Source of Pollen, Distance from Pollinator, and Time of Pollination Affect Yields in Block-type Pecan Orchards

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Abstract. Inadequate cross-pollination of pecan (Carya illinoinensis (Wangenh.) K. Koch) occurred in block-type orchards generally thought exempt from pollination-related crop losses because of an abundance of nearby potential pollinizers. “Off-genotypes” appeared to be potentially major assets in such orchards due to their role as backup pollinizers; hence, their presence assures against crop losses due to poor pollination. Fruit-set in ‘Desirable’ main crop rows declined sigmoidally as distance from ‘Stuart’ pollinizer rows increased. For 15.4-m row spacings, rate of decrease was maximum between 49 and 78 m, depending on crop year. Maximum fruit-set was in rows immediately adjacent to the pollinizer. Tree age/size and spring temperature influences on the characteristics of flower maturity windows are probably primary factors contributing to pollination-related fruit-set losses in block-type orchards relying upon pollen from a single complementary pollinizer or from neighborhood trees. For example, flower maturity was earlier in older/larger trees, and higher spring temperatures accelerated catkin development relative to that of pistillate flowers. Maximum fruit-set occurred when pistillate flowers received pollen around 1 day or less after becoming receptive, whereas no fruit-set occurred when they were pollinated around four or more days after initial receptivity. These findings indicate that many block-type orchards in the southeastern United States are exhibiting pollination-related crop reductions and that future establishment of such orchards merits caution regarding the spatial and temporal distribution of pollinizers.

Pecan is a wind-pollinated monocious crop that exhibits heterodichogamy (Adriance, 1930; Grauke and Thompson, 1996). Synchrony of pollen release and of pistillate flower receptivity varies from none to complete and is influenced by genotype and environmental conditions (Wood et al., 1997; Worley et al., 1992). Pollination dynamics can therefore cause fruit-set problems. Crop losses can be due to poor pollination, excessive self-pollination, or xenia (Romberg and Smith, 1946). Environmental and age factors potentially influence crop loads in certain tree fruit crops (Buban, 1996) and may do so in pecan. The general dearth of pollinizers in the vicinity of southwestern U.S. orchards led to research establishing that pollination-related crop losses occur in that region when pollinator distance is two or more rows from the main crop cultivar (Marquard, 1988). Conversely, the relative abundance of diverse cultivars and feral germplasm proximate to most orchards in the region has contributed to the commonly accepted belief that pollination is not a problem. This has been partially based on observations of the great abundance (Woodroof, 1924, 1930) and mobility (Woodroof and Woodroof, 1927) of pollen produced by trees and its potential availability to regional orchards. Nevertheless, the detection of extensive self-pollination in many southeastern block-type orchards by Wood and Marquard (1992) suggests that pollinator-related crop reductions are occurring. Block-type plantings, where a single cultivar typically occurs in sections 5–50 or more rows across, are common within the region and are becoming increasingly popular because of managerial advantages.

This report documents the occurrence of pollination-related crop reductions in block-type orchards within the southeastern United States. It also provides insight into how pollinizer distance, spring temperatures, tree age/size, and the timing of the exposure of pistillate flowers to pollen influence pollination.

Materials and Methods

Experiment 1. Documentation of crop loss in block-type plantings due to lack of pollination. Pecan orchards are typically established in blocks composed of the main cultivars and one pollinizer (or rarely two). The pollinizer is often positioned on the orchard’s perimeter or in rows ≥75 m from many main crop rows. ‘Desirable’ and ‘Stuart’ is one such cultivar pair commonly planted in the southeastern United States. Because ‘Desirable’ is protandrous (Type I) and ‘Stuart’ is protogynous (Type II), their floral maturity has been commonly assumed to be reciprocally complementary. Therefore, a large block-type orchard of ‘Desirable’ and ‘Stuart’ was selected for study. A second orchard was also studied in which a small block of a single cultivar (‘Pawnee’) was planted next to a large orchard of supposedly complementary cultivars.

The first test orchard was located in the central zone of the Georgia Pecan Belt (near Albany, Ga.), and is near the primary U.S. production area. Trees were 15 years old and were managed according to standard commercial practices (Crocker, 1995). The =150-ha orchard was arranged in repeating blocks of 19 rows of ‘Desirable’ and four rows of ‘Stuart’. Rows were N–S oriented and 15.4 m apart (50 feet) with 10.8 m (35 feet) between trees within the row. Orchard composition was almost exclusively these two cultivars, but with an occasional seedling tree (i.e., escaped rootstock) or “off-genotype” (“OG,” another cultivar or an escaped rootstock). Other genotypes were fairly abundant within 1 km of the test orchard and many were present in the general vicinity.

