

# Effects of postharvest relative humidity and various re-cutting on vase life of cut rose flowers

Article

Accepted Version

Chamani, Esmaeil and Wagstaff, Carol (2019) Effects of postharvest relative humidity and various re-cutting on vase life of cut rose flowers. International Journal of Postharvest Technology and Innovation, 6 (1). pp. 70-82. ISSN 1744-7569 doi: https://doi.org/10.1504/IJPTI.2019.104207 Available at http://centaur.reading.ac.uk/87817/

It is advisable to refer to the publisher's version if you intend to cite from the work. See <u>Guidance on citing</u>.

To link to this article DOI: http://dx.doi.org/10.1504/IJPTI.2019.104207

Publisher: Inderscience

All outputs in CentAUR are protected by Intellectual Property Rights law, including copyright law. Copyright and IPR is retained by the creators or other copyright holders. Terms and conditions for use of this material are defined in



the End User Agreement.

## www.reading.ac.uk/centaur

### CentAUR

Central Archive at the University of Reading

Reading's research outputs online

Effects of postharvest relative humidity and various re-cutting on vase life of	1
cut rose flowers	2
Esmaeil Chamani <sup>a*</sup> and Carol Wagstaff <sup>b</sup>	3
<sup>a</sup> Department of Horticultural Sciences, Faculty of Agriculture and Natural Resources, University of Mohaghegh Ardabili, Ardabil 56199-11367, IR	4 5
<sup>b</sup> School of Food Biosciences, University of Reading, Whiteknights, PO Box 226, Reading RG6 6AP, UK	6
*Coresponding author. E-mail address: echamani@uma.ac.ir	7
	8

#### Abstract

9

Studies were conducted to evaluate the effects of different relative humidity levels (60, 75, and 10 90%) and recutting (0, 1, 2, 3, 4, and 5 cm recutting end of flower stem) treatments on vase life of 11 cut rose flower. Two separate experiments (bucket and vase experiments) were conducted based 12 on completely randomized design with factorial arrangement with 8 replications in bucket 13 experiment and 5 replications in vase experiment. Analysis of variance revealed that two ways 14 effects of various RH and recuts did not significantly (P  $\leq 0.05$ ) affected flower vase life, relative 15 fresh weight, solution uptake, and bacterial populations. Cut rose flower stored in chamber with 16 90% relative humidity had the longest vase life, while those one kept in 60% showed the shortest 17 longevity. The result of mean comparisons revealed with increasing relative humidity from 60% 18 to 90%, bacterial populations was increased too. 19

Keywords: bacterial count, flower diameter, relative fresh weight, solution uptake

#### Introduction

21

20

Control the uptake of  $CO_2$  for photosynthesis and prevention of water vapor is done via 22 stomata functioning, which are on the leaf surface. Guard cells bounded to the stomata and 23 equipped for autonomous ABA synthesis to control stomata opening and closure (Bauer *et al.*, 24 2013). Growing conditions largely effect on stomata responsiveness to closing during postharvest 25 life of flowers (Fanourakis *et al.*, 2016). Flowers longevity is limited at an early stage of 26 postharvest due to a range of factors mainly happen during preharvest. The shortened longevity 27 of cut roses is primarily related to water loss for their large leaf area and unfavorable growth 1 conditions affect the stomatal response (In et al., 2016b) and phenotype, which is determined by 2 genotype (In et al., 2016a). Further, it's also reported that the variation in rose flower longevity 3 has been associated mainly with vascular obstruction through the stem, which affect water status 4 (van Doorn, 1997). In addition, some other factors such as limitation by fungi infection, vascular 5 occlusion and pedicel bending, can be resulted to early flower senescence in cut roses. To 6 increase vase life of cut rose flowers, it's necessary to prevent or retard flower wilting (Rasouli et 7 al., 2015). 8

