Environmental Factors Affecting the Release and Dispersal of Ascospores of *Mycosphaerella citri*  

S. N. Mondal, T. R. Gottwald, and L. W. Timmer

First and third authors: University of Florida, Citrus Research and Education Center, and Department of Plant Pathology, 700 Experiment Station Road, Lake Alfred 33850; and second author: U.S. Department of Agriculture, Agricultural Research Service, Horticultural Research Laboratory, Ft. Pierce, FL 34945. 

Accepted for publication 21 March 2003.

**ABSTRACT**


Greasy spot, caused by *Mycosphaerella citri*, produces a leaf spot disease affecting all citrus species in Florida and the Caribbean Basin. *M. citri* produces pseudothecia and ascospores, which are considered the principal source of inoculum, in decomposing leaves on the grove floor. In studies using a computer-controlled environmental chamber, a single rain event triggered release of most mature ascospores beginning 30 to 60 min after the rain event. Additional rain events did not bring about further release. High relative humidity without rain triggered release of low numbers of ascospores, but vibration and red/infrared irradiation had little or no effect on ascospore release. After three to four cycles of wetting and drying of leaves, all pseudothecia had matured and released their ascospores. In the field, ascospores were detectable starting about 2 h after the beginning of a rain or irrigation and most ascospores were released within 16 h. Ascospore release was greatest following rain events and somewhat less following irrigations, and low numbers of ascospores were detectable on days without precipitation. Ascospore numbers declined linearly with horizontal distance from the source and as a function of the logarithm of ascospore numbers with vertical distance. Low numbers of ascospores were detected 7.5 m above the ground and 90 m downwind from the grove. Ascospore release can be advanced by irrigating frequently during dry, nonconducive conditions to stimulate ascospore release when environmental conditions are unfavorable for infection, but the eventual effects on disease severity are uncertain.

Greasy spot, caused by *Mycosphaerella citri* Whiteside, is a leaf spot disease that produces premature leaf drop accompanied by reduced tree vigor, yield, and fruit size (22). *M. citri* also infects fruit causing greasy spot rind blotch, which reduces the acceptability of fruit for the fresh market. The disease is most severe on grapefruit and its hybrids, but affects all citrus. It is prevalent in Florida and most of the Caribbean Basin, and other greasy spot-like diseases occur elsewhere (8,23).

*M. citri* is a loculoascomycete fungus, and ascospores are produced in pseudothecia in decomposing leaf litter on the grove floor. Most of the leaf drop in Florida citrus groves occurs in January and February. Whiteside (26) demonstrated that alternating wetting and drying was needed for pseudothecial development in dead, infected leaves. Mondal and Timmer (18) found that wetting infected leaves three times a week for 30 to 60 min resulted in maximal production of pseudothecia. The optimum temperature for pseudothecial production was 28°C. Once pseudothecia and ascospores are mature, wetting the leaves triggers the release of ascospores. Most of the ascospores are released in a few hours after the leaves are wetted (7). In Florida, peak ascospore release under grove conditions previously occurred in June and July at the beginning of the rainy season (26). Spore release now peaks in April-May (21,25) because microsprinkler irrigation moistens leaf litter two to three times per week and accelerates leaf decomposition (25). Peak release in Costa Rica occurs in June shortly after the rainy season begins (8). In Texas, where rains begin later, peak release occurs in July-August (24). Ascospores are deposited on the leaf surface where they germinate and mycelium grows epiphytically. In Florida, the epiphytic growth begins to develop in late June or early July and continues through September (17). Hyphal tips penetrate the leaves through stomata. *M. citri* grows slowly in the mesophyll and symptoms of chlorosis and necrosis develop after several months, usually in November-December, in Florida.

The effects of environmental factors on the release and dispersal of ascospores of *Venturia inaequalis*, *V. pirina*, *Apiospora morbosa*, and *Anisogropa anomala* that cause apple scab, pear scab, black knot of cherry, and eastern filbert blight, respectively, have been extensively studied (3,6,10,12,14,20). Other than the fact that ascospore release follows the wetting of decaying leaves with mature pseudothecia of *M. citri* (23,26), little is known about the factors that trigger release, the duration of ascospore production from leaves, and dispersal of ascospores.

