Physiological, Biochemical, and Molecular Changes in *Pelargonium* Cuttings Subjected to Short-term Storage Conditions

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Abstract. The purpose of this study was to evaluate physiological, biochemical, and molecular changes that occur in unrooted *Pelargonium ×hortorum* cuttings during storage. *Pelargonium* cuttings of ‘Sincerity’ (good shipper), ‘Wendy Ann’ (moderate shipper) and ‘Snowmass’ (poor shipper) were stored at 25 °C and evaluated over a 5-day period. Following removal from storage, cuttings of all cultivars exhibited steady and significant decline in photosynthesis, respiration, carbohydrate, starch, and protein over time. However, no significant differences were observed among cultivars for all of these parameters. Ethylene levels produced by ‘Sincerity’ and ‘Wendy Ann’ began to increase 3 days following storage; whereas, ‘Snowmass’ showed an increase after 1 day, reaching a peak at 3 days, and then declined. When unrooted cuttings of ‘Snowmass’ were stored for 5 days at temperatures ranging from 4 to 25 °C, it was observed that those stored at 4 °C had a significantly higher visual rating, chlorophyll content, and root and shoot weight than at higher temperatures tested. As temperature increased from 10 to 25 °C, quality of cuttings declined. Changes in gene expression of two ACC synthases and an ACC oxidase were evaluated in ‘Snowmass’ cuttings stored at 4 and 25 °C. Correlations between ethylene and ACC levels with gene expression were observed. Chemical name used: 1-amino-1-cyclopropane-1-carboxylic acid (ACC).

Production of *Pelargoniums* for use as cut flowers, bedding plants, and potted plants is a worldwide industry. Wholesale value of vegetatively and seed-propagated *Pelargoniums* increased by 60% from 1979 to 1990. In the United States, *Pelargoniums* account for 17.3% of all wholesale bedding plant sales. In 1990, *Pelargoniums* produced from cuttings were valued at almost $86.5 million, far more than seed or flat *Pelargoniums* (Berninger, 1993). To meet the increasing market demand, production and shipment of healthy cuttings and plants are necessary. However, current shipping and storage techniques have had deleterious effects on *Pelargonium* cuttings. Wilting, chlorosis, necrosis, and abscission of leaves typically follows most shipping and storage procedures (Carrow and Bahnemann, 1980; Kaltaler, 1966; Marousky and Harbaugh, 1981). Chlorotic leaves reduce acceptability of cuttings and promote susceptibility to *Botrytis*, which readily spreads within shipping containers (Carrow and Bahnemann, 1980). As a result, regrowth of cuttings is also jeopardized following damage in storage (Carrow and Bahnemann, 1980).

Several biochemical changes occur during leaf senescence, including changes in chlorophyll, protein, nucleic acids, respiration, and plant hormones. The most visible evidence of leaf senescence is loss of chlorophyll. *Pelargonium* cuttings have been shown to become chlorotic when packed and exposed to simulated storage conditions (Kaltaler, 1966; Krebs and Zimmer, 1977). Reports have indicated that proteins are rapidly degraded during senescence before leaf chlorosis (Martin and Thimmann, 1972; Mizrahi et al., 1970; Osborne, 1962). It is well documented that many forms of stress stimulate ethylene production, which results in an acceleration of senescence (Abeles et al., 1992). *Pelargoniums* are ethylene-sensitive flowering plants (Nell, 1993). Storage conditions to reduce ethylene production and maintain quality of plant material, including storage under low pressure, low temperature, and supply of ethylene biosynthesis or action inhibitors such as aminooxyacetic acid, (AOA), aminoethoxyvinylglycin (AVG), and silver thiosulfate, (STS) have been tested (Eisenberg et al., 1976; Paton and Schwabe, 1987; Purier and Mayak, 1987; Whalen and Wulster, 1991). However, these techniques have produced limited success in maintenance of quality of *Pelargoniums* during storage. Although applying STS inhibits ethylene action and reduces petal shattering, it is only effective during early storage. Moreover, STS renders plants susceptible to *Pythium* root rot (Nell, 1993) and reduces root formation (Whalen and Wulster, 1991), and it has been recently banned for commercial use (Oglevee, personal communication). AOA and AVG inhibit ACC synthase activity, while Co²⁺ inhibits ACC oxidase activity, but these are not used due to phytotoxicity problems (Wang and Arteca, 1992). Temperature control appears to be the most effective method to avoid leaf chlorosis in cuttings and plugs (Nell, 1993), but it is costly and in some cases not practical.

