Papaya Fruit Growth, Calcium Uptake, and Fruit Ripening

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Abstract: The uptake of Ca by ‘Sunset’ papaya (Carica papaya L.) fruit and its role in ripening was studied. The highest mesocarp Ca uptake rate occurred in fruit that were <40 days postanthesis when fruit transpiration was probably highest. Ca uptake rate by the mesocarp was low, from 60 to 80 days postanthesis when fruit fresh and dry weight increased. Mesocarp Ca uptake rate increased again from 100 to 140 days postanthesis when fruit fresh weight growth rate declined and dry weight growth rate increased. Mesocarp Ca concentration did not significantly differ from the peduncle to the blossom end, although Ca was significantly higher in the outer than inner mesocarp at the fruit equator. Mesocarp Ca concentration fluctuated significantly throughout the year ranging from 68 to 204 µg·g⁻¹ fresh weight (FW). Soil Ca application did not always increase fruit mesocarp Ca concentration, while K and N fertilization decreased mesocarp Ca concentration. Attempts to increase mesocarp Ca concentration by spraying CaCl₂ onto papaya fruit during growth and by postharvest vacuum infiltration and dipping of the cut peduncle into CaCl₂ were unsuccessful. Mesocarp Ca concentration was positively correlated to the firmness of ripe papaya fruit and the rate of softening of mesocarp plugs. Less correlation was found between fruit firmness and the ratio of Ca concentration to K or Mg concentration, or to Mg plus K concentrations. Mesocarp Ca concentration of 130 µg·g⁻¹ FW or above was associated with slower fruit softening rate than fruit with a lower concentration.

Increased fruit Ca levels delay softening and maintain quality in apple (Conway and Sams, 1987; Pooovaiah, 1986), tomato (Rigney and Wills, 1981), and avocado (Eaks, 1985). Physiological disorders associated with inadequate Ca levels occur in many plants (Bangerth, 1979). Inadequate Ca often occurs in those plant parts that are naturally low in Ca, such as fruit and storage organs (Bangerth, 1979). The best known disorders in fruit are bitter pit of apple (Ferguson and Watkins, 1989) and blossom end rot of tomatoes (Evans and Troxler, 1953). Treatment of mangoes (Tirmazi and Wills, 1981), pears and bananas (Wills et al., 1982), sweet cherry (Facteau et al., 1987), and strawberry (Cheour et al., 1991) delay fruit ripening and senescence.

Inspected shipments of Hawaii papaya fruit in the New York market had overripe and abnormally soft fruit (Cappellini et al., 1991) delay fruit ripening and senescence. Sporadically, commercial papaya-packing sheds in Hawaii have also reported batches of fruit that ripened very rapidly. This soft fruit disorder leads to disruption of marketing and higher losses. The inability to predict the occurrence of this disorder limits the range of options available to deal with this problem. Initial results suggest that low fruit Ca may be related to the problem (Paull, 1987). The objective of this study was to determine the uptake of Ca by papaya fruit and the role of Ca in papaya fruit ripening.

Materials and Methods

Plant materials. ‘Sunset’ papaya fruit grown at Poamoho Experiment Station, central Oahu, were used for preharvest CaCl₂ spray and postharvest CaCl₂ dip treatments. The commercial variety ‘Kapoho Solo’ was used for the field fertilization experiment on the Island of Hawaii. The fruit were harvested and returned to the laboratory in Honolulu on the same day. Fruit were sorted by skin color and stored at 22°C until they reached the desired stage of ripeness.

The skin and mesocarp color were subjectively evaluated by estimating the percent area of yellowing of the fruit surface and of a mesocarp vertical section, respectively. Deformation force was measured with a penetrometer fitted with a 1.6-cm-diameter tip pressed 2 mm into the flesh at the fruit equator.

Fruit from 19-month-old ‘Sunset’ trees were harvested to determine the uptake patterns of Ca, Mg, and K by fruit of different maturities. Two to three phyllotaxic spirals can exist on a papaya tree, and three trees having two spirals each were used in this study. Fruit from each tree was divided into two groups based on the phyllotaxic position on the spiral. Fruit number was marked according to the leaf number starting from the apex. Fruit maturity was determined by dividing the days after anthesis from tagged fruit on companion trees by the leaf number. The fruit were weighed and the volume was measured by water displacement. Fruit, with -20 days difference in maturity, were chosen from each spiral on the three trees. Each fruit was divided into skin, mesocarp, and seeds, and fresh weight (FW) determined. Ten grams of the skin and seeds and 20 g of the mesocarp were dried at 60°C for 7 days to determine dry weight (DW) percentage.

