EFFECT OF PULSING WITH VARIOUS PRESERVATIVES ON POSTHARVEST PERFORMANCE OF CUT Polianthes tuberosa L. ‘SINGLE’ SPIKES

Muhammad Asif¹*, Iftikhar Ahmad¹, M. Qasim¹ and Rashid Ahmad²

¹Institute of Horticultural Sciences, University of Agriculture, Faisalabad, Pakistan.  
²Department of Agronomy, University of Agriculture, Faisalabad, Pakistan.  
*Corresponding author’s e-mail: roymasif@yahoo.co.in

In Punjab-Pakistan, growing tuberose is a profitable business as returns are double than cost despite of certain bottlenecks such as fluctuations in the prices and unawareness of farmers regarding its post-harvest handling. Keeping in view high market demand in local flower markets, the present study was conducted to extend the postharvest life of cut tuberose spikes by pulsing cut spikes with various preservatives. Among various pulsing treatments, 10% sucrose plus 50 mg L⁻¹ salicylic acid (SA) for 24 hours proved best with the longest vase life (10.2 d), more number of days required to open 50% florets (5.0 days), maximum dry weight percentage (13.2%) and superior quality spikes (7.6). Moreover, longest florets (7.0 cm) with greater floret diameter (4.2 cm), highest percentage of opened florets (90.0%) and minimum ion leakage of florets (140.0%) was recorded for spikes pulsed with 10% sucrose plus 50 mg L⁻¹ salicylic acid for 24 hours. On the other hand, highest water uptake (106.0 mL) and the maximum relative fresh weight percentage (109.9%) were observed in cut tuberose spikes treated with 50 mg L⁻¹ salicylic acid without sucrose solution. In summary, pulsing of cut tuberose spikes with 50 mg L⁻¹ SA with or without 10% sucrose proved effective for extending vase life and improving postharvest performance of cut tuberose spikes.

Keywords: Ascorbic acid, floriculture, relative fresh weight, salicylic acid, sucrose, vase life

INTRODUCTION

Tuberose (Polianthes tuberosa L.) is native to Mexico and a member of family Amaryllidaceae. Tuberose spikes have 10-20 pairs of florets, which open from base to upwards and make excellent cut flowers and are popular for their fragrance (De Hertogh and Le Nord, 1993). Moreover, its flowers are also used to extract essential oil, which is used for the production of cosmetic and perfumery products (Hussain, 1986). It is very popular cut flower in Punjab-Pakistan and in fact is the only major cut flower available in the florist shops during summer and very much appreciated by the consumers for its fragrance (Ahsan et al., 2012; Ashfaq et al., 2015).

In Punjab-Pakistan, growing tuberose is a profitable business as returns are double than cost (Usman and Ashfaq, 2013) despite certain bottlenecks such as fluctuations in the prices and unawareness of farmers regarding its post-harvest handling. Previous studies have demonstrated that cut flower growers in Pakistan are generally unaware of modern postharvest handling of cut flowers (Ahmad, 2009). Their packaging and transportation practices are not up to the standard, which result in poor quality produce in the market with shorter vase life. All this results in to heavy postharvest losses, which may range between 10-30% (Hayat et al., 2012) and sometimes up to 40 percent in the absence of floral preservatives (Hutchinson et al., 2003). Tuberose is heat loving plant and can successfully be grown with little care, which makes it the only choice for the cut flower growers during summer season in Punjab, Pakistan. Therefore, more often local florist markets become glutted with it during peak production time, which results in price drop and ultimately cause economic losses to the growers.

