Evaluation of a Sprayable Formulation of 1-Methylcyclopropene in Floriculture

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Abstract. We conducted a series of studies to determine the efficacy of a sprayable formulation of 1-methylcyclopropene (1-MCP; AFxRD-038) to inhibit ethylene-mediated flower abscission in Impatiens walleriana. Exposing Impatiens plants to 1.0 μL·L⁻¹ ethylene for 18 hours caused complete abscission of open flowers and most buds. Sprays of the novel 1-MCP formulation at concentrations >2.5 mg·L⁻¹ protected plants from ethylene. At 5 and 10 mg·L⁻¹, the efficacy of 1-MCP increased as spray volume increased from 102 mL·m⁻² to 306 mL·m⁻². 1-MCP was rainfast with no decrease in efficacy resulting from heavy overhead irrigation within 1–2 minutes of application. Prepared 1-MCP solutions (10 mg·L⁻¹) remained effective up to 2 weeks after mixing if held in airright containers. The sprayable 1-MCP formulation provided protection against exogenous ethylene for a maximum of 4 days and reduced stress-related abscission from 3 days of darkness (in the absence of exogenous ethylene) at 20 °C or 40 hours darkness at 28 °C.

Postproduction losses are an important financial constraint in the floriculture industry. Loss estimates of 5% to 30% per year have been made (Armitage, 1993; Healy, 2009; Jones, 2002; Staby et al., 1978). Losses can occur at many levels from the moment plants leave the production facility until plants are sold to the consumer. Floricultural crops, including many potted plants and bedding plants, are often shipped significant distances in dark trucks that may or may not have adequate temperature control. Losses can be attributed to water stress (especially in the retail environment), damage by mechanical means (vibration, abrasion, breakage) and by ethylene, which may be present exogenously in the shipping chain or could be produced directly by plants and flowers during the stress of transport. In the absence of adequate temperature control, warm temperatures during transport will potentially increase endogenous ethylene production and will increase sensitivity to ethylene whether exogenous or endogenous in nature. In many floriculture crops, ethylene results in flower, bud, or leaf abscission, epinasty, and hastening of senescence (Gibson et al., 2000; Reid and Jiang, 2012).

Skog et al. (2001) measured ethylene levels in midsize to large wholesale and retail businesses in Ontario, Canada. Detectable levels of ethylene (>0.01 μL·L⁻¹) were measured in 63% of air samples and these levels ranged from 0.01 to 10 μL·L⁻¹. Sources of ethylene included fresh or rotting produce, propane exhaust, smoke, gas-fired heating equipment, leaks from ethylene rooms used for gassing produce, and recently gassed produce. Areas that contained only flowers and had no produce on the premises exhibited ethylene concentrations ranging from undetectable (<0.01) to moderate (1.44 μL·L⁻¹). Loading areas should be kept separate from active growing plants and within these confined areas there is a need for ample ventilation, proper maintenance of equipment, and removal of plant debris to reduce ethylene contamination.

As it is unlikely to completely remove ethylene from the postproduction chain, a significant effort has been placed into technologies to reduce the effect of ethylene on floriculture crops. 1-MCP binds to ethylene receptor sites on the plant and with proper application will render plants unsusceptible to endogenous and exogenous ethylene for a period depending on plant growth (Blankenship and Dole, 2003). Current use of 1-MCP in floriculture is limited to a gaseous application sold under various names (EthylBloc; Florallife Inc., Walterboro, SC and Ethylene Buster; Chrysal Intl., Naarden, The Netherlands). With this technology, plants are enclosed in a sealed chamber (treatment room or sealed delivery truck) and the gaseous 1-MCP is released from the commercial powder with a wetting agent. The area must remain sealed for a specified time. This treatment method can be limiting to postproduction efficiency and convenience because plants or flowers must be enclosed in the 1-MCP atmosphere from 4 to 10 h to achieve full protection from ethylene (Jones and Edelman, 2013; Reid and Çelikel, 2008). Other release mechanisms include 1-MCP sachets that are dipped in water and placed into shipping boxes, a technique especially useful for boxed cut flowers or potted plants (Jones and Edelman, 2013).