Recently, tomato fruit storage problems have been reduced by transferring RNA antisense ACC synthase or ACC oxidase or RNA sense ACC deaminase into tomato plants (Hamilton et al., 1990; Klee et al., 1991; Oeller et al., 1991). ACC synthase and ACC oxidase genes have been identified in many plants (Kende, 1993), and ACC deaminase was found in bacteria (Klee et al., 1991). Since ACC is the immediate precursor of ethylene, ACC deaminase can remove the amino group of ACC, thereby inhibiting the conversion of ACC to ethylene.

In this study, we evaluated changes in the physiology and biochemistry of unrooted *Pelargonium* cuttings during storage. We also evaluated changes in two *Pelargonium* ACC synthase genes (*GAC-1, GAC-2*) (Wang and Arteca, 1995) and a *Pelargonium* ACC oxidase gene (*GEFE-1*) (Wang et al., 1994) following storage. Our ultimate goal is to use antisense technology to reduce shipping injury and storage stress in *Pelargonium*.

Materials and Methods

Plant materials. Unrooted *Pelargonium ×hortorum* ‘Sincerity’ (good shipper), ‘Wendy Ann’ (moderate shipper), and ‘Snowmass’ (poor shipper) were subjected to short-term storage. Unrooted cuttings of all cultivars were stored at 4 °C for 1 day, and thereafter at 25 °C for 4 days. Following removal from storage, cuttings of all cultivars exhibited steady and significant decline in photosynthesis, respiration, carbohydrate, starch, and protein over time. However, no significant differences were observed among cultivars for all of these parameters. Ethylene levels produced by ‘Sincerity’ and ‘Wendy Ann’ began to increase 3 days following storage; whereas, ‘Snowmass’ showed an increase after 1 day, reaching a peak at 3 days, and then declined. When unrooted cuttings of ‘Snowmass’ were stored for 5 days at temperatures ranging from 4 to 25 °C, it was observed that those stored at 4 °C had a significantly higher visual rating, chlorophyll content, and root and shoot weight than at higher temperatures tested. As temperature increased from 10 to 25 °C, quality of cuttings declined. Changes in gene expression of two ACC synthases and an ACC oxidase were evaluated in ‘Snowmass’ cuttings stored at 4 and 25 °C. Correlations between ethylene and ACC levels with gene expression were observed. Chemical name used: 1-amino-1-cyclopropane-1-carboxylic acid (ACC).
(poor shipper) cuttings were obtained from virus-indexed stock from Oglevee Ltd. (Connellsville, Pa.). Three cuttings were rooted in a 15-cm-diameter standard plastic pot containing growing media and placed on a mist bench in a greenhouse under natural photoperiod. The growing medium contained 184 g calcitic lime (CON-LIME, Bellefonte, Pa.), 31.4 g KNO₃ (Hummert, St. Louis), 21.2 g triple super phosphate 0–46–0 (available P.O.; Hummert), 83.4 g gypsum 83% (Hummert), and 4.03 g fritted trace elements no. 555 (W.R. Grace & Co., Fogelsville, Pa.) in 100 L of 4 peat : 2 perlite : 1 soil. The medium was steam-sterilized for at least 4 h before use. Pots were subjected to a mist (6 s mist every 6 min) for 20 h·d⁻¹ for the first 3 d then gradually decreased (by 4 h/week) to 8 h·d⁻¹ by the fourth week. After 4 to 6 weeks, pots were transferred to a bench without mist, fertilized at every watering with 250 ppm N of 15–16–17 (15N–7.04P–14.1K) fertilizer (W.R. Grace & Co.) and leached once a week with tap water. Fungicides and pesticides were used as needed. The greenhouse was vented when day temperature reached 26 °C and was kept at 18 °C during the night. Plants were grown for another 4 to 6 weeks before use in our experiments.

