Influence of Plant Bioregulators on Pecan Flowering and Implications for Regulation of Pistillate Flower Initiation

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Abstract. Mitigation of alternate bearing (AB) through regulation of floral initiation of pistillate flowers is central to improving cropload management of pecan [Carya illinoiensis (Wangenh.) K. Koch] trees and orchards. The present study examines the influence of key bioregulators {i.e., an auxin [as B-napthalenecarboxylic acid (NAA)], a cytokinin [6-benylamino purine (6-BA)], an ethylene generator (ethephon), and an auxin transport inhibitor [2,3,5-triodobenzoic acid (TIBA)]} on subsequent season pistillate flowering. Gibberellic acid (i.e., GA₃) and NAA inhibited, whereas prohexadione–calcium (P-Ca; calcium 3-oxido-5-oxo-4-propionylcyclohex-3-enecarboxylate), ethephon, and BA + TIBA promoted floral initiation when topically applied to canopies before the kernel filling stage of seed development. These bioregulators exhibit potential for integration into a bioregulator-based strategy to mitigate AB by selective and timely use in “off” or “on” cycle years, depending on the bioregulator. Field studies provide evidence that a “cytokinin–gibberelin balance” with partial modulation by auxin and ethylene, acts in the endogenous primordial environment of floral meristems as a second-level signal regulating a key step in a three-step process for initiation of pistillate flowers in pecan. This establishes a new model for explaining pistillate flower initiation in pecan and a basis for designing future research on the control and management of pistillate flowering and AB.

Natural selection operating over evolutionary time has produced pecan as an economically important species that exhibits pronounced biennial-like alternations in seed production as a strategy for ensuring long-term reproductive success. This year-to-year variation in pistillate flowering, and subsequent cropload, is termed AB. Although AB-linked variation in pistillate flowering likely increases individual fitness in natural habitats, it is also a major impediment to greater horticultural domestication and is the primary biology-based impediment to horticultural enterprises (Wood, 1991). Excessive year-to-year variability in pistillate flowering limits tree and orchard profitability, thus adversely affecting producers, processors, and consumers through instabilities in nutmeat supply, quality, and price (Sparks, 1974, 1975; Wood, 1991). Although the specific processes regulating AB in pecan remain ambiguous, the trait tightly links to floral initiation processes occurring within bud meristems within the tree’s canopy. Horticultural manipulation of pistillate flowering and mitigation of AB in commercial pecan orchards currently targets minimization of tree stress with orchard management strategies directly or indirectly targeting key exogenous biotic and abiotic stressors (Wood et al., 2003). These include sunlight, nutrient elements, and water as essential resources and pathogens, arthropods, and weeds as potentially harmful pests. Cropload thinning before, or at the time of, inception of kernel (i.e., primarily cotyledon) filling of developing seeds also acts to moderate AB by increasing subsequent year pistillate flowering (Smith and Gallott, 1990; Smith et al., 1993; Wood, 1995). This fruit/seed association implicates one or more seed-associated phloem mobile phytohormones in regulation of floral initiation (i.e., the production of meristems of clearly recognizable flower primordia and includes all preceding reactions that are required if flowers are to be initiated).

Regulation of floral initiation in trees depends on processing of environmental and/or endogenous cues (Amasino, 2010) with initiation in most large-seeded temperate woody perennial angiosperms being primarily controlled by endogenous cues consistent with processing through an autonomous flowering pathway involving phytohormones (Wilkie et al., 2008). Floral initiation in pecan is therefore likely to involve an autonomous flowering pathway as a key step in its floral initiation process (Wood et al., 2003). Like with many other tree-fruit species (Schmidt et al., 2009), florally induced bud primordia on heavy cropload trees (i.e., “on” year of AB cycle) are likely exposed to different phytohormonal environments than are primordia of induced buds on light cropload trees (i.e., “off” year of AB cycle). This raises the possibility that timely application of phytohormones or bioregulators to tree canopies might alter the phytohormonal environment of primordia in such a way as to enable control of pistillate flowering by pecan farmers.

The efficacy and horticultural potential of bioregulators to control the “on” and “off” flowering phases of pecan trees has not been reported despite considerable circumstantial evidence that endogenous phytohormones are involved in floral initiation processes (Barnett and Mielke, 1981; Rohla et al., 2007a, 2007b; Wood, 1982, 1984a, 1984b, 1991, 2003; Wood and McMeans, 1981; Wood et al., 2003). A variety of natural and synthetic bioregulators are efficacious for control of floral initiation processes in several polycarpic perennial crops and involve timely use of floral promoters [generally ethephon and P-Ca and NAA or GA₃ in certain situations (Looney et al., 1985)] in “on” years to promote return flowering the following “off” year and use of floral inhibitors [gibberelic acids (GA₃,₄,₅) (Greene, 2000) and auxin analogs (e.g., NAA in certain cases) in “off” years to decrease subsequent year flowering. It is unknown whether these promoters and inhibitors similarly affect pecan flowering in “on” and “off” years.

