Postharvest Development and Transmission of Watermelon Fruit Blotch

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Additional Index Words. *Acidovorax avenae* subsp. *dtrulli*, *Citrullus lanatus* storage

Summary. Watermelon fruit blotch (WFB) symptoms did not appear on healthy watermelon (*Citrullus lanatus* (Thunb.) Matsum. and Nakai) fruit placed in contact with the diseased surface of other fruit and stored at either 52 or 68 °F (11 or 20 °C) for 1 week. After 3 weeks in storage, some WFB transmission was observed and the frequency of transmission was greater at 68 than 52 °F. Surface abrasion of either the healthy fruit, diseased fruit, or both fruit did not promote transmission of WFB compared with unabraded controls. Some healthy fruit harvested from a field with diseased fruit developed very minor symptoms of WFB in postharvest storage, but the symptoms were not severe enough to cause market problems. Harvesting appeared to halt the spread of WFB symptoms on individual fruit with less than ~10% of the fruit surface affected at harvest. Care is taken during harvest and grading to exclude diseased fruit, and if proper precooling and subsequent temperature management is implemented for marketable fruit, WFB does not appear to be of concern for the marketing of watermelons.

Watermelon fruit blotch (WFB) first appeared in commercial fields in the United States in 1989 (Hopkins, 1989), although a similar disease had been described elsewhere earlier (Webb and Goth, 1965). The causal organism of WFB is a bacterium, *Acidovorax avenae* subsp. *dtrulli* (Schaad et al., 1992), and the epidemiology of the disease has been reviewed in detail (Latin and Hopkins, 1995). Earlier research has been focused on the behavior of WFB in the field because of its devastating impact on watermelon production (Hopkins et al., 1993; Hopkins and Elsom, 1995). As the disease became more prevalent in the United States, industry leaders began to express concerns about the fact that entire loads of watermelons were sometimes rejected by receivers because one or a few fruit with WFB symptoms were found in the load. These concerns about the impact of WFB on marketing watermelons led to the initiation of this project on the postharvest behavior of the disease.

Our objectives were to determine if 1) WFB increases in severity on individual watermelons in storage; 2) apparently healthy fruit harvested from an affected field will develop symptoms of WFB in postharvest storage; 3) WFB can be transmitted from a diseased fruit to an apparently healthy fruit in storage; 4) abrasion of the fruit surface, which can be transmitted from a diseased fruit to an apparently healthy fruit in storage; 4) abrasion of the fruit surface, which can occur in postharvest handling systems, would facilitate transmission of the WFB; and 5) postharvest disease development and transmission is influenced by temperature in the storage environment. An understanding of the postharvest behavior of WFB will be the basis for recommendations for the control of WFB in the marketing chain.

Materials and methods

Seedling production. In 1996 and 1997, 'Royal Charleston' watermelon seeds that tested positive for infection with *A. avenae* subsp. *dtrulli* were obtained from Petoseed Company (Saticoy, Calif.). Additional 'Starbrite' seeds without *A. avenae* subsp. *dtrulli* infection were obtained from Asgrow Seed Co. (Kalama, Wash.). All seeds were planted in Speedling trays (Sun City, Fla.) with 1-inch cells using Scotts Mixture mix 300 as medium (Scotts-Sierra, Marysville, Ohio). Trays were placed in a temperature-controlled room at 86 °F (30 °C) until seedling emergence had begun, then transferred to a greenhouse. Plants were irrigated as needed and fertilized about every 10 d with Peters Professional 9-45-15 (Grace-Sierra, M Ilipitas, Calif.) according to label directions.

Field production. About 1 month after planting, seedlings of ~3 inches (7.5 cm) in height were transplanted to a field at the Coastal Research and Education Center (CREC), Charleston, S.C. The two varieties were evenly spaced throughout the field and no attempt was made to avoid transmission of WFB from one variety to the other during transplanting. Recommended practices for commercial production were followed (Cook et al., 1995), except that frequent light irrigation was implemented to facilitate disease development. The size of the planted area was ~1.2 acres (0.5 ha). The same field was used in 1996 and 1997 to produce plants from both varieties. An additional field (1.2 acres), remote from the WFB infested field, was used in 1997 to produce healthy 'Starbrite' watermelons.