Fruit-set of ‘Desirable’ trees, within the ‘Desirable’/‘Stuart’ orchard, was atypically low in 1995. In the central zone of the block, the region most distant from the ‘Stuart’ pollinizer, fruit-set was evaluated on OG trees (classified based on morphological traits) proximate to other OG trees dispersing pollen during the receptivity period of ‘Desirable’. These trees were selected from a low population randomly dispersed in the orchard’s interior. Fruit-set was then assessed for five ‘Desirable’ trees at each “tree location,” as trees radiated in all directions from these OG trees. Set was assessed by counting the number of fruit from 100 randomly selected fruiting clusters on terminal shoots from the mid exterior canopy of ‘Desirable’ trees, on the side facing the OG tree. This was repeated at five locations within the orchard. Data were then analyzed using PC-SAS by PROC-CONTOUR (SAS, 1990) to derive a contour plot of fruit-set.

The second test orchard was a pure 8-ha block of 9-year-old ‘Pawnee’ trees (protandrous) planted adjacent to a 40-ha orchard made up mostly of 85-year-old ‘Stuart’ (protogynous) and ‘Schley’ (protogynous), but also with randomly dispersed “seedling” or OG trees (≥5%). The ‘Pawnee’ block was configured in 17 × 18 rows with trees spaced at 9 × 9 m. Putative pollinizers were in a contiguous U-shaped orchard immediately adjacent (20 m) to the ‘Pawnee’ block on the west and south sides and slightly set back (200 m) on the north side. There was also 800 or more other genotypes growing within 0.5 km. All trees in the ‘Pawnee’ block were evaluated for fruit-set in Oct. 1995 and 1996 by noting the number of nuts per tree. Fruit-set data were indexed to the cross-sectional area of the tree trunk and analyzed by PC-SAS for linear, quadratic, and cubic relationships. A contour map was then plotted using SAS G-CONTOUR (SAS, 1990).

Experiment 2. Fruit-set as a function of distance from pollinizer. The 15-year-old ‘Desirable’/‘Stuart’ block was evaluated for the influence of pollinator distance on fruit-set of...
the 'Desirable' 'main crop.' Fruit-set at ripening (October) was assessed, as described above, on four 'Desirable' trees in each row between the 'Stuart' pollinator rows in both 1995 and 1996. The mean number of fruit per fruiting cluster on terminal shoots on the lower half of the southern side of the tree was then determined in each of five transects across the orchard to the next row of 'Stuart' pollinizers. Fruit-set per tree was then regressed against distance from the 'Stuart' pollinizers using PC-SAS regression analysis to determine linear and quadratic relationships.

**Experiment 3. Influence of cultivar and tree age/size on relative flower maturation.** Flowering patterns, as related to tree age or size, were evaluated for six protogynous ('Stuart', 'Moneymaker', 'Moore', 'Mahan', 'Schley', and 'Delmas') and two protandrous ('Desirable' and 'Dependable') cultivars located in adjacent orchards. Trees were commercially managed, irrigated, and well spaced. This ensured that the canopy was well exposed to sunlight and wind. Thus, sampled flowers were likely to be exposed to essentially the same air temperatures and radiant energy. Trees of these cultivars were segregated into two age classes: <15 years old or >80 years old. Because flower maturity varies by location within the tree, flowers were repeatedly selected from the lower exterior of the canopy (2 to 3 m aboveground for both age classes) and monitored daily for maturation of male and female flowers. Pistillate flowers were judged mature when pollen adhered to the stigmatic surface, whereas staminate flowers were mature when pollen could be released by gently shaking four randomly selected lower exterior limbs on the southern side of the tree. Experimental design was completely randomized, with two single-tree replicates of eight cultivars of two age classes (<15 and >80 years old) for a total of 32 trees.

**Experiment 4. Influence of cultivar and temperature on relative maturation of flowers.** The influence of temperature on flower maturation was assessed on five cultivars: three protandrous ('Desirable', 'Pawnee', and 'Caddo') and two protogynous ('Stuart' and 'Summer'). Trees were <15 years old and under commercial management. The effect of temperature was estimated by exposing developing flowers to two temperature environments: 1) ambient and 2) elevated (3 to 10 °C above ambient; thus, the maximum ranged from 16 to 37 °C). Elevated temperature was attained by enclosing a terminal shoot in a transparent plastic bag tied around the 2-year-old portions of the branch. Corners were removed to leave 10-cm-diameter holes for gas exchange and temperature moderation. Terminals were bagged 3 weeks before flower maturity and temperature monitored using a thermocouple thermometer (Omega Engineering, Stamford, Conn.). Flowers were evaluated for date of maturation based on the above described criteria. The experimental design was a split-split plot arranged in a completely random manner with two replicates of five cultivars (three protandrous and two protogynous) as the main plot and two temperature treatments (ambient and above ambient) per tree as the subplots (n = 10 trees). Each temperature/treatment subplot was replicated five times per tree per main plot.

