Effect of Temperature on Sporulation of *Botryosphaeria dothidea*, *B. obtusa*, and *B. rhodina*

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**ABSTRACT**


Three *Botryosphaeria* spp. were grown on autoclaved apple and peach stems in cotton-plugged tubes with constant moisture at 6, 12, 18, 24, and 30°C to determine the effect of temperature on sporulation. Number of conidia per pycnidium was determined weekly from 4 to 10 weeks after inoculation. The experiment was repeated three times. Maximum sporulation occurred at 24°C with *B. dothidea* and at 18 and 24°C with *B. obtusa*. Spore production of both fungi showed a quadratic curvilinear response to temperature. Pycnidia were erumpent, typical of their habit in nature. Maximum sporulation of *B. rhodina* occurred at 12, 24, and 30°C instead of at a distinctive peak. Of the three fungi, *B. rhodina* produced the greatest number of conidia per pycnidium at all temperatures. Mycelia and pycnidia of *B. rhodina* grew on top of the bark, which is atypical of their habit in nature. For spore production by *B. dothidea*, there was a significant interaction between temperature and time. Maximum sporulation over the 10-week period occurred in week 4 and/or 6 for *B. dothidea* at 12, 18, and 24°C, with a linear response at 12 and 24°C ($P \leq 0.05$). Conidial maturation of *B. obtusa* and *B. rhodina* had a quadratic curvilinear response due to temperature, with a maximum maturation at 12, 18, and 24°C with *B. obtusa* and at 24°C with *B. rhodina*. Spore maturation would affect longevity of conidial viability. Maximum spore production over time and percent pigmented spores over time by *B. obtusa*, and spore maturation over time by *B. rhodina* occurred in weeks 8, 9, and 10 with a significant linear response ($P \leq 0.05$). All three *Botryosphaeria* spp. produced conidia over the 6 to 30°C range and over the 7-week period (weeks 4 to 10), with maximum sporulation or spore maturation at 18 to 24°C.

Additional keywords: black rot, bot canker, dieback

**Botryosphaeria dothidea** (Moug.:Fr.) Ces. & de Not, *B. obtusa* (Schwein.) Shoemaker, and *B. rhodina* (Cooke) Arx are causative agents of dieback on many fruit, nut, and ornamental woody plants. Ascospores are considered to be a large percentage of the primary inoculum in cooler areas of the United States, such as the Upper Piedmont and mountains of North Carolina and further north, but a small percentage of the primary inoculum in most of the southeastern United States, including the mountains of Georgia (3,5,9,19). In the southern areas, conidia serve as primary and secondary inoculum of all three pathogens (5).

Temperature ranges for conidial germination and mycelial growth are similar but not identical for *B. dothidea*, *B. obtusa*, and *B. rhodina*. Conidial germination for *B. dothidea* occurred from 12 to 39°C, with the peak from 24 to 36°C, and no germination at 9°C in one study (14) and from 8 to 32°C, with the peak from 20 to 28°C, and no germination at 4°C in another study (20). Conidial germination for *B. obtusa* occurred from 12 to 32°C, with the peak from 16 to 28°C, and no germination at 8°C in one study (1), from 8 to 36°C, with the peak from 16 to 32°C, and no germination at 4°C in a second study (7), and from 12 to 32°C, with the peak from 27 to 32°C, and no germination at 8°C in a third study (8). The temperature range for germination has not been determined for *B. rhodina*. Mycelial growth for *B. dothidea* has been reported to occur from 10 to 35°C, with a peak from 20 to 32°C, and no growth at 0, 5, and 40°C (14,18,24). Mycelial growth for *B. obtusa* occurred from 8 to 36°C, with a peak from 20 to 26°C, and no growth at 4°C (7,8). Voorhees (22) found that mycelial growth for *B. rhodina* occurred from 15 to 35°C, with a peak from 25 to 35°C, and little to no growth at 10 and 40°C.