The purpose of this study was to determine the effect of rainfall events, relative humidity (RH), red/infrared (R/IR) radiation, and vibration on the release of ascospores of *M. citri* under laboratory conditions. Under field conditions, the effect of rain events and irrigations on ascospore release, as well as the vertical and horizontal dispersal of ascospores was investigated.

**MATERIALS AND METHODS**

Computer-controlled environmental chamber studies, General. Mature leaves with symptoms of greasy spot were collected from grapefruit (*Citrus paradisi* Macf.) trees that had not been treated with fungicide. Leaves were air dried in the laboratory at ambient temperature 23 to 27°C and stored in paper bags until used. Dried leaves were wetted for 10 to 30 min per day, 5 days per week at 28°C, and dried between wettings to induce formation of pseudothecia (18). After pseudothecial development, about 10 to 15 pseudothecia were dissected periodically from the leaf tissue, squash-mounted in lactophenol-acid cotton blue (5), and examined microscopically to determine pseudothecial maturity. Pseudo-
thecia matured in about 40 days. Leaves selected for ascospore release studies had pseudothecia distributed over more than half of the leaf surface and more than 50% of the pseudothecia matured with no or few pseudothecia that had already ejected ascospores.

A computer-controlled environmental chamber (CCEC), designed and described by Gottwald et al. (7), was used to investigate the effects of environmental factors affecting release of ascospores. The device was used to control, manipulate, and monitor rainfall, temperature, RH, R/IR radiation, leaf wetness, vibration, and air flow (wind speed). For each experiment, four leaves were placed in the sample holder and subjected to the treatments of the above variables as described for each experiment. Spores released from the samples were trapped on a rotating spore trap to allow correspondence of spore release with environment variables and their changes. The spore trap was calibrated to rotate at a constant rate, functioning on the same principle as a Burkard volumetric spore trap (Burkard Scientific Sales, Ltd., Rickmansworth, Hertfordshire, UK), and used the same trapping surfaces prepared in the manner as described below.

Effect of rain. Three experiments were conducted to determine the effect of a single rain event on ascospore release. Because the chamber can flood with excess rain, prolonged simulated rains were not tested. A single rain event was composed of five rains of 5 s each at 1-min intervals. Leaves were exposed to a single rain event at time 0, and ascospores were collected for 180 min. The temperature was maintained at 23 to 24°C, and RH was held at 95 to 100% for the first 80 min and then reduced to 20 to 40% for the duration of the experiment. The number of ascospores trapped during each 30-min period was averaged for the three experiments, transformed to log10, and the data were regressed against time. The first 30-min period was omitted from the regression analysis because no ascospores were released.

Three experiments were conducted to determine the effect of multiple rains on ascospore release. Leaf samples were exposed to six rain events, as defined previously, at about 1-h intervals, and ascospores were collected for 360 min. The RH was held at 95 to 100% for the first 30 min following each rain event and then reduced to 50% until the next rain event. Temperature was maintained at 23 to 25°C. The number of ascospores collected for 180 min during and after each rain event was averaged across the

Fig. 1. Release of ascospores from pseudothecia of *Mycosphaerella citri* in response to a single rain event in a computer-controlled environmental chamber, under the relative humidity (RH), leaf wetness (LW), and temperature (Temp) described.

Fig. 2. Release of ascospores from pseudothecia of *Mycosphaerella citri* in response to multiple rain events in a computer-controlled environmental chamber, under the relative humidity (RH), leaf wetness (LW), and temperature (Temp) described.

Fig. 3. Effect of successive wetting and drying of leaves on the release of ascospores from pseudothecia of *Mycosphaerella citri* in a computer-controlled environmental chamber. Leaves were dried under laboratory conditions between exposure periods, under the relative humidity (RH), leaf wetness (LW), and temperature (Temp) described.
three experiments. The average number of ascospores was transformed to log10 and was regressed against the number of the rain event, i.e., 1 to 6.

