**Inducible Calcium Sinks and Preferential Calcium Allocation in Leaf Primordia of Dracaena sanderiana Hort. Sander ex M.T. Mast. (Dracaenaceae)**

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Additional index words. ribbon plant, calcium oxalate crystals, calcium oxalate monohydrate, raphides, periplasmic cuticular crystals

**Abstract.** The effect of 0, 3, and 7 mM Ca²⁺ on the allocation and deposition of Ca²⁺ into intracellular and sub-cuticular periplasmic calcium oxalate (CO) crystals was examined in leaf primordia of rooted cuttings of **Dracaena sanderiana** Hort. Sander ex M.T. Mast. Crystal development was monitored in two types of cuttings, those rooted in deionized water for 18 months and those rooted in Metro Mix 500 for 6 weeks. Response differed remarkably depending on the type of cutting. Cuttings rooted in deionized water deposited sub-cuticular crystals at the expense of intracellular crystals (raphides). The number of sub-cuticular crystals in leaf primordia of cuttings rooted in deionized water grown in solutions supplemented with either 0, 3, or 7 mM Ca²⁺ was similar, but crystals were considerably smaller in plants grown in 0 mM Ca²⁺. Sub-cuticular crystals appeared developmentally earlier in leaf primordia of all cuttings grown in either 3 mM or 7 mM Ca²⁺ than in cuttings rooted in deionized water grown in 0 mM Ca²⁺. This finding supports the premise that deposition of sub-cuticular crystals is modulated by Ca²⁺ levels and could be induced at an earlier ontogenetical stage by raising rhizospheric Ca²⁺ levels or by lowering rhizospheric Ca²⁺ levels. The total number of sub-cuticular crystals per epidermal cell did not differ significantly between treatments implying that crystal nucleation sites are predetermined and finite in number. In contrast, the formation of intracellular raphides was highly variable and depended on Ca²⁺ concentrations. In terms of Ca²⁺ prioritization, sub-cuticular CO crystals took precedence over intracellular CO raphides.

The occurrence of crystalline calcium oxalate (CO) is ubiquitous in higher plants (Franceschi and Horner, 1980). Plants form calcium oxalate (CO) crystals as means of coping with excess calcium in the cytosolic environment (Webb, 1999). Although the presence of CO crystals has been extensively recorded (Metcalfe and Chalk, 1994), the influence of varied rhizospheric Ca²⁺ levels on crystal development has not been clarified.

**Materials and Methods**

Stock **D. sanderiana** plants were grown in 3.8-L containers in Metro Mix 500 (Grace Sierra, Milpitas, Calif.) in a polycarbonate-covered greenhouse under maximum light intensity of 250–350 µmol·m⁻²·s⁻¹ [photosynthetically active radiation (PAR) during daytime, and 100–150 µmol·m⁻²·s⁻¹] during winter. Temperature was maintained between 20 to 24 °C (night) and 28 to 32 °C (day). The stock plants were top-dressed with 5 g per pot 20N–8.7P–16.6K controlled-release fertilizer (Grace Sierra) every 12 weeks.
To clarify the effect of different levels of rhizospheric calcium on development of the two crystal types, rooted cuttings that exhibited suppressed crystal development were used. We used the following procedure, which achieved this goal. Six-leaf shoot cuttings were taken from stock plants and placed into 20-L leakproof glass containers filled with 5 L of deionized (d.i.) water in the same greenhouse as the stock plants. The cuttings were held upright by insertion into polyurethane-coated test tube racks. Deionized water was added as needed to maintain volume at 5 L.

Because normal raphide development begins when the leaf primordia reaches 1.0% to 1.5% of mature leaf size (Pennisi et al., 2001a), primordia of this developmental age were periodically examined. Four weeks after root initials were visible on the cuttings, one plant was removed from the 20-L glass container monthly and the designated leaf primordia was examined with a microscope equipped with polarizing optics to determine the presence of CO crystals. A total of 18 months after cutting removal from the stock plants was required before raphides were undetectable in the designated primordia. During this time the cuttings formed roots and produced an average of three to four new leaves. These new leaves were smaller than older leaves and the time period between emergence of new leaves increased. However, no other indication of mineral deficiency appeared. Previous research had shown that Dracaena sanderiana has a low nutrient requirement and grows slowly (Pennisi and McConnell, 1997). These rooted cuttings were considered raphide deficient and the term “deficient” is subsequently applied to them.