Five mature fruit were used to determine Ca, Mg, and K distribution. The fruit was divided into exocarp, mesocarp, endocarp, and seeds. The equatorial mesocarp was further divided horizontally into outer and inner tissue. The mesocarp was also divided transversely into peduncle, equatorial and blossom end tissue. The tissues were stored at -20°C until analyzed.

Fruit surface transpiration was measured with a LI-COR model LI-700 transient automatic diffusion porometer (Paull and Chen, 1989). The time taken (seconds) to cycle through the 35% to 55% range was used to determine fruit water diffusive resistance (s·cm⁻¹). Analysis of Ca, Mg, and K. For young fruit, the whole fruit was used. Otherwise, 10 g FW of skin or seeds and 50 g mesocarp was used. Tissue was homogenized in a blender with 100 ml deionized
water. An aliquot (10 ml) was mixed with an equal volume of 12 M HCl and heated at 60°C for 30 min. The solution was filtered through a Whatman #42 ashless filter paper into a 50-ml volumetric flask and made up to the volume with deionized water. For Ca and Mg determination, lanthanum chloride (5% w/v La, 1 M HCl) was added during further dilution for a final La concentration of 0.5% w/v. Leaf petiole, total Ca, Mg, and K concentrations were determined on dry-ashed, ground tissue. Total Ca and Mg concentrations were determined by atomic absorption spectroscopy and K by emission spectroscopy.

Fruit mesocarp and recently matured petiole N were determined on one occasion during the field trial to confirm that applied fertilizer was increasing tissue content. Total N of mesocarp and leaf petiole was measured as NH₃-N in the acid digest.

Fertilization and Ca chloride sprays. ‘Kapoho Solo’ papaya seedlings were transplanted into a commercial field on the Island of Hawaii in November 1987. The soil type belonged to the Papai series of Typic Tropofults. The five fertilizer treatments were applied in a randomized complete-block design and consisted of control, Ca, Ca plus K, K, and N. Commercial fertilization rates were used as the control level, and applied to the whole 10.5-ha field including the experimental block. The control rate of fertiliser (14N-14P-14K) was: 6.1 kg·ha⁻¹ per month in 1988, 14.6 kg·ha⁻¹ per month in 1989, and 9.7 kg·ha⁻¹ per month in 1990. For the other treatments, extra fertilizer was applied when the trees were 6 months old and had begun to flower. Trees were side dressed with CaCO₃ (48.5% CaO) at the rate of 192 g/tree per month, KCl (61% K₂O) at the rate of 192 g/tree per month, and urea (46% N) at the rate of 158.7 g/tree per month. Monthly harvesting began in September 1989 and fruit were air-shipped on the same day to Honolulu for evaluation. Fruit with color break to 15% skin yellow were used and stored at 22°C until the skin color of the majority of fruit in the control treatment reached 85% yellow. Data were collected over 11 months.

The fruit on five trees per treatment were sprayed with 21 of deionized water (control) or 2% (w/v) CaCl₂ every 2 weeks and harvested (10% skin coloring) following six spray applications. Additional fruit were harvested with a knife to leave as long a placenta as possible, and exposed to a high airflow for 24 h at 22°C. Plug firmness was measured by pressing a penetrometer with 0.5 s·cm⁻¹ when the fruit were 120 days old and to 18.6 s·cm⁻¹ at 140 days old (data not shown), a 37-fold increase. Fruit surface transpiration is an important motive force for water and Ca ion flow into fruit (Bangerth, 1979; Ferguson and Watkins, 1989). As the proportion of water imported to tomato fruit via the xylem decreases and that via the phloem increases, fruit Ca accumulation rate also decreases (Ho et al., 1987). The competition between fruit and leaf for water may also reduce the Ca transport to the fruit. The high rate of papaya Ca uptake during the first 60 days postanthesis may therefore be partially associated with the higher rate of transpiration from the fruit surface at that time. The highest Ca uptake rate on a fruit basis occurred 80 to 140 days postanthesis.

