Bending Shoots Stimulates Flowering and Influences Hormone Levels in Lateral Buds of Japanese Pear

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Abstract. Flower development of the lateral buds was accelerated in Japanese pear [Pyrus pyrifolia (Burm.) Nak.] when vertical shoots were bent at a 45° angle in late June. The indole-3-acetic acid (IAA) concentration in lateral buds on vertical (control) shoots increased in mid-July, while remaining nearly constant in bent shoots. The abscisic acid (ABA) concentration of buds in bent shoots rose between 4 July and 15 Aug., whereas control shoots exhibited an increase in concentration followed by a decline. Gibberellin(a7, GA4+7) concentration was high on 16 June, and then declined by 4 July, with the decline being greater in bent shoots. Gibberellin(a7) concentration was higher in the buds on vertical shoots than in those on bent shoots for much of July. The concentrations of zeatin-type cytokinins (CKs) in lateral buds were higher in bent shoots than in vertical shoots. Bending of pear shoots may weaken competition between buds and other organs through altering hormone levels in lateral buds, resulting in acceleration of flower development.

Flower initiation in Japanese pear occurs in late June after shoot elongation ceases (Banno et al., 1986). Floral primordia appear in apple (Malus ×domestica Borkh.) buds only when a critical number of nodes has formed on the axis (Abbott, 1970; Fulford, 1966). If the critical node number is not reached by the end of the growing season, the bud remains vegetative. This suggests that every bud can become a flower bud, but not all buds form flowers because of unsuitable conditions (Tsujikawa et al., 1990).

In Japanese pear, flower buds usually are formed on apices of spurs, and on terminal and lateral buds of shoots. The degree of flower bud formation differs among cultivars. ‘Kosui’, the leading cultivar in Japan, has few flower buds because of unsuitable conditions (Tsujikawa et al., 1990).

Hormone extraction and purification. Buds were collected randomly on 16 June, 4 and 17 July, and 2 and 15 Aug. from the tagged shoots of each subgroup (seven trees). Immediately following collection, the samples were frozen in liquid N2 and stored at –80 °C until analysis. The length of 15 shoots per plot on the trees for anatomical study were measured weekly from 5 June to 17 July. Bud developmental stage was determined microscopically in buds of five uniform shoots per plot. The nine stages of flower bud development as described by Peng and Iwahori (1994) were used: 0 = vegetative; 1 = early stage of initiation; 2 = final stage of initiation; 3 = lateral flower bud apparent; 4 = bract and lateral flower buds beginning to develop; 5 = sepal primordia developing; 6 = petal primordia forming; 7 = stamen primordia forming; 8 = pistil primordia forming; 9 = petal and pistil well-developed. Each of the selected shoots bore between 19 and 23 lateral buds. Apical buds, and lateral buds located at the fifth node below the apex, the center node, and the fifth node from the base were also examined. The lateral buds were located near the center of the distal third, central third, and proximal third of the shoot, respectively, and will be referred to as apex, bud node 15–19, bud node 10–12, and bud node 5, respectively. Shoots were still growing at the beginning of bud observations; thus, the stage of development of bud node 15–19 differed over time. Buds were sampled four times: on 21 June, 21 July, 21 Aug., and 18 Sept.

Hormone extraction and purification. Buds were analyzed for indole-3-acetic acid (IAA), abscisic acid (ABA), gibberellin (GA1, GA4, GA7, GA4+7), trans-zeatin (Z), and trans-zeatinriboside (9RZ). Hormonal analysis was performed according to Takahashi and Yamaguchi (1986) with a slight modification. Each hormone was extracted from frozen buds and fractionated with high-performance liquid chromatography (HPLC, ModelLC-6AD, Shimazu Corp., Kyoto, Japan) equipped with a UV detector. Indole-3-acetic acid and ABA were estimated by gas chromatography–mass spectrometry (GC-MS), and GAs and cytokinins (CKs) by enzyme-linked immunosorbent assay (ELISA). Gibberellin and GA4 and GA7 cannot be separated by the HPLC method used, so these GAs were estimated and...
evaluated as GA_{1+3}, and GA_{4+7}, respectively.