**Successive wetting and drying.** Three experiments were conducted to determine the effect of repeated wetting and drying on ascospore release. In each experiment, a leaf sample was placed in the chamber, exposed to a single rain event, and then ascospores were captured for 180 min on day 1. The RH was maintained at 95 to 100% following the rain events and the temperature was 23 to 24°C. Then, the sample was dried under laboratory conditions at 23 to 27°C and about 40 to 70% RH, and the process was repeated on days 3, 5, and 7. The number of ascospores trapped each time was averaged for the three experiments, transformed to the log10, and regressed against the day of exposure to rain.

**RH.** Two experiments were conducted to determine the effect of constant high RH on ascospore release. Leaf samples were exposed at 95 to 100% RH at 25 to 27°C and the ascospores released were collected for 360 min. Means were calculated for each 30-min period, averaged across the two experiments, transformed to log10, and regressed against time for the constant RH experiments.

**R/IR irradiation and vibration.** The effect of a single R/IR exposure in the absence of vibration was evaluated in two experiments. The leaf sample was exposed to a single rain at time 0. The sample was exposed to R/IR during the first 15 min of the experiment. The RH was maintained at 80 to 100% and the temperature at 22 to 23°C. The effects of R/IR irradiation and of vibration to simulate wind movement of leaves were determined in two additional experiments. After a rain event at time 0, samples were exposed to two periods of R/IR irradiation at 60 and 120 min, each 15 min in duration. Momentary vibrations were imposed at 30, 60, 90, and 120 min. The length of the experiments was 180 min, during which time the RH was maintained at 95 to 100% and the temperature at about 24°C.

**Field studies. Rain and irrigation effects.** The effect of rain events and microsprinkler irrigations on the release of ascospores was determined with a Burkard volumetric spore sampler (Burkard Scientific Sales). The sampler was placed in a grove of mixed citrus at the Citrus Research and Education Center in Lake Alfred, FL. Coatings for the spore trap tape, handling of tapes, and counting of ascospores were as described previously (25). The spore sampler was operated for 24 h following rain events on 17 to 18 May, 27 to 28 May, and 7 to 8 June 2002 to determine the release pattern of ascospores in response to wetting under field conditions. The spore sampler was operated continuously from 1 May to 15 June 2000, 15 April to 15 June 2001, and 15 April to 10 June 2002, during the peak ascospore release period to determine the effect of irrigations and rain events. Rainfall data were collected from an automated weather station located within 500 m of the grove. Dates of irrigations and the amount of water applied were recorded during the above periods. Analysis of variance was used to compare the log10 of the number of ascospores released on days with rain events or irrigations with those with no precipitation.

**Dispersal.** Experiments were conducted in 2002 to determine the horizontal and vertical distribution of ascospores adjacent to a mixed planting of citrus species of about 5 ha from Lake Alfred. Four Burkard spore samplers were operated continuously from 29 April to 10 June 2002 at 0.5, 30, 60, and 90 m downwind from the grove edge. Total ascospores captured each week were square-root transformed and regressed against distance. Four weekly samplings were conducted.

Vertical dispersal was determined by mounting AGI-30 liquid impinger air samplers (Ace Glass, Inc., Vineland, NJ) on rotating platforms so the orifice was always oriented into the wind (20). The samplers were connected to a manifold and then to a vacuum pump using 10-mm-diameter tygon tubing. Each sampler drew air at 10 liters per minute. One spore sampler was located at 0.5, 2.5, 5.0, and 7.5 m above the ground on a tower 5 m from the edge of the grove. Sampling was conducted during and following rain events on 27 May, 29 May, and 31 June 2002. A 50% glycerol solution (10 ml) was placed in each sampler, the vacuum pump was operated for about 5 h, and the liquid was collected. The number of ascospores per 10-ml sample was determined by a hemacytometer. Nonlinear regression analysis was used to relate ascospore number and height above the ground.
RESULTS

Computer-controlled environment chamber studies. Effect of rain. A single rain event usually triggered release of large numbers of ascospores about 30 min later as shown in a typical experiment (Fig. 1). Peaks usually occurred within the first 60 min, after which release declined with occasional small peaks. Across all three experiments, 57% of the ascospores were captured in the first hour following a rainfall event. The number of spores trapped was negatively related to time after the rainfall event ($R^2 = 0.84$, $P = 0.03$).

