Storage Temperature and Duration Affect Quality and Post-storage Recovery of Vegetative Dendranthema × grandiflorum

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Abstract. Experiments were conducted to evaluate Dendranthema × grandiflorum (Ramat.) Kitamura cv. Bright Golden Anne quality and post-storage growth following storage in the range of 5 to 35°C, initial soil water levels (60%, 80%, 100%), and durations (0 to 8 days). Transpiration rate showed a quadratic relationship with storage temperature. Initial soil water content had little effect on transpiration rate in dark storage environments. The lowest transpiration rate was observed in plants stored at 15 or 20°C. Amino acid (AA) leakage and post-storage growth were well-correlated. Plants stored at or above 25°C became etiolated during storage, while storage at 15°C or below did not cause etiolation. Temperatures at or below 15°C did not affect subsequent growth rate of chrysanthemum plants. Storage at 20°C and above caused a reduction in post-storage growth rate following 2 days of storage.

Due to the availability of inexpensive labor and low production costs, many large-scale floriculture and foliage plant producers have established production centers in developing countries in tropical and subtropical regions. The development of international markets for floricultural crops has increased the necessity for information on long-distance shipping and storage conditions before reaching the consumer.

Adverse storage environments, especially high temperature and low relative humidity, enhance water loss from plants due to an increase in the water potential gradient between the leaf and surrounding environment (Lorenzo-Minguez et al., 1985; Stanley et al., 1982, 1983). High transpiration rate during shipping/storage can induce water stress and enhance ethylene production and leaf abscission (Graves and Gladon, 1985). Peterson et al. (1980) reported that water stress is a major factor in leaf abscission of Ficus benjamina during storage. Prolonged exposure to adverse shipping/storage environments can deteriorate plant quality and reduce growth after storage or delay recovery from storage stress. Buck and Blessington (1982) reported that foliar damage was higher when Ficus benjamina plants were exposed to high temperatures (37°C). Tropical plants are susceptible to chilling damage when they are exposed to low temperatures (<7 to 10°C) (Buck and Blessington, 1982). Poole and Conover (1983) reported that Ficus benjamina dropped more leaves as storage duration increased, but shipping durations up to 21 days could be obtained without significant loss in quality if proper temperature was maintained. It is, therefore, essential to provide proper shipping/storage conditions to avoid any detrimental effects on plant quality.

The major objectives of this study were to evaluate the quality and post-storage growth of Dendranthema × grandiflora plants after storage at various temperatures and for several durations and to determine the best storage conditions without affecting plant quality.

Materials and Methods

Pretreatment and culture. Rooted D. × grandiflorum cv. Bright Golden Anne shoot cuttings (Yoder Brothers, Barberton, Ohio) were planted (one plant per pot) in 10-cm plastic pots (=0.5 liter) containing 50 to 60 g (dry weight) 1 peat:1 perlite (v/v) medium amended with (all per m³) 4.8 kg each of dolomite, gypsum, and superphosphate (0N-20P-0K); 1.1 kg MagAmp (W.R. Grace, Fogelsville, Pa.); 0.8 kg KN0₃ and 13.2 g g fritted trace element. Plants were watered, as needed, with N, P, and K at 200, 87, and 165 mg·liter⁻¹, respectively, and allowed to establish in the greenhouse for 2 weeks under natural photoperiod. Environmental conditions during the growth period were: day/night temperatures, 27 ± 3°C/20 ± 2°C; day/night relative humidity, 38% ± 15%/70% ± 10%; light intensity of 300 to 1000 µmol·s⁻¹·m⁻².

Storage treatments and data collection. Two days before the storage experiment, plants were hand watered to runoff and excess water was allowed to drain overnight. The following morning, 100 plants were randomly selected and their weight at container capacity (CC) was recorded. Plant plus pot weight at 80% and 60% of CC and the amount of water needed to retain 100%, 80%, and 60% of CC after drying were calculated from the weight data. The plants were then allowed to dry until the desired percentage of CC was reached. The pots were covered with clear plastic film to prevent direct evaporation from the soil surface. Plants were then transferred to a storage chamber maintained at 5, 10, 15, 20, 25, 30, or 35°C. Relative humidity in the chamber was maintained at 75% ± 10% for each temperature, thus, the vapor pressure deficit ranged from =2.2 to =15 mb. Air inside the storage chamber was circulated using a fan fixed onto the ceiling of the chamber. Plants were kept in darkness, except for use of a low-intensity, green safelight during the data collection. Plants were not watered during the 8 days of storage.

