Pollen Viability of Selected Diploid Watermelon Pollenizer Cultivars

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Abstract. In the Spring and Fall 2006, the pollen viability of four diploid watermelon pollenizers was evaluated in Quincy, FL. Triploid watermelon plants [Citrullus lanatus (Thunb.) Matsum. & Nakai.] do not produce sufficient viable pollen to pollinize themselves and a diploid cultivar must be interplanted as a pollen source. Recent studies have illustrated differences in triploid watermelon yields as a result of the pollenizer cultivar used. The viability of the pollen produced by pollenizer cultivars may greatly influence the fruit set and fruit quality in the triploid watermelon crop. Pollen samples were taken from ‘Companion’, ‘Jenny’, ‘Mickylee’, and ‘SP-1’ and were stained to determine their viability. There were no significant differences in pollen viability among cultivars and all cultivars had high average viability. Pollen viability was never lower than 95% for any cultivar. This study indicates that pollen viability of the cultivars evaluated should not influence their effectiveness as pollenizers.

Seedless watermelons account for 78% of the watermelons sold in the United States (U.S. Department of Agriculture, 2006). Triploid watermelon plants do not produce sufficient viable pollen to pollinize themselves and a diploid cultivar must be interplanted as a pollenizer (pollen source) (Maynard and Elmstrom, 1992). Growers have traditionally planted every fourth or fifth row in the field with a diploid cultivar and harvested both seeded and seedless watermelons.

Diploid cultivars have recently been developed to be planted in-row as pollen sources between triploid plants without altering the triploid spacing’s or affecting marketable yield. Freeman et al. (2007) observed significant differences in the performance of in-row pollenizers based on seedless watermelon yield. Triploid plants pollenized by ‘Companion’ yielded less than those pollenized by ‘Jenny’, ‘Patron’, ‘SP-1’, and ‘Sidekick’. Fiacchino and Walters (2003) demonstrated that ‘Crimson Sweet’ was a more effective pollenizer than ‘Fiesta’ based on greater seedless watermelon yields and less incidence of hollow heart. Studies on ‘Crimson Sweet’ and ‘Fiesta’ by Stanghellini and Schultheis (2005) illustrated that there were no statistical differences in pollen production between the two cultivars. This suggests that there may be other factors that contribute to a pollenizer’s performance.

Pollen viability could be a determining factor in the performance of watermelon pollenizers. Pollen flow from a pollenizer would be of little value if the viability was low, because pollen tube growth and ovule fertilization are necessary for seedless fruit maturation (Maynard and Elmstrom, 1992; Rhodes et al., 1997; Robinson and Decker-Walters, 1997). Poor pollination in watermelon can affect fruit shape and thus its marketability and may be a contributing factor to the physiological disorder termed hollow heart (Fiacchino and Walters, 2003; Maynard, 1992). Research in other plant families has shown that there can be substantial differences in pollen viability within a species and between closely related species (Fortescue and Turner, 2004; Lavi et al., 1996; Nikkanen et al., 2000; Parzies et al., 2005). Fortescue and Turner (2004) reported significant differences in pollen viability between cultivars of banana (Musa acuminata Colla). Studies conducted in mandarin orange (Citrus reticulata Blanco) have shown that pollen source can affect yield and quality characteristics such as soluble solids (Vithanage, 1991; Wallace and Lee, 1999). Differences in pollen viability have been reported in rabbiteye blueberry; however, all cultivars examined had high average viability and probably had no effect on reproductive success (Brevis et al., 2006).

Cultivars used in this study were chosen because previous research has illustrated differences in their performance and pollen viability may be a contributing factor (Freeman et al., 2007). The objective of this research was to determine the pollen viability of four diploid watermelon pollenizer cultivars.

Materials and Methods

On 3 Apr. and 1 Aug. 2006, 4-week-old watermelon seedlings were transplanted into raised beds covered with black polyethylene mulch in the spring and white on black polyethylene mulch in the fall. Experiments were performed at the North Florida Research and Education Center (NFREC) in Quincy, FL. Soil type at NFREC is Norfolk loamy sand (fine-loamy, kaolinitic, thermic Typic Kandiudults). The experimental design was a randomized complete block with four replications. The pollen viability of four diploid pollenizer cultivars, ‘Companion’, ‘Jenny’, ‘Mickylee’, and ‘SP-1’, was determined. Experimental plots were 4.57 m long with in-row spacing of 0.91 m and between-row spacing of 2.43 m. Three pollenizer seedlings were planted at even spacing in the center of each plot. Fertilization, irrigation, and pesticide application practices were followed using recommendations provided by the University of Florida Institute of Food and Agricultural Sciences (Olson et al., 2006).

In the spring, sampling was initiated on 17 May with additional pollen samples taken on 24 May and 31 May. For the fall, sampling was initiated on 31 Aug. with other pollen samples taken on 7 Sept. and 14 Sept. The sampling period was scheduled to coincide with the fruit-setting period of adjacent triploid plants transplanted on the same date.