The initial six leaves on the cuttings when placed in d.i. water senesced at the rate of about one leaf every three months. These leaves were removed at experiment initiation, and the rooted deficient cuttings had four fully expanded leaves. Another group of six-leaf shoot cuttings were taken from the stock plants six weeks prior to experiment initiation. They were placed into Metro Mix 500 (Grace Sierra, Inc.) under mist in a greenhouse (Fig. 1). After 6 weeks, the potting mix was carefully rinsed from the roots. After positive microscopic evaluation for raphides these cuttings were considered raphide nondeficient and the term “nondeficient” is subsequently applied to them.

At experiment initiation, all cuttings had four fully expanded leaves. Nine deficient and nine nondeficient cuttings were randomly selected. Each rooted cutting was placed in a 100 mL beaker filled with 80 mL of d.i. water containing 100 ppm of nitrogen (KNO₃), supplemented with 0, 3, or 7 mM Ca²⁺ using calcium acetate as the calcium source. (Fig. 1). The Ca²⁺ concentrations used were based on experiments used to induce intracellular raphide formation in Lemna minor (Franceschi, 1989). Other nutrients (e.g., phosphorus, magnesium, etc.) were excluded to avoid potential interference with the uptake of Ca²⁺ and CO crystal formation. Potassium has not been shown to be incorporated in CO crystals (Franceschi and Horner, 1980), therefore potassium nitrate was used as the nitrogen source.

Beakers were covered with plastic film to minimize evaporation. Liquid levels were kept at 80 mL with addition of d.i. water as needed. The beakers were arranged in a completely randomized design with three replicates per Ca²⁺ treatment (Fig. 1). Plants were grown for four months in a growth chamber under 12-h photoperiod, 350 μmol·m⁻²·s⁻¹ (light sources Sylvania VHO Full White lamps and Incandescent 25W rough surface lamps), and temperatures of 20 °C day/15 °C night (12 h cycle).

Statistical analysis was performed using analysis of variance (ANOVA) procedure in Microsoft Excel®. Dracaena sanderiana primordia were observed with a Nikon Optiphot-Pol research microscope (Nikon Nippon Kogaku K.K., Tokyo) equipped with polarizing optics. Detailed cellular measurements were made with an ocular micrometer. Crystal counts were taken from micrographs. Photographs were taken with an automatic Nikon UFX-II camera attachment (Nikon Nippon Kogaku K.K., Tokyo).

Results and Discussion

Two crystal maturation zones occur in the elongating leaf primordia; the raphide idioblast zone and the sub-cuticular pericellular crystal zone (Fig. 2A). Dracaena sanderiana exhibits typical monocot leaf primordia growth. Consequently, the distance between the base of the leaf primordia and the sub-cuticular crystal maturation zone is correlated to the physiological age of the cells where sub-cuticular crystal initiation has begun. The shorter this distance, the younger the cells; the longer this distance, the older the cells.

Deficient rooted cuttings. The sub-cuticular crystal maturation zone occurred physiologically 1.5% of mature leaf size (Pennisi et al., 2001a), while the raphide idioblast zone occurred physiologically 3.0% of mature leaf size (Franceschi and Horner, 1980). The idioblast and sub-cuticular crystal zones were visible on the cuttings, one plant was removed from the 20-L glass container monthly and the designated leaf primordia were examined with a microscope equipped with polarizing optics. Detailed cellular measurements were made with an ocular micrometer. Crystal counts were taken from micrographs. Photographs were taken with an automatic Nikon UFX-II camera attachment (Nikon Nippon Kogaku K.K., Tokyo).

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ologically and anatomically later in deficient cuttings grown in 0 mM Ca\textsuperscript{2+} (Fig. 2B) compared with deficient cuttings grown in 3 mM Ca\textsuperscript{2+} and 7 mM Ca\textsuperscript{2+} (Table 1; Fig. 2D–E). In contrast, the sub-cuticular crystal maturation zone occurred physiologically and anatomically earliest in deficient cuttings grown in 7 mM Ca\textsuperscript{2+} compared with deficient cuttings grown in 0 mM Ca\textsuperscript{2+} and 3 mM Ca\textsuperscript{2+} (Table 1). Minute crystals were observed in the sub-cuticular crystal maturation zone of deficient cuttings grown in 0 mM Ca\textsuperscript{2+} (Fig. 2C). These crystals did not exceed 3 \mu m in size. In contrast, cuttings grown in 7 mM Ca\textsuperscript{2+} developed some large (9–10 \mu m in length) sub-cuticular crystals, while the largest crystal size in cuttings grown in 3 mM Ca\textsuperscript{2+} was 6–8 \mu m (Table 1). Sub-cuticular crystals between 9 and 10 \mu m were seldom observed in earlier examinations of leaf primordia of container-grown plants (Pennisi, 1999). Although crystal size was affected by Ca\textsuperscript{2+} regimes, the total number of sub-cuticular crystals per cell was not (Table 1).