Water deficit stress during postharvest handling is one of the most important factors to 9 determine cut rose flower longevity. Abnormal flower opening, flower wilting, bent neck, and 10 failure to open are the results of water deficit stress (Jin et al., 2006). Hence, correct postharvest 11 handling and preventing dehydration as well as controlling temperature and relative humidity 12 during postharvest storage is essential to maximize flower vase life and quality (Reid, 2001). 13 Among the postharvest factors effecting on flowers longevity, RH is strongly correlated with cut 14 rose's longevity (In et al., 2016a). Carvalho et al. (2015) reported that preharvest high RH ( $\geq$ 15 85%), hampered stomata functioning and adversely affected cut rose longevity. Arve et al. (2017) 16 reported that plants developed at high RH during leaf development, when exposed to daily 17 change in RH and temperature, showed improved stomata functioning. Moreover, different cut 18 rose cultivars showed different patterns of reaction during postharvest, where in cultivar 'Akito' 19 decreased stomata functioning adversely affected the vase life. However, 'Grand Prix' cultivar 20 did not significantly affect by stomata functioning (Woltering & Paillart, 2018). 21

Some physical treatments such as splitting or crushing stems and also removing bark at the 22 base of the stem increased water uptake and 25% increase in fresh weight which resulted in 23 enhancing flower longevity compared to control (Milner, 2009). Similarly, it's reported that bark 24 removal and stem-end splitting when applied after short term storage for 24 h at 4°C, increased 25 the vase life of cut rose and acacia. However, crushing stems had no effect on the vase life of 26 fresh-cut rose (Ahmad et al., 2011). It's reported to improve water uptake and maintain water 27 balance for cut flowers and foliage, water loss should be decreased by reduction in leaf area and 28 store them in low temperature and high RH and using some pulse treatments such as sucrose in 29 vases to improve their vase life (Ahmad et al., 2011). In the ambient air, relative humidity was 30 affected water loss in harvested crops. With consideration that various crops need different 1 relative humidity, however, harvested crops keep their nutritional quality, appearance and weight 2 at very high RH (Hong *et al.*, 1999). 3

Influence on overseas shipping on flowers longevity has not been well understood. 4 Furthermore, Ahmad et al. (2011) reported that high RH decreases water loss and maintain 5 flower longevity. However, In et al. (2016 b) reported that high RH lead to the attenuated 6 stomatal responsiveness and increase water loss. Due to study the effect of shipping influence and 7 some contradictory results on effect of RH on flowers longevity, the aim of present experiments 8 was to examine the relationship between different relative humidity, various recuts and bacterial 9 population's effects on cut rose flowers longevity. Here, we have shown that various postharvest 10 conditions can significantly affect cut roses longevity. 11

#### MATERIAL AND METHODS

#### **Plant material**

Cut H3O rose flowers produced in Ethiopia greenhouses were obtained at the commercial 14 stage of bud opening (petals starting to reflex) from MM Flower Factory in Cambridge and 15 transferred to the Reading University and were held in a cold room (4°C) and then transported 16 within 3 days to the experiment chambers (phytotrons). No symptoms of botrytis were observed 17 in flowers. Two separate experiments (bucket and vase experiments) were conducted based on 18 completely randomized design by factorial arrangement with 8 replications in bucket experiment 19 and 5 replications in vase experiment .

#### **Bucket experiments**

Cut rose flowers were recut under the tap water to 1, 2, 3, 4 and 5 cm and placed in various 22 relative humidity (RH) at rates 60, 75, and 90%. No recut flowers were used as a control. Cut 23 rose flowers place in buckets. Three buckets were used for each recut and each bucket was 24 containing 8 cut rose flowers (exactly 24 flowers were used for each recut). Experiment 25 replicated to confirm the results of first experiment at room condition [Temp:  $22\pm2^{\circ}$ C and light 26 intensity:10 µmol. m<sup>-2</sup>. s<sup>-1</sup>]. 27

#### Vase experiment

28

3

13

12

In vase experiment, 10 cut rose flowers were used for each RH. Half of them 5 cm recut 1 and half of them without recut placed in various RHs at rates 60, 75, and 90%. Each cut rose 2 flowers place in a vase. Experiment replicated to confirm the results of first experiment at room 3 condition [Temp:  $22\pm2^{\circ}$ C and light intensity:10 µmol. m<sup>-2</sup>. s<sup>-1</sup>]. 4

#### Assessments

#### Vase life

Vase life was recorded as the time in days after treatment (day 0) that flowers reached the 7 end of their longevity due to bent neck or advanced signs of fading on all petals (Liao *et al.*, 8 2000; Chamani *et al.*, 2006).

