Ethephon Residual Catalysis on Unrooted Impatiens hawkeri Cuttings and Stock Plants

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Abstract. Ethephon [(2-chloroethyl) phosphonic acid] is used to increase stock plant cutting productivity through increased flower and flower bud abscission and branching. However, ethylene evolution resulting from ethephon application is suspected to cause leaf abscission of unrooted cuttings during shipping. It was the objective of this study to assess ethylene evolution from ethephon-treated cuttings during storage and shipping of unrooted cuttings. Impatiens hawkeri W. Bull ‘Sonic Red’ and ‘Sonic White’ stock plants were treated with 0, 250, 500, or 1000 mg L\(^{-1}\) ethephon. Cuttings were harvested from 1 to 21 days later and each harvest was stored at 20 °C in sealed jars for 24 h before ethylene measurement. Higher ethephon doses resulted in greater ethylene generation. Cuttings harvested 1 day after treatment with 0, 250, 500, or 1000 mg L\(^{-1}\) ethephon evolved 0.07, 1.3, 1.7, or 5.8 μL L\(^{-1}\) g\(^{-1}\) (fresh weight) ethylene in the first 24 h of storage at 20 °C, respectively. Twenty-one days after treatment, cuttings from the same plants evolved 0.05, 0.05, 0.15, or 0.14 μL L\(^{-1}\) g\(^{-1}\) (fresh weight) ethylene in the first 24 h of storage at 20 °C, respectively. As cuttings were harvested from Day 1 to Day 21, ethylene concentrations evolved within the first 24 h of storage decreased exponentially. Rinsing cuttings, treated 24 h earlier with 500 mg L\(^{-1}\) ethephon, by gently agitating for 10 s in deionized water reduced ethylene evolution to 0.7 μL L\(^{-1}\) g\(^{-1}\) (fresh weight) as compared with 1.7 μL L\(^{-1}\) g\(^{-1}\) for unrisned cuttings. Cuttings harvested 24 h after treatment with 500 mg L\(^{-1}\) ethephon stored at 10, 15, 20, and 25 °C for 24 h evolved 0.37, 0.81, 2.03, and 3.55 μL L\(^{-1}\) g\(^{-1}\) (fresh weight) ethylene. The resulting mean temperature coefficient (Q\(_{10}\)) for the 10 to 25 °C range from all replications was 5.15 ± 0.85. Thus, ethylene continues to evolve from ethephon-treated Impatiens hawkeri stock plants for up to 21 days and can accumulate to high concentrations during cutting storage.

Ornamental plant producers use cuttings shipped from around the world for finished plant production. The value of unrooted cutting imports into the United States totaled $61 million in 2006 with the bulk coming from Central America, South America, Mexico, and Asia (Foreign Agricultural Service, 2007). In the top 15 floricultural crop-producing states, flats, hanging baskets, and pots of geraniums (Pelargonium L’Hér. ex Ait.), poinsettias (Euphorbia pulcherrima Willd. ex Klotzsch), and New Guinea impatiens (Impatiens hawkeri W. Bull) had a wholesale value of $330 million and were produced from ≈138 million cuttings (National Agriculture Statistics Service, 2007). In this case, the loss of a single cutting is $2.38 lost in potential revenue. Understandably, reducing losses of cuttings during shipping is a critical research interest for cutting producers, plant plug producers, and finished plant producers.

Ethephon [(2-chloroethyl) phosphonic acid] is used on New Guinea impatiens stock plants and many other herbaceous species to increase branching and abscise sink tissues. Ethephon is lipophilic and, on entering the plant, undergoes a degradation reaction releasing ethylene, Cl\(^{-}\), and H\(_2\)PO\(_4\)\(^{-}\) (Warner and Leopold, 1969; Yang, 1969). Application of ethephon results in the abscission of flower buds and flowers, likely as a result of ethylene-mediated promotion of abscisic acid (ABA) synthesis (Goren et al., 1993; Hansen and Grossmann, 2000; Klee, 2002) and subsequent cellulase and polygalacturonase activity (Mishra et al., 2008), thus reducing sink tissue carbohydrate demands. Ethylene-related inhibition of indole-3-acetic acid (IAA) synthesis (Chadwick and Burg, 1970; Weber and Osborn, 1969) is likely cause of increased branching in both young liners and stock plants (Faust and Lewis, 2005; Hayashi et al., 2001) treated with ethephon.