Each sample was divided in half; one half was used for GA, IAA, and ABA analyses and the other for CK analyses. Samples for GAs, IAA, and ABA were homogenized and extracted in 40 mL of 80% aqueous acetone containing butyl hydroxy toluene (BHT) at 0.1 g L^{-1}. At this point 50 ng of [^{13}C_6]-IAA and 100 ng of [^{2H}(D)3]-ABA were added as internal standards for quantification. The solution was then filtered and the residue was re-extracted twice with 40 mL of 80% acetone. The pooled extracts were then evaporated in vacuo to an aqueous solution, and solvent-partitioned as shown in Fig. 1 to obtain an acidic ethyl acetate–soluble (AE) fraction. Samples for CK analysis were similarly extracted and concentrated to give an aqueous solution which was subjected to solvent partitioning (Fig. 2) to obtain a fraction soluble in n-butyralcohol (n-BuOH).

The AE and n-BuOH fractions were dried in vacuo and then dissolved separately in a small amount of methanol. Each methanol solution was loaded on a Pegasil ODS HPLC column (10 mm i.d. × 150 mm; Senshu Scientific Co. Ltd., Tokyo) equilibrated with 5% acetonitrile, and eluted with a gradient of water–acetonitrile containing 0.5% acetic acid. The elution conditions were as follows: 0–5 min 5% acetonitrile; 5–30 min linear gradient from 5% to 30% acetonitrile; 30–50 min linear gradient from 30% to 80% acetonitrile; 30 min 80% acetonitrile wash. The flow rate during the entire elution program was 1.5 mL min^{-1}. The retention times of IAA (36 min), ABA (47 min), GA_{1+3} (31 min), GA_{4+7} (60 min), Z (9 min), and [9R]Z (20 min) were determined by running authentic standards under the same conditions.

GC-MS analysis for IAA and ABA. Indole-3-acetic acid and ABA were identified and estimated by GC-MS. Prior to the analysis, the putative ABA fractions were methylated with ethereal diazomethane, and the putative IAA fractions were first methylated, then trimethylsilylated with N-methyl-\(N\)-\((trimethylsilyl)\)trifluoroacetamide. For GC-MS a JOEL (Nippon Denshi Co. Ltd., Tokyo) JMX-DX303 GC-MS equipped with a multiple ion detector was used. The column (0.53 mm i.d. × 15 m) was packed with OV-1. Injection port, separator, and chamber temperatures were 230, 250, and 200 °C, respectively. The ionization voltage was 70 eV. The temperature program was as follows: 0–1 min at 120 °C; 1–11 min linear gradient from 120–280 °C, followed by 10 min at 280 °C. Selected ion monitoring was conducted at m/e 202/208 and 261/267 for IAA/[^{13}C_6]-IAA, and 134/137, 162/165, and 190/193 for ABA/[D_3]-ABA, respectively. The relative intensities of the ions m/e 202/208 and 190/193 were traced to identify IAA and ABA, respectively. Hormones were quantified from a standard curve using authentic IAA and ABA.

Identification and semi-quantification of GA and CK. GA and CK were identified and estimated based on their retention times on HPLC and immuno-reactivities to specific antibodies.

The GA_{1+3} fractions were analyzed with polyclonal antibodies raised against GA_{1+3} bovine serum albumin (BSA) conjugate via the C-7 carboxyl group of the GA, as reported by Atzorn and Weiler (1983), and the GA_{4+7} fractions were analyzed with monoclonal antibodies against GA_{4} prepared by Nakajima et al. (1991). The cross-reactivities of the antibodies raised against GA_{1+3} to GA_{4} and against GA_{1+3} to GA_{4} were 70% and 72%, respectively, on a molar basis. The ELISA assay was performed as described by Atzorn and Weiler (1983) for GA_{1+3}, and by Nakajima et al. (1991) for GA_{4+7}.

Zeatin and [9R]Z were analyzed by ELISA with polyclonal antibody against [9R]Z, which is highly specific for Z and [9R]Z, as described by Weiler (1980). Synthesis of [9R]Z-immunogens is reported in Weiler and Spanier (1981). The quantified values were evaluated as Z-equivalent for Z and [9R]Z. The cross-reactivity of antibody raised against [9R]Z to Z was 44%, on a molar basis.

Statistical procedures. Data on stage of flower development were analyzed using the analysis of variance (ANOVA) procedure, and means were separated with Duncan’s new multiple range test. Data for hormone concentrations were compared using Tukey’s honestly significant difference.

Results

Shoot length. Shoots elongated until early July (Fig. 3). The bending treatment was initiated when shoot elongation had almost ceased; thus, the elongation rates and final lengths of the vertical vs. the bent shoots were not significantly different.

