Calcium Uptake and Whole-plant Water Use Influence Pod Calcium Concentration in Snap Bean Plants

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ADDITIONAL INDEX WORDS. Phaseolus vulgaris, nutrition, minerals, mineral requirements, green beans, vegetables, human nutrition, bioavailability, health, postharvest

Abstract. Understanding the mechanisms that regulate xylem transport of calcium (Ca) to snap bean (Phaseolus vulgaris L.) pods could allow approaches to increase pod Ca concentration and enhance the nutritional value of edible pods. Using the snap bean cultivars Hystyle and Labrador, which exhibit high and low pod Ca levels, respectively, we wished to determine whether there were differences between the two cultivars in stem xylem-sap Ca concentration and whether any differences in sap Ca concentration were related to differences in whole-plant water uptake or Ca import between the cultivars. Well-watered greenhouse-grown plants were placed in a growth chamber at a constant light intensity for an equilibration period. Put weight loss was measured to determine whole-plant water use and stem xylem exudate was subsequently collected from the severed base of the shoot at flowering and at two stages of pod development. ‘Hystyle’ displayed an exudate Ca concentration that was 50% higher than ‘Labrador’ during pod development. ‘Labrador’ showed 35% greater total water transport through the stem than ‘Hystyle’. ‘Labrador’ plants also showed a significantly larger leaf area than ‘Hystyle’ plants. Additional plants were used to determine total, long-term Ca influx. No difference was observed between cultivars in total Ca influx into the aerial portion of the plant. With whole-shoot Ca influx being equivalent and pod transpiration rate identical in the two cultivars, our results suggest that the higher whole-plant water uptake in ‘Labrador’ led to a dilution of Ca concentration in the xylem stream and thus less total Ca was transported to developing pods, relative to that in ‘Hystyle’. Increased transpiration efficiency, enhanced root uptake of Ca, or reduced Ca sequestration in the xylem pathway of the stem could lead to an enhancement in pod Ca concentration in future cultivars of snap bean.

Calcium is an essential element in human nutrition due to its structural role in bone development and in regulating cellular metabolism (Linder, 1991). Guidelines suggest that humans should consume ≈1000 to 1300 mg of Ca per day, the equivalent of 1 L of milk (Institute of Medicine, 1997). Adolescents should consume at least 1300 mg of Ca per day in an effort to reduce the risk of osteoporosis later in life (Johnston et al., 1992). Adequate Ca nutrition may also decrease the risk of colon cancer (Bruce, 1987). Although dairy products provide a majority of the Ca needs of humans, milk intake in the United States has declined in recent years. Dairy products represent 55% and 46% of the total Ca intake in adolescent and young adult females, respectively (Penninton and Young, 1991). Snap bean pods are high in bioavailable Ca, being low in compounds that can inhibit absorption (e.g., phytate, oxalate) upon digestion (Heaney et al., 1988). When compared to other fruits and vegetables, snap beans display a high Ca concentration (≈5 mg·g⁻¹ dry weight) (Quintana et al., 1996). Snap beans are well liked by children, as well as adults, thus making this food an ideal candidate for nutritional improvement (Pao et al., 1982). Understanding mechanisms that regulate partitioning of Ca within the snap bean plant could allow for enhancement in the Ca concentration of the edible pods by breeding, biotechnological, or production approaches.

Within the snap bean plant, regulation of Ca partitioning to the pods is incompletely understood. Transpiration, as well as growth induced water uptake by developing plant organs, appears to drive the transport of Ca in the xylem pathway to vegetative tissues of the snap bean plant (Bell and Biddulph, 1963; Biddulph et al., 1959, 1961; Mix and Marschner, 1976; Nonami and Boyer, 1987). When atmospheric relative humidity is increased by covering the whole plant (leaves and pods) in a plastic enclosure, pod Ca concentration is lowered, indicating that whole-plant and/or pod transpiration are important determinants of Ca partitioning to pods (Mix and Marschner, 1976).