Stock plant sink tissue management has been critical in lowering cutting losses resulting from carbohydrate depletion during shipping. Rapaka et al. (2008) demonstrated an inverse relationship between cutting carbohydrate content at harvest and subsequent postharvest leaf senescence. Quality of subsequent rooting is also directly proportional to carbohydrate content (Rapaka et al., 2005). Other authors have suggested a correlation between cut flower petal senescence and sugar depletion (van Doorn, 2004).

Problematically, excess ethylene can damage unrooted cuttings by promoting apical meristem necrosis, leaf yellowing, and abscission. Thus, cuttings harvested from ethephon-treated stock plants still evolving ethylene may be exposed to damaging ethylene concentrations during packaging and shipping. Because ethephon-treated plants do not rapidly release the abscised sink tissues, it may be possible to postpone cutting harvest after ethephon application until ethylene concentrations return to normal. The objective of this study was to investigate the persistence of residual ethephon activity on stock plants and unrooted cuttings. We applied a range of ethephon concentrations and subsequent storage treatments to show how temperature and rinsing of cuttings may affect ethylene evolution.

Materials and Methods

Plant material. Five unrooted cuttings of each Impatiens hawkeri W. Bull ‘Sonic Red’ and ‘Sonic White’ were propagated into 30-cm diameter, 13.85-L plastic pots containing a soilless growing substrate (Fafard 3P; Conrad Fafard, Inc., Agawam, MA). Once rooted, these plants were maintained as stock plants and irrigated manually with water alone, 600 mg L\(^{-1}\) nitrogen liquid fertilizer (20N–4.37P–16.6K; Ultrasol, SQM North America, Atlanta, GA), or flouvable lime (Limestone F; Cleary Chemical Inc., Dayton, NJ) to maintain a target electrical conductivity of 1.0 to 1.5 dS m\(^{-1}\) and a pH of 5.8 to 6.2 using the pourthrough monitoring method. Temperature set points were 24/18 °C (day/night), and average daily light integral was 6.1 mol m\(^{-2}\) d\(^{-1}\).

Ethephon application and cutting storage. Ethephon (Florel; Monterey Lawn and Garden Products, Inc., Fresno, CA) treatments were applied at 1600 HR with a 2-L compression hand sprayer (Hudson Manufacturing Company, Chicago, IL) at 0, 250, 500, or 1000 mg L\(^{-1}\) a.i. to three plants each. Cuttings were harvested at 0900 HR the next day and every 48 h thereafter for 21 d for a total of 11 harvests. Two node cuttings of
similar size and age with three to four fully expanded leaves were arbitrarily selected from all parts of the stock plant. Cutting fresh weights were recorded before six each were placed in sealed 0.95-L glass jars. The jars were stored in the dark at 20 ± 1 °C for 7 d and they remained closed for the storage duration. Ethylene concentrations in the jars were measured by gas chromatography every other day beginning 24 h after each harvest and ethylene evolved from the cuttings was calculated on a fresh weight basis. For the rinsing experiment, an additional group of 500 mg·L⁻¹ ethephon-treated cuttings was gently rinsed for 10 s in 100 mL of deionized water and gently blotted dry before storage.

For the temperature treatment experiment, four additional sets of 500 mg·L⁻¹ ethephon-treated cuttings were harvested 24 h after treatment and stored at 10, 15, 20, or 25 °C, respectively, in sealed 0.95-L glass jars for 7 d.

Ethylene determination and cutting condition. A gas chromatograph (Varian 3400; Varian Inc., Walnut Creek, CA) fitted with a glass column (Porapak Q, 80–100 mesh, 183 cm × 2 mm; Sigma Aldrich, Inc., St. Louis, MO) running at 120 °C injector, 120 °C column, and 130 °C detector (flame ionization) temperatures was used to measure ethylene concentration during storage. Flow rates for the He carrier, H₂, and O₂ were 30, 16, and 90 mL/min, respectively. Ethylene quantification was based on a response factor generated using a 1 μL·L⁻¹ ethylene standard. Injection volume was 1 mL of headspace gas drawn through a neoprene port on the jar lid.

In addition, evidence of phytotoxicity was recorded daily as the number of yellow or necrotic leaves, the percentage of leaf area affected, the number of abscised leaves, and turgidity, which was ranked on the following scale: 1 = all leaves turgid; 2 = one or two leaves wilting; or 3 = all leaves wilted.