The differences in seasonal ranges for conidial production among these *Botryosphaeria* spp. are greater than the differences in the in vitro germination and mycelial growth, Weaver (23,24) recovered *B. dothidea* conidia from 22 March to 30 December, with peak number of conidia collected in late July and early August. Creswell and Milholland (6) recovered high numbers of conidia of *B. dothidea* from blighted blueberry stems from June through August and low numbers from February through May. Some researchers have found viable pycnidia of *B. dothidea* produced throughout the year on dead stems of elm (11), pistachio (12,13), and peach (15). A similar but slightly earlier summer trend has been documented for *B. obtusa*, with high numbers of conidia recovered from mid-March through August from apple twigs killed by fire blight in North Carolina (2). Yet in Georgia, conidial production during winter and spring months was attributed to *B. obtusa* (3,4,21). Pycnidia of *B. obtusa* found on dead apple stems contained mature conidia in January and February, with 90% of the pycnidia empty by the end of April (21). *B. obtusa* conidia were isolated from apple buds from October through the silver tip stage in March and from peach buds from August to March, with peak recovery in January and February (3,4). Conidia of *B. rhodina* were recovered from peach trees starting in mid-March, with peak recovery occurring sporadically from early June to late July, and limited or no recovery during winter months (4,16). *B. rhodina* is the least common of the three species isolated from diseased peach stems.

Small geographic differences in pathogen and fruit tree phenology can be important in developing and timing control measures. Sporulation has been predominately studied in vivo, often in the field using spore trapping techniques (15). The objective of this study was to measure in vitro sporulation for *B. dothidea*, *B. obtusa*, and *B. rhodina* from 6 to 32°C at 6-degree increments and determine if sporulation was affected by nutrient differences between apple and peach stems as the substrate.

**MATERIALS AND METHODS**

**Experiment design.** The experiment was conducted in incubators to test the effect of temperature (6, 12, 18, 24, and 30°C) on conidial production. The experiment was conducted three times. A Latin square design pattern with two squares allowed testing for the effect from individual incubators across the three repetitions of the experiment, hereon referred to as periods. One Latin square consisted of 6,
12, and 18°C, and the second Latin square consisted of 12, 24, and 30°C. Within a Latin square, each temperature was tested in each of the three incubators over the three periods. One temperature (12°C) was repeated in both Latin squares to test significance between Latin squares, with the possibility of combining squares into one data set. Within one period, all incubator tests of both Latin squares were run concurrently. A period lasted 10 weeks from the time inoculated stems were placed in incubators until the final sampling.

**Inoculation and stem culture maintenance.** Straight, 4- to 7-mm-diameter water sprouts were obtained from apple and peach trees. The stems were cut into segments 1 cm long, and a single stem segment was placed in a 12 × 75 mm Kimax disposable borosilicate glass tube. The tubes were plugged with nonabsorbent cotton and autoclaved at 121°C for 1 h.

Isolates of *B. dothidea*, *B. obtusa*, and *B. rhodina* were obtained from apple and peach orchards prior to each period and grown on potato dextrose agar for 14 to 21 days. Cultures were covered with 0.01% Tween 20, and pycnidia were scraped free with a rubber policeman for *B. dothidea* and *B. rhodina* and an aluminum spatula for *B. obtusa*. Pycnidia were crushed with a ceramic mortar and pestle and filtered through four layers of sterile cheesecloth. Conidial concentrations were determined with a hemacytometer and adjusted to 4 × 10⁴ conidia per ml. Under a laminar air flow hood, 0.2 ml of conidial suspension was applied to each stem segment with a hypodermic needle.

A tray contained three tubes (replications) of every fungus–host combination for a total of 18 tubes, which were completely randomized in two rows in the tray. Trays were constructed of cardboard and covered with aluminum foil with the shiny side up. Glass tubes containing stem segments were set at opposing angles of 25° with the bases of tubes touching in the middle of the tray. Stem cultures were 10 to 12.5 cm below two fluorescent and two 25-watt incandescent bulbs, which provided approximately 75 to 45 µE·m⁻²·s⁻¹ of intensity from the middle to the corner of the incubator, respectively. Seven trays of stem cultures, representing 7 weeks of sampling, were completely randomized in two rows in the incubator at zero time. All incubators were initially set at 25°C for 5 days to allow for spore germination and initial colonization of the stem, then incubators were adjusted to the appropriate temperature, which took approximately 4 h.

Sterile deionized water was added to tubes as needed in a laminar air flow hood. Water was added up to twice a week to tubes in the 30°C incubator and as little as every 2 weeks to tubes in the 6°C incubator.

