Environmental Control of Moisture Content and Viability in Schlumbergera truncata (Cactaceae) Pollen

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Abstract. The objective of this study was to determine the effects of temperature, relative humidity (RH), and storage duration on moisture content and viability of Schlumbergera truncata (Haworth) Moran (Thanksgiving cactus) pollen. Pollen viability was assayed by the fluorochromatic reaction (FCR) test, percentage fruit set, and numbers of viable seeds per fruit. Pollen moisture content varied from 10% to 12% until flowers senesced. Pollen viability was 85% on the day of anthesis, remained at 80% from 1 to 3 days after anthesis, and decreased to 65% when flowers began to senesce. When pollen was stored for 4 days at 21 °C, moisture content decreased to 1% for pollen stored at 1% RH but increased to 33% for pollen kept at 100% RH. Fruit and seed yields obtained with pollen stored for 4 days at 21 °C and 15%, 28%, or 52% RH did not differ significantly from those obtained with fresh pollen. Pollen stored for 4 days at 21 °C and 1%, 68%, or 90% RH produced fewer fruit and seeds than fresh pollen. Fruit and seed yields obtained with pollen stored 20 days at 21 °C and 15% or 28% RH were similar to yields obtained with fresh pollen. Pollen with ≤23% moisture content that was stored at –18 °C for ≥24 hours retained its original viability (≈85%) whereas pollen with a high (32% to 34%) moisture content exhibited low (<22%) viability after storage at –18 °C for ≥24 hours. Pollen stored for 140 days at 4 or –18 °C and 13% to 51% RH yielded as many viable seeds per fruit as fresh pollen. Although S. truncata pollen is tricellular, it tolerates desiccation to a low (4%) moisture content and can be stored at temperatures below 0 °C.

Interspecific hybridization is commonly employed in breeding floricultural crops to broaden the genetic base (Ewart, 1981; Langton, 1985). Difficulties may arise when attempting to cross species with asynchronous flowering times. Breeders can either manipulate the environment (photoperiod and/or temperature) to synchronize the flowering times of different species or devise methods for collecting and storing pollen.

The genus Schlumbergera Lemaire contains six species of succulent perennials that are native to southeastern Brazil (Barthlott and Taylor, 1995). Schlumbergera truncata is grown as a flowering potted plant, but the other five Schlumbergera sp. are rare in cultivation and have not been exploited extensively for breeding commercial cultivars (McMillan and Horobin, 1995). A Schlumbergera breeding project has been underway at the University of Massachusetts and one of its principal aims is to broaden the germplasm base via interspecific hybridization. Schlumbergera sp. vary in their flowering times when plants are cultivated in greenhouses under natural photoperiods. For example, at lat. 40° to 50°N, the primary flowering period for Schlumbergera sp. is as follows: September and October for S. orssichiana Barthlott & McMillan; November for S. kautskyi (Horobin & McMillan) N.P. Taylor and S. truncata; February and March for S. russelliana (Hooker) Britton & Rose; and March and April for S. opuntioides (Löfgren & Dusén) D. Hunt and S. microsphaerica (Schumann) Hövel (Barthlott and Rauh, 1977; McMillan and Horobin, 1995; T. Boyle, unpublished data). Time of flowering for S. truncata can be controlled by manipulating photoperiod and temperature (Boyle, 1997), but there have been no reports on environmental regulation of flowering for the other five Schlumbergera sp. Consequently, it would be desirable to store Schlumbergera pollen for crossing species with asynchronous flowering times.

Few studies have examined the storage requirements of cactus pollen. Pollen of Cereus flagelliformis Miller [currently Disocactus flagelliformis (L.) Barthlott] and Cereus grandiflorus [currently Selencereus grandiflorus (L.) Britton & Rose] remained viable for 13 and 17 d (respectively) when stored at 75.5 to 21 °C and 30% relative humidity (RH) (Pfundt, 1909). Cullmann et al. (1987) reported that cactus pollen would remain viable for 1 or 2 weeks if stored at 5 to 10 °C, but no data were presented. Pollen of Hylocereus undatus (Haworth) Britton & Rose and H. polyrhizus (Weber) Britton & Rose lost viability within 1 d when stored at room temperature, but pollen could be stored at –18 to –196 °C for at least 9 months if dehydrated to 5% to 10% moisture content before storage (Metz et al., 2000).