When leaf samples were exposed to rain events at hourly intervals, the release pattern did not change appreciably compared with a single rain event. A typical experiment is shown in Figure 2. Over all three experiments, 46, 35, 10, 5, 3, and 1% of the total ascospores were captured following the first to sixth rain events, respectively. The number of ascospores released was negatively related to the number of the rainfall event ($R^2 = 0.82$, $P = 0.01$).

Successive wetting and drying. When leaves were wetted to release the ascospores and then dried and rewetted on days 1, 3, 5, and 7, the majority of ascospores were released on the first day with successively fewer released after repeated wettings (Fig. 3). Averaged across the two experiments, 60, 18, 14, and 8% were recovered on days 1, 3, 5, and 7. The numbers released were negatively correlated with the day of exposure to a rain event ($R^2 = 0.91$, $P = 0.04$). In one of the three experiments, substantial numbers were observed on day 5, indicating that more pseudothecia had matured.

Relative humidity. Low numbers of ascospores were released from leaves with pseudothecia exposed to 95 to 100% RH for 360 min (Fig. 4). Low numbers were released early in the exposure period with a peak at about 200 to 220 min. The number of ascospores released was positively related to the time of exposure to high humidity ($R^2 = 0.81$, $P = 0.01$) if the first hour of the experiment was omitted.

R/IR irradiation and vibration. Irradiation with R/IR did not have obvious effects on the pattern of spore release. When applied during and after the initial rain event, a small spike did occur in the middle of the period (Fig. 5), but that was not observed in a second experiment. The slope of the regression line for ascospore release with time in Figure 5 was significantly different from that in Figure 1 when the two were compared by the $t$ test ($t = 3.6$, $P < 0.05$).

Fig. 6. Release of ascospores from pseudothecia of *Mycosphaerella citri* in response to a single rain event and two exposures to red/infrared or to momentary vibrations (V) in a computer-controlled environmental chamber, under the relative humidity (RH), leaf wetness (LW), and temperature (Temp) described.

Fig. 7. Pattern of ascospore release following rain events on three dates in 2002, as determined by a Burkard volumetric spore sampler located in a grove of mixed citrus near Lake Alfred, FL.

Fig. 8. Relationship of rainfall and irrigation events to ascospore release of *Mycosphaerella citri*, as determined by a Burkard volumetric spore sampler located in a grove of mixed citrus near Lake Alfred, FL.
The IR irradiation may have slightly delayed the release of ascospores (Fig. 5). When leaves were exposed an hour or two after the rain event, no obvious change occurred in the course of ascospore release (Fig. 6). When the slope of the regression line for ascospore release was compared with that for data in Figure 1, there was no significant difference ($t = 1.25, P \geq 0.05$).

In two experiments, vibrations were applied to simulate wind movement of leaves. In no case was there any apparent deviation in the pattern of ascospore release due to vibration (Fig. 6).

**Field studies. Rainfall and irrigation effects.** Ascospore release was detected 1 to 2 h after a rain began in the three events that were followed (Fig. 7). Total release was less by 7 to 8 June because many of the spores had already been released. Peak release occurred 6 to 8 h after the rain began and continued for 12 to 16 h. No differences were noted in the ascospore release pattern during daytime and nighttime hours (data not shown).

Wetting of the leaf litter on the floor of the grove by rainfall or microsprinkler irrigation induced release of ascospores (Fig. 8). Only low numbers of ascospores were released on days with no irrigation or rainfall. The average number of ascospores released was substantially higher on days with rain than on days with irrigation in all 3 years (Table 1). Ten percent or fewer of ascospores were released on days with no rain or irrigation.