**Sampling.** Sampling began the fourth week after inoculation and continued for 7 weeks. One tray was removed from each incubator every week. Two pycnidia were cut from the center region of each stem and placed in separate 7-4-ml glass vials. Sterilized water (0.1 ml) was added to each vial. Each pycnidium was crushed with the flat end of a metal rod, and the contents of the vial were stirred. Two 10-µl volumes were removed and injected into two counting wells. Counting wells were custom made with two hemacytometer coverglasses, each with a 6-mm-diameter hole sandblasted in the center, that had been glued with epoxy to a glass slide. All conidia within both wells were counted, added, and multiplied by five to calculate conidia per pycnidium.

**Statistical analysis.** Analysis of variance was preformed to test the fixed effect of temperature (6, 12, 18, 24, and 30°C), time (weeks 4 to 10), interaction between temperature and time, with the largest reductions in CN/P at 18 and 24°C (Fig. 2). Sporulation decreased over time, with the largest reductions in CN/P at 18 and 24°C (Fig. 2).

For *B. obtusa*, conidial counts were only available for two periods because contamination with other fungi, particularly an unidentified fungus hyperparasitic on *B. obtusa* and an *Alternaria* sp., detrimentally affected spore production in one period. Temperature significantly affected sporulation and the percent conidia with dark pigmentation of *B. obtusa* (*P* ≤ 0.05) with a quadratic curvilinear response (Table 2). CN/P was highest at 18 and 24°C with the least number produced at 6 and 30°C (Fig. 1). The percent conidia with dark pigmentation was highest at 12, 18, and 24°C (Fig. 3). CN/P and the percent conidia with

**RESULTS**

At all temperatures, pycnidium development began at the ends of the stem segments during the second week and formed in the middle of stem segments during the third week. All three *Botryosphaeria* spp. produced conidia at the five temperatures tested. The mycelium of *B. dothidea* and *B. obtusa* grew below the bark, and the pycnidia erumped through the bark. The mycelium and pycnidia of *B. rhodina* grew above the bark, atypical of its habit in nature.

Latin square based on comparison of the data at the 12°C temperature was not significantly different (*P* = 0.0765, 0.1012, and 0.4185 for *B. dothidea*, *B. obtusa*, and *B. rhodina*, respectively). Therefore, temperatures from the two Latin squares were treated as one data set, and data from the two 12°C treatments were combined. Host was not significantly different (*P* ≤ 0.05) for *B. dothidea*, *B. obtusa*, and *B. rhodina*. Therefore, replications from weekly conidial counts from both hosts were combined.

For *B. dothidea*, CN/P was significant (*P* ≤ 0.05) as affected by temperature, time, and the interaction of temperature and time (Table 1). Sporulation peaked at 24°C (Fig. 1). Sporulation decreased over time, with the largest reductions in CN/P at 18 and 24°C (Fig. 2).

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* Data were analyzed using PROC MIXED (SAS Institute, Cary, NC) and linear and quadratic responses determined by contrast of least square means.

**Fig. 1.** Mean conidial number per pycnidium averaged over a 7-week sampling period produced at 6, 12, 18, 24, and 30°C for *Botryosphaeria* *dothidea* (■ and solid line), *B. obtusa* (▲ and long-dashed line), and *B. rhodina* (▲ and solid line). Bars are standard deviation of least square means based on PROC MIXED analysis (*P* ≤ 0.05).
dark pigmentation of *B. obtusa* significantly increased over time (*P* ≤ 0.05) with a significant linear response (Table 2). Maximum sporulation occurred in weeks 8 and 10 (Fig. 4). The percent conidia with dark pigmentation was highest at weeks 7, 8, 9, and 10 (Fig. 5).

For *B. rhodina*, CN/P was significantly (*P* ≤ 0.05) different due to temperature, but not significant for linear or quadratic response patterns (Table 3). CN/P were highest at 12, 24, and 30°C, but means at 24 and 30°C were not different from means at 6 and 18°C (Fig. 1). The mean CN/P were higher at all five temperatures for *B. rhodina* than at the temperature with the highest mean for *B. dothidea* and *B. obtusa* (Fig. 1). For *B. rhodina*, CN/P were not significantly different (*P* ≤ 0.05) over time (Table 3, Fig. 4).