The moisture status of pollen has a significant effect on its longevity (Towell, 1985). Two critical moisture levels exist for pollen. The upper level can be defined as the minimum moisture content at which freezable water is present, and pollen that reaches or exceeds the critical upper moisture level loses viability when exposed to freezing temperatures (Hanna and Towill, 1995). The lower level can be defined as the minimum moisture content at which viability is retained. Critical moisture levels vary among species and among genotypes within species (Hanna and Towill, 1995; Stanley and Linskens, 1974; Towill, 1985) and have not been defined for any Cactaceae taxa (Cullmann et al., 1987; Metz et al., 2000; Pfundt, 1909). The objective of this study was to determine the effects of temperature, RH, and storage duration on the moisture content and viability of S. truncata pollen.

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Materials and Methods

Experiments were conducted from 1996 to 2000 on diploid (2n = 22) S. truncata plants grown in greenhouses at the Univ. of Massachusetts, Amherst (lat. 42°22.5’ N). Plants were grown using standard greenhouse practices and flowering was induced by natural short-day photoperiods occurring from mid-September to April (Boyle, 1997).

**Pollen viability.** Pollen viability was assessed by the fluorochromatic reaction (FCR) test, percentage fruit set, and numbers of viable seeds per fruit. Fluorescein diacetate (2 mg·mL⁻¹ in acetone) was added dropwise to an aqueous solution of 0.8 M sucrose until the solution became persistently cloudy (Heslop-Harrison and Heslop-Harrison, 1970; Heslop-Harrison et al., 1984). Pollen samples were hydrated for 1 h at 100% RH and 21 °C (Shivanna and Heslop-Harrison, 1981) then dispersed on a microscope slide in a drop of stain. Slides were incubated for ≈15 min at 21 °C and examined with a microscope (Standard 16, Carl Zeiss, Inc., Thornwood, N.Y.) equipped with a 100-W high-pressure mercury lamp and a filter set for blue waveband excitation (exciter filter BP 450-490, dichromatic beam splitter FT 510, and barrier filter LP 520). Grains that were brightly fluorescing were scored as a positive FCR whereas grains that were weakly fluorescing or nonfluorescing were scored as a negative FCR. Intracellular accumulation of fluorescein (= positive FCR) indicates the presence of a semipermeable plasmalemma and esterase activity (Heslop-Harrison and Heslop-Harrison, 1970). Fruit and seed set data were collected in three experiments to ascertain if stored pollen could produce viable seed.

**Effect of flower age on moisture content and pollen viability (expt. 1).** Flowers of ‘Dark Marie’ were examined several times daily for five consecutive days and a tag was applied to flowers on the day of anthesis. All flowers were harvested on the fifth day. Flowers were held above preweighed scintillation vials (with the anthers projecting into the vial) and pollen was dislodged from the anthers using a vortex mixer (model 232; Fisher Scientific Co., Pittsburgh, Pa.) set at the highest speed. Pollen was collected in the greenhouse under ambient temperature and RH and vials were capped immediately after collection. Each vial contained pollen from two flowers (=20 mg fresh weight (FW)). After transfer to the laboratory (21 °C and =50% RH), vials were uncapped and immediately weighed on an analytical balance. Vials were placed in a 85 °C forced-air oven for 24 h and reweighed. Moisture content (percentage FW) was determined for each sample. Pollen collection commenced at 1100 HR and all samples were weighed and placed in the oven by 1200 HR. The experiment consisted of five time intervals (0, 1, 2, 3, or 4 d after anthesis) with six replications (vials) per time interval.

Pollen collected 0, 1, 2, 3, or 4 d after anthesis was assayed for viability using the FCR test. Additional pollen samples collected at the same time as those described above were used for FCR tests. Pollen was hydrated and stained as described previously. Five samples were analyzed per time interval and ≥300 grains were scored per sample. FCR tests were conducted on the day of pollen collection between 1330 and 1940 HR. Before hydration, pollen samples were held in capped scintillation vials on a laboratory bench (21 °C and ≈15 μmol·s⁻¹·m⁻² irradiance from cool-white fluorescent lamps).

**Effect of RH on pollen moisture content (expt. 2).** Pollen was collected from five different clones using the method described in Expt. 1. One sample was used to determine pollen moisture content and the remaining samples were stored for 4 d at 1%, 15%, 28%, 48%, 68%, 90%, or 100% RH. RHs were controlled with anhydrous CaSO₄ (1% RH), glycerol–water solutions (15% to 90% RH), or deionized water (100% RH). Glycerol–water solutions were prepared according to Forney and Brandl (1992). Actual RHs generated by the humidity control agents were measured with a Vaisala HMP38 data processor equipped with a HMP35E humidity and temperature probe (Vaisala Inc., Woburn, Mass.).

