Annual Bloom-time Phosphorus Fertigation Affects Soil Phosphorus, Apple Tree Phosphorus Nutrition, Yield, and Fruit Quality

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Abstract. A randomized, complete block, split-plot experimental design with six replicates was established and maintained annually for the first five fruiting seasons (1999 to 2003) in a high-density apple [Malus sylvestris (L.) Mill var. domestica (Borkh.) Mansf] orchard on M.9 rootstock planted in Apr. 1998. Main plot treatments involved eight different nutrient regimes, each containing three tree subplots of each of five different cultivars (Ambrosia, Cameo, Fuji, Gala, and Silken). This report compares a +phosphorus (P) treatment, involving annual fertigation at bloom time of 20 g P/tree as ammonium polyphosphate (10N–15P–0K), to a –P treatment. Both treatments also received nitrogen, potassium, and boron nutrients through fertigation. Drip fertigation of P increased 2 M KCl-extractable P to 0.4-m depth within 0.5-m distance of the drippers. Leaf and fruit P concentrations were consistently increased by the +P treatment with few differences among cultivars. P-fertigated trees also had a 20% increase in cumulative yield overall cultivars during the first five fruiting seasons. Standard fruit quality measurements, including fruit size, soluble solids concentration, titratable acidity, and red coloration were unaffected by P application. However, reductions in incidence of water core at harvest, increased resistance to browning, and elevated antioxidant content of harvested fruit measured in some years imply a role for P in apple membrane stability. The cumulative results indicate that applications of 20 g P as ammonium polyphosphate annually at bloom would be advantageous for apples receiving adequate fertigated applications of nitrogen, potassium, and boron. Best apple performance was associated with leaf P concentrations above 2.2 mg g⁻¹ dry weight and fruit P concentrations between 100 and 120 mg kg⁻¹ dry weight.

Phosphorus fertilization of orchard apple trees has received much less attention than nitrogen (N) and potassium (K) fertilization as a result of limited reports of positive responses to phosphorus (P) fertilization in the historical literature (Boyton and Oberly, 1966). Furthermore, unit area whole tree P nutrient demands have been calculated to be low, ≈20 kg ha⁻¹ for 14-year-old ‘Golden Delicious’ apple trees yielding 90 t ha⁻¹ spaced at 500 trees ha⁻¹ in contrast with much higher N (≈120 kg ha⁻¹) and K (≈200 kg ha⁻¹) requirements (Neilsen and Neilsen, 2003). Few nutrient estimates are available for high-density plantings, which can approach 4000 trees ha⁻¹ for ‘super spindle’.

More recently, several conditions have been identified when apples respond to P fertilization. These include times when apple root length is limited, as when trees are newly planted (Taylor and Goubran, 1975) and when replant disorders further inhibit root growth (Neilsen and Yorston, 1991), or when low soil P levels limit P availability to roots (Cripps, 1987; Raese, 1998). Responses to P application have included increased vigor and accelerated flowering of newly planted trees (Neilsen et al., 1990; Taylor and Goubran, 1975) and increased cropping and increased P concentrations in leaf and fruit tissue of mature trees (Cripps, 1987; Raese, 1998). Foliar sprays of soluble P compounds have increased fruit P concentrations and reduced susceptibility to low temperature breakdown (LTB) in ‘Cox’s Orange’ (Johnson and Yagorovtjan, 1978) and LTB and firmness of ‘Mcintosh’ apples (Webster and Lidster, 1986), but it is not known whether soil P applications would result in similar improvements in fruit quality. Responsiveness to P fertilization has often been made difficult by the absence of unique leaf P deficiency symptoms at low tissue P concentrations (Benson and Covey, 1979). Little is known about the seasonal requirements for P, although these cumulative responses suggest that apple P demand may peak at a time of intense cell division and meristematic activity as might occur in spring when simultaneous strong fruit, root, and shoot growth occurs.