Data analysis. Within the study, each of the three experiments was designed as a randomized complete block design of three blocks with treatments of ethephon concentration, storage temperature, and rinsing of residual ethephon. Experimental results (i.e., ethylene evolved as a result of ethephon dosage, storage temperature, or rinsing) were tested for significance and best fit linear, exponential, quadratic, or higher-order models using the GLM procedure. Model terms were evaluated for significance by comparison of F values at α = 0.05. Subsequent regression analysis using the best fit model was performed using PROC REG (SAS Institute, Cary, NC). The equation \( R^2 = R_1^2 \left( \frac{T_2 - T_1}{T_2} \right) \) was used to calculate Q₁₀ values for ethylene evolution during the first 24 h of storage 1 d after treatment for temperature ranges 10 to 15°, 15 to 20°, and 20 to 25 °C in each replication. The resulting data were tested using Scheffe’s procedure (PROC GLM) with an experiment-wise Type I error rate at α = 0.05.

Results

During storage, the quality and appearance of cuttings from all treatments were similar. Additionally, no statistically significant
differences were observed between cultivars in the initial experiments; thus, cuttings were pooled during subsequent experiments. Higher ethephon doses resulted in greater ethylene generation and ethylene accumulated to very high concentrations (Fig. 1) over the 7-d storage period in all experiments. Cuttings harvested 24 h after treatment with 500 or 1000 mg L^{-1} ethephon produced 1.7 or 5.8 μL L^{-1} g^{-1} ethylene, respectively, within the first 24 h after harvest. Control cuttings produced 0.06 μL L^{-1} g^{-1} ethylene in the same period. We reported ethylene accumulation within the first 24 h of storage for the rinsing, residual activity, and temperature experiments to simplify the results presentation and reduce duplication.

Ethylene evolved from ethephon-treated plants for at least an additional 21 d after treatment depending on application rate (Fig. 2A). Cuttings harvested 1 d after treatment with 0, 250, 500, or 1000 mg L^{-1} ethephon produced 0.07, 1.3, 1.7, or 5.8 μL L^{-1} g^{-1} ethylene in the first 24 h of storage, respectively. As subsequent sets of cuttings were harvested, ethylene produced within the first 24 h of storage decreased exponentially Day 1 through 21. Cuttings harvested 21 d after being treated by the same ethephon concentrations produced 0.05, 0.05, 0.15, or 0.14 μL L^{-1} g^{-1} ethylene in the first 24 h of storage, respectively.

Rinsing 500 mg L^{-1} ethephon-treated cuttings with water did not reduce evolved ethylene concentrations compared with that of the untreated control in the first 24 h after treatment (Fig. 2B). Cuttings harvested 24 h after treatment with 500 mg L^{-1} ethephon and rinsed in distilled water produced 0.7 μL L^{-1} g^{-1} ethylene in the first 24 h of storage compared with 1.7 μL L^{-1} g^{-1} ethylene from unrinseed cuttings and 0.05 μL L^{-1} g^{-1} for untreated cuttings.

Lower temperatures resulted in significantly less ethylene generation from ethylene-sensitive species packed alongside ethephon-treated cuttings (Fig. 3) within the first 24 h of storage. Cuttings harvested 24 h after treatment with 500 mg L^{-1} ethephon and stored at 10, 15, 20, and 25 °C evolved 0.37, 0.85, 2.59, and 3.56 μL L^{-1} g^{-1} of ethylene, respectively. The relationship between storage temperatures and ethylene concentrations evolved within the first 24 h of storage 1 d after ethephon application is best described by the linear equation, 0.21x – 1.97 (R^2 = 0.88). Additionally, these cuttings had a highest temperature coefficient (Q_{10}) of 10.25 from 15 to 20 °C, and 4.70 and 2.18 from 10 to 15 °C and 20 to 25 °C, respectively. Although the Q_{10} for 15 to 20 °C differed significantly from the Q_{10} for 10 to 15 °C and 20 to 25 °C, the latter two did not differ significantly from one another.