Commercial pecan production enterprises need better horticultural tools for managing flowering and AB. Successful development and exploitation of such tools depend on acquiring better understanding of floral initiation processes operating in pecan. This study assesses certain promising bioregulators for activity and/or influence on pecan flowering and examines how their interactions influence pistillate flower initiation. It reports that several synthetic bioregulators possess potential as horticultural tools for controlling pistillate flowering and AB in pecan and, based on observed influence of bioregulators on flowering, presents a “three-level signaling” model explaining regulation of pistillate flower initiation in pecan trees.

Materials and Methods

General orchard characteristics

A series of experiments were conducted between 2002 and 2010 using orchards located at Byron, GA, a humid climatic zone (lat. +32°39’54” N, long. +83°44’31” W) with an average elevation of ≈156 m (509 ft). The dominant orchard soil is a Faceville sandy loam soil (fine, Kaolinitic, thermic Typic Kandiudult soil). The site has a freeze-free growing period of ≈280 d with annual precipitation of ≈1.3 m (51 inches). The test cultivars were Summer, Oconee, or Pawnee, grafted to an open-pollinated ‘Elliott’ seedling rootstock.
Orchards were managed according to the Georgia Extension Services guidelines for pests, fertilizers, etc. (Hudson et al., 2007). Trees also received subsurface drip irrigation with drip emitters rising to the soil surface at 1-m intervals to supplement tree water needs. Parallel irrigation lines ran the length of the tree rows, positioned ≥1.2 m on either side of tree trunks. Drip emitters delivered water at 3.78 L·h⁻¹ for ≥12 h·d⁻¹, depending on water needs, throughout the growing season. Irrigation lines were within an herbicide strip maintained in a vegetation-free state using glyphosate (Monsanto, St. Louis, MO). Trees received annual broadcast applications of nitrogen, potassium, phosphorous, magnesium, and calcium as needed, based on previous July leaf analysis, at principal leaf development growth stage “11” (Finn et al., 2007). Canopies also received foliar sprays of micronutrients (zinc, nickel, copper, manganese, iron, and boron) during canopy expansion (growth stage “11-17”) each spring. Orchard management included routine foliar sprays of pesticides to control pecan scab [Fusicladosporium effusum (Winter)] disease and various arthropod pests. Pest management practices produced fruits with little pecan scar damage and little or no fruit drop or fruit damage by arthropod pests.

Study 1: Influence of gibberellic acid, prohexadione-calcium, and ethephon on flowering

This experiment tested the hypothesis that flowering in pecan can be manipulated by timely applications of GAs, P-Ca (an inhibitor of GA synthesis) and ethephon (a generator of ethylene). Efficacy testing used either 25-year-old ‘Pawnee’ trees or 5-year-old ‘Ocoee’ trees in an “on” year of crop production. Expt. 1: Influence of bioregulators and partial defoliation on return flowering of ‘Pawnee’ shoots. This study used individual ‘Pawnee’ trees exhibiting an overall light cropload (less than ≈5% of terminals fruiting). Treatments were 1) non-treated control [i.e., the control included Silwet L-77®, an organosilicone surfactant, at 0.025% v/v used in all plant growth regulator treatments at this rate; Helena Chemical Co.]; and 2) GA₃ (ProGibb® at 50 mg a.i./L; Valent BioSciences Corporation). Treatment application involved three biweekly foliar sprays beginning the second week of June and extending to early July (i.e., six applications). Treated trees were in their fourth leaf. The study consisted of the two aforementioned treatments structured as a RCB consisting of 10 single-tree experimental units (n = 20). Measured parameters were mean number of pistillate flowers or young fruit per tree at anthesis the next spring. Statistical analysis was by least squares ANOVA at P ≤ 0.05.

Phase 2: Gibberellins, and return flowering of ‘Oconee’ trees. This study assessed the effect of GA₃ on pistillate flower production of ‘Pawnee’ trees in their “off” year when the overall orchard cropload was very light (less than ≈5% terminals fruiting), thus likely ensuring a relatively large “on” year for pistillate flower production at anthesis the next spring. Experimental design was a RCB using two treatments: 1) nontreated control (i.e., included Silwet L-77® as described previously, being in both treatments); and 2) GA₃ (ProGibb® at 50 mg a.i/L). Influence on subsequent year (i.e., the “on” year) flowering was assessed in consequence of spraying (through an air-blast sprayer) tree canopies with GA₃ during early morning to achieve leaf soak. The first application was the first week of August, ≈12 weeks after full bloom (AFB), and repeated biweekly for a total of three applications ending in early September, ≈21 to 31 d before fruit ripening.

Treatments were arranged as a RCB with eight single-tree blocks (n = 24). Experimental units consisted of single ‘Pawnee’ trees blocked by relative cropload. Measured parameters were 1) percentage of terminal shoots displaying pistillate flowers at anthesis; 2) average number of pistillate flowers per terminal; and 3) mean number of pistillate flowers per flowering cluster. A randomly selected population of 20 terminal shoots per tree per treatment was used for measurement of relevant parameters. Measurements were made at anthesis the following spring. Statistical analysis was performed on parameters by two-way least squares ANOVA at P ≤ 0.05.