**Ascospore dispersal.** The total precipitation during the three trapping periods investigated for vertical distribution ranged from 3.4 to 9.1 mm, and wind speed varied from 1.6 to 2.3 m s$^{-1}$. Low numbers of ascospores were captured at 7.5 m, the greatest height tested. Ascospore capture was strongly negatively related to trapping height on all three sample days (Fig. 9A) and best fit a negative logarithmic pattern.

During the four 1-week periods when horizontal distribution was studied, the precipitation ranged from 0.9 to 54.5 mm per week and wind speeds averaged 1.3 to 3.4 m s$^{-1}$. Wind direction was $187 \pm 67^\circ$ for the line of spore traps that were aligned straight west of the grove. The number of ascospores captured was greatest from 13 to 20 May when the number of mature pseudothecia was the highest. The number of ascospores captured declined linearly in distance from the grove, and the slopes were negative ($-0.0033$ to $-0.0062$) in all 4 weeks, and all were significantly different from 0 ($P \leq 0.05$).

**DISCUSSION**

Exposure of leaf litter bearing mature pseudothecia of *M. citri* to free moisture triggered release of high numbers of ascospores. Many ascomycotina, including *V. inaequalis* (9,12), *V. pirina* (10), *Anisogramma anomala* (20), *Guignardia citricarpa* (15), *M. fijiensis* (4), and *Apiosporina morbosa* (14), behave similarly. Initial ascospore release by *M. citri* occurred about 30 min after a simulated rain event in laboratory studies and was detectable after 2 h in the field in our studies. This difference probably represents the time required for pseudothecia and asci to be fully hydrated under field conditions, to swell and force ascii through the ostiole where ascospores are forcibly ejected (1). Hydration under field conditions is probably not as uniform as in the CCEC because it is dependent on the amount of rain, canopy density, and depth of the leaf litter. Detection of ascospores in the field may also reflect the time required for sufficient spore release to occur to provide populations large enough for detection.

In our studies, a single rain event was sufficient to stimulate release of all of the ascospores from mature pseudothecia. Additional rain events did not trigger further release, such as that observed with *V. inaequalis* (7). High RH alone was sufficient to trigger release of low numbers of ascospores of *M. citri* in the CCEC as occurs with *V. inaequalis* (9,16). Ascospores were detectable in the field even on days with no rain or irrigation, suggesting that dew triggers ascospore release under grove conditions. There was no indication that vibration increased ascospore release. RIR radiation increases release of ascospores of *V. inaequalis* (3,12) and *V. pirina* (10), and there is a strong diurnal effect on ascospore release with these species. There were only minor effects of vibration and irradiation using the CCEC, and there was no indication of diurnal effects in ascospore recovery in the field.

**Fig. 9. A, Relationship between number of ascospores captured per liquid-impinger sampler and height above ground. Equations for the best fit lines were May 27 – $y = 250.2 – 108.7 \ln x$; May 30 – $y = 322.4 – 137.6 \ln x$; and June 8 – $y = 273.6 – 132.4 \ln x$. B, Relationship between total ascospores captured per week by a Burkard volumetric spore sampler located in a grove of mixed citrus near Lake Alfred, FL, and horizontal distance for four 1-week periods in 2002. In both cases, ascospore counts were square-root transformed prior to regression analysis. ** = significant at $P \leq 0.01$; * = significant at $P \leq 0.10$.**

<table>
<thead>
<tr>
<th>Class</th>
<th>2000 No. of days</th>
<th>Total precip. (mm)</th>
<th>Ascospores$^a$</th>
<th>2001 No. of days</th>
<th>Total precip. (mm)</th>
<th>Ascospores</th>
<th>2002 No. of days</th>
<th>Total precip. (mm)</th>
<th>Ascospores</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rain</td>
<td>7</td>
<td>48</td>
<td>750 ± 295 (66)</td>
<td>14</td>
<td>46</td>
<td>556 ± 115 (68)</td>
<td>11</td>
<td>139</td>
<td>705 ± 164 (66)</td>
</tr>
<tr>
<td>Irrigation</td>
<td>9</td>
<td>153</td>
<td>329 ± 81 (29)</td>
<td>7</td>
<td>119</td>
<td>201 ± 32 (25)</td>
<td>6</td>
<td>102</td>
<td>299 ± 93 (28)</td>
</tr>
<tr>
<td>No water$^b$</td>
<td>24</td>
<td>0</td>
<td>63 ± 9 (5)</td>
<td>42</td>
<td>0</td>
<td>55 ± 10 (7)</td>
<td>39</td>
<td>0</td>
<td>57 ± 6 (6)</td>
</tr>
</tbody>
</table>