In the last few years, a sprayable formulation of 1-MCP (AFxRD-038; Rohm and Hass, Philadelphia, PA) has been registered for in-orchard use on fruit (apples, pears, and kiwifruit) and field crops (sunflower, wheat, and rice) as Harvist™ and Invinsa™. While the sprayable formulation has gained a significant presence in these crops, it is not currently registered for use on floriculture crops, and information on its use in greenhouse settings is limited.

Impatiens plants are highly sensitive to ethylene and show rapid flower abscission (overnight) when exposed to very low concentrations of ethylene (Dostal et al., 1991). Both Impatiens x hawkeri (Dostal et al., 1991; Han, 2003) and I. walleriana can benefit from 1-MCP protection to improve postharvest quality (Han, 2003; Skog, 2001). Impatiens, therefore, can serve as an excellent model plant to investigate the use and efficacy of this novel 1-MCP formulation on floral crops.

The objective of this work was to investigate parameters associated with the use of a sprayable formulation of 1-MCP (as AFxRE-038) on Impatiens plants. Concentration, timing of application, spray volume, pH of spray solution, mixing and storage, overhead irrigation, and duration of protection against exogenous ethylene were investigated.

Materials and Methods

Plant production. Impatiens walleriana ‘Super Elfin Rose Improved’ and ‘Super Elfin White’ (Ball Horticultural Co., West Chicago, IL) and Impatiens hawkeri ‘Tamarind Purple’ (Fides Oro, Santa Paula, CA) were grown from seed or vegetative cuttings. Individual plants were transplanted into 10- or 14-cm containers filled with a commercial greenhouse substrate (LM-111; Lambert Moss Inc., Riviere-Ouelle, Quebec, Canada). Plants were irrigated as needed with 150 mg·L⁻¹ N from (Jack’s L®X™ 21–5–20; J.R. Peter’s Inc., Allentown, PA) and grown in a glass greenhouse at constant air temperature set point of 18 °C and ambient light. Plants were pinched to encourage branching. Before experimentation, plants were cleaned to remove old or damaged flowers so that only young flowers and unopened buds were present when experiments started.

1-MCP application and ethylene treatment. 1-MCP powder (AFxRD-038, 3.8% w/w a.i.; AgroFresh, Inc. Springhouse, PA) was gently mixed with reverse osmosis (RO) water until all powder was dissolved and the solution was clear (<1 min). Unless otherwise indicated, all solutions were prepared with 0.5 mL·L⁻¹ Capsil (Aquatrols, Paulsboro, NJ). The 1-MCP solution was transferred to a 1-gallon pump sprayer and pressurized within 5 min of mixing. The solution was sprayed onto plants until runoff
and allowed to dry for 1 h. Following this, plants were challenged with 1.0 μL·L⁻¹ ethylene flowing at 10 L·min⁻¹ into 0.4-m³ plexi-glass boxes fitted with two small 12-V circulation fans. Control plants were prepared in similar boxes with fresh air at the same flow rate. Ethylene streams were prepared with mass flow controllers and a microprocessor (Aalborg Inc., Orangeburg, NY). In some specific experiments (see below), plants were treated with ethylene by enclosure into a 4-m³ polyethylene tent and establishing static ethylene concentrations by injection of pure ethylene. In all cases, ethylene treatments were at 20 °C in darkness for 18 h. Ethylene concentrations were verified by gas chromatography and were maintained within 5% of desired set point.

Abscession data. Data were recorded on abscised flowers and buds immediately after the plants were removed from ethylene treatment. Data on percentage-abscised flowers were arcsin-transformed analysis of variance (ANOVA) tests were conducted. The Tukey–Kramer honestly significant difference method was used to conduct pairwise comparisons of all treatments, and ANOVA was performed to test for significance of linear and quadratic fit of 1-MCP treatments (JMP Pro v 10; SAS Institute, Cary, NC).

Expt. 1: Determination of suitable ethylene concentration for flower abscission. Impatiens walleriana ‘Super Elfin Rose Improved’ plants were treated with 0, 0.25, 0.75, and 1 μL·L⁻¹ ethylene (flowing) as described above.