Drip fertigation has improved the availability of P to apple trees (Neilsen et al., 1999) by allowing mass flow delivery of high P concentrations directly to the root surface and thereby proving as effective at increasing first-year tree P uptake as P application in the planting hole (Neilsen et al., 1993). P fertigation has therefore become a standard first-year recommendation for growers in southern interior British Columbia (BC) (BC Ministry of Agriculture and Lands, 2007). However, little is known of the persistence of the effect when P is fertigated annually.

This study was undertaken to measure the effect of a single annual P application through fertigation at bloom on performance of a range of potentially important, nontraditional apple cultivars planted at high density on a dwarfing rootstock. Assessment of soil and tree P status and apple yield and quality response was emphasized.

Materials and Methods

In Apr. 1998, an experimental block of five apple [Malus sylvestris (L.) Mill var domestica (Borkh.) Mansf] cultivars (Ambrosia, Cameo, Fuji, Gala, and Silken) were planted at high density on the dwarfing rootstock M.9 in rows separated by 3 m with 1 m between trees in each row. In the first year, all trees were fertigated with the industry standard NP regime involving daily application of calcium nitrate (15.5N–0P–0K; cumulative 47 g N per tree) within 8 weeks of planting and 20 g P per tree as ammonium polyphosphate (10N–15P–0K) soon after planting on 4 June 1998. Beginning in 1999, a randomized complete block, split-plot experimental design was imposed and maintained for the next five growing seasons. Main plots comprised of eight different fertigation regimes, whereas subplot consisted of the five apple cultivars, previously indicated, randomly planted in three-tree plots in each of the six replications for each fertigation treatment. Two border apple trees separated each main plot fertigation treatment and border rows were planted to completely surround the experimental block. Pertinent to this article were two fertigation treatments, which compared the presence (+P) or absence (–P) of P in otherwise comparable fertigation treatments involving annual applications totaling ≈0.17 g boron (B) per tree (except 1999 when ≈0.5 g B was applied per tree) and N (application of 160 mg L⁻¹ N as calcium nitrate) both applied daily from 0 to 4 weeks post full bloom (pfb). Potassium as potassium chloride was also fertigated daily from 4 to 8 weeks pfb supplying ≈20 g K per tree per year. The +P treatment involved a single P application...
of 20 g P per tree of ammonium polyphosphate, annually soon after bloom (based on the ‘Ambrosia’ cultivar) on 21, 25, 24, 16, and 16 of May 1999 to 2003, respectively. Each fertigation treatment had a separate irrigation line with emitters installed on the appropriate line according to the experimental randomization. Each treatment tree was irrigated by two 4-L-h⁻¹ pressure-compensating emitters (The Toro Company, El Cajon, CA) placed 0.3 m either side of the tree in the tree row. To minimize water stress, irrigation was applied daily in response to the previous day’s evaportranspiration demand and automatically scheduled by atmometer so that water application varied directly in response to evaporative demand (Parchomchuk et al., 1996).

Trees were trained to a slender spindle system supported by posts and grown in a 2.0-m-wide herbicide strip maintained by applications of 1 kg·ha⁻¹ glyphosate each year in early May, midsummer, and early fall. Conventional foliar application of zinc and insect and disease control procedures followed standard commercial recommendations (BC Ministry of Agriculture and Lands, 2007).

The experimental site was located on a Skaha loamy sand (Wittneben, 1986), an Aridic Haploxeroll, extensively planted to orchards or vineyards in southern British Columbia. Coarse-textured soils such as these have limited nutrient and waterholding capacities and have previously been shown to be susceptible to the development of K and B deficiency under drip irrigation (Neilsen et al., 1995).