The inability to remobilize Ca from terminal organs, such as snap bean leaves, requires continual root uptake and xylem transport of Ca for continued delivery to mature and developing organs (Hose et al., 2001; Raven, 1977; White, 2001). Increasing soil Ca levels with gypsum does not result in an increase in Ca concentration of snap bean pods (Miglioranza et al., 1997), suggesting that net root Ca influx is saturated at low soil Ca levels and may be regulated by the plant. Differential Ca uptake by snap bean roots (Grusak et al., 1996a), as well as sequestration along the xylem pathway in cells as Ca-oxalate crystals (Zindler-Frank, 1995), may represent limiting steps in the transport of Ca to the pods.

Quintana et al. (1996) evaluated 64 snap bean breeding lines and cultivars in two environments and found a 2-fold range in Ca concentration in pods; this variation had a significant genetic basis. Two of the commercial cultivars from this evaluation, ‘Hystyle’, which displays a high pod Ca concentration, and ‘Labrador’,...
which exhibits a low pod Ca concentration, have been further examined in an effort to identify a physiological basis for the pod Ca differences. Both cultivars have similar net-influx rates of Ca into the aerial portion of the plant (Grusak et al., 1996a). Grusak and Pomper (1998) found that although developing pods of ‘Hystyle’ have twice the stomatal density of ‘Labrador’, pod transpiration rates measured on intact pods were similar for both cultivars under equivalent environmental conditions. Interestingly, when developing pods of ‘Hystyle’ and ‘Labrador’ were placed under identical water-saturated environments, differences in pod Ca concentration were maintained between the two cultivars, suggesting that xylem sap delivered to ‘Hystyle’ had a higher concentration of Ca (Grusak and Pomper, 1998).

Based on these physiological examinations demonstrating similar root Ca uptake and pod transpiration, it was hypothesized that differences in Ca concentration of the xylem sap delivered to pods of ‘Hystyle’ and ‘Labrador’ could explain the cultivar differences in pod Ca concentration (e.g., higher xylem-sap Ca concentration would lead to a higher pod Ca concentration in ‘Hystyle’). Thus, the objectives of this study were to 1) determine whether the Ca concentration of xylem sap delivered was higher than that of ‘Labrador’, and 2) determine whether any observed differences in Ca concentration of stem xylem sap were related to differences in water or Ca import rates into the aerial portion of the snap bean cultivars.

### Materials and Methods

**Plant material.** Greenhouse-grown snap bean plants of the cultivars Labrador and Hystyle were grown during the winter and spring months in 1.5-L plastic pots in Metro Mix (Scotts-Sierra Horticultural Products Co., Marysville, Ohio). The greenhouse was maintained at 25 °C day/18 °C night and metal halide lamps provided supplemental irradiance [minimum photosynthetic photon flux (PPF) of 200 μmol·m⁻²·s⁻¹ at the top of the canopy] with a 12-h photoperiod (lights on at 0800 hr). An automatic watering system delivered 200 mL of a nutrient solution to each pot twice daily. This volume of solution was sufficient to saturate the soil, with excess solution allowed to drain from the pot. Plants were watered daily and fertilized with 3 mM KNO₃, 0.275 mM CaCl₂, 0.25 mM MgSO₄, and 0.25 mM KH₂PO₄. Micronutrient minerals were not added daily as these were presumed to be adequately provided from the soil mix for the short duration of the experiments (no signs of micronutrient deficiency were observed). Soil pH varied from pH 5.0 (initial) to pH 6.0 (final).