In 1996, Clemson University Extension staff reported that a commercial planting of 'Royal Sweet' watermelon in Bamberg County, S.C., was affected with WFB. This field was used to harvest fruit with WFB. In 1997 there were no reports of WFB in commercial fields in South Carolina.

Harvesting. On the morning of 19 June 1996, ~200 'Royal Sweet' watermelons infected with WFB were harvested from the commercial field in Bamberg County. The percentage of fruit surface showing WFB symptoms was visually estimated and fruit within the range of 10% to 25% diseased surface area were selected for the postharvest evaluation. On the afternoon of the same day, ~200 'Starbrite' fruit with no visible symptoms of WFB were selected from the plant at the CREC. All of the harvested fruit were placed in a shaded area and the storage trial was initiated the following morning.

In 1997, healthy 'Starbrite' watermelons produced in the isolated field at the CREC were utilized for the disease transmission component of the study. On 8 July, healthy 'Starbrite' watermelons were harvested, carefully examined for the presence of WFB, and 22 of the best-quality melons without WFB symptoms were placed at each storage temperature (52 and 68 °F) to be used later for the disease transmission experiment. An additional 22 fruit were placed at...
each temperature as a control.

Development of WFB on ‘Royal Charleston’ in the infested field at CREC was substantially delayed in 1997 compared with 1996, possibly due to the unusually cool temperatures that prevailed during the 1997 growing season (Hopkins, 1997). During the period of 15 to 29 July, the infested field was monitored daily and a total of 44 ‘Royal Charleston’ fruit with WFB symptoms affecting less than =10% of the fruit surface were harvested to be utilized for the disease transmission experiment.

**Postharvest Transmission of Disease.** In 1996, 70 healthy ‘Starbrite’ watermelons were placed at each storage temperature [52 and 68 °F (11 and 20 °C)]. Infected ‘Royal Sweet’ fruit, based on the presence of visual symptoms, were utilized to determine if WFB could be transmitted from the infected fruit to the healthy fruit. Healthy watermelons, not in contact with diseased watermelons, were utilized as the control and designated treatment A. In treatment B, healthy fruit were placed in contact with the diseased area of an infected fruit with neither fruit surface abraded.

In a series of abrasion treatments, the fruit surface was abraded with sandpaper to cause superficial injury similar to that which is often observed in normal handling systems. Medium grade sandpaper was cut into pieces =1.25 inches² (3 cm²) and a clean piece of sandpaper was used to lightly score the surface of selected melons before they were placed in contact with each other as follows: treatment C, healthy unabraded paired with diseased unabraded; D, healthy unabraded paired with diseased unabraded; and E, healthy and diseased fruit both abraded. Abraded areas and diseased areas of different melons were placed in direct contact.

Two unabraded ‘Starbrite’ melons were stored at each temperature as controls. All other treatments had three pairs of fruit per replication. This design was repeated five times at each temperature. Data were analyzed as a split-unit design with temperature as the main unit and treatments as split units.

In 1997, abrasion treatments were not included in the WFB transmission study because there had been no apparent effect of the abrasion treatments in 1996 and we had a limited number of fruit with WFB in 1997. Infected ‘Royal Charleston’ fruit were used as the source of WFB. Healthy ‘Starbrite’ watermelons, produced in the noninfested field, were used to determine if postharvest WFB transmission could occur. The disease transmission study included treatments A and B as described for 1996. ‘Royal Charleston’ fruit harvested on each of seven dates were treated as replications, which resulted in seven replications with one to six pairs of fruit per replication. There were a total of 22 pairs of fruit at each temperature.