Composite samples of 30 leaves from the midportion of extension shoots of the current year’s growth from each tree of the three tree plots were collected 15 July, 20 July, 18 July, 11 July, and 7 July 1999 to 2003, respectively, ≈8 weeks after full bloom (based on the ‘Ambrosia’ cultivar). For the ‘Fuji’ cultivar, similar leaf samples were collected 4 weeks (2000 to 2003) and 12 weeks (1999 to 2003) after full bloom. All samples were oven-dried at 65 °C and ground in a stainless steel mill. From 1999 to 2001, a 250-mg subsample was digested for 0.75 h on a block steel mill. From 1999 to 2001, a 250-mg solution (of chilled dH₂O (chilled to 4 °C) and homogenized with a model PT 10/35 Polytron homogenizer (Brinkmann Instruments, Rexdale, Ontario, Canada)) at a speed setting of four for 30 s. The resultant slurry was centrifuged at 15,000 × g for 15 min and the supernatant decanted into a clean test tube. This was then diluted 1:9 in dH₂O. A 0.5-mL aliquot of the resultant diluted extract was added to 1 mL of 0.2 mL dH₂O in 1 mL of 0.2 mL DPPH and the absorbance response calibrated against standard solutions of ascorbic acid.

To measure water-soluble antioxidant content, 5 g powdered, frozen tissue was added to 20 mL of chilled dH₂O (chilled to 4 °C) and homogenized with a model PT 10/35 Polytron homogenizer (Brinkmann Instruments, Rexdale, Ontario, Canada) at a speed setting of four for 30 s. The resultant slurry was centrifuged at 15,000 × g for 15 min and the supernatant decanted into a clean test tube. This was then diluted 1:9 in dH₂O. A 0.5-mL aliquot of the resultant diluted extract was added to 1 mL of 0.2 mL dH₂O in 1 mL of 0.2 mL DPPH and the absorbance response calibrated against standard solutions of ascorbic acid.
Before the termination of the experiment (and removal of trees), soil sampling was undertaken in Spring 2005 at three different distances (0, 0.5, and 1.0 m) perpendicular to the tree rows from one of the drip emitters for the centrally located ‘Fuji’ tree for five replicates of each of the two P treatments (+P). At each distance, soil samples were collected at 0.1-m depth increments beginning at the soil surface (0 to 0.1 m) and extending to the 0.5- to 0.6-m soil layer. Soil samples were placed in cold storage at 4 °C until analyses. Twenty grams of moist soil was extracted in 100 mL of 2.0 M KCl for 1 h before filtration and determination of NO$_3$-N, NH$_4$-N, and PO$_4$-P using a flow injection analyzer and manufacturer’s procedures (O.I. Analytical, College Station, TX). pH was determined on a 10-g soil sample after equilibration in 20 mL of water for 30 min.

Analysis of variance was performed on all leaf and fruit nutrients, yield, and fruit quality characteristics according to the experimental design (SAS, 1989). Data were analyzed as a split-plot design with six replicates and fertigation as main plot treatments with individual degree of freedom contrasts comparing +P and –P treatments. Subplots were random three-tree plots of each of the five different cultivars. Percent data (red color, incidence of water core) were arcsine-transformed before analyses. As expected, cultivar (subplots) effects were significant for nearly all years and parameters, indicating a significant genetic influence on plant characteristics. Of interest to this discussion were significant treatment (P) and treatment × cultivar interactions. Data were analyzed separately by year as a result of the transition of the plots from primarily vegetative to fruiting growth over the 5-year experimental period. Soil nutrients (samples only from the ‘Fuji’ plots, a single time) were analyzed separately as a randomized complete block with two treatments (+P) and five replicates.