**Xylem sap Ca concentration.** Using a completely randomized design, a greenhouse experiment was conducted to determine the Ca concentration of xylem exudate collected from the cultivars Hystyle and Labrador at three stages of development. There were four replicate plants in each treatment combination for a total of 24 plants (2 × 3 × 4 = 48). Plants of each cultivar were grown at the same time and were randomly assigned positions on benches in the greenhouse. Pots were spaced approximately 0.4 m apart on the benches (6 plants/m²). Plants were removed from the greenhouse and placed in a growth chamber (model PG2V; Controlled Environments Ltd., Winnipeg, Manitoba, Canada). The environmental conditions in the chamber were 25 ± 0.5 °C, with a relative humidity of 50% ± 5% and a constant PPF of 550 μmol·m⁻²·s⁻¹ at the top of the canopy. The pots then were watered with deionized water and covered in plastic bags (including the soil surface). After a 2-h equilibration period in the growth chamber, the plants were weighed, and weighed again after an additional 3 h. After the final weighing, stem xylem exudate was collected from plants and assayed for Ca concentration as described above. All leaves were removed from each plant and leaf area was determined using the Student t test.

**Statistical analysis.** Exudate Ca concentration, whole-plant water-use, and total Ca content, plant dry weight, and leaf area were subjected to analysis of variance (ANOVA) or general linear model of analysis of variance (GLM-ANOVA) using the statistical software CoStat (CoHort Software, Monterey, Calif.). Statistical significance of differences between mean values was determined using the Student t test.

### Results

**Xylem sap Ca concentration.** Large quantities of exudate were easily obtained from severed snap bean stems. Initially, consecutive 100-μL aliquots of exudate were collected from plants of each cultivar at 42 DAP in the greenhouse in mid-afternoon under sunny conditions. For both cultivars, 600 μL of exudate was collected by the end of 1 h; 2 to 3 min was required to collect the first 100 μL-aliquot (Fig. 1). The first 100-μL aliquot
collected from each plant displayed the lowest Ca concentration. The sixth 100-µL aliquot of exudate was 5-fold higher in Ca concentration than the first 100-µL aliquot. Flow rates (µL·h⁻¹) of stem exudate were similar for the two cultivars over the course of the 1-h collection period (data not shown).

The first 100 µL of stem exudate was arbitrarily chosen to best represent the xylem-sap concentration in the plant and this quantity was collected from plants in the greenhouse in mid-afternoon from each cultivar at flowering (32 DAP) and at two stages of pod development (42 and 50 DAP). ‘Hystyle’ plants displayed a higher exudate Ca concentration than ‘Labrador’ throughout flowering and pod development (Table 1). The Ca concentration of the first 100 µL of stem exudate of ‘Hystyle’ was 20% higher than ‘Labrador’ at flowering (32 DAP), and 30% higher than ‘Labrador’ at 42 and 50 DAP (Table 1). There was a developmental decline in exudate Ca concentration of 37% in ‘Hystyle’ and 42% in ‘Labrador’ from flowering (32 DAP) until mature green pods were present (50 DAP).

To determine whether plant transpiration influenced xylem-sap Ca concentration, stem exudate was collected from plants in darkness, 3 h after sunset. No difference in Ca concentration was observed between the two cultivars for exudate collected in darkness at 42 DAP (Table 1). However, the Ca concentration was 3.7-fold higher in ‘Hystyle’ and 5-fold higher in ‘Labrador’ in exudate collected in darkness, relative to exudate collected from plants of each cultivar in the afternoon.

Whole-plant water loss and xylem sap Ca concentration.

To determine if there was a difference in whole-plant water-use between ‘Hystyle’ and ‘Labrador’, plants of both cultivars were placed in a growth chamber under constant environmental conditions to compare transpiration rates. At all stages of development, ‘Labrador’ displayed a 35% higher water-use rate, or flow rate through the stem, than ‘Hystyle’ (Table 2). Water-use by each individual cultivar was similar at all three stages of plant development. Both cultivar and developmental main effects significantly affected leaf area; there was not a significant interaction between the main effects. ‘Labrador’ plants had a significantly larger leaf area (2398 cm²) than ‘Hystyle’ (2037 cm²). Leaf area was significantly greater in plants at 32 DAP compared to plants at 42 and 50 DAP; leaf areas were 1975, 2447, and 2370 cm² at 32, 42, and 50 DAP, respectively. The average leaf area for ‘Labrador’ and ‘Hystyle’ plants at each developmental stage can be found in Table 3.