Pulsing with sucrose applied before storage or shipping increased the postharvest longevity of many cut flowers, probably by replacing the carbohydrates during cold storage (van Doorn et al., 1991; Khan et al., 2015), or preventing leaf desiccation (Silvanda et al., 2011). Sucrose (1-5%) have significant effect on quality parameters, viz. vase life, flower quality, water uptake, leaf chlorophyll contents, fresh and dry weight of cut rose flowers and can maintain the vase life of flowers for a longer period (Younis et al., 2006; Mirjalili, 2015). Similarly, pulsing of cut tuberose flower with 10% sucrose before transfer to deionized water improved vase life by 4 days and improved floret opening by 21% relative to control probably by improving water balance (Hutchinson et al., 2003). However, sucrose alone cannot be used without germicides, as sucrose promotes bacterial proliferation leading to shortening of vase life (Ichimura et al., 2002). Pulse treatment of freesia flowers, with 20% sucrose for 24 h prior to three days of simulated shipping, improved subsequent vase life (Woodson, 1987). Abdulrahman et al. (2012) reported that 0.5 g L⁻¹ of sucrose significantly increased time to stem bending, fresh and dry weight, percentage change in fresh weight and total carbohydrates,
and vase life of snapdragon cut flower. Kim and Lee (2002) suggested pulsing snapdragon flowers in sucrose vases resulted in higher longevity period, which indicated that sucrose played a critical role in promoting water absorption and metabolic processes within the flower. Salicylic acid (SA) is a natural, cheap, safe, and biodegradable compound which is suitable alternative for conventional chemical treatments in order to prolong vase life of cut rose flowers (Abdolmaleki, 2015). It is a signaling molecule which regulates plant growth and development together with local and endemic disease resistance in plants in response to various pathogenic attacks and environmental stresses. Moreover, SA treatments extended vase life in association with inhibition of ethylene production (Leslie and Romani, 1988). In free state, SA has a pH of 2.4 and acidic solution inhibits bacteria growth and proliferation (Enyedi et al., 1992). However, SA can modulate plant responses to a wide range of oxidative stresses and prevent cell wall degradation and thereby reduction in water loss (Shirasu et al., 1997). Moreover, addition of 300 ppm salicylic acid to distilled water extended the vase life of Alstroemeria peruviana, Gerbera jamesonii, Lilium asiaticum, Polianthes tuberosa and Rosa hybrida by 30-55% relative to control. Besides, SA treatments increased relative water content, petal water content and initial fresh weight in cut flowers, over control. The beneficial effects of salicylic acid are associated with the plant regulating and anti-stress properties of salicylic acid. Researchers strongly suggest SA as a vase solution additive being natural, cheap, safe and biodegradable compound for extending the postharvest longevity of cut flower species susceptible to vascular blockage of bacteria and ethylene (Tehranifar et al., 2013). SA, a vase solution additive for extending the postharvest longevity of flower species, susceptible to vascular blockage of bacteria and ethylene, helps maintaining relative water constants (RWC). It was observed that SA (100-300 ppm) treated plants showed 50% higher RWC compared to control which extended the postharvest life of cut flowers (Tehranifar et al., 2013). Moreover, SA treatment at 1.5 mM showed the best effect on fresh weight, water uptake and vase life of cut flowers (Marandi et al., 2011). Kazemi et al. (2011) elaborated that SA treatment decreased microbial population and increased water uptake in carnation cut flowers. Ascorbic acid (AsA) (vitamin C), a novel enzyme, is a product of D-glucose metabolism in higher plants. It affects plant growth and development and plays a role in electron transport system (El-Kobisy et al., 2012). Ascorbic acid has been associated with several types of biological activities in plants. For instance, as enzyme co-factors, antioxidant, and as a donor/acceptor in electron transport at the plasma membrane or in the chloroplast (Conklin, 2001). Moreover, a high level of endogenous ascorbic acid is necessary to maintain the antioxidant system that protects plants from oxidative damage (Cheruth, 2009). Azizi et al. (2015) investigated various continuous treatments of ascorbic acid, viz. 0, 50, 100 and 200 mg L⁻¹ to lisanthus cut flowers and reported that longest vase life, maximum solution uptake, highest dry matter and minimum loss of fresh weight were obtained 200 mg L⁻¹ ascorbic acid. Moreover, they concluded that ascorbic acid, as a natural antioxidant which helped prolonging keeping quality of cut lisanthus flowers when applied at suitable concentration.