Expt. 2: Concentration of sprayable 1-MCP to prevent abscission. Two experiments were conducted. In the first, I. walleriana ‘Super Elfin Rose Improved’ plants were sprayed with 1-MCP at concentrations of 0, 0.1, 0.5, 1, and 2 mg·L⁻¹. In the second, sprays of 0, 2.5, 5.0, 10, and 15 mg·L⁻¹ 1-MCP were used. After sprays dried, plants were placed into plexi-glass boxes for treatment with 1 μL·L⁻¹ flowing ethylene. Five single-plant replicates were used per treatment. One additional set of plants treated with 0 mg·L⁻¹ 1-MCP and was kept in the growth chamber at the same temperature and darkness to represent a nonethylene, non-1-MCP control.

Expt. 3: Concentration and volume of sprayable 1-MCP to prevent abscission. Impatiens walleriana ‘Super Elfin Rose Improved’ plants were sprayed with two concentrations of 1-MCP (5 and 10 mg·L⁻¹) and three volumes (102, 204, and 306 mL·m⁻²) in a factorial design and allowed to dry before overnight ethylene treatment. Five single-plant replicates were used per treatment. Two additional sets of control plants were not treated with 1-MCP. One set was placed into 1.0 μL·L⁻¹ flowing ethylene overnight and the second set was kept in an ethylene-free-grown chamber in darkness.

Expt. 4: Water pH and sprayable 1-MCP efficacy. Solutions of 10 mg·L⁻¹ 1-MCP were mixed using municipal tap water, RO, or Milli-Q (Milli-Q Advantage A10 System; Millipore Corporation, Billerica, MA) water.

The pH of the source water was 7.87, 6.45, or 5.75, respectively. Six additional 1-MCP treatments were made with RO water adjusted to pH 4, 5, 6, 7, 8, and 9. The pH was adjusted using 1.0 N NaOH or 0.1 N HCl. Impatiens walleriana ‘Super Elfin Rose Improved’ plants were sprayed with 1-MCP solutions and allowed to dry before overnight flowing ethylene treatment. Three single-plant replicates were used per treatment.

The experiment was repeated using I. hawkeri ‘Tamarinda Purple’ with 1-MCP treatments made with RO water adjusted to pH 7, 8, 9, and 10. Plants were sprayed with 1-MCP solutions and allowed to dry. Plants were then placed into tents for static ethylene treatment (2.5 μL·L⁻¹) overnight. Four single-plant replicates were used per treatment.

Expt. 5: Length of 1-MCP effectiveness for prevention of abscission. Individual groups of I. walleriana ‘Super Elfin White’ plants were sprayed with freshly made 1-MCP at 10 mg·L⁻¹ daily, for 6 d, and placed under optimum greenhouse conditions, ultimately leading to plants that had been sprayed with 1-MCP 0–6 d earlier. On day 6, all plants were treated with 1.0 μL·L⁻¹ flowing ethylene overnight.

Expt. 6: Length of time between 1-MCP treatment and overhead irrigation to prevent abscission. Impatiens walleriana ‘Super Elfin Rose Improved’ plants were sprayed with 10 mg·L⁻¹ 1-MCP solution. Plants were allowed to stand for 1, 2, 3, 4, 5, or 10 min before each pot was irrigated with ≥100 mL tap water directly overhead through a water breaker (Model 400AL; Dramm Corporation, Manitowoc, WI). Three single-plant replicates were used per treatment. One set of plants was sprayed with 1-MCP and not irrigated. One set of control plants were not treated with 1-MCP or irrigated. After 1-MCP and irrigation treatment, plants were allowed to dry completely and were treated with 1.0 μL·L⁻¹ flowing ethylene overnight.

Expt. 7: Duration of overhead irrigation after 1-MCP treatment to prevent abscission. Impatiens walleriana ‘Super Elfin Rose Improved’ plants were sprayed with 10 mg·L⁻¹ 1-MCP solution. Plants were allowed to dry for 1 or 2 min before they were irrigated with tap water for 1 or 5 s (0.45 or 2.25 L, respectively) directly overhead using a hose fitted with a water breaker as above. Three single-plant replicates were used per treatment. One set of plants was allowed to dry immediately after 1-MCP treatment without an irrigation treatment. After 1-MCP and irrigation treatment plants were allowed to dry completely and were treated with 1.0 μL·L⁻¹ flowing ethylene overnight.