#### **Relative fresh weight**

Following formula was used for calculation of relative fresh weight of stems: RFW (%) = 11  $(Wt/Wt=0) \times 100$ ; where Wt = weight of stems (g) at t = days 0, 2, 4, 6, etc. and Wt=0 = weight 12 of the same stem (g) on day 0. 13

#### **Solution Uptake**

Vase solution uptake was determined by using the formula: Solution uptake (ml day-1 g-1 15 fresh weight) = (St-1-St)/Wt=0; where, St = solution weight (g) at t = days 1, 2, 3, etc. St-1 = 16 solution weight (g) on the preceding day, and Wt=0 = fresh weight of the stem (g) on day 0. 17

#### Microbial count analysis

Preparation of nutrient agar and Maximum recovery diluent (MRD) for total plate count 19 was done as described previously by Chamani and Wagstaff, (2018). However, For the bacterial 20 count, 5'cm from the basal end of flower stems removed and then sterilized by careful blotting 21 with ethanol (98% v/v). After that, further cut the stem into 2 mm segments and weighted and 22 then were added to 90 ml of MRD in a stomacher bag and shaken for 60 seconds with 230 rpm 23 which this will create 10-1 dilution (w/v). 1 ml of the homogenized/inoculum was sampled from 24 the bag and it was serially diluted in 9 ml MRD to obtain 10-2, 10-3, 10-4 until 10-7. Then 1 ml 25 of the respective solution was placed on 15 ml nutrient agar temperature between 45 °C to 50 °C 26 on petri dish using pour plate technique and the plates were swirled to mix evenly. The 27 inoculated plates were allowed to cool at room temperature until the liquid solidified. The plates 28

14

18

5

6

then were incubated at 30 °C in inverted condition. After  $24 \pm 1$  h of incubation, number of 1 colonies per plate was counted using a colony counter. Plates with colonies more than 300 2 colonies are labelled with TNTC (too numerous to count) and plates with colonies less than 30 3 colonies were discarded . 4

#### **Statistical analysis**

All experiments were done in completely randomized design based on factorial 6 arrangement. Bucket experiments were done by 8 replications. Vase experiments were done by 5 replications in each trial for morphological traits and 3 replications for microbial count. Data 8 were subjected to analysis of variance (ANOVA) using Statistical Analysis System Ver. 9.2. 9 (SAS Institute, Cary, NC, USA). Mean differences between treatment were compared using 10 Duncan's Multiple Range Test at P<0.05. Graphs were then plotted using Excel spread sheet . 11

**Results** 

#### **Bucket experiment**

#### Vase life

Analysis of variance revealed that various RH and recuts significantly ( $P \leq 0.05$ ) affected 15 flower vase life, relative fresh weight, and solution uptake in both of the experiments. However, 16 no significant ( $P \leq 0.05$ ) difference was found in interaction effects of RH and recuts. Mean 17 comparison of results showed that different recut significantly ( $P \leq 0.05$ ) affected flower vase life. 18 Cut flowers with No recut significantly ( $P \leq 0.05$ ) had the lowest vase life. However, cut flowers 19 which recut 3, 4, and 5 cm had the highest vase life compared to no recut and 1 cm recuts. No 20 significant difference was found between 1 and 2 cm recuts. It's found that at least 2 cm recut is 21 necessary to get the highest vase life (Figure 1A & Figure 4). The result also revealed that 22 different relative humidity levels significantly ( $P \le 0.05$ ) affected flower vase life. The highest and 23 significant vase life was found in flowers placed in 90 % RH compared to 60 and 75% RH. 24 However, 75% Relative humidity significantly increased flower vase life compared to 60% RH 25 (Figure 1B & Figure 4). 26

