Anatomical Features Facilitating Supercooling of the Flower within the Dormant Peach Flower Bud

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Abstract. Excision of the flower from the peach [Prunus persica (L.) Batsch.] flower bud raised the 50% injury temperature of flowers (cooled at 1°C/hour) from -18 and -20°C to -10 and -13°C on two test dates, 26 Feb. 1988 and 5 Dec. 1990, respectively. Ice inoculation of the excised flowers at -3°C further raised the 50% injury temperature to -7 and -8°C for the two dates, respectively. These observations suggest that supercooling is a true mechanism for avoiding freezing injury. Low temperature scanning electron microscopy of freeze fractured cells verified that the flower froze intracellularly, whereas the subtending tissue froze extracellularly. Ice inoculation of the flower and the flower bud axis from which the scales were removed demonstrated that a barrier to ice propagation (effective to -11°C) from the flower bud axis to the flower was present. This barrier may involve the provascular strands and a cell zone at the flower base (BZ) that were devoid of intercellular spaces. These tissues also had smaller cells, smaller vacuoles, greater ratio of cell wall thickness to cell size than tissue just below the BZ, which may result in greater cell rigidity and restriction of extracellular freezing. The cells outside the provascular strands at the base of the flower were also lacking in intercellular space, were smaller in size, and had a higher ratio of cell wall thickness to cell size compared to cells near the base of the scales. In the intact flower buds in which the flowers supercool below -11°C, the presence of the first and second scales was important to full expression of supercooling because their removal raised the supercooling point, whereas the removal of lower scales did not. Sequestration of ice by the first two subtending scales during the early stages of freezing may be important to the creation of a dry region at the flower base that prevents ice propagation into the flower at temperatures below -11°C.

Supercolling of flowers within dormant buds of several genera and woody plant species appears to be a mechanism for avoiding freezing injury. This supercooling is often referred to as deep supercooling as it occurs below the freezing point of the subtending tissue. Deep supercooling has been observed in flower buds of rhododendron spp. (George et al., 1974; Graham and Mullin, 1976), blackberry (Warmund et al., 1988), blueberry (Biermann et al., 1979), flowering dogwood (Sakai, 1979a), forsythia (Nus et al., 1981), Rubus spp. (Warmond et al., 1988), and several Prunus spp. (Burke and Stushnoff, 1978; Quamme, 1974), some conifers (Sakai 1978, 1979b), and in multiple buds of grape (Pierquet and Stushnoff, 1980). Low temperature injury to buds of these species is associated with sudden freezing of a supercooled fraction of water within the bud that was detected by thermal analysis as a low-temperature exotherm (LTE).

The supposition is that the LTEs of buds are produced by intracellular freezing of the flower because of their sudden occurrence and lethal effects but this has not been verified. Extracellular ice crystals have been seen in the tissue subtending the flower in rhododendron (Ishikawa and Sakai, 1981), peach (Dorsey, 1934; Quamme, 1978) and plum (Dorsey and Strausbaugh, 1923) with aid of a light microscope. However, it has not been possible to determine if intracellular ice is present in the flower with aid of a light microscope (Quamme, 1978).

The dormant peach flower bud usually contains a solitary flower. Studies of seasonal changes in supercooling of peach flowers indicated that deep supercooling is maintained throughout the season, from flower bud formation in autumn until bud burst in spring (Quamme, 1974, 1983).

It is not clear what prevents the ice propagation from subtending tissue into the peach flower, but several observations suggest that a structural barrier is involved. Killing the flower by freezing it below the supercooling point (Quamme, 1978) or by heating it to 65°C for 30 min (Ashworth, 1982) raised the supercooling temperature but did not eliminate it. Disruption of the flower base by cutting also raised the supercooling point of the flower from below -20 to -10°C (Quamme, 1978). Ashworth (1984) observed that in the spring when the xylem elements differentiated, supercooling of the flower was lost. He suggested that ice propagates into the flower through the xylem.

The structure of the tissue subtending the flower also appears important to the full development of supercooling (below -10°C) in peach. Ice masses have been observed in the base of the scales and in the pith of the flower bud axis below the flower (Dorsey, 1934; Quamme, 1978). Ice formation in the scales was associated with a decrease in water content at the base of the flower (Quamme, 1978). It was proposed that water movement to the flower bud scales in the initial stages of freezing created a dry region below the flower that prevented ice propagation into the flower, but it is unknown which, if any, of the scales are essential to full supercooling of the flower.