Expt. 8: Efficacy of aged 1-MCP solutions. Solutions of 10 mg·L⁻¹ 1-MCP were prepared over a period of 7 weeks. Solutions were kept in airtight glass bottles in darkness. Freshly made and 1-, 2-, 4-, 5-, and 7-week-old solutions of 1-MCP were sprayed on I. hawkeri ‘Tamarinda Purple’ and allowed to dry. Plants were placed in plastic tents for a static 2.5 μL·L⁻¹ ethylene treatment overnight. Six single-plant replicates were used per treatment. Two additional sets of plants were treated with 0 mg·L⁻¹ 1-MCP, one set was placed into ethylene overnight and the other kept in an ethylene-free area at the same temperature and darkness.

Expt. 9: 1-MCP to prevent abscission in prolonged darkness. Impatiens hawkeri ‘Tamarinda Purple’ plants were sprayed with 0 or 10 mg·L⁻¹ 1-MCP and allowed to dry. Plants were then placed into plexi-glass boxes with 0 or 1.0 μL·L⁻¹ flowing ethylene at 28 °C for 40 h. Five single-plant replicates were used per treatment.

Expt. 10: 1-MCP to prevent abscission at high temperatures. Impatiens hawkeri ‘Tamarinda Purple’ plants were sprayed with 0 or 10 mg·L⁻¹ 1-MCP and allowed to dry. Plants were then placed into plexi-glass boxes with 0 or 1.0 μL·L⁻¹ flowing ethylene at 32 °C for 48 h. Five single-plant replicates were used per treatment.

Results and Discussion

Expt. 1: Determination of suitable ethylene concentration. Plants showed 50%, 98%, and 100% flower abscission after overnight exposure to 0.25, 0.75, and 1 μL·L⁻¹ ethylene respectively. This is similar to Dostal et al. (1991) who found that a 6-h treatment with 1 μL·L⁻¹ ethylene caused 100% flower abscission in Impatiens xhawkeri. Therefore, our further experiments were conducted using 1 μL·L⁻¹ exogenous ethylene overnight in darkness. Also, preliminary experiments showed no difference in flower abscission whether plants were enclosed by plastic plant sleeves or loose polyethylene bags (data not shown), thus

| 1-MCP pretreatment (mg·L⁻¹) | Ethylene treatment (μL·L⁻¹) | Flower abscission (%) |
|-----------------------------|-----------------------------|----------------------|
| 0                           | 1                           | 96 ± a               |
| 0.1                         | 1                           | 73 ± b               |
| 0.5                         | 1                           | 47 ± c               |
| 1                           | 1                           | 43 ± b               |
| 2                           | 1                           | 25 ± c               |

| Expt. 2 | 0 (control) | 0 | 4 ± a |
|---------|-------------|---|------|
| 0.1     | 1           | 93 ± b |
| 0.5     | 1           | 10 ± a |
| 1       | 1           | 14 ± a |
| 5       | 1           | 5 ± a  |

Data are means of five replicates per treatment. For each experiment, letters after values in each column represent means separation using Tukey–Kramer honestly significant difference at \( P < 0.05 \). Means followed by the same letter are not significantly different.
Table 2. Flower abscission (%) of Impatiens walleriana ‘Super Elfin Rose Improved’ sprayed with varying volumes and concentrations of 1-methylcyclopropene (1-MCP; AFxRD-038). After 1-MCP treatment, plants were exposed 1 µL·L⁻¹ ethylene or kept in an ethylene-free atmosphere for 18 h in darkness.

| 1-MCP concn (mg L⁻¹) | Ethylene treatment (µL·L⁻¹) | Spray volume (mL·m⁻²) | Flower abscission (%) |
|-----------------------|-----------------------------|-----------------------|----------------------|
| 0                     | 0 (control)                 | 306                   | 6⁺                   |
| 0                     | 1                           | 306                   | 95                   |
| 5                     | 1                           | 102                   | 34                   |
| 5                     | 1                           | 204                   | 10                   |
| 5                     | 1                           | 306                   | 9                    |
| 10                    | 1                           | 102                   | 17                   |
| 10                    | 1                           | 204                   | 11                   |
| 10                    | 1                           | 306                   | 7                    |

1-MCP concn (M) Spray volume (V) ** M x V

Data are means of three replicates per treatment.