Flower development. Since in no case was the interaction of treatment and position significant by ANOVA, only main-effect means are presented (Table 1). In both treatments, flower development in the apical buds occurred earlier than in the lateral buds. Variabili-
it was large within each plot, so the effect of bending shoots on flower development was nonsignificant on most dates except for 18 Sept., when bending treatment significantly accelerated flower development regardless of bud position.

**Hormonal changes.** Indole-3-acetic acid concentration in lateral buds of control shoots rose in early- to mid-July, before declining in mid-August (i.e., the concentration rose after shoot elongation ceased) (Fig. 4A). In contrast, the IAA concentration in lateral buds on bent shoots remained relatively constant throughout the measurement period.

Abscisic acid concentration in both treatments increased until early July, then declined in the control (Fig. 4B). However, it continued to increase in buds on bent shoots and was significantly higher than in the controls on 15 Aug.

The concentration of GA4+7 was high initially, then declined in both treatments (Fig. 5A) as shoot elongation ceased (Fig. 3). The decline was greater in bent than in control shoots. Concentrations of GA1+3 were much lower than those of GA4+7 in both treatments, and changed less throughout the measurement period (Fig. 5B).

The concentration of Z declined throughout the period of measurement in control shoots (Fig. 6A), but bending increased the concentration, which was significantly higher than that of the control on 4 and 17 July and 2 Aug. Bending had a similar effect on [9R]Z concentration, and was higher in the bent than in the vertical shoots on 17 July and 2 Aug. (Fig. 6B).

**Discussion**

Japanese pear growers bend vertical shoots in late June when shoot elongation has almost ceased (Fig. 3). The treatment accelerates lateral flower development regardless of the bud position (Table 1). Tromp (1967) discovered that horizontally grown apple trees form more flower buds and terminate growth sooner than do vertical trees. Other researchers (Luckwill, 1970; Owens, 1991) reported a similar relationship between number of flowers and cessation of shoot elongation. In our experiment, however, bending accelerated bud development without affecting shoot elongation, probably because the treatment was applied near the end of the shoot elongation period. A similar observation was made by Tromp (1972).

Bud development was more rapid on bent than on control shoots regardless of bud position. The buds for hormonal analysis were sampled regardless of position. Therefore, the results are meaningful only for the general (regardless of bud position) comparison of hormones and flowering. Another experiment must be conducted to establish a specific relationship between hormones and flowering at specific positions.

Previous studies involving applications of plant growth regulators or estimations of endogenous hormonal levels have shown relationships between flower development and levels of plant hormones, although there are many conflicting or uncertain results (reviewed by Buban and Faust, 1982; Hoard, 1984; Owens, 1991). The effect of auxin on flowering is unclear (Buban and Faust, 1982; Jackson and Sweet, 1972), since it has increased flowering in some studies (Southwick and Weeks, 1950), but not others (Luckwill, 1970). On the contrary, inhibitors of auxin efflux reportedly stimulate flowering (Edgerton and Hoffman, 1965; Luckwill, 1970; Owens, 1991).

Gibberellins inhibit flowering of apple and pear (Banno et al., 1985b; Griggs et al., 1970; Higashiuchi et al., 1990; Hoard, 1978; Huet, 1973; Owens, 1991). Auxin and GAs may affect flowering by increasing the sink strength of apical buds (i.e., vegetative shoot elongation) (Cline, 1994; Ho, 1994; Sachs, 1977, 1993). However, Looney et al. (1985) reported that GA may increase flowering in apple. These conflicting reports may indicate

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**Table 1. Effects of bending and position on seasonal change in developmental stage in buds of ‘Kosui’ Japanese pear.**

| Date       | 21 June | 21 July | 21 Aug. | 18 Sept. |
|------------|---------|---------|---------|----------|
| Treatments (T) | Control | Bent    |         |          |
|             | 0.0     | 0.7     | 2.0     | 1.9      |
|             | 0.0     | 1.0     | 2.2     | 3.6      |
| Position (P)x |         |         |         |          |
| Apex       | 0.0     | 1.9     | 4.2     | 4.0      |
| Node 14–18 | 0.0     | 0.5     | 2.0     | 3.5      |
| Node 10–12 | 0.0     | 0.4 b   | 1.5 bc  | 3.0 a    |
| Node 5     | 0.0     | 0.4 b   | 0.7 c   | 0.9 b    |

**Analysis of variance**

| Main effect | Treatments (T) | Position (P) |
|-------------|----------------|--------------|
|             | NS             | NS           |
|             | NS             | NS           |
|             | NS             | NS           |
|             | NS             | NS           |