**Postharvest Increase in Severity of WFB on Individual Watermelons.** In 1996, 30 ‘Starbrite’ and 10 ‘Royal Charleston’ watermelons were selected with diseased areas ranging in diameter from =1.25 to 4 inches (3 to 10 cm). The area showing symptoms on each fruit was outlined in indelible ink, then 15 ‘Starbrite’ and 5 ‘Royal Charleston’ fruit were placed at each of the two storage temperatures to observe if the area with symptoms increased during the 3-week storage period.

**Postharvest Development of WFB Symptoms on Fruit That were Symptomless at Harvest.** In 1996, the control fruit described earlier as Treatment A served as a means of observing the development of WFB on fruit that did not have symptoms of WFB at the time of harvest but were grown in an infested field. In 1997, two approaches were taken. The control fruit identified earlier as treatment A consisted of symptomless ‘Starbrite’ fruit harvested from a noninfested field.

Additionally, on 11 July, ‘Starbrite’ and ‘Royal Charleston’ fruit without WFB symptoms were harvested from the infested field. These fruit served as another control to observe if WFB symptoms develop postharvest on symptomless fruit harvested from an infested field. Twenty-five ‘Starbrite’ and seven ‘Royal Charleston’ were placed at 52 °F. Eighteen ‘Starbrite’ and seven ‘Royal Charleston’ were placed at 68 °F.

**Results**

**Occurrence of WFB Symptoms on Transplants and Fruit in the Field.** In 1996, visual symptoms of WFB as described by Latin and Hopkins (1995) were apparent on leaves of ‘Royal Charleston’ within 1 week after seedling emergence, but were absent from ‘Starbrite’. Within 1 month after transplanting, WFB symptoms were observed on some plants of both cultivars in the field.

In 1997, WFB symptoms on transplants developed in a manner similar to the 1996 study. Once the seedlings were transplanted to the field, additional WFB symptoms on the foliage appeared substantially later (=15 June) and were less severe than observed in 1996, possibly due to the prevailing cool, dry weather in the 1997 season (Hopkins, 1997).

In 1996, nearly all of the fruit of ‘Royal Charleston’ at the CREC developed symptoms of WFB as they neared harvest maturity. Fruit quality deteriorated drastically before ‘Starbrite’ had reached harvest maturity. For this reason, diseased ‘Royal Sweet’ fruit were harvested from the commercial farm for use in the study on postharvest transmission of the disease. About 25% of ‘Starbrite’ fruit in the CREC field developed symptoms of WFB as they neared harvest maturity in the field, but there were sufficient numbers of symptomless fruit to conduct the postharvest studies. In 1997, the first WFB symptoms were noted on ‘Royal Charleston’ fruit on 15 July 1997.

**Postharvest Increase in WFB Severity on Individual Fruit that Had Symptoms at Harvest.** In 1996, all fruit were evaluated weekly for 3 weeks to observe increases in the diseased areas that had been delineated with indelible ink. After 1 week, none of the fruit at 52 °F showed an increase in the affected area. At 68 °F, affected areas on ‘Starbrite’ fruit had not increased and only one of the ‘Royal Charleston’ fruit had an estimated increase of 20% in the diseased area.

After 2 weeks, blotch symptoms on fruit at 52 °F still had not worsened. At 68 °F, 3 of the 15 ‘Starbrite’ had an estimated increase of blotch symptoms of 5%. ‘Royal Charleston’ fruit had not changed since the first week’s evaluation, with only one fruit showing an increase in severity. In sharp contrast, fruit with blotch symptoms in the field rapidly worsened and decayed within a few days.

After 3 weeks, fruit at 52 °F had minor increases in the area of blotch symptoms with two of the ‘Royal Charleston’ and two of the ‘Starbrite’ showing increases of =10%. ‘Starbrite’ fruit had no additional ‘Starbrite’ fruit had increased blotch symptoms, but the three fruit that were worsening in the 2-week evaluation continued to develop a larger area of blotch. One of the three had =100% more area affected than at harvest time and the other two had increased to =50% greater area than at harvest time. None of the ‘Royal Charleston’ had an increase in diseased area. Removing the watermelons from the vine when the blotted area was no greater than 4 inches (10 cm) in diameter dramatically
reduced the further development of symptoms compared to fruit that were not harvested.