Table 3. Flower abscission (%) of Impatiens walleriana ‘Super Elfin Rose Improved’ sprayed with 10 mg·L⁻¹ 1-methylcyclopropene (1-MCP; AFxRD-038) and exposed to 1 µL·L⁻¹ ethylene (for 18 h in darkness) after 0 to 6 d.

| Days between 1-MCP spray and ethylene exposure | Flower abscission (%) |
|-----------------------------------------------|-----------------------|
| 0                                             | 4⁺                    |
| 1                                             | 8                     |
| 2                                             | 12                    |
| 3                                             | 15                    |
| 4                                             | 28                    |
| 5                                             | 54                    |
| 6                                             | 65                    |

Significance: L *** Q **

Data are means of four replicates per treatment. Significance of linear (L) or quadratic (Q) regression: **, *** Significant at P ≤ 0.01 or 0.001, respectively.

Table 4. Flower abscission (%) of Impatiens walleriana ‘Super Elfin Rose Improved’ sprayed with 1-methylcyclopropene (1-MCP; AFxRD-038). After pretreatment, plants were held for 0 to 30 min, then irrigated (overhead) with 100 mL tap water. After irrigation plants were allowed to dry for 2 h, then exposed to 1 µL·L⁻¹ ethylene for 18 h in darkness.

| 1-MCP treatment (mg·L⁻¹) | Time elapsed before irrigation (min) | Flower abscission (%) |
|--------------------------|--------------------------------------|----------------------|
| 0 (control)              | —                                    | 97                   |
| 10                       | —                                    | 3                    |
| 10                       | 1                                    | 1                    |
| 10                       | 2                                    | 2                    |
| 10                       | 3                                    | 2                    |
| 10                       | 4                                    | 2                    |
| 10                       | 5                                    | 0                    |
| 10                       | 10                                   | 3                    |
| 10                       | 30                                   | 2                    |

Significance: NS

Plants were not irrigated after 1-MCP treatment.

Table 5. Flower abscission (%) of Impatiens walleriana ‘Super Elfin Rose Improved’ sprayed 10 mg·L⁻¹ 1-methylcyclopropene (1-MCP; AFxRD-038) and exposed to 1 µL·L⁻¹ ethylene for 18 h in darkness.

| Time elapsed before irrigation (min) | Volume of irrigation (L) | Flower abscission (%) |
|--------------------------------------|--------------------------|----------------------|
| 1                                    | 0.45                     | 5                    |
| 2                                    | 2.25                     | 7                    |

Significance: Time (T) NS Volume (V) NS T x V NS

Plants were not irrigated after 1-MCP treatment.

Expt. 2: Concentration of sprayable 1-MCP to prevent abscission. Sprays of 1-MCP at concentrations from 0.1 to 2.0 mg·L⁻¹ significantly reduced ethylene-induced flower abscission (Table 1). There was a significant difference between plants treated with 2.0 and 0.1 mg·L⁻¹ 1-MCP. The linear and quadratic regressions of 1-MCP concentration and flower abscission were highly significant at P ≤ 0.001 and 0.01, respectively. However, the 25% abscission seen with 2.0 mg·L⁻¹ 1-MCP would still render plants unmarketable. In a second experiment, there was no difference in flower abscission when 1-MCP was sprayed at rates of 2.5–15 mg·L⁻¹ (Table 3). These results suggest that 2.5 mg·L⁻¹ 1-MCP spray concentration is the lowest concentration for protection against an overnight exposure to 1.0 µL·L⁻¹ ethylene.

Expt. 3: Concentration and volume of sprayable 1-MCP to prevent abscission. In this experiment, only 1-MCP spray volume was significant (P = 0.004), 1-MCP concentration and the concentration x volume interaction were not significant (P = 0.1983 and 0.0829, respectively) (Table 2). Within 1-MCP concentrations, the percentage of flowers that abscised decreased as the spray volume increased. MacKinnon et al. (2009) saw similar results in tomato with higher spray volume leading to higher efficacy of 1-MCP in preventing epinasty after ethylene exposure. This is likely due to better coverage of plants from the greater spray volumes. It was suggested that 1-MCP in the aqueous form may have limited translocation in tomatoes (MacKinnon et al., 2009) and as seen with Impatiens, it appears important to fully cover all foliage and flowers with spray solution to achieve 1-MCP binding to open ethylene receptor sites.

