit quality, plant habit, and resistance to leaf scorch [caused by *Diplocarpon earlium* (Ellis & Everh.) F.A. Wolf], leaf spot [caused by *Myosphaerella fragariae* (Tul.) Lindau], powdery mildew [caused by *Sphaerotheca macularis* (Wallr.:Fr.) Jacz. f. *fragariae* Peries], and two-spotted spider mites (*Tetranychus urticae* Koch).

Fifteen hundred fifteen (1515) anthracnose-resistant selections were made from the seedlings field-tested in Mississippi. Four of these anthracnose-resistant strawberry clones were released as breeding lines (Galletta et al., 1993) and have been used as the source of anthracnose resistance in several breeding programs. Smith et al. (1998) released the cultivar Pelican, which is highly resistant to both anthracnose crown rot caused by *C. fragariae* and anthracnose fruit rot caused by *C. acutatum*. ‘Pelican’ is also resistant to five races of red stele caused by *Phytophthora fragariae*. Evaluation of strawberry seedlings and advanced breeding lines from state and private breeding programs is an ongoing project. The USDA anthracnose-resistance screening procedure has effectively identified resistant genotypes in seedling progenies from the North Carolina State University breeding program, with >32,000 resistant strawberry seedlings identified between 1998 and 1999 (Ballington et al., 2002).

Smith and Black (1987) found that resistance to *C. fragariae* was influenced by environmental conditions after inoculation and that plants incubated at a high temperature (35°C) for 48 h in a dew chamber (relative humidity near 100%) had higher disease severity ratings when compared with plants incubated at 25 or 30°C. Plants maintained in a greenhouse at 32°C after dew chamber incubation developed more severe symptoms than did those held in a greenhouse at 25°C. Two- to 4-week-old strawberry seedlings (age after transplanting at the first true-leaf stage) were more susceptible to *C. fragariae* than 14- to 18-week-old seedlings when spray inoculated with a conidial suspension (Smith et al., 1990). Tissue culture-induced (somaclonal) variation is another strategy being pursued.
for generating disease-resistant genotypes. Hammerschlag et al. (2006) screened shoots regenerated from leaf explants of six commercially important cultivars for resistance to a virulent isolate of *C. acutatum*. Somaclones with higher levels of anthracnose resistance were identified for all the cultivars, and the greatest increases in disease resistance were observed for somaclones of Chandler, Peli-can, and Sweet Charlie, where resistance increased 12-fold. These studies provide evidence that in-vitro screening can be used to evaluate strawberry germplasm for anthracnose resistance and that somaclonal variation is influenced by strawberry genotype.

**INHERITANCE OF RESISTANCE TO ANTHRACNOSE**

Gupton and Smith (1991) determined that the estimates of dominance genetic variance were six to 10 times higher than those for additive genetic variance. The frequency distribution of disease severity ratings was bimodal suggesting major gene action. Narrow-sense heritability estimates of 0.37 and 0.26 were probably sufficient to produce gains because of the high (0.87 and 0.85) broad-sense heritability estimates. Giménez and Ballington (2002) also found that nonadditive effects were more important than additive effects in the inheritance of resistance on runners. The epistatic nature of anthracnose resistance on runners appears to be supported by results of crosses between susceptible parents that result in up to 20% resistant seedlings (Ballington et al., 2002). Giménez and Ballington (2002) found that the frequency distribution of lesion lengths within progenies suggests that resistance to *C. acutatum* on runners is quantitative. Results of Garcés et al. (2002) also support the theory that inheritance of resistance to anthracnose is quantitative in nature.

Increased losses due to anthracnose fruit and crown rots may be related to the shift from matted-row culture to the annual plasticulture production system in the United States. Breeding for genetic resistance to anthracnose and development of resistant cultivars are primary means for reducing economic losses due to this disease. This is environmentally sound because it results in reduced use of fungicides.

**CONCLUSIONS**

As our knowledge of the anthracnose pathogens and the epidemiology of anthracnose diseases has increased, so has our ability to control these diseases. Changes in cultural practices have resulted in reduced levels of disease. At the same time, development of more effective fungicides and their registration for use on strawberries have greatly reduced losses due to both anthracnose crown rot and fruit rot. Anthracnose-resistant cultivars also have reduced economic losses due to these diseases. Even so, growers may sustain severe losses when environmental factors are highly favorable for anthracnose development.

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