Inoculation of the flower from the exterior of the flower bud with ice through an excision in the scales at different temperatures raised the killing point of the flower from -25 to -8°C (Quamme, 1978). Wounding the flower directly with a needle also destroyed the supercooling capacity (Ashworth, 1982). Quamme (1978)
suggested that the cuticle and epidermis prevented nucleation from the exterior but this hypothesis has not been tested.

The objectives of this study were to determine if freezing injury to the flower within the peach flower bud is avoided by supercooling, if there is an anatomical barrier to ice propagation into the base of the flower, and if the flower bud scales and flower cuticle play a role in maintaining the flower in a supercooled state.

**Materials and Methods**

Effects of ice inoculation on flower injury and supercooling. All studies were conducted during the winter months of 1987 to 1992 on flower buds produced on shoots from mature ‘Redhaven’ *Prunus persica* (L.) Batsch. peach trees in orchards at the Summerland Research Centre. The shoots were collected and used immediately or stored at 0°C or -3°C until used.

Survival of the excised flower, uninoculated and inoculated with ice at -3°C, was compared to the survival of the intact flower within the flower bud. The excision was carried out at 22°C under a stereoscopic microscope, and the tissue was immediately transferred to a petri dish containing moist filter paper. The whole flower bud was cut to include a piece of the shoot attached to the base. The flower was excised by cutting away the scales to expose the flower and then cutting the flower just above the flower bud axis with a sharp scalpel. Ice inoculation of the excised flower was carried out by cooling it to -3°C in a cold room, placing a small droplet of ice water on the surface of the tissue, and introducing a piece of ice into the water droplet with a pair of tweezers. Whole flower buds were cooled to -3°C, and the base of the flower bud axis was inoculated by dipping the cut surface into ice water and allowing the water to freeze before further cooling. A sample (10) of each inoculated, and uninoculated, excised flowers and the whole flower buds were each placed in separate metal containers at -3°C. A set of samples was prepared for each test temperature. The sets of samples were cooled in the cold room to preselected temperatures 2.5°C apart in the range of injury at 1°C/h. At each test temperature, a set of samples was removed from the freezer, thawed at 0°C, and then incubated over moist filter paper in petri dishes for three days at 3°C. After incubation, the flowers and flower buds were sectioned and observed for browning with a stereoscopic microscope. Flowers that were brown were classified as injured, whereas those that remained normal green were classified as uninjured. The experiment was repeated three times with material collected on each of two dates, 26 Feb. 1988 and 5 Dec. 1990. Micrographs of injured flowers in the bud were made to show the location of injury.

To determine whether there was a barrier to the ice propagation from the flower bud axis into the flower, the flower was examined for the presence of supercooling after the flower bud axis, from which the scales were removed, was inoculated with ice. Exotherm analysis was also performed on inoculated and uninoculated excised flowers, inoculated flowers on flower bud axes, and inoculated whole flower buds as controls. Excised flowers were inoculated on the uncut and the cut surface where the excision was made to determine whether the cuticle and epidermis restricted the ice propagation. Excision and ice inoculation were carried out as described previously. The flower buds were collected for this study on 24 Feb. 1988.

Exotherm analysis of the flower buds and flower bud parts was conducted by noting sudden deflections observed on temperature recordings during cooling. The method used was similar to that described previously (Quamme, 1974) except that the thermocouple (20 gauge, copper-constantan) was attached with silicon grease to the surface of the base of the flower bud or that of the flower. After attachment to the thermocouple, the tissue was placed in a Dewar flask at -3°C and cooled to below -30°C at 5°C/h. The tissue temperature was determined with a 3974A data logger interfaced with a computer (model 300; Hewlett-Packard, Rockville, Md.).

Effect of scale removal on LTE. The importance of the flower bud scales to supercooling of the flower was determined by scale removal from the flower bud base. The LTE temperature was measured on buds from which varying numbers of scales were removed. The number of scales remaining on the bud axis was counted after the exotherm analysis was run. The collection date for the flower buds used in this study was 7 Feb. 1989. The sample size was 14 to 18, except for flower buds with 5 scales, for which exotherms were determined on 3 flower buds. Analysis of variance was performed on the data using the General Linear Models procedure of SAS Institute (1985).