Studies of aqueous 1-MCP on plums (Manganaris et al., 2007), tomatoes, and avocados (Choi et al., 2008) suggest that complete coverage of target plant material is necessary to achieve full protection from ethylene-mediated senescence. When tomatoes and avocados were partially immersed in an aqueous 1-MCP solution, there was uneven protection of the fruit showing limited diffusive capacity of aqueous 1-MCP on the surface of treated plants (Choi et al., 2008).

Volumes of 204 mL·m⁻² (equal to 2 qts/100 ft²) are generally recommended for plant growth regulator applications (Latimer and Whicker, 2012) and this experiment confirmed that current greenhouse industry spray protocols should provide sufficient coverage of 1-MCP when used at rates of 5 and 10 mg·L⁻¹.

Expt. 4: Water quality and sprayable 1-MCP efficacy. There was no difference in 1-MCP efficacity when prepared with municipal, RO, or Milli-Q water; abscission averaged 8% across all treatments (data not shown). As pH of the spray solution increased from 4 to 9 (in RO water), abscission increased from 4% to 14%. In the second experiment, there was no difference in abscission as pH increased from 7 to 10 (average of 11%). Although pH 4 and 9 were statistically different in the first experiment, we concluded that within the pH range studied, there is no commercially significant effect of water source pH on activity of this formulation.

Expt. 5: Length of 1-MCP effectiveness for prevention of abscission. Impatiens walleriana ‘Super Elfin White’ plants challenged with ethylene within 0 to 3 d of 1-MCP treatment exhibited significantly less flower abscission than plants challenged with ethylene four or more days after 1-MCP treatment (Table 3). Plants with four or more days delay between 1-MCP treatment and ethylene challenge showed more than 25% flower abscission and plants were unmarketable. Linear and quadratic regressions were significant at P < 0.001 and 0.01, respectively; flower abscission was affected by length of delay between 1-MCP treatment and ethylene challenge. Pelargonium peltatum pretreated with 1.0 µL·L⁻¹ gaseous 1-MCP for 2 h completely inhibited ethylene-induced petal abscission; however, the half-life of 1-MCP activity was not <2, 3, and 6 d after 1-MCP treatment at 25, 20.7, and 12 °C, respectively, with no residual effect of 1-MCP protection after 4 or 5 d at 25 or 20.7 °C, respectively (Cameron and Reid, 2001). These results suggest that it is necessary to apply the sprayable formulation of 1-MCP a maximum of 3 d before...
plants will be exposed to exogenous ethylene. The likely reason for this transient effect is due to creation of new ethylene receptors in the actively growing plant, which are not protected by the previous 1-MCP treatment (Blankenship and Dole, 2003; Cameron and Reid, 2001).

Expt. 6: Length of time between 1-MCP treatment and overhead irrigation. There was no significant difference in flower abscission between plants treated with 1-MCP only and those receiving irrigation 1 to 30 min after 1-MCP treatment (Table 4). The linear and quadratic regressions were not significant. All 1-MCP and irrigation treatments protected plants from external ethylene at a market acceptable level showing that complete 1-MCP protection is conferred within one minute of 1-MCP application. MacKinnon et al. (2009) challenged tomatoes with irrigation 15 to 240 min after 1-MCP pretreatment and found that there was no significant difference from untreated controls suggesting that 1-MCP is rainfast on tomatoes as early as 15 min after application. This would allow practitioners the opportunity to apply sprayable 1-MCP without concern that future rain or irrigation will negatively affect the application.

Expt. 7: Duration of overhead irrigation after 1-MCP treatment to prevent abscission. There was no significant difference between I. walleriana ‘Super Elfin Rose Improved’ plants treated with 1-MCP solutions only and those receiving an overhead irrigation treatment 1 or 2 min after the 1-MCP spray. The time between 1-MCP application and irrigation, irrigation volume, and the interaction of time × volume were all nonsignificant (Table 5). The U.S. Label for AFxRD-038 suggests that 1-MCP sprays should be applied under drying conditions, in early morning or at night to allow for adequate absorption, irrigation should not be used for at least 2 h following application and spray should be used 1 h before expected rain (EPA, 2010). These results, with the results of Expt. 5, show that irrigation or rainfall 1 min after 1-MCP treatment, regardless of irrigation volume, will not reduce 1-MCP efficacy in Impatiens plants.