Eight cuttings (10 to 15 cm long) were collected from the three cultivars from stock plants grown above, the cut base was wrapped in moist Kim Pak (Seedburo Equipment Co., Chicago), and cuttings were placed in a partially sealed plastic bag packed in shipping boxes and stored at 25 °C unless designated otherwise. For all parameters tested in this study, four individual cuttings were used per replication at each sampling date. All experiments were repeated three times.

**Measurements of photosynthesis and respiration.** A Hansatech LD-2 system (Decagon, Pullman, Wash.) was used in the study. This system measures leaf photosynthesis or respiration by monitoring oxygen production or consumption within a sealed chamber containing a leaf disk. Carbon dioxide concentration was maintained at saturating levels to keep stomates open. Temperature was maintained at 25 °C in the chamber. Oxygen concentration was measured using a Clark electrode (Decagon), which can detect minute changes in oxygen.

**Ethylene and ACC determinations.** Ethylene was analyzed with a Hewlett Packard (Valley Forge, Pa.) 5830 dual-column gas chromatograph equipped with 3.175-mm-od x 1.2-m columns packed with activated alumina. Injector port, column, and flame ionization detector temperatures were 90, 90, and 300 °C, respectively (Wang and Arteca, 1992). Three fully expanded leaves (3 to 4 g fresh weight) were collected from each cutting, placed in a 50-mL vial containing 1 mL distilled water, sealed with a serum cap, and kept at 4 or 25 °C in the dark for 2 h before measuring ethylene.

For ACC analysis, 1 g of leaf tissue was extracted in 85% ethanol. The extract was dried, resuspended in 1 mL of water, and centrifuged at 4500 x g for 15 min. An aliquot (300 μL) was mixed with 0.5 mL ice-cold 20 mm HgCl₂ in a 13 x 100-mm test tube and sealed with a serum cap. For converting ACC to ethylene, 0.2 mL of an ice-cold mixture of 5% NaOCl and 15 m NaOH (2:1 v/v) was injected. The tube was vortexed for 15 s, allowed to incubate for 5 min, and ethylene in the test tube headspace was determined by gas chromatography (Tsai et al., 1988). Remaining leaves were frozen in liquid N₂ and stored at –80 °C until analyzed.

**Total protein, carbohydrate, starch, and chlorophyll determination.** Total protein was determined according to Bradford (1976) using bovine serum albumin as a standard. Sucrose, glucose, fructose, and starch were determined according to the manufacturer’s protocols using UV-based kits (Boehringer Mannheim Biochemicals, Indianapolis). Chlorophyll was determined according to Witham et al. (1971).

**Temperature treatments.** ‘Snowmass’ cuttings were stored at 4, 10, 20, and 25 °C for 5 d. Following storage, four cuttings from each of the treatments were used for chlorophyll extraction, determining leaf weight per plant, and visual rating, with 5 = best (dark green and turgid) and 1 = worst (chlorotic and/or not turgid). The remaining four cuttings from each treatment were planted in soil and grown for 10 d, and then measurements were recorded for leaf weight per plant, root weight, and chlorophyll content. Visual ratings were also made. Two sets of controls were used where one set was sampled before rooting and the other was planted and grown for 10 d.

**RNA extraction.** RNA was extracted according to Wang and Arteca (1995). Briefly, 15 g of leaves were ground to a fine powder in liquid N₂ and then homogenized in 30 mL of homogenization buffer 0.2 m Tris, pH 9, containing 5 m guanidium thiocyanate, 1% polyvinyl pyrrolidone, 0.62% sodium Sarkosyl, and 1% b-mercaptoethanol. The homogenate was centrifuged at 3000 x g, at 4 °C for 20 min. The supernatant was filtered through two layers of MiraCloth (Calbiochem) into a polycarbonate tube and centrifuged at 47,000 x g at 4 °C for 30 min. The supernatant was layered on a 10-mL 5.7 m CsCl cushion and centrifuged at 112,000 x g in a SW-28 rotor at 4 °C for 24 h. After centrifugation, the RNA pellet was collected from the bottom of the tubes as described by Sambrook et al. (1989).