As in the initial greenhouse study, the first 100 µL of stem exudate collected from ‘Hystyle’ plants in the growth chamber displayed a higher Ca concentration than ‘Labrador’ throughout flowering and pod development (Table 2). Specifically, ‘Hystyle’ showed an exudate Ca concentration that was 47% higher at flowering (32 DAP), and 59% higher midway through pod development (42 DAP) and 27% higher at harvest (50 DAP) than exudate collected from ‘Labrador’ (Table 2). During plant development, there was a decline in exudate Ca concentration of 34% in ‘Hystyle’ and 24% in ‘Labrador’ from flowering (32 DAP) until mature green pods were present (50 DAP).

Plant growth and long-term whole-shoot uptake of Ca.

Plant growth, as determined by dry weight (DW) measurements of the three plant organs (leaves, stems and pods) at the three time points, was similar for the two cultivars (Table 3). The only significant differences were that the total mean DW of ‘Labrador’ plants (14.6 g) was significantly larger than that of ‘Hystyle’ (12.3 g DW) at 42 DAP, and that the mean stem DW was significantly larger at all three time periods in ‘Labrador’ as compared to ‘Hystyle’ (32, 42, and 50 DAP). Almost all the weight gain from 42 to 50 DAP was attributable to pod growth in plants of both cultivars.

No significant differences in total whole-shoot Ca content were seen between cultivars at flowering (32 DAP) or during pod development (42 and 50 DAP) (Table 4). The only significant difference between ‘Hystyle’ and ‘Labrador’ in Ca content of the leaves, stems, and pods or flowers, was in significantly larger amounts of Ca in pods of ‘Hystyle’ at 42 and 50 DAP than ‘Labrador’ (Table 4). About 74% of the total Ca transported in the aerial portion of 50 DAP plants of both cultivars was accumulated in the leaves. By 50 DAP, ‘Hystyle’ had partitioned 45.2 mg and ‘Labrador’ 31.1 mg of Ca to the pods.

The Ca concentration of pods, on a DW basis, was significantly higher at 42 DAP in ‘Hystyle’ as compared to ‘Labrador’, being (mean ± se) 11.4 ± 0.5 and 7.6 ± 0.7 mg·g⁻¹ DW, respectively. Pods of ‘Hystyle’ at 50 DAP also displayed a significantly higher Ca concentration at 8.3 ± 0.8 mg·g⁻¹ DW as compared to ‘Labrador’ pods at 5.8 ± 1.2 mg·g⁻¹ DW. In a separate experiment, ‘Hystyle’ and ‘Labrador’ plants grown using one-third the rate of nutrient supply also did not display a significant difference in total whole-shoot Ca content between the two cultivars at 50 DAP (Student t test level of significance of 0.05, n =16). However, pods of ‘Hystyle’ still showed a significantly higher Ca concentration at 10.5 ± 0.5 mg·g⁻¹ DW compared to ‘Labrador’ pods at 7.8 ± 0.2 mg·g⁻¹ DW (data not shown).
Table 2. Whole-plant water use measured gravimetrically over a 3-h period in a growth chamber under constant environmental conditions and calcium concentration of exudate (first 100 µL) collected from the severed stem of ‘Hystyle’ and ‘Labrador’ snap bean plants.*