Expt. 8: Efficacy of aged 1-MCP solutions. Solutions of 1-MCP lost effectiveness as the age of the solution increased beyond 2 weeks and effectiveness was especially lost as solutions aged from 2 to 4 weeks (Table 6). This finding is at odds with recommendations that AFxRD-038 be used very soon after mixing (EPA, 2010) in that efficacy was maintained for at least 14 d after mixing. One explanation for this is that the concentration of the prepared solution, 10 mg-L⁻¹, was 4-fold greater than the minimum effective concentration of 2.5 mg-L⁻¹. We hypothesize that the solution was releasing gaseous 1-MCP more or less continually after mixing, and between weeks 2 and 4 enough had been lost to cause a failure in protection from ethylene. If this is true, mixing protocols could be extremely important in experiments and applications where the concentrations involved are very similar to those needed for protection from ethylene.

Expt. 9: 1-MCP to prevent abscission in prolonged darkness. In the absence of ethylene, dark storage at 22 °C increased flower abscission in Impatiens ×hawkeri ‘Tamarinda Purple’ and pretreatment with 10 mg-L⁻¹ 1-MCP inhibited abscission (Table 7). When plants were kept in darkness for less than 3 d, there was no difference in flower abscission with or without 1-MCP treatment. The effects of 1-MCP treatment, days in darkness and the 1-MCP × days in darkness interaction were all highly significant (P < 0.0001) (Table 7).

Shipping conditions are generally dark (not allowing for photosynthesis) and warm (increasing respiration rates). During shipping, carbohydrate levels decrease and plants may show symptoms of senescence such as lower leaf chlorosis, internode elongation, and abscission of buds and flowers (Jones, 2002). Holding Begonia ×hiemalis plants in the dark caused more bud abscission than vibration or nonfreezing low temperature treatments and pretreatment with 1-MCP reduced abscission and increased display life (Kim et al., 2007). Skog et al. (2001) also showed that Impatiens ×hawkeri ‘Paradise Aglia’ and I. walleriana ‘Accent White’ and ‘Seashell’ were improved with gaseous 1-MCP pretreatment in the presence or absence of ethylene during postproduction. Impatiens ×hawkeri, I. walleriana, and double Impatiens plants placed in cardboard boxes for 3 or 5 d showed significantly increased bud abscission, which was reduced by gaseous 1-MCP pretreatment (Han, 2003). From this, we suggest that Impatiens plants that will be subjected to postproduction stresses for more than 2 d may also be pretreated with sprayable 1-MCP.

Expt. 10: 1-MCP to prevent abscission at high temperatures. Impatiens hawkeri ‘Tamarinda Purple’ plants kept in darkness at 28 °C for 40 h exhibited significantly less flower abscission if treated with 1-MCP than untreated plants, regardless of the presence of exogenous ethylene (Table 8). Effects of 1-MCP, ethylene, and 1-MCP × ethylene were highly significant (P < 0.0001). Flower abscission damage to plants without a 1-MCP pretreatment was seen within 48 h in the absence of ethylene in warm temperatures (28 °C) (Table 8), whereas damage was seen in 3 d when plants were kept at cooler temperatures (22 °C) (Table 7).

Table 6. Flower abscission (%) of Impatiens hawkeri ‘Tamarinda Purple’ pre-treated with 10 mg-L⁻¹ 1-methylocyclopene (1-MCP; AFxRD-038) prepared over a 7-week period. After 1-MCP pretreatment, plants were exposed 2.5 μL-L⁻¹ ethylene or kept in an ethylene-free atmosphere for 18 h in darkness.

| 1-MCP treatment (mg-L⁻¹) | Ethylene treatment (μL-L⁻¹) | Age of 1-MCP solution (wk) | Flower abscission (%) |
|-------------------------|---------------------------|---------------------------|----------------------|
| 0 (control)             | 0 (control)               | N/A                       | 12 c                 |
| 0                       | 1                         | N/A                       | 100 a                |
| 1                       | 0                         | 0                         | 13 c                 |
| 1                       | 0                         | 1                         | 12 c                 |
| 1                       | 1                         | 2                         | 13 c                 |
| 1                       | 4                         | 5                         | 97 a                 |
| 1                       | 7                         | 7                         | 100 a                |
*Data are means of five replicates per treatment. Means followed by the same letter are not significantly different using Tukey–Kramer honestly significant difference at P = 0.05.