Despite the small differences between CN/P due to temperature for *B. rhodina*, the percent conidia with dark pigmentation and percent pigmented conidia with a septum were significantly different (*P* ≤ 0.05) due to temperature and time for *B. rhodina* (Table 3). Percent conidia with dark pigmentation was lowest at 6°C and highest at 24°C, with significant linear and quadratic curvilinear responses (*P* ≤ 0.05) (Table 3, Fig. 6). Percent pigmented conidia with a septum were lowest at 6 and 30°C and highest at 12 and 24°C, with a significant quadratic curvilinear response. Percent conidia with pigmentation and percent pigmented conidia with a septum increased over time (*P* ≤ 0.05), with a significant linear response (Table 3). Maximum percent conidia with dark pigmentation and percent pigmented conidia with a septum occurred in weeks 8, 9, and 10; and weeks 9 and 10, respectively (Fig. 7).

**DISCUSSION**

Conidia of *B. dothidea*, *B. obtusa*, and *B. rhodina* were produced at all temperatures tested. Conidial production occurred at 6°C, which is below or near temperatures at which conidial germination or mycelial growth does not occur. A hypothesis, not tested in this study, is that conidial production could be initiated at

**Fig. 2.** Mean conidial number per pycnidium per week from 4 to 10 weeks after inoculation for *Botryosphaeria dothidea* at 6 (□ and solid line), 12 (■ and short-dashed line), 18 (○ and solid line), 24 (● and long-dashed line), and 30°C (▲ and long-dashed line). Bars are standard deviation of least square means based on PROC MIXED analysis (*P* ≤ 0.05).

**Fig. 3.** Percent conidia of *Botryosphaeria obtusa* with dark pigmentation (■) averaged over a 7-week sampling period at 6, 12, 18, 24, and 30°C. Bars are standard deviation of least square means based on PROC MIXED analysis (*P* ≤ 0.05).

**Fig. 4.** Mean conidial number per pycnidium per week averaged over five temperatures (6, 12, 18, 24, and 30°C) from 4 to 10 weeks after inoculation for *Botryosphaeria obtusa* (□ and solid line) and *B. rhodina* (○ and dashed line). Bars are standard deviation of least square means based on PROC MIXED analysis (*P* ≤ 0.05).

**Fig. 5.** Percent conidia of *Botryosphaeria obtusa* with dark pigmentation (■) averaged over five temperatures (6, 12, 18, 24, and 30°C) from 4 to 10 weeks after inoculation. Bars are standard deviation of least square means based on PROC MIXED analysis (*P* ≤ 0.05).

**Table 2.** Significance of temperature (6, 12, 18, 24, and 30°C), time (from week 4 to 10 after inoculation), and interaction between temperature and time on the number of conidia per pycnidium and percent conidia with dark pigmentation for *Botryosphaeria obtusa*

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* Data were analyzed using PROC MIXED (SAS Institute, Cary, NC) and linear and quadratic responses determined by contrast of least square means.
low temperatures during winter so that mature conidia are available for release when temperatures are suitable for conidial germination. Particularly in the southeastern United States, temperatures fluctuate between cold and warm periods throughout the winter and spring seasons. For _B. dothidea_, the in vitro temperature range found in this study corresponded with the temperatures that would occur during the months that Weaver (24) collected conidia in central Georgia. Weaver found no correlation between air temperature and the number of conidia collected. However, his collection methods depended on release of conidia, and release usually is in response to wetting (9,13,17). While we found that conidia were produced over the 6 to 30°C range, peak sporulation occurred at 24°C, with a moderate number of conidia produced at 18°C. This suggests that _B. dothidea_ may have a relatively narrow temperature range for peak conidia production.