**Northern analysis.** Total RNA was fractionated on a 1.5% agarose gel containing 2.2 m formaldehyde. Following electrophoresis, RNA was transferred to a nylon membrane by capillary transfer in 2× SSC overnight, air dried for 1 h, and fixed by UV exposure. Filters were prehybridized at 42 °C with a solution containing 6× SSC, 5× Denhardt’s reagent, 0.1% SDS, 100 mg·mL⁻¹ denatured fragmented salmon sperm DNA, and 50% formamide for 4 h. Hybridization was performed overnight at 42 °C using 1 x 10⁶ cpn·mL⁻¹ of GAC-1, GAC-2, and GEEF-1 probes were labeled with ³²P-dCTP using random priming. Membranes were washed at room temperature in 2× SSC plus 0.1% SDS for 15 min, 2× SSC plus 0.1% SDS at 62 °C for 15 min, and 0.2× SSC plus 0.1% SDS at 62 °C for 15 min. The blot was exposed to Kodak XAR5-X-ray film with two intensifying screens at –80 °C for 24 to 48 h. Blots were rehybridized using a pea ribosomal gene (Jorgenson et al., 1982) to ensure that equal amounts of total RNA were present in each lane.

**Results and Discussion**

All three cultivars have exhibited a decline in photosynthetic rate, total protein, starch, sucrose, glucose, and fructose 1 d following dark storage at 25 °C (Figs. 1–3). These changes occurred before a reduction in chlorophyll content or respiration rate and before induction of high ethylene levels. This gradual decline for all parameters and cultivars tested, except for ethylene, continued for 5 d (Figs. 1–3). These findings agree with earlier reports on rapid protein degradation before visible leaf chlorosis (Martin and Thimann, 1972; Mizrahi et al., 1970; Osborne, 1962). In addition, this work agrees with other reports indicating that, when Pelargonium cuttings are packed and exposed to simulated storage conditions, they turn chlorotic (Kaltaler, 1966; Krebs and Zimmer, 1977). ‘Snowmass’ shows a significant increase in ethylene production 3 d after storage followed by a decline to undetectable levels by the fifth day. This is in contrast to the observed storage response of ‘Sincerity’ and ‘Wendy Ann’ (Fig. 3). It is well documented that many forms of stress stimulate ethylene production, which results in senescence (Abeles et al., 1992). Exogenous applications of ethylene to Pelargoniums during simulated storage conditions have been shown to promote leaf chlorosis (Purer and
Our results agree with others using different *Pelargonium* cultivars, whereby a 3- to 4-fold increase in ethylene production has been obtained following simulated storage (Purer and Mayak, 1987).

When ‘Snowmass’ cuttings were stored at different temperatures from 4 to 25 °C, a gradual decline in visual rating was observed with increasing temperatures (Fig. 4). Leaf weight and chlorophyll content were unchanged at 4 and 10 °C; however, at 20 and 25 °C, leaf weight and chlorophyll content were significantly

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**Fig 1.** Changes in photosynthesis, respiration, and chlorophyll in *Pelargonium* ‘Sincerity’, ‘Wendy Ann’, and ‘Snowmass’ cuttings following removal from storage at 25 °C in the dark. The error bar in the upper right hand corner of each graph corresponds to standard error of the mean.

**Fig 2.** Changes in glucose, fructose, and sucrose in *Pelargonium* ‘Sincerity’, ‘Wendy Ann’, and ‘Snowmass’ cuttings following removal from storage at 25 °C in the dark. The error bar in the upper right hand corner of each graph corresponds to standard error of the mean.
reduced (Fig. 4). When ‘Snowmass’ cuttings were stored for 5 d at different temperatures and then planted and allowed to grow for 10 d before evaluation, there was no difference in visual rating, leaf weight, chlorophyll, and root weight between the 4 °C treatment and control. At 10 °C, a reduction in visual rating and root weight was observed; however, there was little difference in leaf weight and chlorophyll content. When cuttings were stored at 20 to 25 °C, there was a significant reduction in all of the parameters tested (Fig. 4). These results were similar to those reporting reduced temperatures were effective in maintaining quality in various cultivars (Nell, 1993).