The mean papaya fruit surface diffusive resistance was 0.5 s·cm⁻¹ when the papaya fruit were 60 days old and increased to 11.3 s·cm⁻¹ when the fruit were 120 days old and to 18.6 s·cm⁻¹ at 140 days old (data not shown), a 37-fold increase. Fruit surface transpiration is an important motive force for water and Ca ion flow into fruit (Bangerth, 1979; Ferguson and Watkins, 1989). As the proportion of water imported to tomato fruit via the xylem decreases and that via the phloem increases, fruit Ca accumulation rate also decreases (Ho et al., 1987). The competition between fruit and leaf for water may also reduce the Ca transport to the fruit. The high rate of papaya Ca uptake during the first 60 days postanthesis may therefore be partially associated with the higher rate of transpiration from the fruit surface and the lower surface area to weight ratio of the fruit.

Papaya mesocarp Ca uptake (Table 1) did not parallel mesocarp growth (Fig. 1A) suggesting that Ca uptake was not determined by fruit growth alone. Sucrose in papaya mesocarp increases 4-fold, 110 to 140 days postanthesis (Chan et al., 1979), a period of steady...
Ca uptake (Table 1). High sucrose accumulation in mesocarp during this period reflects increased phloem importation or decreased osmotic potential, either or both of which could increase xylem water flow to the mesocarp. Alternatively, root pressure may also provide a certain amount of Ca to papaya fruit, as suggested for tomatoes (Banuelos et al., 1985). In addition, Ca import into tomato fruit is favored at night when the humidity is normally highest, although the rate of Ca absorption and translocation from root to shoot does not differ between the day and night (Tachibana, 1991).

Calcium concentration was lowest in the mesocarp (Fig. 1D) and was not significantly different among the peduncle, equatorial, and blossom regions (Table 2). However, Ca concentration was significantly higher in the outer mesocarp (187 µg·g⁻¹ FW) than the inner mesocarp (127 µg·g⁻¹ FW). In contrast, a longitudinal gradient in Ca is found in avocado (Chaplin and Scott, 1980) and tomato (Ehret and Ho, 1986). The vascular network in tomato fruit has increased branching from the proximal to the distal half and a decline in the ratio of xylem to phloem area, increasing the resistance to xylem water movement leading to lower Ca content in the distal half (Ehret and Ho, 1986). The vascular bundles in papaya mesocarp extends longitudinally throughout the mesocarp and therefore may distribute Ca relatively evenly from the peduncle to the blossom end. The high levels of K at the peduncle (Table 2) may be related to the active sugar uptake via the phloem occurring in mature fruit (Chan et al., 1979). While the Mg distribution (Table 2) may be a function of the higher uptake 80 to 110 days postanthesis (Fig. 1E) and lower rate in mature fruit.

Spraying papaya fruit six times over 12 weeks with CaCl₂ (2% w/v) during fruit growth and development did not significantly

![Graphs](image_url)
Table 1. The changes in papaya fruit dry weight percentage and Ca concentration of mesocarp tissue during fruit growth and development.

| Days after anthesis | Dry wt (%) | C a (µg·g⁻¹FW) | C a (µg·g⁻¹DW) |
|--------------------|------------|----------------|----------------|
| 30                 | 8.9        | 405a           | 4506a          |
| 60                 | 7.7        | 256 b          | 3340 ab        |
| 80                 | 6.9        | 135 c          | 1899 b         |
| 100                | 7.0        | 180 c          | 2511 b         |
| 120                | 7.1        | 163 c          | 2748 b         |
| 140                | 8.6        | 158 c          | 1677 b         |

Data were analysed with Duncan’s multiple range test and means with some letters in the same column were not significantly different at 5% level, n = 6.

FW = fresh weight.

Table 2. Longitudinal distribution of Ca, Mg, and K in papaya mesocarp from fruit at the 10% yellow stage.

| Position           | Ca     | Mg     | K      |
|--------------------|--------|--------|--------|
| Peduncle end       | 143 a  | 92 c   | 3084 a |
| Equatorial zone    | 117 a  | 202 a  | 1221 b |
| Blossom end        | 133 a  | 143 b  | 1213  b|

Data were analysed with Duncan’s multiple range test and means with same letters in the same column were not significantly different at 5% level. n = 5.