**Results and Discussion**

**Soil effects.** Extractable soil P concentrations were elevated by the annual ammonium polyphosphate treatment (+P) when measured before removal of the experimental trees in 2005. Soil P was significantly increased directly beneath the drip emitters for each 0.1-m depth increment to 0.4-m depth (Fig. 1A). A similar pattern was measured at 0.5 m from the emitters perpendicular to the tree row, except that the increase was not significant at 0.1- to 0.2-m depth (Fig. 1B). There were no differences in profile P concentration at 1-m distance from the emitter location (Fig. 1C). In general, extractable soil P concentration increased above 0.5-m depth at all distances from the emitter. There were no significant effects of treatments on soil NO$_3$-N and NH$_4$-N concentrations measured in the same extract (as P) at any distance or depth relative to the emitters (data not shown). Soil NH$_4$-N concentrations were near zero, whereas soil NO$_3$-N concentrations averaged 0.20, 0.45, and 0.50 mg kg$^{-1}$ across all depths at 0, 0.5, and 1.0 m from the drip emitter. Similarly, soil pH changes associated with treatments were minimal with a single occurrence of a significant pH decrease from 7.2 (–P) to 7.0 with the P treatment directly beneath the emitter at 0.1- to 0.2-m depth. pH values also increased from near 7 (at the surface) to near 8 at 0.4- to 0.6-m depth.

Mobility of annual bloom time applications of fertigated ammonium polyphosphate within 0.5-m radial distance of the drip emitters is indicated by the elevated 2 M KCl-extractable P concentrations to 0.4-m depth. Increased mobility of P dissolved in irrigation water has been previously reported for sandy soils (O’Neill et al., 1979). It is likely that this P would have been highly available to the apple trees because previous research on this soil series indicated a clustering of roots near and immediately below drip emitters (Neilsen et al., 1997). For example, after 5 years, the average root location occurred at ≈0.25 m depth and lateral distance from the drip emitter location for ‘McIntosh’ receiving N-P fertigation and growing on the same M.9 rootstock. The modest increases in 2 M KCl-extractable soil P (not exceeding an average of 2.5 mg kg$^{-1}$) resulting from the +P treatment nevertheless exceeded soil solution P concentrations previously measured for this soil when it had not received fertilizer P applications (Neilsen et al., 1993). These measurements were made at the end of the study and represented the situation after the cumulative application of 80 g P per emitter (8 years × 10 g P/emitter) or ≈125 kg ha$^{-1}$ P assuming uniform wetting within an 0.5-m radius of each emitter. Previous research has indicated that the Langmuir adsorption maxima for this soil is typical of loamy sands in the region (158 mg kg$^{-1}$ P) and that solution P concentrations after single-dose incubations remain elevated for 12 weeks (Neilsen et al., 1993). Thus, the method of P fertigation in this study should have sufficed to significantly increase annual, early-season P uptake for these apple trees.

Application of ammonium polyphosphate in the +P treatment resulted in coapplication of 13.3 g N per tree in the NH$_4$-N form at bloom. The N regime was otherwise similar for the two treatments, involving maintenance of a NO$_3$-N concentration of 168 mg L$^{-1}$ in the fertigating solution daily for 4 weeks p.h. Hence, the amount of N applied per tree varied with the amount of water applied but, on average, in 1999 to 2003, was 46.7 g N per tree per year. Thus, although the P treatment received 28.5% more N per tree as NH$_4$-N, its effectiveness was limited by the short duration of its application. Previous research has indicated that when NH$_4$-N is fertigated, soil solution NO$_3$-N concentration increase is delayed (Neilsen et al., 1999). Furthermore, only low amounts of NH$_4$-N are measurable in the soil after fertigating all N as NH$_4$-N (Klein and Spieler, 1987). Also, the amount of applied irrigation water has a direct effect on root zone soil solution NO$_3$-N concentrations, which are rapidly reduced to low values after cessation of fertigation and continued irrigation (Neilsen et al., 1998).

Rapid acidification of sandy soils beneath drip emitters in response to fertigation with ammonium-based fertilizers has been long recognized (Edwards et al., 1982) and was the rationale for use of calcium nitrate as the primary N fertigation source in this study. The limited decline in soil pH associated with eight annual, bloom-time applications of ammonium polyphosphate indicates this method of P fertigation should be sustainable with minimal consequences to soil acidification.