To characterize the effects of storage on ethylene production in

![Graph](image-url)

**Fig. 3.** Changes in starch, protein, and ethylene in *Pelargonium* ‘Sincerity’, ‘Wendy Ann’, and ‘Snowmass’ cuttings following removal from storage at 25 °C in the dark. The error bar in the upper right hand corner of each graph corresponds to standard error of the mean.

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**Fig. 4.** Changes in visual rating, chlorophyll (Chl), root and leaf weight in *Pelargonium* ‘Snowmass’ cuttings after storage in the dark for 5 d at 4, 10, 20, and 25 °C. A control sample was taken before storage. Following storage, half of the cuttings were evaluated immediately (top) and the other half was planted, grown for 10 d, and then evaluated (bottom). The error bar in the upper right hand corner of each graph corresponds to standard error of the mean.
When ‘Snowmass’ cuttings were stored at 4 and 25 °C, GAC-1 mRNA was not detected at 4 °C, while it increased 3 to 4 d following storage at 25 °C (Fig. 5). One day following storage at 4 °C, there was an increase in GAC-2 mRNA, while days 3 and 4 were similar to control and correlated with an increase in ACC. GAC-2 mRNA levels from cuttings stored at 25 °C increased 1 d following storage and to higher levels after 3 to 4 d in storage, which correlated with increased ethylene and ACC levels. GEF-E-1 mRNA levels increased in cuttings stored at either 4 or 25 °C 1 d following storage and to higher levels 3 to 4 d correlating with increased ethylene and ACC levels (Fig. 5). Since GAC-1 and GEF-E-1 have also been induced by water stress (Wang and Arteca, 1995) it is likely that they were involved in the production of ethylene and decline in quality following storage.

Worldwide demand for virus-indexed Pelargonium plants and cuttings has increased dramatically over the past 10 years. However, obtaining uniform plants from cuttings and minimizing petal shattering in Pelargoniums that have been shipped long distances or stored for long periods is a major problem, as they are exposed to darkness, transit vibration, water stress, and temperature extremes. Some cultivars are not severely affected and perform well following shipping or storage; whereas, many desirable cultivars are adversely affected resulting in petal shattering, leaf chlorosis, disease developments, and poor rooting. While ethylene in many cases has been implicated in decline in quality of cuttings and petal shattering during short-term shipping or storage, there are no methods available to overcome these adverse effects. Current methods include reduced temperatures, hypobaric storage, ethylene scrubbers, or use of ethylene action blockers or ethylene biosynthesis inhibitors.

The use of antisense technology has been successful in delaying fruit ripening in tomato, which is very sensitive to ethylene (Hamilton et al., 1990; Oeller et al., 1991). Climacteric fruit can be induced to ripen by treatment with ethylene at concentrations above 0.1 mL of ethylene/L air. It has been shown that the use of antisense technology using the ACC synthase gene can reduce ethylene levels to below 0.1 nL ethylene/g fruit mass per hour (Oeller et al., 1991). We have shown that Pelargoniums are similar in sensitivity to tomato; therefore, it is likely that this approach will be also useful for Pelargoniums. In this study we have shown that a Pelargonium ACC synthase cDNA (GAC-1) and an ACC oxidase (GEFE-1) are turned on under storage conditions, thereby enabling us to regulate a specific gene that is turned on by shipping and storage stress while allowing other genes required for the production of ethylene during normal plant growth and development to function normally. While the use of antisense technology is no longer novel for use in fruit ripening, it is novel for delaying leaf and flower senescence in Pelargoniums to produce plants with reduced problems from injury, thereby, increasing the longevity of plants in the home environment.

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