Expt. 3: Influence of prohexadione–calcium on flowering of whole trees. The second study assessed the effect of P-Ca (Apogee®) on pistillate flower production. It was initiated when ≈25-year-old AB ‘Pawnee’ trees were in a moderate “on” year (≥60% terminals fruiting), thus ensuring an “off” year for pistillate flower production at anthesis the next spring. Trees were treated three times at weekly intervals with one of five P-Ca concentrations (i.e., 0, 62.5, 125, 250, and 500 mg a.i/L). Timing of treatment application incorporated two distinctly different seed development stages. These were during the liquid endosperm stage before initiation of kernel filling versus cotyledon development after initiation of kernel filling. Initiation of kernel filling was considered to be at the gel stage of seed development when the inner surface of the liquid-filled endosperm coenocyte (the large liquid-filled central vacuole) exhibits a noticeable gelatinous layer of cells [i.e., the stage in which endosperm cellularization is initiated by the formation of radial microtubular systems at the surface of the endosperm nuclei, just before cell wall formation of the first aleuroid structures that eventually grow in a centripetal manner, producing cell files oriented toward the center of the central vacuole of the endosperm until it is completely cellularized and seed cotyledons are filled (Olsen, 2001)].

Pre-kerfilling treatments began ≈4 weeks (mid-July) before initiation of kernel filling followed by retreatment at 3 weeks and again at 2 weeks. Post-kerfilling treatments began at the time of initiation of kernel filling (i.e., ≈13 Aug.), with applications again 1 week and 2 weeks later.

Experimental design was a RCB with four blocks of five continuous variable P-Ca treatments at two fruit development stages, thus giving 10 treatments (n = 40). Treatment application was through an air-blast sprayer (at 935 L·ha⁻¹) to all portions of the tree canopy during early morning to ensure leaf soak and bioregulator uptake. Assessment of treatment effects on pistillate flower production was the next spring at anthesis, an “off” year, by assessing the percentage of terminal shoots displaying pistillate flower clusters. Data were analyzed using curvilinear regression.

Expt. 4: Influence of prohexadione–calcium and ethephon on flowering of whole trees. Results from the described Phase 2 study led to a study testing efficacy of 1) P-Ca [a GA metabolic inhibitor; calcium 3-oxido-5-oxo-4-propionylcyclohex-3-ene-2-carboxylate (Apogee); BASF Corporation, Research Triangle Park, NC, at 500 mg a.i./L]; 2) ethephon ([2-chloroethyl]phosphonic acid, Ethrel® at 100 mg a.i/L; Buyer CropScience, Inc., Calgary, Alberta, Canada); and 3) “P-Ca plus ethephon” (i.e., “Apogee + Ethrel”) as potential floral promoters in young pecan trees. Foliar applications were made on 1 June, 21 June, and again 14 July (i.e., 3, 6, and 9 weeks post-full bloom). Treatment of ‘Oconee’ trees was through an
air-blast spray when trees were in their fifth leaf.

The experimental design consisted of the four aforementioned treatments structured as a RCB containing 10 blocks (n = 40). Experimental units consisted of single-tree experimental units blocked by cropload. Measured parameters were 1) percentage of terminal shoots exhibiting pistillate flowers (based on a random sampling of 20 shoots); and 2) average number of pistillate flowers per terminal shoot. Measurements were made at anthesis the next spring. Statistical analysis was performed on parameters by two-way least squares ANOVA at P ≤ 0.05. Mean separation for main effects and interaction using Student’s t test.

**Study 2: Influence of cytokinin and auxin on flowering**

Expt. 1: Influence of benylaminopine, triiodobenzoic acid, and naphthalenacetic acid on flowering of individual terminal shoots when applied in the “on” phase. This study was initiated when overall orchard cropload was relatively heavy (≈60% terminals flowering) for ≈25-year-old ‘Sumner’ trees. Treatments were 1) nontreated control (i.e., the control included Silwet L-77 at 0.025% v/v used in all plant growth regulator treatments at this rate); 2) 6-BA [or N-(phenyl-methyl)-1H-purine-6-amino, at 50 mg a.i./L]; 3) NAA (K-Salt Fruit Fix) at 10 mg a.i./L; 4) TIBA (at 100 mg a.i./L); and 5) TIBA + BA. Application of treatments occurred as three bieweekly foliar sprays beginning the first week of July and extending to mid-August. It consisted of the five aforementioned treatments structured as a RCB of five single-tree blocks (n = 25). A set of 15 san-exposed terminal shoots served as the experimental unit. Measured parameters were 1) percentage of new terminal shoots displaying pistillate flowers; and 2) mean number of pistillate flowers per flower cluster at anthesis the next spring. Statistical analysis was by least squares ANOVA at P = 0.05. Mean separation is by Student’s t test at P = 0.05.