Abdulrahman et al. (2012) investigated various levels of ascorbic acid (50, 100, 150 mg L⁻¹) and suggested that 150 mg L⁻¹ (AsA) significantly increased vase life, fresh weight and percentage of total carbohydrates in cut snapdragon flowers. Moreover, ascorbic acid at 200 ppm improved growth and keeping quality by delaying flower opening and stimulating accumulation of carbohydrate, probably due to a fact that vitamins like ascorbic acid are considered as a bio-regulator compounds, which in little concentration exerted profound influence upon quality and long vase life (Bedour and Rawia, 2011). Sheikh et al. (2014) suggested ascorbic acid as cheap, safe, biodegradable compound and suitable alternative for chemical treatments in order to prolong vase life of cut flowers of Eustoma as it exhibited the longest vase life and higher relative water contents and petal water contents at 300 mg L⁻¹ ascorbic acid due to its acidic and anti-stress properties.

Keeping in view high market demand in local flower markets and role of pulsing with various chemicals to improve the postharvest life of cut flowers, present study was conducted with an aim to extend the postharvest life of cut tuberose spikes.

MATERIALS AND METHODS

**Plant material:** Cut spikes were obtained from a commercial grower at Pattoki, Distt. Kasur, Punjab. Flowers grown in open field were harvested early in the morning before 10:00 A.M., packed in cardboard boxes and immediately transported in air-conditioned vehicle to Postharvest and Floriculture Laboratory, Institute of Horticultural Sciences, University of Agriculture, Faisalabad, Pakistan, within 5 hours of harvest.

**Spikes handling in laboratory:** On arrival, flowers were recut to 60 cm length for the removal of any air embolism. Except upper two, all leaves were removed from lower side of spikes and placed in separate jars according to the treatments having 500 mL of pulsing solution for 24 hours at room temperature. Afterwards, spikes were placed in jars with three spikes each according to the treatments having 500 mL of distilled water, in a vase life evaluation room. Vases (glass jars) were arranged in a vase life evaluation room at 23±2°C temperature, 60±10% R.H. and a photosynthetically active photon flux of 12 μmol m⁻² s⁻¹ with 12h photoperiod from cool...
white florescent tubes. Experiment was arranged in a completely randomized design with five replications in each treatment having three cut spikes in each experimental unit (glass jar).

**Pulsing treatments:** The concentrations of chemicals were selected on the basis of preliminary experiments (data not shown). There were seven treatments and a control, viz. 10% sucrose, 50 mg L\(^{-1}\) salicylic acid, 100 mg L\(^{-1}\) acetyl salicylic acid, 100 mg L\(^{-1}\) ascorbic acid, 10% sucrose plus 50 mg L\(^{-1}\) salicylic acid, 10% sucrose plus 100 mg L\(^{-1}\) acetyl salicylic acid and 10% sucrose plus 100 mg L\(^{-1}\) ascorbic acid. Data regarding following postharvest indices were recorded using standard procedures:

**Spike Characteristics:**

**Vase life (days):** Stems were observed daily for visual appeal. Vase life was considered ended if, either half of the florets were wilted or bent neck symptoms appeared or the visual quality of the spike did not meet the commercial standards (Joyce et al., 2000; Ahmad et al., 2011).

**Water uptake (mL):** Volume of water absorbed during first five days of vase life was measured by recording remaining volume of vase water on day 5 of vase life from each jar containing three spikes. Water uptake was calculated as:

\[
\text{Water uptake (mL)} = (S_0 - S_5)
\]

Where, \(S_0\) is the amount of vase water on day 0 and \(S_5\) is the amount of vase water on the day 5 (Kazemi and Ameri, 2012).

**Relative fresh weight (% of initial FW):** Relative fresh weight (RFW) on day 5 of vase life (Kazemi and Ameri, 2012) was calculated as:

\[
\text{Relative fresh weight of spike (RFW)} = \left(\frac{FW_5}{FW_0}\right) \times 100
\]

Where \(FW_5\) is the fresh weight (g) of stem on day 5 and \(FW_0\) is the fresh weight (g) of the same stem on day 0 (He et al., 2006; Ahmad et al., 2011). Relative fresh weight of one spike was measured from each replication and average was worked out.

**Days to open 50% florets (days):** Number of days was counted starting from the day 0 to the day when 50% of the total florets were completely opened on the spike.

