Self-incompatibility in *Cornus florida*

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Additional index words: breeding, pollen tube growth

Abstract. Low seed set has been reported following self-pollinations of flowering dogwood (*Cornus florida* L.). The objective of this study was to verify the presence of self-incompatibility in *C. florida*. ‘Cherokee Princess’ stigmas and styles were collected 1, 2, 4, 8, 12, 24, 48, and 72 hours after cross- and self-pollinations, stained with aniline blue and observed using a fluorescence microscope. Pollen germinated freely following self-pollinations, but self-pollen tubes grew slower than those resulting from cross-pollinations. By 48 hours after cross-pollination, pollen tubes had reached the bottom of the style while pollen tubes in self-pollinated flowers had only penetrated the upper third of the style. Evidence of reduced pollen tube growth rate in self-pollinations of ‘Cherokee Chief’ and ‘Cherokee Brave’ was also obtained. This study provides evidence of a gametophytic self-incompatibility system in *C. florida*. It was also determined that stigmas of *C. florida* ‘Cherokee Princess’ are receptive to pollen from 1 day prior to anthesis to 1 day after anthesis.

The genus *Cornus* consists of ≥50 species, many of which are cultivated as ornamentals (Dirr, 1998; Eyde, 1988). The most popular member of the genus is *C. florida*, or flowering dogwood, which is highly valued as a landscape tree because of its white, pink or red bracts in the spring, autumn foliage color, and fruit display. According to the 1998 Census of Horticultural Specialties, annual sales of *C. florida* exceeded $26 million, ranking it third among flowering deciduous trees for U.S. sales. Many cultivars are available in the marketplace, but most have been the result of grower selections of sports or superior seedlings rather than organized plant breeding efforts (Santamour and McArdle, 1985).

In recent years, disease problems have plagued native populations, nursery stock and landscape plantings of *C. florida* (Sherald et al., 1996; Sinclair et al., 1987) and have prompted a search for disease resistant germplasm. A plant with resistance to anthracnose (*Discula destructiva*) was discovered in the wild and released as the cultivar Appalachian Spring (Windham et al., 1998). Three powdery mildew resistant cultivars have recently been released (Windham et al., 2000) and other resistant individuals identified (Mmbaga and Sheng, 2001); all these plants were selected from seedling populations growing at nurseries. While progress in developing disease resistant cultivars has been made through selection of open-pollinated seedlings, development of cultivars with multiple disease resistance or with a combination of disease resistance and specific ornamental characteristics will most likely require controlled pollinations between selected parents.

Received for publication 23 Dec. 2002. Accepted for publication 21 Apr. 2003. Mention of trade names of commercial products in the publication is solely for the purpose of providing specific information and does not imply recommendation or endorsement by the U.S. Dept. of Agriculture.

Research Geneticist.

*HortScience* 39(2):335–338. 2004.
Results and Discussion

Evaluation of self-incompatibility. Pollen germination was observed on only a few ‘Cherokee Princess’ stigmas collected 1 to 2 h after pollination; however, germinating pollen was present on at least 80% of the stigmas that were collected four or more hours after either cross- or self-pollination (Table 1; Fig. 1A and B). From 4 to 12 h after pollination, more germinated grains were observed on stigmas that had been cross-pollinated than on those that had received self-pollen. By 24 h after pollination, similar numbers of germinated grains were noted on both cross- and self-pollinated stigmas. The number of germinated pollen grains was difficult to determine by 48 h after pollination due to decreased fluorescence and collapsed appearance of the pollen grains; therefore, counts of germinated grains were omitted for samples collected 48 and 72 h after pollination. No germinated pollen grains were observed in emasculated, unpollinated controls.

Very few ungerminated pollen grains were observed on any of these stigmas, even in the samples collected 1 to 2 h after pollination and those that served as the unpollinated control. It appears that ungerminated pollen washed off the stigmas during the fixation or staining procedures. Therefore, it is not known whether the differences in number of germinating grains between self- and cross-pollinations in samples collected 4 to 12 h after pollination reflect actual differences in pollen germination or were caused by different amounts of pollen being applied to stigmas.

While considerable background fluorescence was observed in C. florid styles, pollen tubes were easily discerned following both cross- and self-pollination. Pollen tubes exhibited fluorescence along their entire length, with bright callose plugs occasionally apparent (Fig. 1). Differences in pollen tube lengths between ‘Cherokee Princess’ cross- and self-pollinations were observed in samples collected from 4 to 72 h after pollination (Fig. 2). From 4 to 24 h after pollination, pollen tubes of cross-pollinated specimens were about twice the length of those from self-pollinations. By 48 h after cross-pollination, pollen tubes were observed emerging from the base of the style (Fig. 1D). In contrast, pollen tubes had only penetrated the top one-third of the style in self-pollinations observed 48 h after pollination. No additional pollen tube elongation was noted in styles collected 72 h after self-pollination.

Considerable differences in pollen tube length following cross- and self-pollinations of ‘Cherokee Chief’ and ‘Cherokee Brave’ were also noted (Fig. 3). As with the ‘Cherokee Princess’ × ‘Cherokee Chief’ pollinations, by 24 h after pollination pollen tubes had grown about halfway down the styles in the ‘Cherokee Chief’ × ‘Cherokee Princess’ and ‘Cherokee Brave’ × ‘Cherokee Princess’ pollinations. At this time, pollen tubes resulting from self-pollinations were only about one-half the length of those from cross-pollinations.