#### Solution uptake

27

12

13 14

The result of experiment revealed that cut flowers placed in 60% and 90% relative humidity 1 had the highest and lowest solution uptake during whole experiment time, respectively. However, 2 75 % relative humidity made intermediate effects on cut flowers (Figure 2A). Mean comparison 3 revealed that recut flowers with 2, 3, 4, and 5 cm significantly ( $P \le 0.05$ ) had the highest solution 4 uptake during whole experiment time. Moreover, cut flowers with No recut (C0) had the lowest 5 solution uptake (Figure 2B). 6

#### **Relative fresh weight**

Fresh weight of cut rose flowers decreased from the second day of vase life, indicating a deterioration of their water status. The result of experiment also showed that cut flowers placed in 9 90% and 60% relative humidity had the highest and lowest relative fresh weight during whole 10 experiment time, respectively. However, 75 % relative humidity made intermediate effects on cut 11 flowers (Figure 3A). Mean comparison revealed that recut flowers with 2, 3, 4 and 5 cm, 12 significantly ( $P \le 0.05$ ) had the highest relative fresh weight during whole experiment. However, 13 cut flowers with no recut had the lowest relative fresh weight during experiment (Figure 3B). 14

#### Vase experiment

#### Vase life

Analysis of variance revealed that various RH and recuts significantly ( $P \le 0.05$ ) affected 17 flower vase life, relative fresh weight and solution uptake. But, no significant ( $P \le 0.05$ ) 18 difference was found in interaction effects of RH and recuts. The result revealed that different 19 relative humidity significantly ( $P \le 0.05$ ) affected flower vase life. The highest and significant 20 vase life was found in flowers placed in 90 % RH compared to 60 and 75% RH. However, 75 % 21 Relative humidity significantly increased flower vase life compared to 60% RH (Figure 5A & 22 Figure 8). In fact, with increasing relative humidity, flower vase life increased too . 23

Cut flowers with no recut significantly ( $P \le 0.05$ ) had the lowest vase life. However, cut 24 flowers which recut 5 cm had the highest vase life compared to no recut. It's concluded that recut 25 the flowers increased its vase life two times compared to No recut (Figure 5B and Figure 8). 26

#### **Relative fresh weight**

27

7

15

The result of experiment revealed that cut flowers placed in 90% and 60% relative humidity 1 had the highest and lowest relative fresh weight during whole experiment time respectively and 2 75% relative humidity made intermediate effects on cut flowers (Figure 6 A). Mean comparison 3 revealed that recut flowers with 5 cm significantly ( $P \le 0.05$ ) had the highest relative fresh weight 4 during whole experiment time compared to no recut (Figure 6 B). 5

#### Solution uptake

The result of experiment revealed that cut flowers placed in 60% and 90% relative humidity 7 significantly had the highest and lowest solution uptake till days 7 respectively. After days 7, 8 solution uptake in flowers placed in 90% RH remained constant (Figure 7A). Whereas, solution 9 uptake in flowers placed in 60 and 75% RH decreased because of flowers drying. It's also 10 revealed that recut flowers significantly had the highest and significant solution uptake during 11 experiment after days 5 compared to No recuts (Figure 7B). 12

#### **Bacteria count**

The result of mean comparisons revealed with increasing relative humidity from 60% to 14 90%, bacterial populations was increased too. But, no significant ( $P \leq 0.05$ ) differences were 15 found among them. However, with recutting the stem ends in cut rose flowers which held in vase 16 solution containing Crysal, bacterial populations significantly ( $P \leq 0.05$ ) decreased in stem ends. 17 It seems Crysal, did not deleted bacterial population. However, it's inhibited bacterial 18 multiplications. In fact, bacterial count in the 5 cm of stem end showed that, bacterial populations 19 were significantly decreased after 4 cm recut compared to recut less than 3 cm recut (Figure 9A). 20 However, in vase experiment, there were significant ( $P \leq 0.05$ ) differences between recut and 21 non-recuts flowers. However, it was found much more bacterial populations in stem ends on non-22 recut flowers compared to recut flowers (Figure 9B). 23

#### Discussion

Range of factors including preharvest growth conditions, shipping overseas, proper harvest
time, and appropriate storage can affect flowers longevity (Fanourakis *et al.*, 2015; In *et al.*,
26
2016, Baker, 2018). To maintain the natural appearance of flowers, quality deterioration should
be delayed. All consumers prefer cut flowers with high longevity (Asghari *et al.*, 2014).
28

