Residual Effects of Ethylene on Tulip Growth and Flowering

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Additional index words. Tulipa gesneriana, 1-methylcyclopropene, 1-MCP, EthylBloc™ sachet, hydroponics

Abstract. Ethylene effects were investigated on two tulip (Tulipa gesneriana L.) cultivars, Markant and Carreria. Pre-cooled bulbs were treated with ethylene (flow-through) for 1 week at 0, 0.1, 1.0, or 10 µL L⁻¹ (± 10%) in a modified hydroponic system. After ethylene exposure, plants were either destructively harvested for root measurements or forced in a greenhouse for flower measurements. Ethylene exposure at concentrations as low as 1 µL L⁻¹ during the first week of growth reduced shoot and root elongation and subsequently increased flower bud abortion. At 10 µL L⁻¹, root growth was essentially eliminated. In a second experiment, bulbs were treated overnight with 1-methylcyclopropene (1-MCP) before a 7-day exposure to 1 µL L⁻¹ ethylene. 1-MCP pretreatment eliminated the harmful effects of ethylene on root and shoot growth. This study illustrates the effects of ethylene exposure during hydroponic tulip production and demonstrates a potential benefit to treating bulbs with 1-MCP before planting.

Tulip (Tulipa gesneriana L.) is an ornamental geophyte prized for centuries as a cut flower, potted plant, and garden favorite. Tulip bulbs are grown in fields and sold to specialized producers who “force” the plants into flower after a cold treatment of 12 or more weeks. In cut flower tulip production, bulbs are grown in fields and sold to specialized producers who “force” the plants into flower after a cold treatment of 12 or more weeks. In cut flower tulip production, bulbs are grown in fields and sold to specialized producers who “force” the plants into flower after a cold treatment of 12 or more weeks. In cut flower tulip production, bulbs are grown in fields and sold to specialized producers who “force” the plants into flower after a cold treatment of 12 or more weeks. In cut flower tulip production, bulbs are grown in fields and sold to specialized producers who “force” the plants into flower after a cold treatment of 12 or more weeks.

During tulip bulb development and storage before cooling, ethylene can cause a number of physiological and morphological disorders, including gummosis (excretion of polysaccharides), flower bud abortion, shortened leaves or flowers, reduced or eliminated roots, deformed anthers, abnormal growth habit, loss of fresh weight during storage, and excessive growth of daughter bulbs (splitting) (Kamerbeek and de Munk, 1976). The degree of ethylene damage depends on a number of factors, including concentration, duration (De Munk, 1972), temperature during exposure (De Munk, 1973), and tulip cultivar (Miller et al., 2004; De Wild et al., 2002). A significant source of ethylene in the tulip industry is the fungal pathogen fusarium (Fusarium oxysporum Schlecht f. sp. tulipae), which produces ethylene when colonizing tulip bulbs (Kamerbeek and de Munk, 1976). These fungal infections therefore create a unique production challenge for cut flower tulip growers given the potential for ethylene exposure to actively growing roots and shoots.

1-Methylcyclopropene (1-MCP) is widely used in postharvest horticulture, gaining popularity for ornamental plants as an ethylene perception inhibitor that blocks ethylene binding sites in plant tissue (Blankenship and Dole, 2003; Watkins, 2006), thus protecting the tissue against ethylene damage for variable periods (Sisler and Serek, 1997). Gude and Dijkema (2005) demonstrated protection from ethylene damage in tulip storage by 24 h 1-MCP applications (0.2 µL L⁻¹) at 12-d intervals. The potential for 1-MCP application immediately before planting has not been investigated and is the subject of this research.

Materials and Methods

Pre-cooled bulbs (16 weeks at 5 °C) of Tulipa gesneriana L. ‘Markant’ were treated with flow-through ethylene at nominal concentrations of 0, 0.1, 1.0, or 10 µL L⁻¹ (± 10%) for 1 week at 20 °C in a modified hydroponic system. Bulbs were impaled on a planting grid and placed in 4.5-L plastic tubs. One sachet per treatment container was placed in a small beaker containing 25 mL distilled water and then immediately enclosed with the bulbs for 24 h at 20 °C. Control bulbs were similarly sealed in a separate container without 1-MCP. The containers were then opened, beakers were removed, 1.5 L tap water was added, and then resealed for ethylene exposure at 0 or 1 µL L⁻¹ (± 10%) in the same hydroponic system as described previously. This experiment also included three replicates of six bulbs per treatment. After 1 week, plants were destructively harvested for the same root and shoot measurements described previously.

All data were subjected to analysis of variance using standard least squares in JMP (SAS Institute, Cary, NC) with regression analysis or a means separation procedure (Tukey’s honestly significant difference) where appropriate. Using a natural log (ln) transformed ethylene concentration provided the best fit for the regression analysis; thus, all values were transformed on the basis of ln (ethylene concentration + 0.1) (because ln 0 is undefined). Analysis of variance and means separation for experiments using 1-MCP were calculated using non-transformed data.