The most striking feature of deficient cuttings was the numerical variability of raphide idioblasts in the leaf primordia (Table 1; Fig. 2B). Cuttings grown in 0 mM Ca\textsuperscript{2+} had no raphide idioblasts while cuttings grown in 3 and 7 mM Ca\textsuperscript{2+} developed large numbers of raphide idioblasts. The greatest number of intracellular raphides formed in cuttings grown in 7 mM Ca\textsuperscript{2+} (Fig. 2E).

**Nondeficient rooted cuttings.** Nondeficient cuttings also exhibited pronounced differences in sub-cuticular crystal deposition and formation of raphide idioblasts (Table 1). The sub-cuticular crystal maturation zone occurred earliest in leaf primordia of cuttings grown in 7 mM Ca\textsuperscript{2+} (Fig. 3D) compared with other treatments (Fig. 3A and B; Table 1). Sub-cuticular crystals up to 10 \mu m were found in all treatments (Fig. 3F), with the greatest number of large crystals found in cuttings grown in 7 mM Ca\textsuperscript{2+} (Table 1). Total number of sub-cuticular crystals per epidermal cell was not significantly different among nondeficient cuttings for any Ca\textsuperscript{2+} concentration. The most pronounced difference between Ca\textsuperscript{2+} treatments in nondeficient cuttings was the number of raphide idioblasts. Highest values were counted in cuttings grown in 7 mM Ca\textsuperscript{2+}.

**Comparison between deficient and nondeficient rooted cuttings.** Deficient and nondeficient cuttings had similar numbers of sub-cuticular crystals at all Ca\textsuperscript{2+} levels (0, 3, or 7 mM Ca\textsuperscript{2+}) but the sub-cuticular crystals deposited in cuttings supplemented with Ca\textsuperscript{2+} were larger. The large crystals (10 \mu m) found in deficient and nondeficient plants grown in 7 mM Ca\textsuperscript{2+} might have occurred because crystal nucleation occurred earlier in leaf development, thus the developmental period for crystal growth was extended. Conversely, the smaller crystal size in deficient plants grown in 0 mM Ca\textsuperscript{2+} may have occurred because developing epidermal cells had differentiated to the stage where deposition of cellulose material in the cell wall had been initiated and the sub-cuticular crystals became spatially separated from Ca\textsuperscript{2+} ions in the living protoplasm, inhibiting crystal growth. These positions are strongly supported by measurements of the sub-cuticular crystal maturation zone from the primordial leaf base (Table 1).

The interaction term (Ca-levels times nutrient status) for sub-cuticular crystals 6 to 10 \mu m was significant, while nutrient status was nonsignificant (Table 1). A subsequent analysis revealed a linear and quadratic relationship for calcium levels in the deficient cuttings but not for the nondeficient cuttings.

With the exception of deficient cuttings grown in 0 mM Ca\textsuperscript{2+}, both deficient and nondeficient cuttings formed raphide idioblasts. However, deficient cuttings developed...
Table 1. Sub-cuticular periplasmic and raphide crystal related characteristics of leaf primordia from deficient and nondeficient *Dracaena sanderiana* cuttings grown in three rhizospheric levels of Ca²⁺.

| Factors | Ca²⁺ levels | Significance (P value) |
|---------|-------------|------------------------|
|         | 0 mM (D) | 3 mM (ND) | 7 mM (D) | 7 mM (ND) | Ca²⁺ status | Nutrient Interaction |
| Sub-cuticular crystal maturation zone (µm from primordium base) | 2940  | 2550  | 2060  | 2090  | 1720  | 1970  | L ³°°, Q ³°° | 0.0005  | 0.0001 |
| Sub-cuticular crystal number (per epidermal cell) | 47  | 9  | 27  | 25  | 31  | 24  | L ³°°, Q ³°° | 0.007  | NS |
| <1–2 µm | 28  | 18  | 14  | 21  | 10  | 23  | NS | NS |
| 3–5 µm | 0  | 4  | 11  | 6  | 15  | 10  | L ³°°, Q ³°° | 0.001  | NS |
| 6–10 µm | 56  | 50  | 52  | 52  | 56  | 57  | NS | NS |
| Total crystal number per epidermal cell (sum of all size crystals) | 56  | 50  | 52  | 52  | 56  | 57  | NS | NS |
| Number of raphide idioblasts ≈ primordium length 1-5 mm | 0  | 2  | 9  | 6  | 11  | 10  | L ³°°, Q ³°° | 0.0002  | 0.0001 |
| Number of raphide idioblasts ≈ primordium length 6-14 mm | 0  | 10  | 45  | 25  | 54  | 32  | L ³°°, Q ³°° | 0.0001  | 0.0001 |