**Dry weight (%):** Dry weight percentage was calculated by using following formula:

\[
\text{Dry weight % age} = \left(\frac{DW}{FW}\right) \times 100
\]

Dry weight of same spike was measured whose fresh weight was measured earlier on day 0 from each replication and averaged. For estimation of dry weight, cut spikes were sun dried for one day, packed in brown paper bags and kept in oven at 80°C for 48 hours to record the dry weight at the end of vase life (Ahmad, 2009). Dry weight percentage was worked out from each replication and average was worked out.

**Spike/flower quality:** Flower quality was rated by three different judges (postgraduate floriculture students) adopting the method described by Dest and Guillard (1987) and average was worked out. Flower quality was rated in numbers using a scale ranging from 1 to 9, where 1 = Poor quality, 5 = Medium quality, 9 = Good quality. Spike quality was judged when 50% of florets were opened on spikes.

**Floret Characteristics:**

**Average life of floret (hours):** It was calculated by counting the numbers of hours starting from the opening of bud till floret lose its freshness. It was calculated from two florets of each spike in each replication and average was computed.

**Floret head diameter (cm):** Head diameter of fully opened two uppermost florets from each spike in each replication was measured on the last day of vase life with digital caliper and then their average was computed.

**Floret length (cm):** Length of two fully opened uppermost florets from each spike in each replication was measured on the last day of vase life with a measuring scale and average was worked out.

**Open floret (%):** Total number of florets (opened and closed) was counted on last day of vase life and number of opened florets was calculated as:

\[
\text{Open florets (%age)} = \left(\frac{\text{florets opened}}{\text{total florets}}\right) \times 100
\]

Open florets percentage was computed from 15 spikes per treatment and average was worked out.

**Ion leakage of floret (%):** Ion leakage of florets was measured on the last day of vase life of each treatment. For this purpose, five fresh florets were randomly selected from each replication and rubbed with silica powder. Rubbed petals were washed thoroughly with distilled water and shifted to test tubes containing 15 mL distilled water. These test tubes were placed on an orbital shaker for 10 min and EC\(_1\) was recorded. Again test tubes were placed on the orbital shaker for another 100 min. and EC\(_2\) was recorded (Ahmad, 2009) and ion leakage was calculated by using following equation:

\[
\text{Ion leakage (%) = } \left(\frac{EC_2}{EC_1}\right) \times 100
\]

**Change in pH and EC (DS m\(^{-1}\)):** Initial and final pH and EC of vase water were recorded using pH and EC meter (Hanna-HI9813-6) to observe changes in pH and EC during the study. For this purpose, three readings were taken from each replication on day 0 and final day of vase life of each cut spike and average was worked out.

**Change in total soluble solids (°Brix) of florets:** Total soluble solids of florets were measured on day 0 and at the end of vase life. For this purpose, six fully open florets were squashed to obtain sap from each replication. TSS in the sap was measured with the hand refractometer with a range of 0 to 30 °Brix by placing one drop on the prism as described by Bayleyegn et al. (2012). Between samples, prism of refractometer was washed with alcohol, rinsed by distilled water and dried using tissue paper. Three readings were taken from each replication and then their average was computed.

**Statistical analysis:** Experiment was arranged in a completely randomized design and data were analyzed using General
RESULTS

Spice Characteristics:
**Vase life (days):** Data revealed highly significant differences (P≤0.0001) among various pulsing treatments for vase life (Table 1). Longest vase life (10.2 d) was recorded in spikes pulsed with combination of 10% sucrose plus 50 mg L⁻¹ salicylic acid for 24 hours followed by spikes pulsed with 50 mg L⁻¹ salicylic acid (9.2 d). Pulsing with 10% sucrose plus 50 mg L⁻¹ salicylic acid extended vase life by 3.6 days compared to spikes pulsed in control (distilled water). All the pulsing treatments increased the vase life of spikes, compared to distilled water which had shorter vase life (6.6 days).