$^a$ Ascospore captured per day ±SE of the mean in a continuously operating Burkard volumetric spore trap (Burkard Scientific Sales, Ltd., Rickmansworth, Hertfordshire, UK). Numbers in parentheses are the percentage of the total ascospores captured.

$^b$ Days with no rain or irrigation.

TABLE 1. Release of ascospores of *Mycosphaerella citri* from a grove of mixed fruit in Lake Alfred, FL, on days with rain, irrigation, or no water during peak release periods in 2000, 2001, and 2002
Rain stimulated release of more ascospores of *M. citri* than irradiation. This was probably attributable to the fact that irradiation wets only a portion of the surface area and thus of the leaf litter, whereas sufficient rainfall wets all of the litter. Irrigation was continued for about 3 h in most cases and was sufficient to thoroughly wet leaf litter as were most rainfall events. By irrigating during dry periods, it may be possible to induce ascospore release during periods unfavorable for infection and reduce disease. Ascospores of *M. citri* are fragile and short-lived and would likely die prior to infection under unfavorable conditions (28). However, epiphytic growth seems to be produced in abundance from the few ascospores produced in the summer rainy season (3). Because considerable infection can occur from relatively few ascospores (17), treatments and cultural manipulations designed to reduce inoculum or cause inoculum to be released when infection is not possible (19, 27) may not be highly effective for this disease.

The life cycles of many ascomycotina appear adapted to the phenology of the host plant. Ascospore production of *V. inaequalis* and *Anisogramma anomala* peaks when the host is in the most susceptible stage (6, 9, 13, 20). In the case of *Monilinia vaccinii-corymbosi* on blueberries, apothecium development of the pathogen has adapted to the phenology of individual cultivars of the host (11). Thus, with these temperate crops, pathogens tend to have life cycles closely adapted to the phenology of the host. However, we saw no evidence of this with *M. citri*. First, leaves of citrus appear to be susceptible even after they are mature (22), and conditions are favorable for infection and disease development during most of the year. The majority of the citrus leaves fall in February-March in Florida and decompose and produce pseudothecia over the next 2 to 3 months (18, 23, 25). Ascospore production peaks in April and May, but some ascospores are produced year round (21, 25). Thus, ascomycete pathogens of tropical and subtropical crops, such as banana (4) and citrus, tend to have overlapping cycles throughout the year (25).

Ascospores of *M. citri* are small and can be dispersed at least 80 m and probably much further, as are those of *V. inaequalis* (2, 13) and *Anisogramma anomala* (20). They are subject to desiccation and are fragile (28). However, if winds are strong and RH high, they could be quickly dispersed and establish new infections at some distance. Because it appears that relatively few ascospores are necessary to establish infections (17), significant disease may result from the few ascospores reaching the leaf from considerable distance. Thus, plot size for any experiments designed to reduce inoculum production would need to be large. Nevertheless, the greater severity of greasy spot on leaves on the lower canopy suggests that most of the infections result from ascospores dispersed short distances from the grove floor.

*M. citri* behaves much in the same manner as many other ascomycetes in that thorough wetting of leaf litter stimulates release of mature ascospores that are dispersed long distances vertically and horizontally by wind. However, ascospore release by this pathogen is not responsive to light as are some other ascomycetes. The host is susceptible throughout the year and the phenology of the pathogen is not synchronized with that of the host.

**ACKNOWLEDGMENTS**

This research was supported by the Florida Agricultural Experiment Station and a grant from the Florida Citrus Production Research Advisory Council, projects 991-36P and 013-16P, and approved for publication as Journal Series No. R-09225. We thank L. Zhang, T. Riley, and E. Taylor for technical assistance.

**LITERATURE CITED**