**Tree nutrition.** Annual fertigation of a single-time application of ammonium polyphosphate at bloom early in the growing season consistently increased midterminal, midsummer leaf P concentration above 2.0 mg g$^{-1}$ dry weight for all cultivars in the first...
fives growing seasons (Table 1). Increases were particularly pronounced early in the growing season (4 weeks post bloom) as indicated by samples taken during the growing season for the ‘Fuji’ cultivar (Fig. 2). In the first 2 years (1999 and 2000), leaf P was unaffected when measured as late as 12 weeks post bloom. Subsequently, samples collected at 12 weeks, 2001 to 2003, also had increased leaf P concentration resulting from bloom time P fertigation. In two of the first five growing seasons, midsummer (8 weeks post bloom) leaf P concentration was significantly increased across all apple cultivars for the +P treatment compared with the –P treatments (Table 1).

At the time of commercial harvest, fruit P concentration was usually increased by P fertigation for all apple cultivars (Table 2). The single exception was for ‘Gala’ in 2001, which, unlike the other four cultivars in this year, did not show a significant increase in fruit P concentration. In contrast, fruit P concentrations were minimally affected by treatments (data not shown). Only in the first year of the study (1999) was fruit P concentration increased from 51.6 mg kg$^{-1}$ fresh weight (fw) (–P) to 60.1 mg kg$^{-1}$ fw (+P); otherwise (2000 to 2003), fruit P concentration was unaffected by P application.

The single-time fertigated application of P generally improved the P nutrition of both vegetative (leaf) and reproductive (fruit) tissue indicating that P applied in this way was readily available to all the apple cultivars tested. The increased tissue P concentrations were consistent with increases in soil-extractable P concentration measured at the end of the study for the P fertigation treatment. It is noteworthy that all leaf P concentrations, regardless of treatment, exceeded 1.5 mg kg$^{-1}$ dry weight (dw), normally considered adequate for apples (Shear and Faust, 1980).

Higher optimum leaf P concentrations of 2.5 mg kg$^{-1}$ dw have been advocated for mature apples (Shear and Faust, 1980). Higher optimum leaf P concentrations of 3.0 mg kg$^{-1}$ dw have been advocated to stimulate flowering, initial vigor, and yield of newly planted trees (Neilson et al., 1990; Taylor and Goubran, 1975). Fruit P concentration of P-fertigated fruit always exceeded 100 mg kg$^{-1}$ fw and often was above 120 mg kg$^{-1}$ fw, whereas, after the first year of the study (1999), fruit not receiving P usually had concentrations $\approx$ 100 mg kg$^{-1}$ fw or lower. Few fruit P thresholds have been developed for apple, although whole fruit P concentrations above 110 mg kg$^{-1}$ fw have been recommended for cultivars susceptible to low-temperature breakdown (Neilson and Neilson, 2003). In an extensive study (3 years, 90 orchards) of traditional apple cultivars (McIntosh’, ‘Spartan,’ and ‘Golden Delicious’) in the fruit-growing region of southern British Columbia, harvest fruit P concentration averaged 100 mg kg$^{-1}$ fw (Wolk et al., 1998). These concentrations were similar to fruit in this study not receiving P.

Table 1. Leaf nitrogen (N) and phosphorus (P) concentration (8 weeks post full bloom) for apple cultivars (Ambrosia, Cameo, Fuji, Gala, and Silken) as affected by fertigation of 20 g P as ammonium polyphosphate (+P) or not (-P) immediately after bloom, 1999 to 2003.