Weight measurements were taken twice daily at 0700 and 1900 hr for 8 days. Transpiration rate was calculated from the weight-loss data.

AA leakage from leaf disks was estimated as described by Leopold et al. (1981). Two leaf disks, each 1.5 cm in diameter, were cut from the fourth leaf from the apex. Disk fresh weights were recorded and they were placed in glass vials containing 20 ml deionized, distilled water. Glass vials were shaken in a water bath maintained at 22°C for 2 hr. Absorbance readings of
the water were measured at 280 nm ($A_{280}$) with a spectrophotometer (Bausch and Lomb Spectronic 21, Rochester, N.Y.). Vials were then frozen overnight at –70°C in an ultra-low freezer to rupture cells, thawed, and shaken again for 2 hr and $A_{280}$ again was measured. Percentage AA leakage was calculated from $A_{280}$ readings of original and frozen samples.

At the end of the experiment, plants were harvested and leaf area was measured with a LI-COR-3050 area meter (LI-COR, Lincoln, Neb.).

Representative plants placed in each storage temperature were taken out after 0, 2, 4, 6, or 8 days of storage and placed back in the greenhouse. Plant height increase was recorded in the greenhouse for 3 weeks at 4-day intervals.

Experimental design and analysis. Plants were arranged in a split-plot design with 10 replications for weight loss measurements and five replications for other characteristics measured. Temperature and initial soil water level were the whole-plot and split-plot factors, respectively. Plants in the post-storage growth experiment were arranged in a split-split plot design. Temperature, initial water level, and storage duration were whole-, split-, and split-split plot factors, respectively. The data were analyzed using analysis of variance procedure. Regression analysis was carried out to test for the temperature and storage duration effects. Least square means were computed and differences in the means were tested using least significant difference procedure, where necessary.

Results and Discussion

The transpiration rate rapidly declined after placing plants in dark storage and leveled off after ≈48 hr (Fig. 1). Plants stored at 30 and 35°C showed an initial decline of transpiration rate followed by a rapid increase after 60 and 108 hr, respectively. Plants stored at these two temperatures showed severe damage at the end of storage.

Slight diurnal fluctuation of transpiration rate was observed in plants stored at 5 and 10°C, probably due to incomplete stomatal closure in the dark at these temperatures. It has been reported that stomata of some species do not close completely in darkness (Kramer, 1983; Rajapakse et al., 1989). Wilson (1983) reported that a low temperature “locks open” the stomata of some plants. Slow stomatal response has also been reported at low temperatures (McWilliam et al., 1982).

At all temperatures, plants with higher soil water levels showed higher average transpiration rates than those held at a lower CC (Fig. 2). The difference in transpiration rates between soil water levels was highest between 15 and 20°C, with a smaller difference at extreme temperatures. In previous studies, a decrease in transpiration rate has been reported with decreasing soil water content under greenhouse conditions (Ceulemans et al., 1979; Morison and Gifford, 1984). In light, stomata are fully open and both stomatal and cuticular transpiration account for the water loss. As water stress progresses, stomata begin to close, reducing the transpiration rate. In the present study, conducted in darkness, the stomata were probably closed or partially open; thus, the effect of soil water content on transpiration was not as drastic as under normal greenhouse conditions. Cuticular transpiration may have accounted for the majority of water loss in the present experiment due to closed or partially open stomata and poor epicuticular wax development in chrysanthemum leaves (Rajapakse et al., 1988).

The lowest transpiration rate was observed in plants stored at 15 and 20°C. The slightly higher transpiration rates at 5 and 10°C could be due to partial opening of the stomata or impaired stomatal movement as was shown at low temperatures for some tropical plants (Wilson, 1983). The relationship between transpiration and storage temperature is explained by a quadratic model.

The increase in transpiration rate with increasing temperature can be partially explained by the increase in water potential gradient (wpg) between the leaf surface and ambient air. Lorenzo-Minguez et al. (1985) reported that transpiration of Schef-
flera increased with increasing wpg. An increase in air temperature without increasing absolute humidity of the air increases the rate of transpiration because of the increase in wpg between leaf surface and surrounding environment (Kramer, 1983).