7

13

24

The results of our experiments showed that various RH and recut treatments significantly 1 affected flower vase life, relative fresh weight, solution uptake, and bacterial populations in both 2 of the experiments. The highest vase life and relative fresh weight was observed in flowers 3 treated with 90% RH. Flowers stored in 60% RH had the highest solution uptake in bucket 4 experiment but in vase experiment in flower stored in 90% RH the amount of solution uptake did 5 not change after day 7 while decreased in 60 and 75%. It can be deduced that 90% RH is ideal 6 condition for prolonged vase life of rose flowers. However, it's reported that low temperature and 7 high RH decreases water loss and maintain flowers quality (Ahmad et al., 2011). The difference 8 between saturation vapor pressure and actual air vapor pressure defines evapotranspiration of leaf 9 and petals and is playing key role in water uptake. Hence, if it happens with high difference, 10 evapotranspiration will be increased too. However, high air relative humidity reduced 11 evaporation of water from the flower petals and leaves, resulted in to high fresh weight and long 12 longevity (Siddiquei, 2015). Our finding is in line with the study that reported the solution uptake 13 was affected strongly by RH compared to sucrose concentration, and greater solution uptake was 14 happened in lower relative humidity condition (Shimizu & Ichimura, 2007). 15

Moreover, Faragher et al (1986) reported that although keeping of cut rose flowers (*Rosa* 16 *hybrid* L. cv. Mercedes) at 65% relative humidity (RH) decreased petal water content by 20% 17 compared to flowers stored at 95% RH, it did not shorten vase life. However, it can be because of 18 cultivar types and some other unknown factors. For reduction of disease development low 19 temperature and higher relative humidity have also been suggested (Harkema *et al.*, 2013). 90% 20 relative humidity has been preferred for keeping of *Anthurium andraeanum* lindl, Strelitzea 21 reginae (Vieira *et al.*, 2014). 22

Results of both bucket and vase experiments also showed that recutting rose flower stem 23 could extent it's vase life, maintain higher fresh weight and uptake more solution in comparison 24 to no recut. Experiment with various recut showed that recutting flowers with 1, 2, 3, 4, and 5 cm 25 significantly increased vase life during whole experiment time compared to control. In fact, to 26 obtain the best results, it's necessary to recut flower stem ends at least 2 cm. Our results are in 27 consistent with an earlier study, where difference in vase life of cut carnation flowers was due to 28 flower stalks height (Chandra et al., 2013). All done experiments to evaluate flower vase life, 29 were conducted by using of various stems lengths which sometimes influenced flowers vase life. 30 In fact, cut flowers have been tested either at the stem length of harvest as little as 12 cm to as 1 much as 75 cm depending upon tested cultivars (Fanourakis et al., 2013). It's found that there was 2 significant negative correlation between vase life and stem length (Mortensen & Fjeld, 1998). 3 Actually, Short stems (i.e. short water transport path) and/or less leaves (i.e. lower water loss in 4 cut flower basis) would reduce loss of water balance resulted in longer vase life. Additionally, 5 some physical treatments such as splitting or crushing stems and also removing bark at the base 6 of the stem increased water uptake and 25% increase in fresh weight which resulted in enhancing 7 flower longevity compared to control (Milner, 2009). It's reported that bark removal and stem-8 end splitting increased the vase life of rose and acacia (Ahmad et al., 2011). 9

Our finding also revealed that with increasing relative humidity from 60% to 90%, bacterial 10 populations were increased too. However, with recutting the stem ends in cut rose flowers which 11 held in vase solution containing Crysal, bacterial populations significantly ( $P \le 0.05$ ) decreased in 12 stem ends. Vase life of cut rose flowers is short which could be related to excessive water loss 13 from the rose leaves, resulting in leaf desiccation and the development of bent necks (Mortensen 14 & Fjeld, 1998). Actually, short vase life in cut flowers is often the results of vascular occlusions 15 that restrict vase solution supply. Water absorption in stem is typically caused by blockage of cut 16 stem ends by microbes and physiological plugging which inhibits water uptake by flower stalk 17 (Hussen & Yasin, 2013). The accumulation of bacteria, in the stem ends may play an important 18 role in reduction of vase life, as a result of decreasing water uptake (van Doorn, 1997). 19