| Treatment | 1999 | 2000 | 2001 | 2002 | 2003 |
|-----------|------|------|------|------|------|
| Leaf N (mg g$^{-1}$ dw) | 26.3 | 26.6 | 24.3 | 27.0 | 22.0 |
| +P | 25.9 | 26.0 | 24.4 | 25.8 | 21.9 |
| -P | 24.8 | 25.3 | 24.4 | 25.8 | 21.9 |
| Significance | NS | * | * | NS | NS |

| Leaf P (mg g$^{-1}$ dw) | 2.6 | 2.7 | 2.3 | 2.2 | 2.2 |
| +P | 2.1 | 1.9 | 1.8 | 1.7 | 1.8 |
| -P | 2.6 | 2.7 | 2.3 | 2.2 | 2.2 |
| Significance | *** | * | ** | * | * |

*Significance NS = nonsignificant or paired means significantly different at P = 0.05, 0.01, 0.001, or 0.0001, respectively.

Table 2. Whole fruit phosphorus (P) concentration at commercial harvest for apple cultivars [Ambrosia (A), Cameo (C), Fuji (F), Gala (G) and Silken (S)] as affected by fertigation of 20 g P as ammonium polyphosphate (+P) or not (-P) immediately after bloom, 1999 to 2003.

| Treatment | 1999 | 2000 | 2001 | 2002 | 2003 |
|-----------|------|------|------|------|------|
| Fruit P (mg kg$^{-1}$ fw) | 158 | 128 | 110 | 129 | 143 |
| +P | 158 | 128 | 110 | 129 | 143 |
| -P | 129 | 94 | 84 | 90 | 78 |
| Significance | NS | *** | *** | *** | NS |

*Significant interaction between P treatment and cultivar only in 2001.

**Significance NS = nonsignificant or paired means significantly different at P = 0.01, 0.001, or 0.0001, respectively.
surface applications of monoammonium phosphate (MAP) fertilizer (Raese, 1998). Application of 150 g of MAP fertilizer within the planting hole, in association with some form of soil disinfection, has improved establishment and initial growth of apple in replant problem orchards (Neilsen and Yorston, 1991). As a consequence of stimulation of second-year flowering by first-year P applications (Neilsen et al., 1990) and the effectiveness of first-year P fertilization (Neilsen et al., 1993), P fertilization of newly planted apple trees in the first year is a general commercial recommendation regardless of soil P and replant status (BC Ministry of Agriculture and Lands, 2007). The results reported here for a soil of low to moderate P status would suggest bloom time annual fertigation of P to trees receiving N and K fertilization should be recommended because cumulative yield over the first five growing seasons has increased by ≈20% (=5.3 Mt/ha/year) for the five tested apple cultivars. It is also noteworthy that an increase in the number of fruit associated with P fertilization was not associated with the decrease in fruit size normally associated with higher crop loads (Goffinet et al., 1995).

**Fruit quality.** Standard quality characteristics of harvested fruit, including SSC, TA, fruit firmness, and percent red color, were unaffected by P fertigation (data not shown). The incidence of water core at harvest was assessed in the last 3 years of the study and in the 2 years of harvest that fruit P concentration the incidence of water core was reduced for fruit fertilized with P (Table 4). In 2001, only ‘Fuji’ and ‘Silken’ exhibited water core, but for both of these cultivars, water core was reduced and for ‘Silken’ completely eliminated. In 2002, the reduction in water core was significant across all cultivars. The effect was not observed in 2003 when crop load was low for all cultivars. In 2001 and 2002, the low P years, membrane leakage of ‘Fuji’ and ‘Silken’ apple, which were most susceptible to water core at harvest, was reduced in both cultivars (Table 5). Water and lipid-soluble antioxidant concentration was higher for all P-treated fruit, except ‘Gala’ in 2002 (Table 6). In 2002, the degree of browning of cut apple slices from stored fruit was inhibited for all apple cultivars in fruit that received P at bloom (Table 7). Fertilization of P had no effects on SSC, TA, firmness, or color at harvest, indicating the increased yield associated with P application did not negatively affect standard fruit quality characteristics. Beneficial effects of elevated apple fruit P concentrations on incidence of low-temperature breakdown (Johnson and Yogoratnam, 1978), including fruit firmness (Webster and Lidster, 1986), have been periodically reported. From these earlier studies, critical P fw concentration thresholds were recommended to exceed 90 mg kg⁻¹ for ‘McIntosh’ and 110 mg kg⁻¹ for ‘Cox’s Orange Pippin’ for maximum apple quality. Although such thresholds are likely to vary with cultivar, P concentrations of fruit not receiving P in our study were usually less than 100 mg kg⁻¹ in 2001 and 2002 (an exception was ‘Fuji’ apple at 106 mg kg⁻¹ in 2001). In these years, trees fertilized with P had fruit P concentrations in excess of these minimum values. These fruit were characterized by improved membrane stability as indicated by reduced incidence of water core at harvest, reduced membrane leakage, and susceptibility to browning and increased antioxidant content after 90 d of cold air storage. These results are sufficient to justify further research to clarify the relationship between P concentration and these indicators of fruit quality over a wider range of cultivars, seasons, and storage conditions.