AA leakage was not different between soil water levels and, hence, the AA leakage data from three soil water levels were pooled for further analysis. In contrast, Leopold et al. (1981) reported that desiccation of cowpea leaf tissues increased the AA leakage in proportion to the level of desiccation. Bewley et al. (1978) also demonstrated that AA leakage increased with the development of water stress.

AA leakage increased as the storage temperature and duration increased (Fig. 3). Plants stored at 30 or 35C showed the highest AA leakage and the rate of AA leakage rapidly increased as storage duration increased. At another temperatures, AA leakage increased at a slower rate over the storage time. Plants stored at 15C showed a higher rate of AA leakage than those at 25, 20, 10, or 5C.

The leakage of solutes and AA from tissues has been used as a measure of membrane damage (Ingram, 1985; McWilliam, 1980). The increase in AA leakage at high storage temperatures and longer durations indicated that membrane damage occurred at these conditions.

Membrane damage during storage may be a major factor in determining quality and recovery of plants in post-storage. The extent of membrane damage depends on duration of exposure, plant species, and temperature. A sigmoidal response of solute leakage to temperature and exposure time has been reported with Pittosporum tobira root cells (Ingram, 1985). At extremely high temperatures, short exposure can enhance the leakage of solute, indicating severe membrane damage. Once the membranes are severely damaged, the plant may not be able to recover from the storage stress. Storage of plants at optimum temperatures minimizes membrane damage and allows better recovery after storage. Chilling temperatures also have been reported to increase electrolyte leakage from chilling-sensitive species (Wang, 1982). Low storage temperatures, in this study, did not damage the membranes, indicating the tolerance of chrysanthemum to low temperatures.

Soil water content did not affect extension growth during storage, and, therefore, data were pooled to calculate rate of stem elongation. Stem elongation in storage increased as the storage temperature increased up to 25C (Fig. 4). The lowest rate of stem elongation was found at 5C while it was highest at 25C. The rate of stem elongation declined as the time in storage at 15C or above increased, with =30% decrease in the plants stored at these temperatures (data not shown). Stem elongation in storage is undesirable because it is due to etiolation and not to dry matter accumulation. Storage at 5 to 15C would be best for chrysanthemum for short durations because the plants did not elongate in this temperature range.

Measurements of post-storage growth of chrysanthemum plants in the greenhouse indicated that storage at 5, 10, or 15C did not affect subsequent growth rate after any storage duration (Fig. 5). However, plants stored at 20C or above showed a reduction in post-storage growth in the greenhouse. The most rapid reduction occurred with 2 days of storage; thereafter, the growth rate remained steady with increasing storage time. Plants stored at 30C for >4 days or the plants stored for >2 days at 35C did not recover after placing in the greenhouse due to severe foliar damage.

AA leakage and recovery from storage stress were closely related. Plants stored at 30 or 35C had a high AA leakage and did not recover in the greenhouse. Plants that had low AA leakage recovered well in the greenhouse. This result indicates the need for minimum storage in order to obtain the best quality plants after storage at ambient temperatures or above and the importance of low-temperature storage if long-term shipping/storage is necessary. Buck and Blessington (1982) observed that Ficus benjamina and Ficus lyrata stored at 4 or 37C had less recovery after storage as the duration of exposure increased.

Our results indicate that the transpiration rate in storage is mainly determined by temperature; soil water content had little influence. Temperatures >20C caused plant etiolation and loss of quality after removal from storage. Plant injury was severe and recovery from storage stress was slow as storage duration and temperature increased. Plants stored between 5 and 15C gave the best quality and post-storage recovery. Selection of a low temperature is important if long-term shipping/storage is necessary before chrysanthemums reach the consumer. How-
Fig. 5. Effect of storage temperature and duration on average post-storage growth of *Dendranthema × grandiflorum* in greenhouse. Each point indicates the average growth of five replicates and three soil water levels. Plant height increase was measured in the greenhouse for 3 weeks at 4-day intervals.

However, it is important to minimize duration of dark shipping/storage to obtain best quality after storage.

**Literature Cited**