Our data show that _B. obtusa_ is capable of producing conidia at low temperatures, but to no greater degree than _B. dothidea_. Yet on apple and peach trees, _B. obtusa_ has been recovered during the late fall, winter, and early spring months and is still active into early summer. Results in this study neither contradict nor provide additional insight to further explain Britton and Hendrix’s (4) and Beisel et al.’s (3) bud infestation data, Taylor’s (21) pycnidial collection data, and Sutton’s (19) spore trap data. For _B. obtusa_, peak sporulation occurred at both 18 and 24°C, with moderate sporulation at 12°C, which suggests that _B. obtusa_ has a broader temperature range for peak conidia production than does _B. dothidea_. _B. rhodina_ produced more conidia in vitro than the other species. However, it also exhibited a habit that would not be found in nature. The mycelium of _B. rhodina_ grew profusely above the bark, and pycnidia formed above the bark also. The unnatural constant moisture maintained in this experiment probably contributed to this unusual growth habit. Voorhees (22) reported a similar growth habit for some _B. rhodina_ isolates. _B. rhodina_ is the least common of the three _Botryosphaeria_ spp. in peach orchards (4). Brown and Britton (5) suggested that its minor role was due to the production of copious amounts of gum by peach trees in response to infection by _B. rhodina_. However, Pusey (16) recovered _B. rhodina_ from the outer bark and infrequently from inner bark layers, even when gumming was not evident. The atypical habit seen in Voorhees’s (22) study and in all three experiments in our in vitro research suggests that _B. rhodina_ may have a lower propensity to penetrate and colonize inner bark layers than the other two species. Environmental and/or host factors that regulate or influence _B. rhodina_’s growth habits still need to be defined.

Despite the high numbers of CN/P produced by _B. rhodina_ over the 6 to 30°C range, conidial maturation was affected by temperature. Immature conidia of _B. rhodina_ are hyaline. The typical maturation proceeds to development of a dark brown pigmentation in the cell wall, followed by production of a single central, horizontal septum. A low percentage of conidia were pigmented at 6°C, with a high percentage having pigmentation at 12 to 30°C (Fig. 5). Of the pigmented conidia, <50% had a septum at 6 to 30°C, whereas >60% formed septa at 12 to 24°C. Based on these results and the general knowledge that hyaline conidia are more susceptible to damage from UV radiation, a higher number of conidia could remain viable for longer at 12 to 24°C. These results need to be documented under natural conditions to understand the extent that temperature impacts prevalence of _B. rhodina_.

The pattern of change in CN/P over time differed with each _Botryosphaeria_ spp. CN/P decreased over a 7-week period with _B. dothidea_, increased over time with _B. obtusa_, and did not change significantly over time with _B. rhodina_. For _B. rhodina_, the percentage of mature conidia (darkly pigmented with a septum) increased over time in a similar linear pattern to CN/P for _B. obtusa_. In our study, natural moisture events that would result in the release and dispersal of conidia did not occur; therefore, conidia potentially remained associated longer with pycnidia than would occur in nature. When water was added to a tube, it was directed to flow down the side of the tube; therefore, conidia would more likely be dispersed at the lower half than at the middle to upper half of the stem.

The study did not demonstrate how CN/P correlated with field measures of the number of conidia per stem area, which would have required additional measurements such as the number of pycnidia per stem area. Nor did we measure the length of time that _Botryosphaeria_ spp. could

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**Table 3. Significance of temperature (6, 12, 18, 24, and 30°C), time (from week 4 to 10 after inoculation), and interaction between temperature and time on the number of conidia per pycnidium, percent conidia with dark pigmentation, and percent pigmented conidia with a septum for _Botryosphaeria rhodina_**

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* Data were analyzed using PROC MIXED (SAS Institute, Cary, NC) and linear and quadratic responses determined by contrast of least square means.

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**Fig. 6.** Percent conidia of _Botryosphaeria rhodina_ with dark pigmentation (● and solid line) and pigmented conidia with a septum (● and dashed line) averaged over five temperatures (6, 12, 18, 24, and 30°C) from 4 to 10 weeks after inoculation. Bars are standard deviation of least square means based on PROC MIXED analysis (P ≤ 0.05).

**Fig. 7.** Percent conidia of _Botryosphaeria rhodina_ with dark pigmentation (● and solid line) and pigmented conidia with a septum (● and dashed line) averaged over five temperatures (6, 12, 18, 24, and 30°C) from 4 to 10 weeks after inoculation. Bars are standard deviation of least square means based on PROC MIXED analysis (P ≤ 0.05).
continue to colonize and produce conidia from a section of stem, which would require longer stem sections to simulate progressive stem colonization.

The data from our study do not directly contribute to improving control strategies of the diseases caused by *Botryosphaeria* spp., but they do provide basic information to be used in combination with spore trapping data, temperature ranges of conidium germination, and seasonal recovery of pathogens from plant tissue to better understand the seasonal behavior of these pathogens. Our study demonstrates that in vitro temperature ranges for sporulation and spore maturation can differ from those for spore germination and mycelial growth.

**LITERATURE CITED**