**Conclusions.** A single, annual application of 20 g P per tree at bloom as ammonium polyphosphate dissolved in irrigation water was beneficial for the performance, over the first five fruiting seasons, of a range of apple cultivars, including ‘Fuji’, ‘Gala’, ‘Ambrosia’, ‘Silken’, and ‘Cameo’ planted at high density on the dwarfing rootstock M9. These trees were also receiving optimum fertigation recommendations for sandy soils, which included daily application of 168 mg L⁻¹ N as calcium nitrate, maintenance B applications (0.17 g B per tree as Solubor) both applied 0 to 4 weeks postbloom, and daily applications of K applied 8 to 12 weeks postbloom as potassium chloride (≈20 g K per tree). The results imply that conventional fertigation recommendations for P, which currently advise P application only in the first year, should be modified to include annual bloom-time P application. A question is also raised as to whether continuous fertigation of P would provide additional benefits. Application of P in this manner, timed to coincide with a period of high shoot, fruit, and root growth around bloom, is highly effective as indicated by its mobility in applied water throughout the rooting zone and its ability to increase leaf P concentration throughout the growing season and fruit P concentration at harvest. Furthermore, cumulative yield of these trees was increased by ≈20% for all tested apple cultivars during the

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**Table 3. Annual and cumulative fruit number, fruit size and per tree yield for apple cultivars [Ambrosia (A), Cameo (C), Fuji (F), Gala (G) and Silken (S)] as affected by fertigation of 20 g P as ammonium polyphosphate (+P) or not (–P) immediately after bloom, 1999–2003.**

| Treatment | 1999 | 2000 | 2001* | 2002 | 2003 | 1999–2003 |
|-----------|------|------|-------|------|------|-----------|
| cultivar  | All  | All  | A     | C    | G    | F         | S         | All  | All  | All    |
| +P        | 8    | 49   | 74    | 52   | 52   | 47        | 98        | 93   | 71    | 49      |
| –P        | 9    | 40   | 61    | 48   | 42   | 81        | 63        | 66   | 28    | 49      |
| Significance | NS  | NS   | NS    | **  | NS   | NS        | NS        | NS   | NS    | NS      |

| Fruit size (g/fruit) | All | All | All | All | All | All |
|----------------------|-----|-----|-----|-----|-----|-----|
| +P                   | 2.0 | 10.3| 12.1| 12.2| 12.2| 14.1 |
| –P                   | 1.9 | 7.7 | 10.8| 11.5| 10.5| 13.0 |
| Significance | NS | NS | NS | NS | NS | NS |

**Table 4. Incidence of water core at harvest for apple cultivars [Ambrosia (A), Cameo (C), Fuji (F), Gala (G) and Silken (S)] as affected by fertigation of 20 g phosphorus (P) as ammonium polyphosphate (+P) or not (–P) immediately after bloom, 2001 to 2003.**

| Treatment | 2001* | 2002 | 2003 |
|-----------|-------|------|------|
| cultivar  | All   | All  | All  |
| +P        | 0.62  | 0.00 | 0.22 |
| –P        | 0.88  | 0.16 | 0.29 |
| Significance | *** | *    |    |