Table 3. Effect of fertilization on Ca, Mg, and K in papaya leaf petiole and mesocarp from color break fruit.

| Mineral content of | Petiole (% dry wt) | Fruit mesocarp (µg·g⁻¹Fresh wt) |
|--------------------|---------------------|---------------------------------|
| Fertilization      | Ca                  | Mg                             | K                        |
| Control            | 0.98 bc             | 0.65 a                         | 1.89 b                   |
| Ca                 | 1.29 a              | 0.47 b                         | 1.93 b                   |
| Ca + K             | 1.15 ab             | 0.29 c                         | 3.80 a                   |
| K                  | 0.84 c              | 0.41 bc                        | 3.74 a                   |
| N                  | 0.99 bc             | 0.76 a                         | 1.21 c                   |
| Ca                 | 138 b               | 119 a                          | 1070 b                   |
| Ca + K             | 145 a               | 180 b                          | 1095 b                   |
| K                  | 151 a               | 197 a                          | 1518 b                   |
| N                  | 120 d               | 186 b                          | 1497 a                   |
|                   | 130 c               | 182 b                          | 904 c                    |

Data were analyzed with Duncan’s multiple range test and means with same letters in the same column were not significantly different at 5% level. n = 165.

increase the mesocarp Ca level (data not shown) such as occurs with sprayed apple fruit (Glenn et al., 1985). Lenticels, cracks, or other surface irregularities are the important pathway of Ca diffusion through the cuticle in fruit such as apple (Glenn et al., 1985); in the papaya, lenticels are absent and stomata of mature fruit are apparently blocked by wax (Paull and Chen, 1980). The cuticle of papaya, although not well developed in immature fruit (Sanxter, 1989), increases in thickness from 15 to 50 µm during fruit growth and development (Quintana and Paull, 1993). Papaya cuticle thickness is almost twice that of apples (Glenn et al., 1985) and may be a more effective barrier to cuticle absorption.

Vacuum infiltration of Ca into papaya was unsuccessful (data not shown) using similar conditions reported to increase Ca in apple tissue by 70% (Ferguson and Watkins, 1989). Water soluble flowerdye was vacuum infiltrated only 1 to 2 mm into papaya fruit. Attempts to draw Ca into the mesocarp via the peduncle using transpiration were also unsuccessful in uniformly increasing mesocarp Ca. Either transpiration by detached mature fruit was not an effective motive force or the peduncle vascular bundle may have come wholly or partially blocked during the 48-h treatment.

Calcium fertilizer treatment significantly increased Ca concentration in fruit and leaf petioles (Table 3) above the control. Calcium soil fertilization also did not always increase mesocarp Ca concentration. The higher mesocarp Ca concentration in all fertilizer treatment of fruit harvested in April (Fig. 2) may be related to the high rainfall during January to February when the fruit were young. There was an overall increase in mesocarp Ca concentration (Fig. 2) but a decrease in Mg concentration in all fertilizer treatment throughout the year. Mesocarp K did not significantly change throughout the year of sampling. Both N and K fertilizer treatments significantly decreased mesocarp Ca concentration (Table 3). Calcium uptake by papaya plants and fruit may therefore be influenced by temperature, relative humidity, levels of other minerals in the soil, and plant age as for other crops (Clarkson, 1984: Shear, 1975).

Potassium fertilization reduced the Ca concentration in leaf petioles and fruit mesocarp (Table 3). This agrees with the finding that K fertilizer decreased leaf petiole Ca concentration (Awada and Long, 1971: Awada et al., 1986). Calcium fertilization did not affect the K concentration in leaf petiole and the fruit mesocarp (Table 3), while Ca plus K fertilizer increased mesocarp Ca (Table 3). Since the application of K fertilizer to papaya plants increases mesocarp total soluble solids (Awada and Long, 1971). it is possible that increases in mesocarp K and sugars may reduce
Fig. 2. Effect of fertilization on mesocarp Ca concentration from color break ‘Kapoho Solo’ papaya mesocarp from December to October. The fertilization treatments were control (●), Ca (Δ), Ca plus K (❍), K (❏) and N (▲). Analysis of variance: month***, Ca***, month × Ca***, n = 16; ***Significant at P ≤ 0.001

The fruit water potential, resulting in water and Ca movement into the mesocarp. There is a strong association between high N fertilization and Ca deficiency disorders in apple fruit (Ferguson and Watkins, 1989; Shear, 1975); this suggests that high N fertilization could induce Ca deficiency in papaya fruit. However, high N fertilization of papaya tended to delay ripening (Table 4) and had no effect on papaya leaf petiole Ca concentration, but decreased mesocarp Ca concentration.