**Fig. 4. Seasonal change in (A) IAA and (B) ABA concentrations (ng·g−1 fresh weight) in lateral buds of control (○) and bent (●) shoots of ‘Kosui’ Japanese pear. Values are means ±SE, n = 3 (SE bars present only when larger than symbol). Vertical bar indicates Tukey’s honestly significant difference (P ≤ 0.05).**

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**Table 2. Effects of bending and position on seasonal change in developmental stage in buds of ‘Kosui’ Japanese pear.**

| Date       | 21 June | 21 July | 21 Aug. | 18 Sept. |
|------------|---------|---------|---------|----------|
| Treatments (T) | Control | Bent    |         |          |
|             | 0.0     | 0.7     | 2.0     | 1.9      |
|             | 0.0     | 1.0     | 2.2     | 3.6      |
| Position (P)x |         |         |         |          |
| Apex       | 0.0     | 1.9     | 4.2     | 4.0      |
| Node 14–18 | 0.0     | 0.5     | 2.0     | 3.5      |
| Node 10–12 | 0.0     | 0.4 b   | 1.5 bc  | 3.0 a    |
| Node 5     | 0.0     | 0.4 b   | 0.7 c   | 0.9 b    |

**Analysis of variance**

| Main effect | Treatments (T) | Position (P) |
|-------------|----------------|--------------|
|             | NS             | NS           |
|             | NS             | NS           |
|             | NS             | NS           |
|             | NS             | NS           |

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|------------|---------|---------|---------|----------|
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|             | 0.0     | 1.0     | 2.2     | 3.6      |
| Position (P)x |         |         |         |          |
| Apex       | 0.0     | 1.9     | 4.2     | 4.0      |
| Node 14–18 | 0.0     | 0.5     | 2.0     | 3.5      |
| Node 10–12 | 0.0     | 0.4 b   | 1.5 bc  | 3.0 a    |
| Node 5     | 0.0     | 0.4 b   | 0.7 c   | 0.9 b    |

**Analysis of variance**

| Main effect | Treatments (T) | Position (P) |
|-------------|----------------|--------------|
|             | NS             | NS           |
|             | NS             | NS           |
|             | NS             | NS           |
|             | NS             | NS           |
that GAs influence flowering without affecting growth, or that the response of the meristem to GA may vary depending on the developmental stage (Bonnet-Masimbert and Zaerr, 1995; Kondo and Gemma, 1993).

In contrast, CK are reported to increase flower bud development in some fruit species (Banno et al., 1985b; Luckwill, 1970; McLaughlin and Greene, 1984; Ramirez and Hoad, 1981), probably by activating cell division within the buds (McGaw, 1995; McLaughlin and Greene, 1991).

The direct role of ABA in flower bud development is thought to be negligible (Hoad, 1981), probably by activating cell division within the buds (McGaw, 1995; McLaughlin and Greene, 1991).

While shoot bending influenced hormone levels, certain of these changes may have been the result of changes in other hormone(s) or alterations in the physical characteristics of the shoot. For example, shoot bending reduces the basipetal transport of IAA (Bangerth, 1990, 1993), and this could reduce IAA content of lateral buds. Van Huizen et al. (1995) reported that auxins stimulate the conversion of GA from inactive to active forms in peas (Pisum sativum L.). Therefore, the decrease in GA observed in this experiment may have been a result of reduced IAA levels. A decrease in the auxin concentration following decapitation can increase the cytokinin concentration either by increasing transport in the xylem (in bean; Bangerth, 1994), or by increasing metabolism in the buds (in pea; King and van Staden, 1987). So, CK levels also may be tied to auxin levels. Additionally, the relation between ethylene production and auxin level is also suggested in apple; ethylene reduces polar auxin transport (Sanyal and Bangerth, 1998), whereas auxin application increases the ethylene level (Robiaille and Leopold, 1974). Tsujikawa et al. (1990) reported that the removal of competing sinks, such as young leaves or fruits, promoted flowering of Japanese pear. Sink strength and/or dominance strongly correlates with cytokinin concentration in the organs and with the auxin transport rate of each organ (Bangerth, 1993; Ho, 1994; Sachs, 1993). Thus, the relationship between auxin transport and flower development after shoot bending merits re-examination.

In conclusion, shoot bending advanced flower development. This advancement was associated with notable changes in hormone levels. We propose that the bending of pear shoots alters hormone levels, thereby increasing the ability of the lateral buds to compete for assimilates.