Low-temperature scanning electron microscopy (LTSEM) of freeze-fractured floral parts. To confirm if cells within the flower froze extracellularly or intracellularly, flower buds (5) were frozen intact on the shoot to -23°C (below the LTE, -18°C) at a cooling rate of 1.2°C/h, rewarmed to a holding temperature of -12°C at a rate of 5°C/h. Flowers and flower bud axes with the scales removed were excised from flower buds, and mounted on a precooled cryo-specimen stub at the holding temperature with the aid of a binocular microscope. The sample on the cryo-specimen stub was quench-frozen in supercooled liquid nitrogen at -209°C with a cryo-apparatus (K-1250; Emitech, Ashford, UK). After quench-freezing, the sample was fractured and sputter-coated in the cryo-apparatus with gold-192C (Dubochet et al., 1985; Steinbrecht and Zierold, 1987). The gold-coated fractured floral parts on the cryo-specimen stub were transferred and maintained at -123°C during examination (from 10 to 25 kV) with scanning electron microscope (SEM) (JSM-840A; JEOL Ltd., Tokyo) fitted with an Emitech LTSEM cryo-stage. Similar observations were made on excised flowers that were quench-frozen from room temperature.

Peach flower bud anatomy. For SEM, the flower buds were stored in the buffered glutaraldehyde solution (5%, 0.07 M, pH 6.8), cut longitudinally along the flower axis and then post-fixed in 2% osmium tetroxide. These fixed samples were dehydrated in ethanol, critical-point dried using CO2 with a CPD 020 critical-point dryer, (Balzer’s Fürstentum, Liechtenstein) mounted on aluminum stubs, and sputter-coated with gold using a sputter-coater (Desk II; Denton Vacuum Inc., Cherry Hill, N.J.). Samples were examined with the SEM, operated at 25 kV.

Flower buds were collected for anatomical investigation on 6 Jan. 1988, preserved with 5% glutaraldehyde in Sorenson’s buffer (0.07 M, pH 6.8), and then vacuum infiltrated with a second change of the same buffered glutaraldehyde solution (Sabatini et al., 1963). The samples were washed in Sorenson’s buffer (six times for 0.5 h each) and stored at 2°C in the same buffer for several days. Several flower buds were dissected in buffered glutaraldehyde to include the flower and flower bud base, and these parts were post-fixed in 2% osmium tetroxide in Sorenson’s buffer at 4°C (Hayat, 1986). After fixation in 2% osmium tetroxide, the flower bud parts were dehydrated in an ethanol gradient, vacuum infiltrated, using propylene oxide as an intermediate fluid, and then embedded in JEMBED 812 epoxy resin for sectioning (Luft, 1961). Longitudinal, 2-µm-thick (semi-thick) sections of the base of a flower and the upper flower bud axis were cut with a microtome (Ultracut E; Reichert-Jung, Vienna) using a glass knife. Serial semi-thick sections (with a glass knife) and serial ultra-thin sections (50-60 nm, with a diamond knife) were cut from another flower bud using the same ultramicrotome.

The parts of the flower that were sectioned are indicated in Fig. 1.
Fig. 1. A scanning electron micrograph image of a longitudinal section of a peach flower bud shows the flower (F), the flower bud scales (Sc), and the flower bud axis (FBA). The line at the base of the flower bud indicates the axis where cross sections were cut and the line from the flower to the second scale indicates the axis where longitudinal sections were made. Bar = 1 mm.

The serial sectioning began about 0.4 mm below the first scale subtending the flower and ended just below the ovary. This sectioned region was 1.2 mm long. Semi-thick sections were also cut with a glass knife longitudinally through the base of the first and second flower bud scales and the flower (Fig. 1).

Semi-thick sections were stained with toluidine blue and basic fuchsin in sodium carbonate solution (Lewis and Knight, 1977) and examined with a photomicroscope (Carl Zeiss, Oberkocken, Germany). The ultra-thin serial sections were mounted on copper grids, stained with uranyl acetate followed by Reynold’s lead citrate, and viewed with transmission electron microscope TEM; (JEOL 100 CX II) at 80 kV.

Cell size measurements were made from micrographs of the semi-thick serial cross sections through the flower base (0.1 mm apart) and longitudinal sections at the base of the first two subtending scales and the flower (four sections). Cell diameter (30 cells) was measured in each of the series of cross or longitudinal sections and expressed as cell cross sectional area. The cell wall thickness of the same cells was measured from micrographs of the sections. The ratio of cell wall thickness to cell cross sectional area was also determined. The percentage of intercellular space was determined by area weight ratios from the micrographs.

Results

Effects of ice inoculation on flower injury and supercooling.

The excised flowers, both inoculated and uninoculated with ice, were injured at a higher temperature than intact flowers in the flower bud (Fig. 2 A and B). 50% injury determined from the spline interpolation occurred to the inoculated excised flower at –8 and –7°C, to the uninoculated excised flower at –10 and –13°C, and to the intact flower in the flower bud at, –18 and –20°C on the first and second collection dates, respectively. The subtending tissue of whole buds did not show browning at all test temperatures.