Expt. 2: Influence of benylaminopine, triiodobenzoic acid, and naphthalenacetic acid on flowering of this tree. This experiment tested the hypothesis that return flowering in pecan is able to be manipulated through treatment of “on”-phase tree canopies with cytokinin, auxin, or an auxin inhibitor. Testing was done using 8-year-old ‘Summer’ trees in an “on” crop year. Experiment initiation was when the overall orchard cropload was moderate (≈40% to 50% of terminals fruiting) and efficacy evaluation was the next spring at anthesis. The experimental design was a RCB of seven discrete variable treatments. Treatments were 1) nontreated control (with Silwet L-77 at 0.025% v/v in all treatments); 2) cytokinin [6-BA or N-(phenyl-methyl)-1H-purine-6-amino; a synthetic cytokinin; formulated as MaxCel® (1.9% BA) and applied at 50 mg a.i./L]; 3) auxin [NAA, a synthetic auxin formulated as K-Salt Fruit Fix (AMVAC, Los Angeles, CA); applied at 10 mg a.i./L]; 4) auxin transport inhibitor TIBA at 100 mg a.i./L; Sigma, St. Louis, MO); and 5) cytokinin plus auxin transport inhibitor (“MaxCel + TIBA”).

Treatment application was with a hand-gun power sprayer during early morning to achieve leaf soak. The first application was 5 June and again for five more applications at 2-week intervals for a total of six applications. Immediately after spraying the entire tree canopy, 15 “on” terminal shoots (i.e., bearing fruit) and 15 “off” terminal shoots (i.e., not bearing fruit) were randomly selected in the canopy periphery and tagged for subsequent year evaluation.

The experimental design consisted of the five aforementioned treatments structured as a RCB with 10 single-tree blocks (n = 50). Experimental units were single trees blocked by cropload. Measured parameters were 1) number of staminate flower groups per previous season terminal shoot; 2) number of pistillate flower clusters arising from both terminal and lateral shoots origination in the previous season’s terminal shoot; and 3) total number of pistillate flowers (i.e., florets) produced by the pistillate flower population associated with “2.” Parameters were measured for both the “on” and “off” shoot populations at anthesis the next spring. Statistical analysis was performed on parameters by two-way least squares ANOVA at P ≤ 0.05. Mean separation was by two-way least squares ANOVA at P = 0.05 using Student’s t test.

Results and Discussion

**Study 1: Influence of gibberellic acid, prohexadione–calcium, and ethephon on flowering**

Expt. 1: Influence of bioregulators and partial defoliation on return flowering of “off”-phase ‘Pawnee’ shoots. Treatment of terminal shoots with certain bioregulators reduced floral initiation based on observation of pistillate flowers at anthesis the next season (Table 1). Shoots previously “off” (also in an “off” tree) flowered at a rate of ≈65% (i.e., nontreated control) with partially defoliated (manually) shoots flowering at only 1%, indicating that foliage contributes one or more floral inducing factors. Shoots treated the previous season with GA3 or GA4+7 plus BA were largely vegetative the next spring, exhibiting only 26% and 29%, respectively, as many flowers as the control. Either BA or P-Ca alone did not influence the percentage of flowering terminal shoots. Bioregulators also influenced the number of florets per floral cluster on those shoots that did flower. Control shoots and P-Ca-treated shoots produced 4.4 florets per floral cluster, but BA slightly reduced florets per cluster, and both GA3 and GA4+7 plus BA substantially reduced florets per cluster. Thus, not only did GAs reduce the percentage of shoots flowering, but also the number of florets per shoot cluster. Taken together, it appears GA3 and GA4+7 in combination, and possibly GA4 or GA3 independently, inhibit floral initiation when applied the previous year during the period of floral induction within pecan buds. This indicates that GAs likely act as floral inhibitors in pecan, and P-Ca exhibits efficacy through inhibition of GA biosynthesis.

It appears that amount of foliage per shoot affects floral initiation. Because treatment trees returned with a heavy “on” crop, and non-structural shoot carbohydrates were not measured, it is unknown whether this foliage effect is the result of energy reserves available to bud meristems or to dependence of phytohormones otherwise produced by foliage. Regardless, leaves produced one or more floral promoters during the early stages of floral initiation, and results are consistent with phytohormone regulation. This indicates that reproductively induced primordia depend on a leaf originating phloem mobile promoter for continuation of floral primordia or to dependence of phytohormones otherwise produced by foliage. Regardless, leaves produced one or more floral promoters during the early stages of floral initiation, and results are consistent with phytohormone regulation.

Table 1. Influence of treating “off”-phase ‘Pawnee’ terminal shoots with foliar sprays of plant bioregulators [gibberellic acid (GA3, ProGibb at 50 mg a.i./L), prohexadione–calcium (P-Ca; Apogee; at 500 mg a.i./L), gibberelins A4+7 and N-benzyladenine (Promalin) at 50 mg a.i./L], and N-benzyladenine (BA, at 50 mg a.i./L), and partial defoliation (only the apical-most leaflet of each compound leaf remaining) of terminals on pistillate flower bloom the next spring. *

| Treatment                  | Percentage of terminal shoots displaying pistillate flowers (%) | Number of pistillate flowers per terminal clusters (no.) |
|----------------------------|---------------------------------------------------------------|---------------------------------------------------------|
| Control                    | 65 aβ                                                        | 4.4 aβ                                                  |
| Partial defoliation        | 1 b                                                          | 0.7 d                                                   |
| GA3                        | 17 b                                                         | 1.4 cd                                                  |
| BA                         | 56 a                                                         | 2.9 b                                                   |
| GA4+7 and BA               | 19 b                                                         | 1.7 c                                                   |
| P-Ca                       | 59 a                                                         | 4.4 aβ                                                  |

*Treated shoots within canopies received three biweekly applications beginning early July.