<table>
<thead>
<tr>
<th>DAP</th>
<th>Aerial parts</th>
<th>Hystyle</th>
<th>Labrador</th>
<th>Hystyle</th>
<th>Labrador</th>
<th>Hystyle</th>
<th>Labrador</th>
</tr>
</thead>
<tbody>
<tr>
<td>32</td>
<td>Leaves</td>
<td>4.62 ± 0.34 NS</td>
<td>5.25 ± 0.19</td>
<td>34.3 ± 2.0*</td>
<td>45.8 ± 2.3</td>
<td>56.3 ± 4.7*</td>
<td>38.4 ± 2.5</td>
</tr>
<tr>
<td></td>
<td>Stem</td>
<td>2.05 ± 0.15*</td>
<td>2.72 ± 0.09</td>
<td>0.20 ± 0.01 NS</td>
<td>0.25 ± 0.01</td>
<td>6.87 ± 0.49*</td>
<td>8.23 ± 0.27</td>
</tr>
<tr>
<td>42</td>
<td>Leaves</td>
<td>7.38 ± 0.59 NS</td>
<td>8.27 ± 0.50</td>
<td>3.43 ± 0.28*</td>
<td>4.65 ± 0.31</td>
<td>12.30 ± 0.84*</td>
<td>14.67 ± 0.82</td>
</tr>
<tr>
<td></td>
<td>Stem</td>
<td>1.49 ± 0.19 NS</td>
<td>1.75 ± 0.27</td>
<td>10.89 ± 0.01</td>
<td>10.92 ± 0.01</td>
<td>10.89 ± 0.01</td>
<td>10.92 ± 0.01</td>
</tr>
<tr>
<td>50</td>
<td>Leaves</td>
<td>6.29 ± 0.78 NS</td>
<td>6.03 ± 0.77</td>
<td>3.43 ± 0.28*</td>
<td>4.26 ± 0.31</td>
<td>15.55 ± 1.89 NS</td>
<td>16.08 ± 1.57</td>
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<tr>
<td></td>
<td>Stem</td>
<td>5.72 ± 0.88 NS</td>
<td>5.78 ± 0.66</td>
<td>5.72 ± 0.88 NS</td>
<td>5.78 ± 0.66</td>
<td>5.72 ± 0.88 NS</td>
<td>5.78 ± 0.66</td>
</tr>
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*DAP = days after planting, n = 8; *indicates that the means are significantly different between ‘Hystyle’ and ‘Labrador’ plant using a t test with P < 0.05.

**Discussion**

**Xylem sap collection and Ca concentration.** Using our collection technique, large quantities of xylem exudate were obtained from the severed stem of the snap bean plants. As with any method that is used to sample and measure the components of the plant xylem-sap, it is difficult to determine which aliquot best represents the in vivo composition of the sap. Although the first aliquot of exudate may have contained contamination from damaged cells of the cut stem surface, this aliquot contained the lowest concentration of Ca, and thus contamination seemed minimal. Using a similar method, Ehret and Ho (1986) reported a Ca concentration of ≈300 µg·mL–1 in stem exudate from tomato (Lycopersicon esculentum Mill.) in a combined sample collected over 6 h, which is similar to the Ca concentration we found in aliquots of exudate from stems of snap bean after 1 h. Since the goal of this study was to determine whether there was a relative difference in Ca concentration in the xylem-sap delivered to the pod of each cultivar, we chose the first 100-µL aliquot of exudate as the best representation of the in vivo situation, and used this in all comparisons. However, comparisons of either the first or later matched aliquots demonstrated that the xylem-sap Ca concentration of ‘Hystyle’ was higher than that of ‘Labrador’ at all developmental stages examined.

First aliquots of xylem exudate collected from plants of each cultivar in the greenhouse had a higher Ca concentration than from those collected from plants in the growth chamber (Tables 1 and 2). This may be due to higher whole-plant transpiration rates in plants in the growth chamber than in the greenhouse. The higher transpiration rate by plants in the growth chamber would have resulted in greater water uptake by the roots in relation to Ca uptake, thereby lowering the stem xylem-sap Ca concentration (growth chamber relative to greenhouse plants). Diurnal changes in Ca concentration in the root pressure exudate of poplar (Populus tremula L. × P. alba L.) have been reported; Ca concentration was highest at night in poplar xylem exudate (Siebrecht et al., 2003). Tanner and Beevers (2001) reported that water transport in the xylem brought about by root pressure and the resulting guttation was sufficient for long-distance Ca transport in sunflower (Helianthus annuus L.). In this study, when plant transpiration was greatly diminished during the night period and exudate was collected from either cultivar, the Ca concentration was much higher than that of exudate collected from plants either during the day in the greenhouse or from plants in the growth chamber (Table 1). However, the Ca concentration was similar for xylem exudate collected during the night for both ‘Hystyle’ and ‘Labrador’.