#### Conclusions

Cut roses longevity closely depends on preharvest growth conditions, genetic background, 21 and cut flowers storage. According to results of present study, high RH (90%) with recutting cut 22 H3O rose flower stems at least 2 cm, extended the cut rose longevity via maintaining the proper 23 stomatal functioning, reduction in bacterial population, developing a normal water balance, and 24 maintaining relative fresh weight Moreover, by increasing recut the stem end from 0 to 5 cm, 25 bacterial population in the stem was decreased too. 26

20

27

#### References

Ahmad L, Joyce DC, John DF, 2011. Physical stem-end treatment effects on cut rose and acacia28vase life and water relations. Postharvest Biol Technol 258-264.29

Arve LE, Kruse OMO, Tanino KK, Olsen JE, Futsæther C, Torre S, 2017. Daily changes in VPD
during leaf development in high air humidity increase the stomatal responsiveness to darkness
and dry air. J Plant Physiol 211: 63–69.

Asghari R, Salari A, Gharehdaghi S, 2014. Effect of pulsing solution and packaging type under
exogenous ethylene on physiological characteristics and post harvesting quality of cut roses
(*Rosa hybrida*). Am Eur J Agric Environ Sci 14: 329-335.

Baker JE, 2018. Preservation of cut flowers, in: Plant Growth Regulating Chemicals.CRC7Press, pp. 177–191.8

Bauer H, Ache P, Lautner S, Fromm J, Hartung W, Al-Rasheid KAS, Sonnewald S, Sonnewald 9
U, Kneitz S, Lachmann N, 2013. The stomatal response to reduced relative humidity requires 10
guard cell-autonomous ABA synthesis. Curr Biol 23: 53–57. 11

Carvalho DRA, Koning-Boucoiran CFS, Fanourakis D, Vasconcelos MW, Carvalho SMP,
Heuvelink E, Krens FA, Maliepaard C, 2015. QTL analysis for stomatal functioning in tetraploid
Rosa× hybrida grown at high relative air humidity and its implications on postharvest longevity.
Mol Breed 35: 172.

Carvalho DRA, Vasconcelos MW, Lee S, Koning-Boucoiran CFS, Vreugdenhil D, Krens FA, 16 Heuvelink E, Carvalho SMP, 2016. Gene expression and physiological responses associated to 17 stomatal functioning in Rosa× hybrida grown at high relative air humidity. Plant Sci 253: 154– 18 163. 19

Chamani E, Irving DE, Joyce DC, Arshad M. 2006. Studies with thidiazuron on the vase life of 20 cut rose flowers. J App Hortic 8: 42-44. 21

Chamani E, Wagstaff C, 2018. Response of Cut Rose Flowers to Relative Humidity and Recut 22 During Postharvest Life. Int J Hortic Sci Technol 5: 145–157. 23

Chandra SA, Kasturi A, Manohar RA, Narender RS, 2013. Effects of harvesting in different24heights on growth and flower yield of carnation. J Hortic 1: 1-4.25

Fanourakis D, Roland P, Andrea S, Andrew JM, Vaia S, Ernst JW, 2013. Sources of vase life26variation in cut roses: A review. Postharvest Biol Technol 78: 1–15.27