**Table 5. Membrane leakage of ‘Fuji’ and ‘Silken’ apples after 90 d storage at 1 °C as affected by fertigation of 20 g phosphorus (P) as ammonium polyphosphate (+P) or not (–P) immediately after bloom, 2001 and 2002.**

| Treatment | 2001 | 2002 | 2002 |
|-----------|------|------|------|
| +P        | 0.19 | 0.19 | 0.17 |
| –P        | 0.28 | 0.33 | 0.28 |
| Significance | ** | *    |    |

**Table 6. Incidence of water core at harvest for apple cultivars [Ambrosia (A), Cameo (C), Fuji (F), Gala (G) and Silken (S)] as affected by fertigation of 20 g phosphorus (P) as ammonium polyphosphate (+P) or not (–P) immediately after bloom, 2001 to 2003.**

| Treatment | 2001* | 2002 | 2003 |
|-----------|-------|------|------|
| cultivar  | All   | All  | All  |
| +P        | 0.62  | 0.00 | 0.22 |
| –P        | 0.88  | 0.16 | 0.29 |
| Significance | *** | *    |    |

**Table 7. Proportion of apple fruit affected by water core.**

| Treatment | 2001* | 2002 | 2003 |
|-----------|-------|------|------|
| cultivar  | All   | All  | All  |
| +P        | 0.62  | 0.00 | 0.22 |
| –P        | 0.88  | 0.16 | 0.29 |
| Significance | *** | *    |    |
Table 6. Water-soluble (WSA) and lipid-soluble antioxidant (LSA) concentrations of ‘Fuji’ (F), ‘Silken’ (S), ‘Ambrosia’ (A), ‘Cameo’ (C), and ‘Gala’ (G) apples after 90 d storage at 1 °C as affected by fertigation of 20 g as ammonium polyphosphate (+P) or not (–P) immediately after bloom in 2001 and 2002.

| Antioxidant | Treatment | F   | S   | A   | C   | G   |
|-------------|-----------|-----|-----|-----|-----|-----|
| WSA (μg·mL⁻¹ ascorbate) | +P | 0.58 | 0.49 | 0.77 | 0.74 | 0.44 | 0.70 |
| | –P | 0.45 | 0.38 | 0.39 | 0.41 | 0.28 | 0.63 |
| Significance | ** | ** | ** | ** | ** | ns |
| LSA (μg·mL⁻¹ tocopherol) | +P | 1.32 | 1.80 | 1.58 | 1.27 | 2.28 | 1.60 |
| | –P | 1.12 | 1.32 | 1.33 | 0.84 | 1.44 | 1.34 |
| Significance | ** | * | * | * | * | ** |

Table 7. L value of ‘Silken’ (S), ‘Gala’ (G), ‘Ambrosia’ (A), ‘Fuji’ (F), and ‘Cameo’ (C) apple slices after 90 d storage at 1 °C as affected by fertigation of 20 g phosphorus (P) as ammonium polyphosphate (+P) or not (–P) immediately after bloom in 2002.

| Treatment | S   | G   | A   | F   | C   |
|-----------|-----|-----|-----|-----|-----|
| +P | 77.4 | 79.0 | 77.3 | 75.5 | 77.9 |
| –P | 76.4 | 78.2 | 76.6 | 73.8 | 77.0 |
| Significance | ** | * | * | * | * |

‘Color of slices was determined 5 d after slicing and storage at 5 °C in zip-lock packages with a Minolta chromometer. A higher L value denotes a lighter color for the slice surface (i.e., less browning). ‘*’Significantly different at P = 0.01 and 0.05, respectively.

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