**Water uptake (mL):** Highly significant differences (P=0.0001) among various pulsing treatments for water uptake on day five of vase life was observed (Table 1). Highest water uptake (106.0 mL) was observed for spikes pulsed with 50 mg L⁻¹ salicylic acid, which was at par with water absorbed by the spikes pulsed with 10% sucrose (104.0 mL) or 10% sucrose plus 50 mg L⁻¹ salicylic acid (101.6 mL). Moreover, water uptake of spikes pulsed with 50 mg L⁻¹ salicylic acid was 26 mL greater than water taken up by the spikes in control (84.0 mL), which absorbed minimum water.

**Relative fresh weight (% of initial FW):** Data depicted significant differences (P=0.0274) among various pulsing treatments for relative fresh weight percentage of spikes on day five of vase life (Table 1). The spikes pulsed with 50 mg L⁻¹ salicylic acid exhibited the maximum relative fresh weight percentage (109.9%) followed by combination of 10% sucrose plus 50 mg L⁻¹ salicylic acid (108.4%), 100 mg L⁻¹ acetyl salicylic acid (107.7%), 100 mg L⁻¹ ascorbic acid (107.5%) and 10% sucrose (110.7%). However, relative fresh weight percentages of all these treatments were statistically similar to each other but greater than other treatments. On the other hand, spikes without pulsing (control) had minimum relative fresh weight percentage (104.6%).

**Days to open 50% florets (days):** Significant differences existed (P=0.0115) among various pulsing treatments for days to open 50% florets on spikes (Table 1). The spikes pulsed with 10% sucrose plus 50 mg L⁻¹ salicylic acid took significantly longer time to open 50% florets (5.0 days) and prolonged the time period to open 50% florets on spikes by 1.6 days as compared to control, followed by spikes pulsed with 50 mg L⁻¹ salicylic acid (4.6 days) and 10% sucrose plus 100 mg L⁻¹ ascorbic acid (4.4 days). However, these differences were non-significant among each other. On the other hand, spikes without pulsing (distilled water) took minimum number of days (3.4) to open 50% florets.

**Dry weight (%):** Highly significant differences (P<0.0001) among various pulsing treatments were recorded for dry weight percentage of spikes (Table 1). Dry weight percentage (13.2%) observed in spikes pulsed with 10% sucrose plus 50 mg L⁻¹ salicylic acid, 10% sucrose (13%), 10% sucrose plus 100 mg L⁻¹ acetyl salicylic acid (12.8%) and 10% sucrose plus 100 mg L⁻¹ ascorbic acid (12.7%) was statistically similar but greater than the rest of treatments. Minimum dry weight percentage (10.7%) was recorded in control which was 2.5% less than spikes pulsed with 10% sucrose plus 50 mg L⁻¹ salicylic acid.

** Spike quality:** Highly significant differences (P=0.0002) in the spike quality were observed among various pulsing treatments (Table 1). Significantly superior quality spikes (7.6) were observed when pulsed with 10% sucrose plus 50 mg L⁻¹ salicylic acid followed by spikes pulsed with 50 mg L⁻¹ acetyl salicylic acid (9.2 d). Pulsing with 10% sucrose plus 100 mg L⁻¹ salicylic acid (9.4 d) and 10% sucrose plus 100 mg L⁻¹ acetyl salicylic acid (12.8) days was statistically similar but greater than the rest of treatments. Minimum spike quality was (5.3) observed in spikes pulsed with distilled water.