**Phase 1: Gibberelins and return flowering of ‘Pawnee’ trees.** Canopy treatment of young “off” year ‘Oconeé’ trees with GA3 during June and early July markedly reduced number of pistillate flowers per tree at anthesis the next spring (Table 2); thus, treatment with GA3 led to production of only 32% as much fruit per tree as the nontreated control. This supports a conclusion from Expt. 1 above that GA3 inhibits pistillate flower initiation in pecan and is especially active

![Image](https://via.placeholder.com/150)
when applied within 3 to 7 weeks after full bloom. It appears to be within this timeframe that pistillate flower induction occurs, being completed when fruit are nearing maximum size (Amling and Amling, 1983; Wetzstein and Sparks, 1983), usually around early August. One or more molecular GA species trigger similar inhibitory activity in other fruit tree crops (Bangerth, 2009; Bertling and Bangerth, 1995) and grape (Vitis vinifera L.; Vasconcelos et al., 2009). It is unclear from the present study whether GA3 acts in pecan to prevent floral induction, to reverse induction processes, or both. It is also possible that it simply stops or retards physiological differentiation processes occurring during the early stages of physiological differentiation from the vegetative state.

**Phase 2: Gibberellin3 and return flowering of ‘Pawnee’ trees.** ‘Pawnee’ trees treated with GA3 during their “off” bearing phase reduced pistillate flower production the next “on” year relative to the non-treated control (Table 2). GA3-treated canopies had fewer terminals displaying pistillate flowers to only 69% of the control, the average number of pistillate flowers per terminal to 47% of the control, and number of pistillate flowers per flower cluster to 69% of the control. Thus, GA3 not only reduces the number of terminal shoots on the tree that displayed pistillate flowers the next spring, but also reduces the number of florets in each flower cluster at anthesis. In apple, GAs inhibits floral bud formation when applied as a single application from before bloom to several weeks post-bloom (Greene, 1989, 2000). It is possible that in pecan that either floral induction or initiation is sensitive to GAs over a period of several weeks with sensitivity depending on molecular GA species, concentration, and possibly crop load (as developing pecan fruit contain gibberellin-like substances and probably export these basipetally) and leaf area (Wood, 1982, 1984a).

**Expt. 3: Influence of prohexadione-calcium on flowering of whole trees.** Application of P-Ca, a metabolic inhibitor of GA synthesis (Evans et al., 1999), to “on”-phase trees increases return pistillate flowering in the next “off”-phase year when P-Ca is applied at

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of "on" phase flowering on floral bud exposed to a high endogenous GA environment during a portion of the pre-kernel filling phase of the previous "off"-phase year. In apple, it appears that GA7 is more inhibitory than is GA3 (Tromp, 1982). Perhaps different GAs vary in efficacy for inhibiting flowering in pecan and GA3 might not be the most active molecular species. These observations indicate that one or more GAs play a major role in regulation of this. AB. This is supported in that 1) GA-like substances are present in pecan fruit (Wood, 1982, 1984a); 2) paclobutrazol, an inhibitor of GA biosynthesis, can increase flowering in pecan (Wood, 1984b); 3) the date of maximum rate of change in the pistillate flower action spectra of pecan is during the dough stage of seed development when GAs are likely present (Wood, 1995); and 4) the efficacious period for mechanical fruit thinning, to ensure return flowering, is about the same time as, or before, the dough stage of seed development (Smith et al., 1993) when GAs are likely to be high. A similar inhibitory activity in other crops, as reviewed by Dennis and Neilsen (1999), and by Bangerth (2009), further supports a floral inhibitory role for GAs.

Study 2: Influence of cytokinin and auxin on flowering

Expt. 1: Influence of benylamino purine, triiodobenzoic acid, and naphthaleneacetic acid on flowering of individual shoots when applied in the on-phase. Treatment of "on"-phase terminal shoots of "on" trees with synthetic cytokinin- or auxin-type bio-regulators potentially influences return flowering (Table 4). Because the next spring was an "off" year, trees produced few flowering shoots with only 13% of control shoots flowering the next spring. The percentage of subsequent season flowering shoots was unaffected by either TIBA or BA alone but greatly reduced by NAA. Surprisingly, shoot exposure to "TIBA + BA" led to 53% of terminal shoots flowering, this being a four-fold increase over the control. Certain bio-regulators also influenced the number of pistillate flowers per flowering shoot with NAA and TIBA reducing and TIBA + BA slightly increasing pistillate florets per cluster. It is somewhat surprising that the inhibitory effect of TIBA, when used alone, is promotive when shoots receive both TIBA and BA. These findings hint that auxins are inhibitory to floral initiation in "on" years and that floral initiation processes within bud meristems of terminal shoots are influenced by a cytokinin-auxin interaction.