Stem xylem-sap Ca concentration in snap bean was also examined by Atkinson et al. (1992), where exudate was collected by placing roots of plants in a pressure chamber. These authors reported Ca concentrations for the first 50 to 100 µL of stem exudate to be 612 µg·mL–1 in ‘Mihoaica-12A3’, 564 µg·mL–1 in ‘Camahuate-72’, and 180 µg·mL–1 in ‘Canadian Wonder’. The exudate Ca concentrations were ≈7.5-fold higher in ‘Mihoaica-12A3’ and ‘Camahuate-72’ and ≈2-fold higher in ‘Canadian Wonder’ than what we found in ‘Hystyle’ exudate collected at 32 DAP from plants in the greenhouse. The higher Ca concentration in exudate...
collected from snap bean plants of Atkinson et al. (1992) may be due to the difference in technique used by these authors, or, it merely adds further support to our observation that xylem-sap Ca concentration varies among snap bean cultivars.

**Whole-plant water-loss and whole-shoot Ca uptake.** There are two possibilities that can explain why ‘Labrador’ would have a significantly lower xylem-sap Ca concentration than ‘Hystyle’. First, if whole-shoot Ca influx is similar between the two cultivars, then a higher whole-plant uptake of water in ‘Labrador’ would lead to a greater dilution of Ca in the xylem sap of ‘Labrador’ relative to ‘Hystyle’. Alternatively, if whole-plant water uptake is similar between the two cultivars, then higher net influx of Ca in ‘Hystyle’ would lead to a higher Ca concentration in the xylem sap of ‘Hystyle’, relative to ‘Labrador’.

To determine whether there was a difference in whole-plant water-use between the two cultivars, plants were placed in a growth chamber under constant environmental conditions in order to maintain steady transpiration rates under equivalent conditions. Although plants of both cultivars were the same size (grams DW), ‘Labrador’ showed a 35% higher whole-plant water-use than ‘Hystyle’ (Table 2). The higher rate of water-use was a result of the larger leaf area (larger transpirational area) of ‘Labrador’, rather than being due to higher transpiration rates per unit leaf area in this cultivar (data not shown).

Cumulative Ca uptake into the aerial portion of the plant was the same in both cultivars at each developmental stage, indicating that the higher concentration of Ca in the xylem-sap in ‘Hystyle’ was not due to a higher net influx of Ca in this cultivar, relative to ‘Labrador’. There was a smaller increase in whole-shoot Ca content in both cultivars from 42 to 50 DAP than from 32 to 42 DAP, which indicated that less Ca was delivered to the aerial portion of the plant later in development. The reduced Ca uptake and transport by both ‘Hystyle’ and ‘Labrador’ late in plant development was mirrored by a reduced xylem-sap Ca concentration in each cultivar from 32 to 50 DAP and demonstrated that net Ca influx does play a role in the determination of xylem sap Ca concentration.

Our water-use data support the scenario that ‘Labrador’ roots take up larger quantities of water than ‘Hystyle’, such that this dilutes the Ca concentration in the xylem stream of ‘Labrador’, relative to ‘Hystyle’. The similar rates of whole-shoot Ca uptake by these cultivars also support this explanation. Ehleringer et al. (1991) found cultivar variation in water-use efficiency (net mol CO₂ fixed per mol H₂O transpired at the leaf level) and transpiration efficiency (net mol CO₂ fixed per mol H₂O transpired at the canopy level) in a range of common North American bean cultivars. Selecting for increased transpiration efficiency in future snap bean cultivars may result in increased xylem-sap Ca concentration and thereby help to increase pod Ca concentrations.