### Table 1. Effect of pulsing with different preservatives on spike characteristics of tuberose.

<table>
<thead>
<tr>
<th>Treatments</th>
<th>*Vase life (days)</th>
<th>*Water uptake (mL)</th>
<th><strong>Relative fresh weight (%)</strong></th>
<th><strong>Dry weight (%)</strong></th>
<th><strong>Spike quality</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>Control (Distilled water)</td>
<td>6.6±0.37 d</td>
<td>87.6±1.6 b</td>
<td>104.6±0.6 b</td>
<td>10.7±0.3 b</td>
<td>5.2±0.49 d</td>
</tr>
<tr>
<td>10% sucrose</td>
<td>8.2±0.37 bc</td>
<td>104.0±3.1 a</td>
<td>110.7±1.4 a</td>
<td>4.0±0.3 bc</td>
<td>13.0±0.1 a</td>
</tr>
<tr>
<td>50 mg L⁻¹ salicylic acid (SA)</td>
<td>9.4±0.24 ab</td>
<td>106.0±3.2 a</td>
<td>109.9±1.0 a</td>
<td>4.6±0.2 ab</td>
<td>10.9±0.1 b</td>
</tr>
<tr>
<td>100 mg L⁻¹ Ascorbic Acid (AA)</td>
<td>8.0±0.63 c</td>
<td>84.0±2.2 b</td>
<td>107.7±2.1 ab</td>
<td>4.0±0.3 bc</td>
<td>10.4±0.4 b</td>
</tr>
<tr>
<td>10% sucrose+50 mg L⁻¹ SA</td>
<td>10.2±0.37 a</td>
<td>101.6±3.3 a</td>
<td>108.4±2.4 ab</td>
<td>5.0±0.3 a</td>
<td>13.2±0.1 a</td>
</tr>
<tr>
<td>10% sucrose+100 mg L⁻¹ SA</td>
<td>8.6±0.46 bc</td>
<td>88.8±4.3 b</td>
<td>105.2±0.3 b</td>
<td>4.2±0.2 b</td>
<td>12.8±0.2 a</td>
</tr>
<tr>
<td>10% sucrose+100 mg L⁻¹ AA</td>
<td>8.6±0.4 bc</td>
<td>89.4±3.8 b</td>
<td>105.1±0.4 b</td>
<td>4.4±0.2 ab</td>
<td>12.7±0.2 a</td>
</tr>
</tbody>
</table>

*Relative fresh weight of spike (RFW) = (FW/FW₀) x 100, where FW₀: Fresh weight at day 0 and FW: Fresh weight at day 5. **Dry weight % age = (DW/FW) x 100, where FW: Fresh weight at day 0 and DW: Dry weight at the end of vase life. *Means separation within columns by Least significant difference test at P ≤ 0.05. **Values were obtained using General Linear Model (GLM) procedures of Statistix 8.1. *Values are averages of 15 replicate spikes. **Values are averages of 5 replicate spikes.
Table 3. Change in pH and EC of vase water, while total soluble solids (TSS) in florets during vase life. Values are averages of 15 readings.

<table>
<thead>
<tr>
<th>Treatments</th>
<th>Initial pH</th>
<th>Final pH</th>
<th>pH change</th>
<th>Initial EC (dS m⁻¹)</th>
<th>Final EC (dS m⁻¹)</th>
<th>EC change</th>
<th>Initial TSS</th>
<th>Final TSS</th>
<th>TSS Change</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control (Distilled water)</td>
<td>6.9</td>
<td>7.1</td>
<td>+0.2</td>
<td>0.65</td>
<td>0.57</td>
<td>-0.08</td>
<td>6.8</td>
<td>6.0</td>
<td>-0.8</td>
</tr>
<tr>
<td>10% Sucrose</td>
<td>6.2</td>
<td>6.3</td>
<td>+0.1</td>
<td>0.57</td>
<td>0.50</td>
<td>-0.07</td>
<td>7.1</td>
<td>7.5</td>
<td>+0.4</td>
</tr>
<tr>
<td>50 mg L⁻¹ salicylic acid (SA)</td>
<td>4.6</td>
<td>4.9</td>
<td>+0.2</td>
<td>0.74</td>
<td>0.81</td>
<td>+0.07</td>
<td>6.5</td>
<td>6.3</td>
<td>-0.2</td>
</tr>
<tr>
<td>100 mg L⁻¹ acetyl salicylic acid (AsA)</td>
<td>4.3</td>
<td>4.4</td>
<td>+0.1</td>
<td>0.25</td>
<td>0.24</td>
<td>-0.01</td>
<td>6.0</td>
<td>5.6</td>
<td>-0.4</td>
</tr>
<tr>
<td>100 mg L⁻¹ ascorbic acid (AA)</td>
<td>5.2</td>
<td>4.6</td>
<td>-0.6</td>
<td>0.88</td>
<td>0.66</td>
<td>-0.22</td>
<td>6.0</td>
<td>5.5</td>
<td>-0.5</td>
</tr>
<tr>
<td>10% sucrose+50 mg L⁻¹ SA</td>
<td>5.5</td>
<td>5.4</td>
<td>-0.1</td>
<td>0.89</td>
<td>0.19</td>
<td>-0.70</td>
<td>6.9</td>
<td>6.9</td>
<td>0.0</td>
</tr>
<tr>
<td>10% sucrose+100 mg L⁻¹ AsA</td>
<td>4.3</td>
<td>4.4</td>
<td>+0.1</td>
<td>0.25</td>
<td>0.24</td>
<td>-0.01</td>
<td>5.6</td>
<td>5.8</td>
<td>+0.2</td>
</tr>
<tr>
<td>10% sucrose+100 mg L⁻¹ AA</td>
<td>4.5</td>
<td>4.5</td>
<td>0.0</td>
<td>0.91</td>
<td>0.11</td>
<td>-0.79</td>
<td>6.6</td>
<td>6.8</td>
<td>+0.2</td>
</tr>
</tbody>
</table>