On the sampling dates, watermelon flowers were removed from the plant before the flowers opened. This was to ensure that pollinators would not remove pollen and an adequate supply of pollen would be available for analysis. Three staminate flowers were removed from each plot and placed into plastic cups and covered to exclude pollinators. Flowers were taken to the laboratory and harvested both seeded and seedless watermelons.

Table 1. Analysis of variance for pollen viability of watermelon pollenizer cultivars tested in Spring and Fall 2006 at Quincy, FL.

| Source            | df | MS     | P value |
|-------------------|----|--------|---------|
| Replication       | 3  | 0.00410| 0.410   |
| Sampling date     | 2  | 0.02827| 0.003   |
| Cultivar          | 3  | 0.00856| 0.124   |
| Date*cultivar     | 6  | 0.00281| 0.669   |
| Error             | 33 | 0.00415|
| Replication       | 3  | 0.00290| 0.408   |
| Sampling date     | 2  | 0.00302| 0.367   |
| Cultivar          | 3  | 0.00281| 0.422   |
| Date*cultivar     | 6  | 0.00257| 0.519   |
| Error             | 33 | 0.00292|

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allowed to open. Samples were analyzed after anther dehiscence was verified using a hand lens. Pollen was removed from the anthers and placed on a slide. Viability was determined using the diaminobenzidine (DAB) test for peroxidase activity in pollen (Dafni et al., 2005). Rodriguez-Riano and Dafni (2000) compared the results of four vital dyes versus pollen germination results and illustrated the superiority of peroxidase tests over other commonly used vital dyes. The DAB test uses a dye that creates a color differential between viable and nonviable pollen. Four 100-pollen grain subsamples were analyzed from each plot using a compound microscope. A pollen grain was considered viable if it turned dark brown or black. All pollen samples were analyzed on the same day pollen was collected. It is important that watermelon pollen be thoroughly mixed with the dye on the slide to have adequate contact between the pollen grains and dye. If large clumps of pollen are not broken up, dye may not infiltrate the pollen and false-negatives may be observed. On each sample date, heat-killed pollen (2 h at 80°C) was analyzed to prevent false-positive readings. New dye was prepared for each sample date. Data were taken as percent viable pollen and square root transformations were performed. Analysis of variance (PROC GLM) and means separation (Duncan’s multiple range test) were accomplished using SAS version 9.1 (SAS Institute, Cary, NC).

Results

There were no significant differences (P ≤ 0.05) in pollen viability observed among the pollenizer cultivars tested (Table 1). In the spring trial, significant differences (P ≤ 0.05) in pollen viability were observed among sampling dates but not cultivars. There were no significant interactions observed between pollenizer cultivar and sampling date (Table 1).

Pollens samples taken on 31 May showed a greater average viability than samples taken on 24 May or 17 May (Table 2). Pollen viability on 31 May was 98.6%, which was greater than 97.4% or 97.0% for 17 May and 24 May, respectively. The average pollen viability over all cultivars and all dates for spring and fall were 97.7% and 97.9%, respectively. There was little variation within the data and the coefficients of variation were never higher than 0.84%.

Discussion

The results of this study illustrate that there was no significant variation in pollen viability between the diploid pollenizer cultivars and that pollen viability changed very little during triploid fruit set. The results of this study are similar to Nepi and Pacini (1993) who studied ‘Greyzini’ (Cucurbita pepo L.) and found average pollen viability to be ≈92% at anthesis. The higher viability of pollen during one sampling date in the spring is more likely the result of environmental conditions and not cultivar characteristics. Nepi and Pacini (1993) also reported that decrease in pollen viability in ‘Greyzini’ was the result of dehydration of the pollen grains. High morning temperature or low relative humidity shortly after anthesis could decrease pollen viability.

The results of this study suggest that pollen viability was not a contributing factor in the varying degrees of performance of these pollenizers. Factors such as floral attractiveness to pollinators, availability and total production of staminate flowers, competitive growth, and quantity of pollen produced may be more important characteristics of pollenizers.

The small amount of variation in pollen viability between the cultivars tested suggests that there may be little variation within common diploid cultivars. Pollen viability appears to play no role in the performance of the pollinizers tested and may not be an important characteristic of pollenizer cultivars.

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Table 2. Pollen viability of four diploid watermelon pollenizer cultivars at Quincy, FL, during the Spring and Fall 2006.

| Pollenizer cultivar | 17 May  | 24 May  | 31 May  | 31 Aug.  | 7 Sept.  | 14 Sept. |
|---------------------|---------|---------|---------|----------|----------|----------|
| Mickylee            | 97.8%   | 98.0%   | 98.7%   | 97.4%    | 98.2%    | 97.8%    |
| Companion           | 97.4    | 97.2    | 99.2    | 97.6     | 98.1     | 98.8     |
| Jenny               | 97.3    | 97.2    | 98.3    | 97.8     | 97.7     | 97.0     |
| SP-1                | 97.2    | 95.5    | 98.1    | 97.9     | 98.7     | 97.6     |
| Date means          | 97.4%   | 97.0%   | 98.6%   | 97.7%    | 98.2%    | 97.8%    |

*p = 0.05; means are compared within the same column.

*n = 0.05; means from the same season are compared within the row.

Nonsignificant.