*Values are means of three replicates rounded to the nearest integer. Measurements were taken on primordia between 10–14 mm in length.

More raphides than the nondeficient cuttings. Deficient cuttings grown in 7 mM Ca²⁺ had 30 more raphide bundles per primordia than nondeficient cuttings grown in 7 mM Ca²⁺ (Table 1, primordium length 6–14 mm).

* Lemma minor roots formed raphides rapidly and reversibly (Franceschi, 1989). Entire new raphide bundles were formed within 30 minutes of induction in *Lemna minor* roots in medium supplemented with 5 and 7 mM Ca²⁺. However, peeled *Gleditsia triacanthos* and *Albizia julibrissin* leaflets required 3 to 5 d to form raphide bundles after an inductive stimulus (elevated rhizospheric Ca²⁺ levels) (Borchert, 1985). Experiments with *Canavalia ensiformis* showed that the number of crystal idioblasts were one-half when plants were grown in low Ca²⁺ levels (0.01 mM Ca²⁺) compared with "normal" Ca²⁺ levels (0.4 mM Ca²⁺) (Frank, 1972).

A plant's ability to utilize stored intracellular Ca²⁺ when subjected to low-Ca²⁺ stress may explain why one tomato line did not exhibit deficiency symptoms compared to another line (Behling et al., 1989). The improved performance of the first line appeared to be associated with a slow movement of absorbed Ca²⁺ allowing for continued growth of the shoot apex and upper lamina under Ca²⁺ deficiency conditions. In contrast, the inefficient second tomato line exhibited little upward movement of Ca²⁺ in the plant after absorption. Franceschi (1989), demonstrated dissolution of CO₂ raphides in *Lemna minor* roots using Ca-ionophore to expedite the process.

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The sub-cuticular crystal maturation zone in leaf primordia 3–4 mm in length of both deficient and nondeficient cuttings grown in 3 and 7 mM Ca²⁺ appeared earlier than in container-grown plants using recommended fertilizer levels (Pennisi, 1999). This finding implies that the deposition of sub-cuticular crystals can be induced at an earlier ontogenetical stage by elevating rhizospheric...
The total number of sub-cuticular crystals per epidermal cell did not differ significantly among cuttings. This finding implies that sub-cuticular crystal nucleation sites are pre-determined and finite in number. The number of intracellular raphide bundles in deficient and nondeficient cuttings increased when rhizospheric Ca²⁺ levels were increased. Highest raphide idioblast numbers were found in deficient cuttings grown in 7 mM Ca²⁺. This elevated rate of CO crystal deposition may have been induced by a decrease and/or a delay in functioning of normal Ca²⁺-utilizing pathways. In terms of Ca²⁺ prioritization, subcuticular CO crystals took precedence over the intracellular CO raphides. This observation was most obvious in deficient cuttings that had been grown in d.i. water for 18 months prior to experiment initiation.

This study represents the first attempt to determine the effects of rhizospheric Ca²⁺ supply on crystal deposition in a plant species that features two or more distinct CO types in different locations of the plant body. Our results support the hypothesis that crystal-forming cells may provide a mechanism for regulation of Ca²⁺ levels in plant tissues and that crystal idioblasts act as a storage depot for Ca²⁺ ions (Franceschi, 1989). They also show that the size of CO crystals can be markedly affected when a plant is subjected to a sudden increase in Ca²⁺ ions and that the nutritional status of a plant has a pronounced effect on the number of raphide idioblasts formed when rhizospheric calcium levels are elevated. This would imply that nutritional programs can markedly increase or decrease the number CO crystals in plants and these changes can affect crop quality (Stebbins et al. 1972). Additional research is needed to advance our understanding of the relationship between rhizospheric calcium levels, CO crystals in plants and crop quality.

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