L⁻¹ salicylic acid (6.8). Inferior spike quality was observed in control (distilled water) (5.2).

Floret length (cm): Highly significant differences were observed (P<0.0001) among various pulsing treatments for floret length (Table 2). Longest floret (7.0 cm) was recorded in spikes pulsed with 10% sucrose plus 50 mg L⁻¹ salicylic acid which was 0.9 cm longer than the florets in control (6.1 cm). No statistical differences were recorded in the floret lengths of the spikes pulsed with 10% sucrose plus 50 mg L⁻¹ salicylic acid, 50 mg L⁻¹ salicylic acid, 100 mg L⁻¹ acetyl salicylic acid and 100 mg L⁻¹ ascorbic acid, viz. 7.0 cm, 6.9 cm, 6.8 cm and 6.9 cm, respectively.

Floret head diameter (cm): Non-significant differences were observed (P>0.05) among various pulsing treatments for floret diameter (Table 2). However, spikes pulsed with 10% sucrose plus 50 mg L⁻¹ salicylic acid and 10% sucrose exhibited numerically greater floret diameter (4.2 cm) each, while minimum diameter (3.8 cm) was recorded in control.

Open florets (%): Highly significant differences were observed (P<0.0001) among various pulsing treatments for open florets percentage (Table 2). Highest percentage were opened in the spikes (20%) pulsed with 10% sucrose plus 50 mg L⁻¹ salicylic acid as compared to control (70%) which had the minimum open florets percentage.

Ion leakage (%): Data collected on ion leakage percentage of florets also exhibited highly significant differences (P<0.0001) among various pulsing treatments. Minimum ion leakage percentage was noted in the florets of spikes pulsed with 50 mg L⁻¹ salicylic acid and combination of 10% sucrose plus 50 mg L⁻¹ salicylic acid, viz. 140.0% and 141.8%, respectively. On the other hand, maximum ion leakage percentage was recorded in the florets of spikes in control and pulsed with 10% sucrose, viz. 175.8% and 171.8%, respectively (Table 2).

Change in EC (dS m⁻¹) and pH: Data regarding change in vase water pH showed variable response. In some treatments it was increased, viz. Control, 10% Sucrose, 50 mg L⁻¹ salicylic acid, 100 mg L⁻¹ acetyl salicylic acid and 10% sucrose plus 100 mg L⁻¹ acetyl salicylic acid but in others it
was decreased. In contrast, EC of all treatment decreased during vase life evaluation period accept in the vase solution containing 50 mg L⁻¹ salicylic acid (Table 3).

**Change in total soluble solids (TSS) of florets (°Brix):** Data regarding change in TSS of florets during the vase life period showed variable trend for various pulsing treatments. TSS remained same or slightly increased in the florets of the spikes pulsed with the pulsing treatment having sucrose while decreased with the pulsing without sucrose (Table 3).