No fertilizer treatment, except for N, had a significant effect on fruit skin color and mesocarp color development (Table 4). The N treatment significantly delayed skin and mesocarp color development. Although Ca fertilizer treatment did not always increase the firmness of the ripe fruit in the monthly harvests, Ca and Ca plus K fertilizer treatments significantly increased fruit firmness with an average deformation force of 76 and 78 N, respectively, vs. 68 N in the control treatment (Table 4). Potassium treatment had no significant effect on fruit firmness (69 N), while fruit from the N fertilizer plots were firmer (81 N) than control fruit (68 N) (Table 4).

Papaya mesocarp Ca concentration in the mesocarp was positively correlated with ripe papaya fruit firmness (Table 5). Other fruit with natural or artificially raised higher Ca concentration also show positive correlations between Ca concentration and fruit firmness and ripening rate of apple (Abbott et al., 1989; Conway and Sams, 1987; Poovaiah et al., 1988), cherry (Facteau et al., 1987), and avocados (Tingwa and Young, 1974). The positive correlation between Ca concentration and ripe fruit firmness was described by the significant quadratic and cubic (firmness ratio = –2.707 + 0.071 Ca – 0.0004Ca$^2$ + 0.0000008Ca$^3$) regression equation (Table 5). The development of skin and mesocarp color during fruit ripening was significantly delayed (Table 4), and fruit softening was significantly slower in fruit from the N treatment. The June data did not follow the same trends as other monthly data. If the data from the N treatment and June data were removed, the significance increased in the cubic (firmness ratio = –2.780 + 0.069Ca–0.0004Ca$^3$ + 0.0000007Ca$^3$) regression equation (Table 5).

When mesocarp Ca concentration was 130 µg·g$^{-1}$FW (≈1.3 mg·g$^{-1}$ DW) or greater, fruit softened at a slower rate than fruit with lower concentrations (Fig. 3). Fruit with < 130 µg·g$^{-1}$FW Ca could not soften at a slower rate than fruit with lower concentrations (Fig. 3). Fruit with < 130 µg·g$^{-1}$FW Ca could not soften at a slower rate than fruit with lower concentrations (Fig. 3).

Table 4. Effects of field fertilization on papaya fruit ripening. Results are means of 11 monthly harvests. Fruit were ripened at 25°C until the skin color of most fruit in the control treatment had reached ≈ 85% yellow.

| Treatment | Initial skin color (%) | Final skin color (%) | Final mesocarp color | Deformation force (N) | Firmness ratio |
|-----------|------------------------|---------------------|----------------------|-----------------------|---------------|
| Control   | 7 a                    | 89 a                | 96 ab               | 68 b                  | - - -         |
| Ca        | 4 b                    | 86 a                | 95 b                | 76 a                  | 1.13          |
| Ca + K    | 4 b                    | 86 a                | 95 b                | 78 a                  | 1.16          |
| K         | 5 b                    | 88 a                | 97 a                | 69 b                  | 1.02          |
| N         | 3 b                    | 80 b                | 91 c                | 81 a                  | 1.20          |

$^1$Data were analyzed with Duncan’s multiple range test and means with same letters in the same column were not significantly different at 5% level. n = 220.

$^2$Firmness ratio = firmness of the treatment/firmness of control treatment.
Table 5. Relationships between mesocarp Ca, Mg, and K concentrations and fruit firmness.

| Repression model | Coefficient of determination |
|------------------|-------------------------------|
|                  | -N and June data               | 11 Months data |
| 1. Firmness ratio = a + bCa + cCa + dCa | 0.429                         | 0.160                     |
| 2. Firmness ratio = a + b(Ca/Mg) + c(Ca/Mg) + d(Ca/Mg) | 0.386**                      | 0.084**                   |
| 3. Firmness ratio = a + b(Ca/K) + c(Ca/K) + d(Ca/K) | 0.255*                       | 0.060**                   |
| 4. Firmness ratio = a + b(Ca/(Mg + K)) + c(Ca/(Mg + K)) + d(Ca/(Mg + K)) | 0.260*                   | 0.177*                   |

Data were analyzed by the procedures of general linear model and nonlinear curve fitting model.