**Literature Cited**

Abbott, D.L. 1970. The role of budscales in the morphogenesis and dormancy of the apple fruit bud. p. 65–80. In: L.C. Luckwill and C.V. Cutting (eds.). Physiology of tree crops. Academic Press, London.

Atzorn, R. and E.W. Weiler. 1983. The immunoassay of gibberellins. II. Quantiﬁcation of GA$_{3}$, GA$_{4}$, and GA$_{5}$ by ultra-sensitive solid-phase enzyme immunoassays. Planta 159:7–11.
Bangerth, F. 1990. Polar auxin transport in fruit trees in relation to fruit drop. Acta Hort. 275:461–468.
Bangerth, F. 1993. Polar auxin transport as a signal in the regulation of tree and fruit development. Acta Hort. 329:70–76.
Bangerth, F. 1994. Response of cytokinin concentration in the xylem exudate of bean (Phaseolus vulgaris L.) plants to decapitation and auxin treatment, and relationship to apical dominance. Planta 194:439–442.
Banno, K., S. Hayashi, and K. Tanabe. 1985a. Effects of SADH and shoot-bending on flower bud formation, nutrient components and endogenous growth regulators in Japanese pear (Pyrus serotina Rehd.). J. Jpn. Soc. Hort. Sci. 53:365–376. (in Japanese with English abstract and figures).
Banno, K., S. Hayashi, and K. Tanabe. 1985b. Effect of application of growth regulators on flower bud formation in Japanese pear (Pyrus serotina Rehd.). Fac. Agr., Tottori Univ. Bul. 38:11–15. (in Japanese).
Banno, K., S. Hayashi, and K. Tanabe. 1986. Morphological and histological studies on flower bud differentiation and development in Japanese pear (Pyrus serotina Rehd.). J. Jpn. Soc. Hort. Sci. 55:258–265.
Bonnet-Masimbert, M. and J.B. Zaerr. 1987. The role of plant growth regulators in promotion of flowering. Plant Growth Regul. 6:13–35.
Buban, T. and M. Faust. 1982. Flower bud induction in apple trees. Wiley, New York.
C R O P  P R O D U C T I O N
Edgerton, L.J. and M.B. Hoffman. 1965. Some physiological responses of apple to N-dimethyl amino succinic acid and other growth regulators. Proc. Amer. Soc. Hort. Sci. 86:28–36.
Edgerton, L.J. and M.B. Hoffman. 1966. The morphogenesis of apple buds. III. The inception of flowers. Ann. Bot. 30:207–219.
Griggs, W.H., G.C. Martin, and B.T. Iwakiri. 1970. The effect of seedless versus seeded fruit development on flower bud formation in pear. J. Amer. Soc. Hort. Sci. 95:243–248.
Higashitani, Y., K. Kido, T. Ichii, T. Nakamishi, Y. Kawai, and T. Ozaki. 1990. The effect of gibberellins and growth retardants on in vitro flowering of the vegetative apex of Japanese pear. Scientia Hort. 41:223–232.
Ho, L.C. 1994. Fruit growth and sink strength. p. 101–124. In: C. Marshall and J. Grace (eds.). Fruit and seed production. Aspects of development, environmental physiology and ecology. Cambridge Univ. Press, U.K.
Hoad, G.V. 1978. The role of seed derived hormones in the control of flowering in apple. Acta Hort. 80:93–103.
Hoad, G.V. 1984. Hormonal regulation of fruit-bud formation in fruit trees. Acta Hort. 149:13–23. Huet, J. 1973. Floral initiation in pear trees. Acta Hort. 34:193–196.
Jackson, D.I. and G.B. Sweet. 1972. Flower initiation in temperate woody plants. Hort. Abstr. 42:9–2.
King, R.A. and J. van Staden. 1987. The metabolism of N6-(4')-isopentenyl)[3H]adenine by isolated organs of Pisum sativum. J. Plant Physiol. 131:181–190.
Kojima, K., Y. Yamada, and M. Yamamoto. 1995. Effects of abscisic acid injection on sugar and organic acid contents of citrus fruit. J. Jpn. Soc. Hort. Sci. 64:17–21.
Kondo, S. and H. Gemma. 1993. Relationship between abscisic acid (ABA) content and maturation of the sweet cherry. J. Jpn. Soc. Hort. Sci. 62:63–38.
Leopold, A.C., K.M. Brown, and F.H. Emerson. 1972. Ethylene in the wood of stress trees. HorticScience 7:175.