Expt. 2: Influence of benylamino purine, triiodobenzoic acid, and naphthaleneacetic acid on flowering of trees. Statistical analysis indicated a significant "bioregulator" treatment effect on return flowering characteristics of 'Pawnee' pecan with flowering being either reduced or increased depending on bio-regulator treatment (Table 5). The number of pistillate flower clusters at anthesis the next spring and, arising from the terminal and lateral shoots originating from the previous growing season's terminal shoot, was unaffected by TIBA or BA treatments but was substantially fewer when treated with NAA and greater when treated with TIBA + BA. This confirms the promotive effect observed in the previously described shoot-based study. The number of pistillate flower clusters was reduced to 21% of the control by NAA and increased 3.5-fold by TIBA + BA. In the case of the total number of pistillate flowers per terminal and associated lateral shoots, TIBA alone had no effect, whereas NAA and BA alone reduced pistillate flowers and TIBA + BA substantially increased the number of pistillate flowers. The number of pistillate flowers was reduced to 16% of the control by NAA and 79% of the control for BA, yet there was a 1.8-fold increase by TIBA + BA. The number of staminate catkin triads per terminal shoot was unaffected by TIBA or BA alone, reduced to 12% of the control for NAA alone, and was slightly increased (16%) by TIBA + BA. Although shoot treatment with a combination of TIBA plus BA (i.e., TIBA + BA) led to several floral clusters per shoot, there were fewer flowers per cluster. The combination of TIBA + BA led to a pronounced positive synergism regarding floral initiation and subsequent flowering on a shoot basis with the ultimate expression of pistillate flowers being potentially influenced by interacting factors (e.g., the carbohydrate environment). This synergism is suggestive of partial regulation of floral initiation processes governed by the relative auxin-cytokinin balance with a high cytokinin:auxin ratio enabling expression of pistillate and staminate flower buds during anthesis the next spring.

Table 3. Influence of foliar sprays of gibberellic acid (ProGibb at 100 mg a.i./L), prohexadione-calcium (P-Ca; Apoge; at 500 mg a.i./L), and ethephon (Ethrel*, at 100 mg a.i./L) on pistillate flower bloom the next spring of young 'Occane' pecan trees.

| Treatment                        | Percentage of terminal shoots displaying pistillate flowering (%) | Avg number of pistillate flowers per terminal (no.) |
|----------------------------------|-----------------------------------------------------------------|---------------------------------------------------|
| Control                          | 0.9 a                                                           |                                                    |
| P-Ca                             | 22 b                                                            | 2.3 c                                             |
| Ethephon                         | 20 b                                                            | 1.5 b                                             |
| P-Ca + ethephon                  | 16 b                                                            | 1.6 b                                             |

*Canopies received three biweekly applications beginning early June.

Analysis of variance by least squares means. Means within a column are not significantly different at P = 0.001 by Student's t test.

Table 4. Influence of treating "on"-phase 'Pawnee' terminal shoots with foliar sprays of plant bioregulators [N'-benzyladenine (BA, at 50 mg a.i./L), 2,3,5-triiodobenzoic acid (TIBA, at 100 mg a.i./L), and 1-naphthaleneacetic acid (NAA, K-Salt Fruit Fix at 10 mg a.i./L)] of terminals on pistillate flower bloom the next season.

| Treatment                        | Percentage of terminal shoots displaying pistillate flowers (%) | Number of pistillate flowers per flowering terminals (no.) |
|----------------------------------|----------------------------------------------------------------|----------------------------------------------------------|
| Control                          | 13 b                                                            | 3.0 b                                                     |
| BA                               | 20 b                                                            | 3.0 b                                                     |
| TIBA                             | 12 b                                                            | 1.9 c                                                     |
| NAA                              | 1 c                                                             | 0.2 d                                                     |
| TIBA + BA                        | 53 a                                                            | 3.9 a                                                     |

*Canopies received three biweekly applications beginning early July.

Analysis of variance by least squares means with means within a column followed by the same letter are not significantly different at P = 0.05 by Student's t test.

Table 5. Influence of foliar sprays of NAA (at 10 mg a.i./L), 2,3,5-triiodobenzoic acid (TIBA at 100 mg a.i./L), and BA (at 50 mg a.i./L) on subsequent season pistillate and staminate flowering of young 'Pawnee' trees.

| Treatment                        | Pistillate floral clusters per previously treated 1-year-old terminal (no.) | Pistillate flowers per previously treated 1-year-old terminal (no.) | Stamine catkin triads per terminal shoot (no.) |
|----------------------------------|-----------------------------------------------------------------------------|---------------------------------------------------------------------|-----------------------------------------------|
| Control                          | 1.9 b                                                                        | 6.8 b                                                               | 10.6 bc                                       |
| BA                               | 1.9 b                                                                        | 5.4 c                                                               | 11.2 ab                                       |
| TIBA                             | 2.1 b                                                                        | 7.0 b                                                               | 9.8 c                                         |
| NAA                              | 0.4 c                                                                        | 1.1 d                                                               | 1.3 d                                         |
| TIBA + BA                        | 6.7 a                                                                        | 12.0 a                                                              | 12.3 a                                        |

*Trees were treated in a moderate "on" year (~40% of terminal shoots fruiting) with six sprays at biweekly intervals beginning early June with production of flowers being assessed at anthesis the next spring.