**Experiment 5. Duration of pistillate flower receptivity.** The stigmatic surface of pistillate flowers was evaluated to assess the effective pollination period. Pistillate flower clusters of five 'Cheyenne' trees (replicates), possessing good seed crops, were covered with gas-permeable pollen bags (cellulose casings) at 1 to 3 d before receptivity. 'Desirable' pollen was dusted onto flowers a predetermined number of days after the initiation of receptivity. Receptivity was based on adherence of pollen to the stigmatic surface of flowers on adjacent flower clusters on the same shoot. Pollination treatments were such that 'Desirable' pollen was applied 1, 2, 4, 6, 8, 10, 15, and 20 d after onset of pistillate flower receptivity. Pollen was lightly dusted onto flowers within the bag using a modified hypodermic syringe, with care being taken to avoid excessive application (Marquard, 1992). Twenty-five flower clusters per treatment on each of the three trees were ultimately pollinated (n = 75 clusters). As controls, open-pollinated flower clusters were tagged and monitored for fruit drop throughout the summer. Developing nuts were evaluated 3 June, 20 June, 31 July, 1 Sept., and 20 Oct. to determine the percentage of fruit retention.

**Results**

**Experiment 1. Documentation of crop loss in block-type plantings due to nonpollination.** Trees located in the center of the 'Desirable' "main crop" blocks, far from 'Stuart' pollinizers, set few fruit unless located near an OG tree (Fig. 1). This trend held in both 1995 and 1996, but 1996 observations were subjective and are not presented. Fruit-set declined as distance from OG trees increased, and declined sharply by the third tree. This produced a plume-like pattern stretching from the southwest to the northeast across the orchard. This plume was aligned with the direction of the prevailing southwesterly winds during the pollination period. These data indicate that crop load was largely reduced by insufficient availability of 'Stuart' pollen at stigma receptivity.

Fruit-set in a second test orchard, an 8-ha 'Pawnee' block, declined in 1995 as the eastward distance from pollinizers increased [Fig. 2, where set (y) = a + bx + cx²; a = 0.69, b = -4.29, c = 1.46 × 10⁻⁶; r² = 0.63]. This gradient was consistent with prevailing southwesterly winds during the pistillate flower maturity window. Conversely, fruit-set increased in an easterly manner (or northeasterly) in 1996 (where y = a + bx² + cx³; a = 1.48; b = 0.00012; c = 5.7 × 10⁻⁷; r² = 0.85). These wind-associated gradients in fruit-set are evidence of insufficient pollen availability for the entire orchard.

**Experiment 2. Fruit-set as a function of distance from pollinizer.** Fruit-set of 'Desirable' in the 'Desirable'/'Stuart' block-type orchard declined sigmoidally as distance from the 'Stuart' pollinizer increased in both 1995 and 1996 (Fig. 3). Crop loads varied such that the 1995 crop was "light" and the 1996 crop was relatively heavy. There was a decline in fruit-set in both 1995 where set (y) = a +
Crop Production

Distance from pollenizer (m)

Pollinating
O=Old tree
Y=Young tree

Protrandrous
(Type I)

Receptive

Moore
Delmas
Mahan
Banjo
Bonanza
Elegant
Moneymaker
Schley
Stuart

Dependent

Protoynous
(Type II)

Day of year

Fig. 3. Fruit-set in a large block-type orchard in 1995 and 1996 of ‘Desirable’ as a function of distance from the pollenizer (‘Stuart’). Orchard configuration was repeating blocks of 19 rows of ‘Desirable’ and four rows of ‘Stuart’. There was a delay in fruit-set (y) in 1995 (y = a + bx + cx2 (a = 1.50; b = -0.002; c = -9.35 x 10^-5; r^2 = 0.96, α ≤ 0.001) and an eastward increase in set in 1996 where y = a + bx + cx2 (a = 1.80; b = -0.003; c = -1.18 x 10^-4; r^2 = 0.95, α ≤ 0.001).

Fig. 4. Effects of cultivar and tree age/size on pollen shedding and pistil receptivity characteristics for protrandrous and protogynous pecan cultivars.