Looney, N.E., R.P. Pharis, and M. Noma. 1985. Promotion of flowering in apple trees with gibberellin A1 and C-3 epigibberelin A1, Planta 165:292–294.
Luckwill, L.C. 1970. The control of growth and fruitfulness of apple trees, p. 237–254. In: L.C. Luckwill and C.V. Cutting (eds.). Physiology of tree crops. Academic Press, London.
Lyndon, R.F. 1994. The environmental control of reproductive development, p. 9–32. In: C. Marshall and J. Grace (eds.). Fruit and seed production. Aspects of development, environmental physiology and ecology. Cambridge Univ. Press, U.K.
Marcelle, R. and C. Sironval. 1963. Effect of gibberellic acid on flowering of apple trees. Nature 197:405.
McGaw, B.A. 1995. Hormonal regulation of apical dominance, p. 98–117. In: P.J. Davies (ed.). Plant hormones. Kluwer Academic, Dordrecht.
McLaughlin, J.M. and D.W. Greene. 1984. Effects of BA, GA10, and daminozide on fruit set, fruit quality, vegetative growth, flower initiation, and flower quality of ‘Golden Delicious’ apple. J. Amer. Soc. Hort. Sci. 109:34–39.
McLaughlin, J.M. and D.W. Greene. 1991. Fruit and hormones influence flowering of apple. I and II. Effects of hormones. J. Amer. Soc. Hort. Sci. 116:450–453.
Mor, Y. and N. Zieslin. 1987. Plant growth regulators in apple. J. Hort. Sci. 62:63–38.
Peng, S.-A. and S. Iwahori. 1994. Morphological and cytological changes in apical meristem during flower bud differentiation of Japanese pear, Pyrus pyrifolia Nakai. J. Jpn. Soc. Hort. Sci. 63:313–321. (in Japanese with English abstract and figures.)
Rakunag, J., H. Gemma, and S. Iwahori. 1995. Flower bud formation in Japanese pear trees under adverse conditions and effects of some growth regulators. Jpn. J. Trop. Agr. 39:1–6.
Ramirez, H. and G.V. Hoad. 1981. Effects of growth substances on fruit bud initiation in apple. Acta Hort. 120:131–136.
Rohilaite, H.A. and C. Leopold. 1974. Ethylene and the regulation of apple stem growth under stress. Physiol. Plant. 32:301–304.
Sachs, R.M. 1977. Nutrient diversion: An hypothesis to explain the chemical control of flowering. HortScience 12:220–222.
Sachs, T. 1993. The role of auxin in plant organization. Acta Hort. 329:162–168.
Sanyal, D. and F. Bangerth. 1998. Stress induced ethylene evolution and its possible relationship to auxin-transport, cytokinin levels, and flower bud induction in shoots of apple seedlings and bearing apple trees. Plant Growth Regul. 24:127–134.
Sax, K. 1962. Aspects of ageing in plants. Annu. Rev. Plant Physiol. 13:489–506.
Southwick, F.W. and W.D. Weeks. 1950. Some attempts to thin apples with naphthaleneacetic acid type materials after calyx. Proc. Amer. Soc. Hort. Sci. 56:70–75.
Takahashi, N. and I. Yamaguchi. 1986. Analyses of endogenous plant hormone levels throughout the life cycle of higher plants. Acta Hort. 176:47–57.
Tromp, J. 1967. Fruit-bud formation and shoot growth in apple in relation to gravity. Naturwissenschaften 54:95.
Tromp, J. 1972. Effects of growth-regulating substances and tree orientation on growth and flower bud formation in apple. J. Hort. Sci. 47:525–533.
Tsujikawa, T., T. Ichii, T. Nakamishi, T. Ozaki, and Y. Kawai. 1990. In vitro flowering of Japanese pear and the effect of GA10. Scientia Hort. 41:233–245.
Van Huizen, R., J.A. Ozga, D.M. Reinecke, B. Twitchin, and L.N. Mander. 1995. Seed and 4-chloroindole-3-acetic acid regulation of gibberellin metabolism in pea pericarp. Plant Physiol. 109:1213–1217.
Waring, P. 1970. Growth and its co-ordination in trees, p. 1–21. In: L.C. Luckwill and C.V. Cutting (eds.). Physiology of tree crops. Academic Press, London.
Weiler, E.W. 1980. Radioimmunoassays for transzeatin and related cytokinins. Planta 149:155–162.
Weiler, E.W. and K. Spanier. 1981. Phytohormones in the formation of crown gall tumors. Planta 153:326–337.