Analysis of variance by least squares means. Means within a column followed by the same letter are not significantly different at P = 0.0001 by Student's t test.

Mean number of pistillate floral clusters produced on the previously treated 1-year-old terminal. Means were 1.5b versus 1.8a for "on" and "off" shoots, respectively. The mean number of pistillate flowers per previously treated 1-year-old terminal when terminal was 4.6a versus 5.0a for "on" and "off" shoots, respectively. The mean number of stamine catkin triads per terminal shoot was 6.5b versus 8.7a for "on" and "off" shoots, respectively. The "on" versus "off" effect was significant at P = 0.001 by analysis of variance for the number of pistillate floral clusters per previous 1-year-old terminal shoot and for the mean number of stamine catkin triads per terminal shoot.
There was a significant effect on the number of pistillate flower clusters per previously treated 1-year-old terminals and the number of stamine catkin triads per terminal shoot but no effect on the number of pistillate flowers per previously treated 1-year-old terminal (Table 5, see footnote). For the two former parameters, values were slightly less for shoots previously in the “on” bearing phase than in the “off” bearing phase. This small difference in return flowering in relation to previous-year bearing phase of treated shoots is likely because trees were young, and differences might have been much greater in older more physiologically mature trees.

Potential as tools for mitigation of alternate bearing in pecan

The present study illustrates that several commercially available bioregulators exhibit potential as horticultural tools for mitigation of AB through regulating pistillate flower initiation and subsequent flowering. It is clear that GA₃, GA₄+7, and NAA possess theoretical use as floral inhibitors and are useful for reducing return flowering in otherwise “on”-phase trees when canopies are properly exposed during the previously “off”-phase season. Conversely, P-Ca, ethephon, and BA + TIBA possess theoretical use as floral promoters and therefore potential for increasing return flowering of otherwise “off”-phase trees when canopies are properly exposed during the previous “on”-phase season. This raises the possibility of bioregulator-based regulation of floral initiation in commercial orchards by orchard managers by using GAs and/or NAA (and probably other auxins as well) in severe “off” years to decrease flowering during the subsequent “on” phase. However, the development of bioregulator-based AB control strategies in orchards is problematic in that success depends on considerable research to identify the appropriate bioregulator mix, rate, and timing in relation to tree cropload and on satisfying regulatory constraints.

Implications regarding flowering in pecan

The ability of NAA to inhibit flowering in pecan is consistent with a “second messenger” role, as presented by Bangert (2006), for auxins (e.g., indole-3-acetic acid) as an inhibitor of flowering in perennial trees with endogenous auxin being substantially controlled by both GA concentration and molecular species within developing fruit. In pome crops, GA-treated fruit are strong auxin excluders and therefore potential for increasing return flowering in the subsequent “off”-phase year. However, the development of bioregulator-based AB control strategies in orchards is problematic in that success depends on considerable research to identify the appropriate bioregulator mix, rate, and timing in relation to tree cropload and on satisfying regulatory constraints.

It is noteworthy that neither BA nor NAA alone positively influence pistillate flower initiation or production but are promotive in combination. This might be because auxin regulates cytokinin concentration and vice versa (Nordstrom et al., 2004), thus enabling relatively homeostatic crosstalk because the two phytohormones regulate plant developmental events. In the absence of TIBA, it might be that enough auxin is produced by pecan organs (i.e., developing fruit, apices, and young foliage) to rapidly metabolize BA (or other cytokinins) to non-bioactive forms, because auxin (e.g., IAA) can rapidly upregulate cytokinin oxidase and block cytokinin biosynthetic pathways, thus reducing the endogenous cytokinin environment to which buds are exposed the previous year when fl orally induced meristems were subject to reversion during the physiological differentiation phase of floral initiation. It is possible that BA alone did not enhance pistillate flowering because other necessary long-distance signals (e.g., ethylene and possibly others) were not quantitatively or qualitatively compatible with increased flowering, or its rapid deactivation prevents an effect on floral initiation processes. The fact that floral initiation increases with application of BA in combination with TIBA indicates that BA is potentially an active cytokinin molecular species for floral initiation in pecan.