**DISCUSSION**

Largely, from aforementioned results, it is evident that pulsing treatments with various preservative solutions including 10% sucrose, 50 mg L⁻¹ salicylic acid, 100 mg L⁻¹ acetyl salicylic acid, 100 mg L⁻¹ ascorbic acid, 10% sucrose plus 50 mg L⁻¹ salicylic acid, 10% sucrose plus 100 mg L⁻¹ acetyl salicylic acid, 10% sucrose plus 100 mg L⁻¹ ascorbic acid and distilled water (control) significantly affected spike and floret characteristics of cut tuberose as presented in Table 1 and 2. It was observed that among various pulsing treatments, 10% sucrose plus 50 mg L⁻¹ salicylic acid for 24 hours reported the longest vase life (10.2 d), more days to open 50% florets (5.0 days), maximum dry weight percentage (13.2%) and the superior quality spikes (7.6) (Table 1) relative to control. Similarly, the longest florets (7.0 cm), greater floret diameter (4.2 cm), the highest percentage of opened florets (90.0%) and the minimum ion leakage (140.0%) was noted in the florets of spikes of cut tuberose pulsed with 10% sucrose plus 50 mg L⁻¹ salicylic acid for 24 hours period as compared to control (Table 4.5.2). On the other hand, the highest water uptake (106.0 mL) and the maximum relative fresh weight percentage (109.9%) were observed in cut tuberose spikes treated with 50 mg L⁻¹ salicylic acid without sucrose solution compared to control (Table 1).

The best performance of sucrose solution in maintaining and prolonging postharvest life might be attributed to supplementing the natural sugar (Nowak and Rudnicki, 1990), preventing leaf desiccation (Silvanda et al., 2011), improving water balance (Hutchinson et al., 2003) and increased osmotic potential of the stem and petals, thus improving their ability to absorb nutrients and maintain their turgidity (Abbasi and Asil, 2011). Our results are also in lines with Bahrehmand et al. (2014) and Srivastava et al. (2015) for tuberose cut flowers and chrysanthemum, respectively. Moreover, Lama et al. (2013) reported extended vase life in cut roses due to sucrose solution are attributed to provision of carbohydrates and reduction in oxidative stress mediated damages during senescence. Similar results were observed in present study as mentioned in Tables 1 and 2 vis-à-vis various spike and floret characteristics studied. However, sucrose alone was not suitable which may be attributed to probable increase of microbial agents in vase solution. Therefore, combining sucrose with a biocide, changing the vase solution or re-cutting the stem end may reverse this effect (Jowkar and Salehi, 2005).

Exogenously applied SA at pre harvest stage increased flower quality, and vase life of cut rose. These results suggest that SA could be used as potential growth promoter to improve postharvest life of roses (Abdolmaleki, 2015). SA has also been found to play a key role in the regulation of plant growth, development and in responses to environmental stresses (Hayat et al., 2010). Further, its role is evident in ion uptake and transport (Harper and Balke, 1981), photosynthetic rate, stomatal conductance and transpiration (Khan et al., 2003). SA can modulate plant responses to a wide range of oxidative stresses (Shirasu et al., 1997). SA and its derivative, acetyl salicylic acid (ASA) also plays a role as an antagonist to ethylene action have been reported to inhibit ethylene production in pear (Leslie and Romani, 1988) and banana (Srivastava and Dwivedi, 2000). Also, the upward gravitropic bending of snapdragon was inhibited using SA (Friedman et al., 2003).

Similar results were observed in present study in which SA alone or in combination with sucrose helped strengthening various spike (Table 1) and floret (Table 2) characteristics of cut tuberose. Besides, SA modulates plant responses to a wide range of oxidative stresses and prevents cell wall degradation and thereby reduction in water loss (Shirasu et al., 1997). The present study results are also in accordance with the results proposed by Marandi et al. (2011), Kazemi et al. (2011) and Tehranifar et al. (2013) vis-à-vis postharvest life of cut flowers. Therefore, SA is used as a vase solution additive being natural, cheap, safe and biodegradable compound for extending the postharvest longevity of cut flower species susceptible to vascular blockage of bacteria and ethylene (Tehranifar et al., 2013).

**CONCLUSION**

It is concluded that pulsing of cut tuberose spikes with 50 mg L⁻¹ SA with or without 10 % sucrose proved effective for extending vase life and improving postharvest performance of cut tuberose spikes.

**REFERENCES**


Postharvest life of tuberose