Table 7. Flower abscission (%) of Impatiens hawkeri ‘Tamarinda Purple’ sprayed with 1-methylocyclopene (1-MCP; AFxRD-038). After treatment, plants were kept in darkness (ethylene-free atmosphere) for 1, 2, 3, or 4 d at 22 °C.

| 1-MCP pretreatment (mg-L⁻¹) | Length of darkness (d) | Flower abscission (%) |
|-----------------------------|------------------------|----------------------|
| 0                           | 1                      | 0 b                  |
| 0                           | 0                      | N/A                  |
| 0                           | 3                      | 19 a                 |
| 0                           | 4                      | 34 a                 |
| 1                           | 0                      | 0 a                  |
| 2                           | 2                      | 1 a                  |
| 3                           | 3                      | 0 a                  |
| 4                           | 4                      | 3 a                  |
| 1-MCP (M)                   |                         | ***                  |
| Darkness (D)                |                         | ***                  |
| M × D                       |                         | ***                  |

*Data are means of five replicates per treatment. Means followed by the same letter are not significantly different using Tukey–Kramer honestly significant difference at P = 0.05.

Table 8. Flower abscission (%) of Impatiens hawkeri ‘Tamarinda Purple’ pretreated with 1-methylocyclopene (1-MCP; AFxRD-038). After treatment, plants were exposed 1 μL-L⁻¹ ethylene or kept in an ethylene-free atmosphere in darkness for 40 h at 28 °C.

| 1-MCP pretreatment (mg-L⁻¹) | Ethylene treatment (μL-L⁻¹) | Flower abscission (%) |
|-----------------------------|---------------------------|----------------------|
| 0                           | 0                        | 6 b                  |
| 1                           | 0                        | 100 a                |
| 1                           | 0                        | 0 d                  |
| 1                           | 1                        | 2 c                  |
| 1-MCP (M)                   |                         | ***                  |
| Ethylene (E)                |                         | ***                  |
| M × E                       |                         | ***                  |

*Data are means of five replicates per treatment. Means followed by the same letter are not significantly different using Tukey–Kramer honestly significant difference at P = 0.05.

*Significant at P ≤ 0.001.
developed as lower leaf chlorosis (81% of cultivars), internode elongation (38% of cultivars), flower senescence (38% of cultivars), and bud abortion (33% of cultivars). Potted *Rosa hybrida* plants subjected to dark storage conditions for various lengths of time and a range of temperatures showed increased flower development (leading to senescence) with storage duration more than 2 d and the interaction of storage duration, temperature, and cultivar were significant with less development at 4 °C than 16 °C and greatest development at 28 °C (Cushman et al., 1998).

**Conclusions**

The present work provides a basis for further experimentation with sprayable 1-MCP on ornamental floriculture crops. Sprays of 1-MCP (AFxRD-038 formulation) at 10 mg·L⁻¹ (204 mL·m⁻²) should be adequate to protect highly sensitive flowers from exposure to 1.0 μL·L⁻¹ exogenous ethylene for 18 h in darkness while lower concentrations will likely provide protection if applied at high volume (above 204 mL·m⁻²). Spray applications of 1-MCP were rainfast and fully efficacious within 1 min of application. The AFxRD-038 formulation was effective across a relatively wide pH range. As general practice, mixed solutions should be used soon after mixing but if held in an airtight container, solutions may show activity up to 2 weeks. Plants should be treated with 1-MCP within 3 d of potential postproduction ethylene exposure for full protection against stressors such as heat, darkness, and ethylene. AFxRD-038 shows great promise for future use in the floriculture industry to provide protection against postproduction stress and mediate ethylene damage in flowering plants.

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