**Discussion**

Cuttings are typically packaged for shipment in polypropylene or polyethylene bags, which offer only a fraction of the headspace of the 0.95-L jars used in this study. Total shipping and handling times rarely exceed 4 d (R. Heins, A. Gerendas, personal communication). Thus, ethylene concentrations in this experiment measured in the first 24 h of storage may be artificially low. The same number of cuttings placed into a polypropylene shipping bag with very little headspace may evolve ethylene concentrations, which are severely damaging for some species. In shipping, problems may occur when cuttings of ethylene-sensitive species are packed alongside ethephon-treated cuttings. Species susceptible to ethylene concentrations as low as 0.1 μL L^{-1} such as Lantana camara L. or Portulaca oleracea L. (Leatherwood, 2008; Rapakka et al., 2007) could suffer leaf abscission when packaged alongside cuttings still releasing ethylene. Safe ethylene concentrations for all herbaceous cuttings are not yet known. However, concentrations of 0.01 μL L^{-1} for 20 h can damage the flowers of a number of cut flower species (Blankenship and Dole, 2003). Reports of ethylene damage during shipping such as leaf abscission have been difficult to confirm because ethylene dissipates once the packages are opened. Additional follow-up experiments should examine this headspace issue and combine various cuttings sensitive to ethylene with cuttings recently treated with ethephon.

Our findings support work by Woodrow et al. (1988) with [14C] ethephon, who reported that 300 mg L^{-1} ethephon degradation on Solanum lycopersicum L. var. cerasiforme (Dunal) Spooner, G.J. Anderson & R.K. Jansen releases ethylene continually for at least 9 d after treatment and is not synthesized de novo. Additionally, Riov and Yang (1982) demonstrated that after exposure to exogenous 12 μL L^{-1} ethylene, ethylene autocatalysis of Citrus sinensis L. Osbeck cv. Washington Navel leaf discs decreases to normal within 12 h of removing the exogenous ethylene source. Woolf et al. (1995) reported a decrease in ethylene evolution from Camellia saluenensis × C. japonica ‘Anticipation’ leaves and vegetative buds treated with 4 mL L^{-1} ethephon a.i. (Ethrel 48; Rhone-Poulenc Ltd., Wellington, New Zealand) as temperatures decreased from 30 to 10 °C. Thus, it appears that ethephon can have residual activity on impatients stock plants for at least 3 weeks, is the primary ethylene source observed in this study, and evolves ethylene more slowly at lower temperatures. Decreasing ethephon concentration results in lower ethylene concentrations evolved from treated cuttings. Thus, it may be possible to establish a minimal effective dose to improve branching and abscise flowers and flower buds while also reducing ethylene evolved. Although rinsing cuttings treated with 500 mg L^{-1} ethephon reduced the amount of ethylene generated from ethephon application, ethylene concentrations were similar to amounts generated by unrinseed cuttings harvested from stock plants treated with 250 mg L^{-1} ethephon. Rinsing most likely removes some of ethephon adhered to the cuticle surface but does not remove ethephon embedded in the cuticle. However, rinsing of cuttings or stock plants may not be
horticulturally feasible considering the costs and disease risks the action would impose.

We have demonstrated that ethephon ethylene evolution decreases over time. By delaying cutting harvests after ethephon application, growers could dramatically decrease the ethylene evolved during shipping. Growers report, however, waiting as little as 2 d before harvesting after ethephon application (B. Troost, personal communication). Awareness of the ethylene status of unrooted cuttings will help producers and growers manage the cold chain and propagation environment more precisely. Storage temperatures less than 15 °C reduced $Q_{10}$ and thus the amount of ethylene generated by treated cuttings within 24 h of harvest. It is possible that at lower temperatures, ethephon’s rate of absorption slows as the cuticle and cell membranes become less permeable. Because ethephon autocatalyzes in the apoplast (Yang, 1969), ethylene synthesizing metabolic activity (Woodrow et al., 1988) can be ruled out. Because ethephon degradation peaks between 15 and 20 °C, storage and handling of cuttings harvested from recently treated stock plants should be conducted at less than 15 °C. Once a cutting shipment leaves the production facility, however, control over temperature is a challenge and peaks of 25 °C are common. Thus, whenever possible, producers should avoid combined boxes of ethephon-treated cuttings and cuttings of ethylene-sensitive species. If combined shipments cannot be avoided, include in the shipment an ethylene blocker such as 1-methylcyclopropene (Blankenship and Dole, 2003).

Regardless of these findings, ethephon alters the carbohydrate distribution in the parent plant through abscission of flowers and flower buds, thus increasing carbohydrate availability to the shoots (Woodrow et al., 1988) and unrooted cutting carbohydrate content. Cuttings with higher carbohydrate content have improved adventitious root formation (Rapaka et al., 2005) and propagation performance. Thus, ethephon is still a valuable tool in cutting production yet should be used with an awareness of these findings. Further research should look into establishing minimal effective ethephon applications, handling and shipping procedures to minimize ethylene evolution and exposure, and delay of harvesting to allow ethephon dissipation.

**Literature Cited**