Results and Discussion

Ethylene concentrations were verified by daily measurements using a Buck 310 gas chromatograph (Buck Scientific, East Norwalk, CT) fitted with an alumina column and a flame ionization detector. After 1 week of ethylene exposure, plants were destructively harvested for the following measurements: plant height (basal plate to the tip of the tallest leaf), root length (mean of three longest roots per bulb), and root fresh weight (including 1-cm basal plate). When the experiment was repeated with the same setup as previously described, instead of destructive harvest, bulbs were forced into flower in a presumed ethylene-free greenhouse, at which time stem length (basal plate to bud tip) and percent aborted flowers (plants showing no flower) were calculated. Days to flower were not calculated, but it was ≈10.

To test the efficacy of 1-MCP pretreatment against ethylene damage, a second experiment was designed in which ‘Carreria’ bulbs were treated with 1-MCP (EthylBloc™ sachets; Floralife, Inc., Walterboro, SC) before ethylene exposure. ‘Carreria’ bulbs were impaled on a planting grid and placed in 4.5-L plastic tubs. One sachet per treatment container was placed in a small beaker containing 25 mL distilled water and then immediately enclosed with the bulbs for 24 h at 20 °C. Control bulbs were similarly sealed in a separate container without 1-MCP. The containers were then opened, beakers were removed, 1.5 L tap water was added, and then resealed for ethylene exposure at 0 or 1 µL L⁻¹ (± 10%) in the same hydroponic system as described previously. This experiment also included three replicates of six bulbs per treatment. After 1 week, plants were destructively harvested for the same root and shoot measurements described previously.

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previous ethylene exposure as did flower bud abortion, respectively, decreasing or increasing linearly as ethylene concentration increased (Figs. 1 and 2).

Our results are consistent with those published by De Munk and de Rooy (1971), in which they measured ethylene effects on tulip growth by planting fusarium-infected bulbs alongside healthy tulips in soil culture. Although our experiments were conducted in a modified hydroponic system, the ethylene-related growth problems could occur in other substrata as well, including sand or soilless mixes (King and Smith, 1987). It is presumed that ethylene concentrations emanating from fusarium-infected bulbs increase as the gas becomes “trapped” as a result of low air movement around soil particles, thick canopy density, or other factors limiting diffusion surrounding individual plants. These influences could therefore contribute to high ethylene concentrations in isolated locations throughout a commercial greenhouse. De Munk (1971) measured ethylene levels as high as 22 \( \mu \text{L}^{-1} \) from enclosed fusarium-infected tulip bulbs; therefore, severe damage caused by our 10\( \mu \text{L}^{-1} \) treatment is entirely plausible in a commercial production system where fusarium-infected tulip bulbs are present.

Deleterious effects of 1 \( \mu \text{L}^{-1} \) ethylene were essentially eliminated with 1-MCP pretreatment (Table 1; Fig. 3). The only exception was with root fresh weight, in which the interaction between 1-MCP and ethylene was nonsignificant. However, the main effects for these data support a similar trend, heavier root mass with 1-MCP treatment than without (3.2 versus 2.7 g, respectively) and lighter root mass with ethylene exposure regardless of 1-MCP treatment (data not shown). In tulip, 1-MCP treatment was previously shown to provide protection from various forms of ethylene damage for up to 12 d after application (Gude and Dijkema, 2005). This is consistent with our data, which indicate 1-MCP was effective at reducing ethylene injury for at least 1 week in the earliest phases of fen.

Table 1. Effect of 1-MCP treatment before 1.0 \( \mu \text{L}^{-1} \) ethylene exposure on plant height (basal plate to longest leaf) and root length (mean of three longest roots per bulb) of *Tulipa gesneriana* ‘Carreria’.

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Plant ht (cm)</th>
<th>Root length (cm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>+MCP + Eth</td>
<td>18.5 a</td>
<td>3.6 a</td>
</tr>
<tr>
<td>+MCP − Eth</td>
<td>16.1 b</td>
<td>3.8 a</td>
</tr>
<tr>
<td>−MCP − Eth</td>
<td>15.9 b</td>
<td>3.4 a</td>
</tr>
<tr>
<td>−MCP + Eth</td>
<td>9.1 c</td>
<td>1.7 b</td>
</tr>
</tbody>
</table>

\( \mu \text{L}^{-1} \)

Source

\( M \) *** ***

\( E \) *** ***

\( M \times E \) *** ***

- Bulbs were exposed to ethylene for 1 week at 20 °C while in the growing environment.
- Means in columns not followed by the same letter are different at \( \alpha = 0.05 \).
- ***Significant at \( P \leq 0.0001 \).
- 1-MCP = 1-methylcyclopropene.
of growth and establishment in the greenhouse.

**Literature Cited**