**Postharvest Development of WFB on Fruit that were Symptomless at Harvest.** In 1996, one of the 10 control fruit (treatment A) at each temperature had developed a diseased area of ~1 inch (2.5 cm) in diameter after 3 weeks in storage. Tissue from these fruit were cultured in the laboratory. A *Acidovorax avenae* subsp. *citrulli* was not isolated from these samples, indicating that if the bacterium must have been present, it was superficial in nature as were the symptoms.

**Postharvest Transmission of WFB from Infected to Symptomless Fruit.** In the 1996 study, none of the 'Starbrite' fruit in any treatment at either temperature had developed symptoms of blotch after 1 week in storage. After 2 weeks in storage, deterioration of the diseased 'Royal Sweet' fruit was so severe at both temperatures that it was impossible to move them and evaluate the 'Starbrite' without jeopardizing the experimental design. None of the 'Royal Sweet' had literally liquefied and could not be handled. Evaluation of the 'Starbrite' fruit for transmission of WFB was postponed until week 3.

After 3 weeks, all of the diseased 'Royal Sweet' fruit were removed from storage and the 'Starbrite' fruit were washed so that an evaluation for WFB transmission could be conducted. In the control groups (Treatment A), one 'Starbrite' fruit at each temperature had developed a small blotch. In both fruit the affected area was ~1.25 inches (3 cm) in diameter and the fruit were otherwise in good condition.

Transmission of WFB to healthy 'Starbrite' fruit that were paired in contact with diseased 'Royal Sweet' varied from 8% to 53% (Fig. 1). There was no significant effect of abrasion treatments on the transmission of WFB from 'Royal Sweet' to 'Starbrite' compared with paired fruits that were not abraded. There was an increase in disease transmission in all four paired-fruit contact treatments compared with WFB symptom development in controls that were not in contact with a diseased fruit (P < 0.05). Storage temperature significantly influenced the rate of WFB symptom development (P = 0.03), with greater frequency of development at 68°F than at 52°F.

In 1997, 'Starbrite' fruit examined after 1 or 2 weeks in contact with diseased 'Royal Charleston' did not exhibit any symptoms of WFB at either storage temperature. At the 3-week examination, none of the 'Starbrite' fruit at 52°F exhibited WFB symptoms, but at 68°F, 3 of the 22 'Starbrite' had very minor (<1 inch) diameter WFB symptoms. A *Acidovorax avenae* subsp. *citrulli* was isolated from two of these fruit. Two of the control fruit not in contact with other fruit also had developed very minor WFB symptoms at 68°F.

**Discussion**

The severity of blotch symptoms did not increase dramatically postharvest on fruit that had ~4 inches (10 cm) diameter symptoms at the time of harvest. The slight increases of disease fruit surface that were noted were more pronounced at 68°F than at 52°F. Symptomless fruit harvested from an infested field had an extremely low incidence of development of WFB symptoms postharvest, and when symptoms did develop, they were minor and of no consequence to the marketing of the fruit.

Watermelon fruit blotch was not transmitted from diseased to healthy fruit in the first week of storage at either temperature, but it was transmitted from diseased to healthy fruit after 3 weeks in storage. The incidence of transmission was greater at 68°F than at 52°F in the 1996 study. In 1997, this pattern was observed again, but the difference between temperatures was not statistically significant. Abrasion of the fruit surfaces does not appear to significantly influence disease transmission.

The fact that storage temperature influences the frequency of disease transmission, with more transmission at the higher temperature, suggests that appropriate precooling of watermelons will reduce postharvest incidence of fruit blotch. Watermelon fruit blotch likely can be controlled in the marketing chain if all fruit with symptoms are excluded from harvest or are eliminated during packing, and if appropriate temperature management is employed for storage and distribution.

**Literature Cited**