Auxins (e.g., NAA) and cytokinins (e.g., BA) are thought to act somewhat in the physiological differentiation stage downstream of activation of key floral induction genes (Horvath, 2009), as activated by florigen (i.e., a leaf produced phloem mobile 20-kDa protein that moves to bud meristems and apices to induce the floral process; Shalit et al., 2009; Zeevaart, 2008). Findings here appear consistent with a positive effect of TIBA, a potent auxin transport inhibitor, observed for flowering in other fruit tree species and a promotive effect of cytokinin on flowering in several perennial tree species (Bangerth, 2009). Because auxin can inhibit cytokinin biosynthesis (Nordstrom et al., 2004), it is possible that flower initiation processes in pecan are not only substantially regulated by GAs through inhibition, as described previously, but also through cytokinins’ promotive effects. If so, then the promotive effect of cytokinin is potentially subject to modulation by auxin from young foliage and fruit. Thus, a cytokinin–GA balance, or equilibrium, appears to regulate “on”-phase pistillate flower initiation with equilibrium potentially regulated by a cytokinin–auxin interaction with either auxin or ethylene acting as quantitative “long-distance” signals (Bangerth, 2009). Several findings support this possibility, including those by Palm et al. (1988) in which increasing endogenous auxin reduces endogenous cytokinin through effects on cytokinin oxidase activity; by Ito et al. (2001) in which decreasing auxin leads to increased bud cytokinin; by Li et al. (2003) in which removing the shoot apex (a rich source of auxin) greatly increases endogenous cytokinin concentration; by Chunjiang and Bangerth et al. (2000) in which they found a mutually antagonizing interaction between cytokinin and auxin; by Nordstrom et al. (2004) in which auxin regulates cytokinin levels and vice versa; and by Rashotte et al. (2005) in which cytokinin can trigger biosynthesis of ethylene. Thus, the present study supports the idea that a cytokinin–GA balance, as thought to operate in Vitis spp. (Vasconcelos et al., 2009), regulates the early stages of floral initiation in pecan, and auxin and ethylene modulates this balance.

Conclusions

Findings here support the theory that the autonomous floral pathway dominates floral initiation process acting between floral induction (i.e., processes required for evocation) and vernalization (inductive process requiring low temperature) or evocation (processes required for irreversible commitment to initiate flower primordia) occurring before actual flower development (processes occurring from after evocation until anthesis). The present study supports conclusions by others, working with different crops, that endogenous factors regulating at least one flowering phase includes GAs, auxin (indole-3-acetic acid), ethylene, and cytokinins. These phytohormones collectively act and interact as a long-distance “level-two floral signal” to regulate flowering through chromatin modification (Bangerth, 2009) after florigen (i.e., florigen as the “level-one floral signal”) triggered chromatin modification that initiates floral induction physiology. Additionally, sugar signaling also plays key roles in developmental processes such as flowering (Gibson, 2005), in which complex interplay between phytohormones and sugars affects each other (Leon and Sheen, 2003). In the case of pecan, it is clear that sugars are intimately involved in one or more processes controlling floral initiation (Smith and Waugh, 1938; Sparks, 1974, 1975; Wood, 1989, Wood et al., 2004; Worley 1979a, 1979b) with their role being expressed in association with successful vernalization and subsequent floral evocation (Wetzstein and Sparks, 1983; Wood, 1989, 1995); however, sugars are not the sole factor driving pistillate flower initiation nor AB (Rohila et al., 2007a, 2007b; Smith et al., 2007). Timely use of bioregulators may provide the means of affecting key downstream floral initiation processes through effects on sugar-based chromatin modulation (Gibson, 2003, 2005; Koch, 1996; Li et al., 2003) such as that occurring during floral vernalization and evocation.

Although the present study does not quantify endogenous phytohormone concentrations, nor presence of associated receptors, in association with floral primordia or their environment, results nevertheless provide evidence that GAs, auxins, ethylene, and cytokinin influence floral initiation in pecan; thus, at least one key process is largely controllable by the action and/or interaction of one or more molecular species of these phytohormones. It is therefore postulated that pecan’s post-induction phase of pistillate flower initiation is largely regulated by the endogenous “cytokinin–GA”
environment to which young bud meristems are exposed during late spring and early summer before kernel filling. This cytokinin–GA balance appears to be subject to modulation by ethylene and auxin exported from foliage and/or fruit and/or influenced by tree and/or organ stress. The cytokinin–GA balance likely affects chromatin-related activities within meristematic cells as well as downstream changes in floral initiation processes needed to prepare pistillate floral primordia for vernalization and evocation.

When the results of the present study are considered within the context of finding by Zeevaart (2008) on the role of florigen an initiating floral signal, Bangerth (2009) on the role of phytohormones in the primordial environment, and others regarding the role of carbohydrate reserves and their effect on primordia formation, it appears that pistillate flower initiation in pecan involves three distinct phases of chromatin modification before new flowers appear in early spring. The following is a proposed model explaining initiation of pistillate flowers in pecan.

Three sequential phases of chromatin modification control pistillate flower initiation, beginning with 1) a foliage produced phloem translocated florigen acting as a first-level signal to initiate phase-one chromatin-modifying inductive processes in young bud primordia; 2) then phase-two chromatin modification regulated by translocated phytohormones, from foliage and/or fruit, acting in the primordia environment during early post-induction as a “cytokinin–GA ratio” based second-level signal subject to modulation by auxin and ethylene; and 3) finally phase-three chromatin modification regulated by concentration of one or more non-structural carbohydrates (e.g., sucrose) acting in the primordia environment during vernalization as a third-level signal enabling floral development in preparation for anthesis.

This model identifies a testable theory possessing several distinct stages as potential candidates for horticultural intervention for controlling pistillate flowering. It also provides a basis for future research toward better understanding of flowering and AB processes and subsequent development of horticultural tools and strategies enabling greater year-to-year stability in nutmeat yield and quality from trees and orchards of pecan and other woody perennial polycarpic species.

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