Self-incompatibility is a genetically controlled mechanism that prevents formation of a zygote after self-pollination of a fertile hermaphrodite plant, and is estimated to be present in almost half of all angiosperm species (Brewbaker, 1959). Two homomorphic self-incompatibility systems, sporophytic and gametophytic, are recognized (de Nettancourt, 1977). In the sporophytic self-incompatibility system, the incompatibility reaction is determined by the diploid genotype of the pollen parent and germination of incompatible pollen is inhibited at the stigmatic surface. Gametophytic self-incompatibility is determined by the haploid genotype of the pollen, and is manifested by an inhibition of pollen tube growth in the style. Incompatible pollen tubes generally reach one-third to three-quarters the length of the style in the time required for compatible tubes to penetrate the entire style (Ascher, 1976). Sporophytic self-incompatibility is associated with plants that produce tricellular pollen, whereas gametophytic self-incompatibility is generally found in plants with bicalcilar pollen (Brewbaker, 1957).

The presence of a self-incompatibility system in C. florid has been suggested by
studies in which self-pollinations have resulted in low or no seed set (Ohta, 1971; Orton, 1983; Reed, 1999). This study confirms this incompatibility and provides evidence of a gametophytic self-incompatibility system. Pollen freely germinated on the stigmatic surface following self-pollination, but pollen tubes grew slower than those resulting from cross-pollination. The slower pollen tube growth and the failure of pollen tubes from self-pollinations to reach the base of the style are indicative of gametophytic self-incompatibility. Based on the presence of bicellular pollen in *Cornus* (Brewbaker, 1967), this is the self-incompatibility system that is predicted for this species. Results of this study indicate that it is not necessary to emasculate *C. florida* flowers before making controlled cross-pollinations.

**Time of stigma receptivity.** Pollen germination was observed on at least 90% of the ‘Cherokee Princess’ stigmas pollinated from 1 d before to 1 d after anthesis (Table 2). Numbers of germinated pollen grains present on stigmas pollinated over this 3-d period were similar. The percentage of stigmas with germinated pollen and the mean number of germinated grains were lower when pollinations were made 2 or 3 d after anthesis. By 3 d after anthesis, pollen germination was observed on only 20% of the stigmas.

Differences in pollen tube lengths were also noted among pollination times. Pollen tubes from flowers pollinated 1 d before to 1 d after anthesis were similar in length to each other, and to pollen tubes from cross-pollinations collected 24 h after pollination in the self-incompatibility experiment (Table 2; Fig. 2). Pollen tube growth was reduced in pollinations made 2 to 3 d after anthesis. pollen tubes from flowers pollinated 3 d after anthesis were about one-half the length of those from flowers pollinated 2 to 4 d earlier.

Since the individual flowers of a *C. florida* inflorescence open over a 2- to 3-week period, obtaining maximum number of seeds may require multiple pollinations of an individual inflorescence. Results of this study indicate that stigmas of *C. florida* ‘Cherokee Princess’ are highly receptive to pollen from the day before anthesis to the day after anthesis. Some stigmas may be receptive 2 d after anthesis, but little seed set is expected from flowers pollinated 3 d or more after anthesis. Stigma receptivity prior to 1 d before anthesis was not tested. For cross-pollinations, stigma receptivity prior to anthesis is irrelevant since self-incompatibility eliminates the need to emasculate flowers.

A study of the effect of multiple applications of pollen on seed set in *C. florida* ‘Cherokee Princess’, ‘Cherokee Brave’, ‘Cherokee Chief’, and Barton determined that seed set was the same when flowers were pollinated every other day as when pollen was applied daily during the first 12 d of flowering (Reed, 1999). Less frequent pollinations resulted in reduced seed set. These findings concur with evidence presented in the current study that stigmas receptivity begins to decrease by two days after anthesis. In the absence of emasculations, it appears that dogwood flowers must

| Maternal parent | Stigmas with germinated pollen (%) | Mean pollen germination mean length of pollen tubes (mm) |
|-----------------|-----------------------------------|------------------------------------------------------|
|                 | 1 d prior to anthesis              | 1 d after anthesis                                   |
|                 | 100                                | 2.3 b                                                |
|                 | Day of anthesis                    | 2.6 ab                                               |
|                 | 90                                 | 2.6 b                                                |
|                 | 1 d after anthesis                 | 4.0 a                                                |
|                 | 100                                | 1.5 b                                                |
|                 | 2 d after anthesis                 | 1.2 b                                                |
|                 | 60                                 | 1.2 b                                                |
|                 | 3 d after anthesis                 | 1.0 b                                                |

Values within a column followed by the same letter do not differ significantly according to Fischer’s least significant difference text (*P* ≤ 0.05).
be cross-pollinated within 1 or 2 d of opening for pollinations to be effective. If maximum seed set from each inflorescence is needed, as when working with young plants with limited number of inflorescences, pollinations should be repeated every other day.

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