Pollen samples (50 to 140 mg each) were placed in uncapped scintillation vials which in turn were placed inside screw top bottles (138 cm³ capacity) containing 25 mL of a humidity control agent. Each bottle contained one pollen sample. Pollen was placed in RH treatments within 1 h of collection. Preliminary experiments indicated that RHs remained constant when pollen samples were added to bottles containing 25 mL of a humidity control agent. Bottles were sealed with parafilm (Amer. Natl. Can, Greenwich, Conn.) and placed in a 21 ± 1 °C incubator. On day 4, the moisture content of each sample was determined by the gravimetric method described previously.

**Effect of pollen storage at 21 °C on fruit and seed set (expt. 3).** A composite sample (14.3% moisture content) was prepared using fresh pollen from four different clones. Pollen was stored for 4 or 12 d in a 21 ± 1 °C incubator at RHs ranging from 1% to 90%. RHs were controlled with anhydrous CaSO₄ (1% RH) or glycerol–water solutions (15% to 90% RH). Viability of fresh and stored pollen samples was assayed by fruit and seed set. Stored pollen was rehydrated (100% RH at 21 °C) for ≈1 h before pollinations (Hanna and Towill, 1995). ‘Eva’ was used as the maternal clone for all crosses. Two control treatments were applied: flowers pollinated with fresh (day 0) pollen (to assess the normal amount of fruit and seed set) and nonpollinated flowers (to determine if flowers would set fruit without pollination). Pollen was applied with a fine artist’s brush and each flower was pollinated once. Fifteen flowers were pollinated per treatment.

Fruit were harvested 7 months after pollination (Boyle et al., 1995) and seeds were cleaned by vacuum extraction (Boyle, 1994). Seeds were sown in 10-cm covered glass petri dishes containing one layer of blue blotter paper (Anchor Paper Co., Charlotte, N.C.) moistened with deionized water. Dishes were sealed with parafilm and placed in an incubator with 12 h days/12 h nights of 25 ± 1/22 ± 1 °C and a 12-h photoperiod of 50 ± 8 μmol·s⁻¹·m⁻² provided by cool-white fluorescent lamps.

The number of germinated seeds (those with protruding radicles) were recorded 30 d after sowing. Percentage fruit set [(no. fruit set/no. flowers pollinated) × 100] and number of viable seeds per fruit were calculated for each treatment. Percentage data were analyzed by the chi-square test and seed viability data were analyzed using analysis of variance procedures of SAS (SAS Inst. Inc., Cary, N.C.). Means for number of viable seeds per fruit were separated by the Waller-Duncan k ratio test (k = 100).

**Effect of pollen storage at 21 °C on fruit and seed set (expt. 4).** A composite pollen sample (11.9% moisture content) was collected from three different clones. Pollen was stored for 20 d at 21 °C and either 15% or 28% RH. RHs were controlled with glycerol–water solutions. The experimental protocol and methods used for data collection and statistical analysis were identical to those described in Expt. 3 except that pollen collected 20 d after the experiment commenced from flowers at anthesis (fresh pollen) was used as a control. Fifteen flowers were pollinated per treatment. ‘Dark Marie’ served as the maternal parent for all crosses.

**Effect of pollen moisture content on survival at –18 °C**
Flowers were harvested on the day of anthesis from five different clones and pollen was extracted as described previously. After extraction, pollen (≈800 mg) was taken immediately to the laboratory (21 ± 1.5 °C and 35% ± 2% RH) in a stoppered flask, mixed briefly in a vortex mixer, and spread in a thin layer in a 15-cm glass petri dish. Samples were collected at 0, 0.25, 0.75, 1, 1.5, 2, 3, 4, or 6 h. Eleven samples were collected per time interval. Six samples (≈12 mg each) were placed in preweighed scintillation vials and dried for 24 h at 85 °C. Moisture content was determined as described previously. Five samples (≈3 mg each) were placed in scintillation vials that were capped immediately afterwards and transferred to a −18 °C freezer for ≥24 h. Moisture content decreased to 1% for pollen stored at 1% RH but increased to 33% for pollen maintained at 100% RH. Variation in pollen moisture content at high RHs has been reported previously (Connor and Towill, 1993; Lanner, Connor and Towill (1993) examined hydration/dehydration characteristics of pollen from six species and found that the moisture content equilibrated within 24 h when the RH was <97% whereas pollen stored over water required >24 h to reach equilibrium. Given the fact that S. truncata pollen was kept in controlled RHs for 96 h, it is reasonable to conclude that the pollen moisture contents reported herein are equilibrium values.

Pollen stored at 100% RH exhibited greater variation in moisture content (sd = 2.1) than pollen stored at lower RHs (sd = 0.4 to 1.3). Variation in pollen moisture content at high RHs has been reported previously (Connor and Towill, 1993; Lanner,