Discussion

Block-type plantings are common in the southeastern United States, a region where orchards are generally assumed to receive an abundance of pollen from nearby orchards, as well as from feral and yard trees (Woodroof and Woodroof, 1927). The present study has documented pollination deficiencies in orchards located in the heart of the southeastern pecan belt and indicates that fruit-set problems can occur due to pollenizer distance, tree aging, and temperature-induced changes in flower maturity windows. The efficacy of pollinizers can therefore be highly variable. This variation indicates that orchards should contain more than one pollinizer if fruit-set problems are to be avoided throughout the life of the planting, and that pollinizers must be planted within a few rows of the main cultivar.

The observed pollination-related crop reductions support a previous conclusion that poor cross-pollination is occurring in many southeastern orchards (Wood and Marquard, 1992). This obviously does not occur in all orchards at all times. Altered flowering windows due to tree aging or to abnormal spring temperatures, as described in the present study, or to changes in relative humidity (Woodroof, 1930; Yates and Sparks, 1993) could easily trigger major pollination-related reductions. Additionally, Woodroof (1930) observed that pollen viability changes significantly (0% to 82%) from year to year and from tree to tree, therefore potentially influencing fruit-set.

Greatly reduced fruit-set occurred when trees were more than ~49 m from a pollinator, and a 30% loss occurred only two rows from the pollinizer during a “light crop” year. This indicates that orchards of similar sized trees, spacings, and configuration should probably be established with main cultivars no more than one row from a pollinator (i.e., main cultivars in double rows flanked by pollinizers). These data also indicate that if main cultivar rows are greater than or equal to about three to four rows (greater than or equal to ~49 m) from pollinator cultivars, large crop reductions can occur in certain years. Even in relatively good years, main cultivars (rows greater than or equal to five to six rows from pollinizers) are likely to exhibit substantial crop reductions. Thus, the main cultivar section of block-type orchards should probably be less than or equal to ~10 to 12 rows, and two to three rows will probably give much better fruit set than 10 to 12 row blocks. Even then, cross-pollination-related crop reductions will probably occur, because observations by Marquard (1988) and Wood (personal observations) on ‘Western Schley’ and ‘Wichita’ block-type planting in the southwestern United States indicate that main culi-
This study also indicates that pecan exhibits reduced fruit-set if pistillate flowers fail to receive pollen within 1-4 d after becoming receptive; if no pollination occurs within 24 d, then there is likely to be little or no fruit-set. The cause of these fruit-drops remains undetermined, although visual observations of the stigmatic surface revealed they had become swollen and distorted and appeared to lose much of their moist appearance 3 to 4 d after initial receptivity. Because pollen grains on the stigmatic surface require rehydration before germination and germ tube penetrate of the stigmatic surface (Wetzstein and Sparks, 1989), insufficient moisture on the aged stigmatic surface may prevent germination and fertilization and therefore fruit-set.

Fig. 6. Influence of the time of pollination on fruit-set of 'Cheyenne' pecan. Percentage fruit retention (%) is described by the equation: y = a + b/x, where x = time interval between initial receptivity and pollination. For June 3, a = 42.6, b = 44.3, \( r^2 = 0.85, \alpha \leq 0.001 \); for June 20, a = 4.6, b = 39.7, \( r^2 = 0.94, \alpha \leq 0.001 \); for July 31, a = 3.2, b = 26.9, \( r^2 = 0.96 \); for Sept. 1, a = -2.7, b = 23.1, \( r^2 = 0.96, \alpha \leq 0.001 \).

The apparent need for pollination soon after stigma receptivity illustrates the vulnerability of orchard systems to shifts in flower maturity windows, especially in orchards composed of two cultivars with a protogynous primary cultivar. Thus, reliance on a single pollen source is risky, for pollination may occur too late. For insurance, orchards should probably contain at least two nearby pollinizers. Woodroof (1924) came to a similar conclusion, but the idea has been largely abandoned in recent years due to the assumed abundance of pollen from pollinizers nearby. Pollinizers could be segregated in rows, or cultivars possessing nutrient shapes nearly identical to that of the main cultivar could be positioned within the orchard. For example, 'Stuart'/ 'Cape Fear' or 'Desirable'/ 'Kiowa' combinations would provide enhanced pollination without a reduction in price due to mixed nuts. A comprehensive listing of flower maturity windows recently developed for southeastern U.S. conditions by Wood et al. (1997) can be used to facilitate pollinator selection.

Extrapolation of these studies suggests that southeastern orchards should be designed such that the main cultivar is pollinated by two or more cultivars possessing good reciprocal complementarit under normal conditions. Many patterns of cultivar placement within orchards are available, based upon relative cultivar value, tree removal strategy, and a variety of cultural or pest management strategies. Whatever the design selected, these data indicate that main cultivar trees should be as close to a pollinizer as is practical and that blocks greater than two to three rows will probably exhibit substantial pollination-related crop reductions.

**Literature Cited**