Fanourakis D, Velez-Ramirez AI, In BC, Barendse H, van Meeteren U, Woltering EJ, 2015. A	1
survey of preharvest conditions affecting the regulation of water loss during vase life. Acta Hortic 1064: 195–204.	2 3
Fanourakis D, Bouranis D, Giday H, Carvalho DRA, Nejad AR, Ottosen CO, 2016. Improving stomatal functioning at elevated growth air humidity: a review. J Plant Physiol 207: 51–60.	4 5
Hussein S, Yassin H, 2013. Review on the impact of different vase solutions on the postharvest life of rose flower. Int J Agric Res Rev 1:13-17.	6 7
In BC, Lee JH, Lee AK, Lim JH, 2016a. Conditions during export affect the potential vase life of cut roses ( <i>Rosa hybrida</i> L.). Hortic Environ Biotechnol 57: 504-510	8 9
In BC, Seo JY, Lim JH, 2016b. Preharvest environmental conditions affect the vase life of winter-cut roses grown under different commercial greenhouses. Hortic Environ Biotechnol 57: 27-37	10 11 12
In BC, Lim JH, 2018. Potential vase life of cut roses: Seasonal variation and relationships with growth conditions, phenotypes, and gene expressions. Postharvest Biol Technol 135: 93–103.	13 14
Jin JS, Shan NW, Ma N, Bai JH, Gao JP, 2006. Regulation of ascorbate peroxidase at the transcript level is involved in tolerance to postharvest water deficit stress in the cut rose ( <i>Rosa hybrida</i> L.) cv. Samantha. Postharvest Biol Technol 40: 236–243.	15 16 17
Kader AA, 2007. Postharvest Technology of Horticultural Crops. 3rd ed. Agriculture and Natural Resources. Publication 3311. University of California, Davis, CA.	18 19
Milner G, 2009. Fresh-Cut Flowers. Jojo Publishing, Melbourne, Australia.	20
Mortensen LM, Fjeld T, 1998. Effects of air humidity, lighting period and lamp type on growth and vase life in roses. Sci Hortic 73: 229–237.	21 22
Rasouli O, Ahmadi N, Behmanesh M, Daneshi Nergi MA, 2015. Effects of BA and TDZ on postharvest quality and expression of laccase and aquaporin genes in cut rose 'Sparkle'. South African J Bot 99: 75-79.	23 24 25
Reid A, 2001. Storage conditions for ornamental crops. The Chief Executive Officer of the Department of Agriculture and the State of Western Australia. No. 71/2001.	26 27

Siddiquei MW, 2015. Postharvest biology and technology of horticultural crops. CRP press. p540.	1 2
Shimizu-Yumoto H, Ichimura K, 2007. Effect of relative humidity and sucrose concentration on leaf injury and vase life during sucrose pulse treatment in cut Eustoma Flowers. Hort Res 6: 301–305.	3 4 5
Van Doorn WG, 1997. Water relations of cut flowers. Horticulture Rev 18: 1-85.	6
van Meeteren U, Aliniaeifard S, 2016. Stomata and postharvest physiology. Postharvest Rip Physiol Crop 1: 157.	7 8
Vieria MRS, Simoes AN, Souza PA, 2014. Recommended temperature and relative humidity for storage of Brazilian tropical flowers, African J Biotechnol 13: 1198-1201.	9 10
Woltering EJ, Paillart MJM, 2018. Effect of cold storage on stomatal functionality, water relations and flower performance in cut roses. Postharvest Biol Technol 136: 66–73.	11 12
	13
	14
	15
	16
	17
	18
	19
	20
	21
	22
	23



Figure 1. Vase life of rose flowers in different RH (A) and various recutting (B)

Different letters indicate significant differences determine using a Duncan's multiple range test (P<0.05). Error bars= SE (n=8)



--

. .

. .





Different letters indicate significant differences determine using a Duncan's multiple range test (P<0.05). Error bars= SE (n=8)



Figure 4. Effects of various RH (A=60%, B=75% and C=90%) with different recuts on cut rose flower vase life



Figure 5. Vase life in different RH (A) and different recutting (B) conditions

Different letters indicate significant differences determine using a Duncan's multiple range test (P<0.05). Error bars= SE (n=5)



----



Figure 6. Relative Fresh weight in different RH (A) and different recutting (B) conditions









Figure 8. Effects of different RH (A=60%, B=75% and C=90%) and recut on flower vase life

Different letters indicate significant differences determine using a Duncan's multiple range test (P<0.05). Error bars= SE (n=5)



Figure 9. Bacterial population in different re-cutting stem ends in cut rose flowers in bucket experiment (A) and vase experiment (B)

Different letters indicate significant differences determine using a Duncan's multiple range test (P<0.05). Error bars= 5 SE (n=5) 6