**, ***Nonsignificant or significant at P = 0.5, 0.1, or 0.01, respectively; n = 20.

Fig. 3. Relationship between mesocarp Ca concentration and ratio between deformation force at full ripe stage for Ca-fertilized fruit and non Ca fertilized fruit.

Therefore be expected to be susceptible to the soft fruit disorder reported for papaya (Paul, 1987). The critical Ca concentration for papaya (1.3 mg·g⁻¹ DW) contrasts with the concentration (0.4 mg·g⁻¹ DW) required to eliminate the development of tissue breakdown found in tipburn of lettuce (Barta and Tibbitts, 1991), tomato blossom-end rot (Cerda et al., 1979), and bitter pit of apple (Ferguson and Watkins, 1989). No correlation was found between ripe papaya fruit firmness and Mg or K concentration (data not shown). There were significant correlations between papaya fruit firmness and the ratio of Ca to K or Mg concentration, or to Mg plus K concentrations (Table 5). The involvement of K or Mg appeared to be related to Ca concentration, rather than an effect of these minerals per se.

The processes characteristic of ripening in intact fruit are basically duplicated in excised pericarp discs or plugs from tomato (Campbell et al., 1991). The ripening pattern of excised papaya mesocarp plugs although X-fold faster is similar to that of intact fruit. The plugs could therefore be a model to study the effect of Ca on mesocarp ripening (Qiu, 1992). Infiltration of plugs with CaCl₂, sodium citrate, and EGTA had no significant effect on mesocarp plug respiration from 10% yellow fruit (Fig. 4A). The pattern of ethylene production in mesocarp plugs (Fig. 4B) was about the same for the control and CaCl₂-treated plugs, although 50% reduced by the Ca treatment. EGTA treatment doubled ethylene production, while sodium citrate reduced ethylene production. Calcium concentration in the mesocarp tissue increased ≈100% when plugs were infiltrated with 50 mM CaCl₂, and was more effective in reducing softening rate in plugs from color break and 10% yellow fruit (Fig. 4C) than in plugs from 30% and 50% yellow fruit (Fig. 4D). EGTA and sodium citrate infiltration increased the softening rate 8 h after treatment in plugs from 10% and 50% yellow fruit. In fruit that contain high levels of cell wall- and middle lamella-bound Ca, removal of Ca²⁺ from pericarp tissue or from recovered cell walls by chelating agents, such as EDTA or citrate, may promote extensive hydrolysis of the pectin molecules (Brady et al., 1985; Buescher and Hobson, 1982). Infiltration of papaya mesocarp plugs with MgCl₂ and KCl did not increase tissue firmness or rate of plug softening: Infiltration with Mg in intact apple fruit (Conway and Sams, 1987) as well as in excised tissue (Stow, 1989) increases the firmness of apples or the fruit tissue to a lesser extent than Ca. The difference in part could be ascribed to papaya having already begun to ripen when treated.

The fractionation of Ca in papaya mesocarp tissue from color break, 50% and 100% yellow fruit indicated that most Ca was in the 80% acetic acid fraction with 10 to 16% in the 0.25 N HCl and <1% in the 1 N HCl fraction. This distribution suggests that most
of the Ca in mesocarp tissue was bound to pectin in the cell wall (Ferguson et al., 1980). It has been suggested that fruit softening occurs either by movement of Ca\(^{2+}\) from the middle lamella or by the loss of Ca\(^{2+}\) attachment sites (Knee and Hartley, 1981). That the Ca concentration in 80% acetic acid fraction did not change during papaya ripening, suggests that the Ca probably stayed in the cell wall. Some loss of Ca\(^{2+}\) cell wall attachment sites may have occurred during papaya ripening as the Ca contents in the 0.25 M and 1 M HCl fractions decreased during ripening.

Calcium concentration in papaya fruit varied significantly with harvest date. There were two periods during fruit growth when there was significant movement of Ca into fruit, and these periods relate in part to FW and DW changes during fruit growth. Fruit mesocarp Ca concentration was related to the rate of softening during papaya fruit ripening. Fruit whose mesocarp Ca concentration was > 130 µg·g\(^{-1}\)FW would not be expected to be susceptible to the soft fruit condition reported by commercial shippers and handlers. However, fruit with mesocarp Ca concentration <130 µg·g\(^{-1}\)FW could be anticipated to occur a number of times during a year. Low mesocarp Ca uptake was related to preharvest environmental conditions and fertilization.

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