Grusak et al. (1996a) reported pod Ca concentrations in hydroponically grown ‘Hystyle’ (3.8 mg·g⁻¹ DW) and ‘Labrador’ (2.5 mg·g⁻¹ DW) that were about half the pod Ca concentrations found in greenhouse, soil-grown plants in the present study. Plants of both cultivars grown in hydroponics by Grusak et al. (1996a) were ≈2-fold larger in both plant dry weight and plant Ca content at 50 DAP as compared to our soil-grown plants. Plants in hydroponics were planted at a much higher plant density (20 plants/m²) than in soil grown plants (6 plants/m²) in this present study. Grusk et al. (1996a) noted that the pod Ca concentration from hydroponically grown growth chamber plants was lower in both cultivars than when plants were grown in the field. These authors suggested that the lower pod Ca concentration in the two cultivars was perhaps due to a denser canopy in hydroponically grown plants and thereby a higher humidity microenvironment around pods that would reduce pod transpiration. Grusk and Pomper (1998) showed that raising the humidity levels around pods lowered pod Ca concentrations in both ‘Hystyle’ and ‘Labrador’ pods. Humidity levels in the greenhouse were generally lower during the daytime than in the growth chamber (data not shown). The lower humidity in the greenhouse would have helped raise pod Ca concentrations higher for plants in this environment, relative to pods of hydroponically grown growth chamber plants.

In the present study, we saw a decline in Ca uptake and in xylem-sap Ca concentration from 30 to 50 DAP with soil-grown plants of both ‘Hystyle’ and ‘Labrador’. Similarly, Grusk et al. (1996a) found a decline in net influx of Ca for these two cultivars over this same time period. We found that soil-grown ‘Labrador’ plants transpired (48 g·h⁻¹) at a significantly higher rate relative to ‘Hystyle’ (35 g·h⁻¹). In contrast, Grusk et al. (1996a) reported that hydroponically grown ‘Hystyle’ and ‘Labrador’ plants both showed similar water-uptake rates per plant for both cultivars of ≈29 g·h⁻¹ from 25 to 50 DAP in a growth chamber. Bean plant density was higher in hydroponically grown plants (20 plants/m²) than in soil-grown plants (6 plants/m²) in this study. The higher planting density of the hydroponically grown plants may have influenced plant water-use rates. Factors other than whole-plant water-use (e.g., crowding of plants, plant size, stem length, and canopy structure) of the plants grown in hydroponics may have resulted in the observed cultivar differences in pod Ca concentration.

**Cultivar development for enhanced pod Ca concentration.** Plant transpiration efficiency, pod transpiration rates, root uptake of Ca, and Ca sequestration in the xylem pathway of the stem are important potential regulatory factors that could impact pod Ca concentration in snap bean. Based on the results of this study, we suggest that if breeders select for cultivars that maintain high xylem-sap Ca concentration (either through higher transpiration efficiency or by increased root Ca uptake), this could lead to higher pod Ca concentrations in snap bean. Utilizing the amplified fragment length polymorphism (AFLP) marker system and 120 F₂₃ segregating individuals of a snap bean cross, Guzman-Maldonado et al. (2003) found two putative quantitative trait loci significantly associated with Ca content of the seed. However, the authors did not examine plant water-use or Ca uptake in the progeny.

Further gains in pod Ca concentration could be achieved if Ca sequestration in the stem was reduced, such that larger quantities of Ca could be made available for delivery to pods. Both cultivars sequestered almost as much Ca in stems as in pods (Table 4). Of the total Ca transported to the aerial portion of the plant by 50 DAP, ‘Hystyle’ and ‘Labrador’ partitioned 15% and 12%, respectively, into pods, whereas ‘Hystyle’ and ‘Labrador’ partitioned 11% and 14%, respectively, into stems. Thus, significant gains in pod Ca concentration could be achieved in both cultivars if Ca sequestration in the stem pathway was reduced.

In conclusion, using the snap bean cultivars ‘Hystyle’ and ‘Labrador’ that exhibit high and low pod Ca levels respectively, we demonstrated that differences in whole-plant water use impacted xylem-sap Ca concentration, and this further impacted pod Ca concentration. It is worth noting that the more water-use efficient cultivar had more Ca-dense pods. Selection for water use efficiency in bean, therefore, would be not only advantageous to the farmer and the environment, but would benefit the consumer as well.
Literature Cited