A LTE occurred on the time-temperature profile of the uninoculated excised flower (Fig. 3A) as it did on that of the whole flower bud (Fig. 3E). The mean LTE temperature determined on the excised flower was 7°C higher than that of the inoculated whole flower bud (Fig. 3A and F). The increase in LTE temperature of the excised flower coincided with the higher killing temperature that occurred in the excised flowers in the first experiment. A slight deflection was sometimes present on the time-temperature profile of the ice inoculated flower at a temperature just below that at which cooling was begun, but it was neither as sharp nor as high as the deflections on the time-temperature profile of the excised uninoculated flower. Inoculation of the excised flower with ice placed on the cut surface of the flower base (Fig. 3B) was as effective in preventing supercooling as inoculation of the excised flower with ice placed on the uncut exterior of the flower (Fig. 3C). The LTE was detected in the intact flower on the inoculated flower bud axis from which the scales had been removed.

Fig. 2. Freezing injury to flowers excised from flower buds and then inoculated with ice at –3°C before freezing ( ), uninoculated flowers excised from flower buds ( ), and flowers intact in the inoculated flower bud ( ) on two collection dates, (A) 26 Feb. 1988 and (B) 5 Dec. 1990. Each value represents the mean ± se of three experiments. Lines were drawn through the points using spline interpolation (SAS/GRAPH, SAS Institute, 1985). The dashed reference line indicates 50% level of injury to the flowers.

removed (Fig. 3D) at the same temperature as that of the uninoculated excised flower (Fig. 3A). Inoculating the flower on the bud axis at the surface of the flower and flower bud axis eliminated the LTE (Fig. 3E) as did with the excised flower (Fig. 3B).

The effect of scale removal on the LTE. Progressive removal of the scales from the base of the flower bud to the second scale below the flower did not affect the LTE temperature (Table 1). Removal of all scales, except for the first scale subtending the flower, significantly raised the LTE temperature from –18°C to –14°C. A significant further rise in LTE temperature did not occur with removal all scales including the first scale subtending the flower.

LTSEM observations on freeze-fractured floral parts. Cell appearance of freeze-fractured flowers subjected to two freezing treatments, quench-freezing from room temperature and quench-freezing from –12°C after freezing to –23°C followed by rewarming, was similar as is shown for floral tube cells (Fig. 4 A and B). The floral tube cells were typical of both treatments and appeared to be neither shrunken nor deformed. Some empty intercellular spaces were observed between the floral tube cells. The cell vacuoles from Table 1. The effect of scale removal on the low temperature exotherm (LTE) temperature.

<table>
<thead>
<tr>
<th>No. of scales remaining on the flower bud axis</th>
<th>LTE (°C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>–12.8 a</td>
</tr>
<tr>
<td>1</td>
<td>–13.7 a</td>
</tr>
<tr>
<td>2</td>
<td>–18.3 b</td>
</tr>
<tr>
<td>3</td>
<td>–17.2 b</td>
</tr>
<tr>
<td>4</td>
<td>–19.2 b</td>
</tr>
<tr>
<td>5</td>
<td>–18.5 b</td>
</tr>
<tr>
<td>Whole flower bud (5–8 scales)</td>
<td>–18.7</td>
</tr>
<tr>
<td>SE</td>
<td>±1.0</td>
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</tbody>
</table>

*Means (n = 3 to 18) within column followed by the same letter are not significantly different by Duncan’s new multiple range test at P ≤ 0.05.
FIG. 4. Low temperature scanning electron micrographs of the fracture surfaces of peach flower buds that were quench-frozen in supercooled liquid nitrogen. Appearance of freeze-fractured cells in the floral tube that were (A) quench-frozen from room temperature and (B) quench-frozen from the -12C holding temperature after freezing to -23C (below the low temperature exotherm), at 1.2C/h. Ice is absent in the intercellular spaces. (C) Extracellular ice in freeze fracture tissue from the base of a flower bud scale after quench-freezing from the -12C holding temperature. Abbreviations: FC = freeze fractured cells, IS = intercellular space, and IC = extracellular. Bars = 20 µm.

Discussion

The nucleation temperature of supercooled, excised flowers was higher than that of intact flowers within the flower bud, which agrees with a previous observation (Quamme, 1978). Initiation of ice formation in the excised flower by ice inoculation further raised the temperature at which injury occurred. The flower appeared to lack resistance to freezing and to be dependent on supercooling for survival. Ishikawa and Sakai (1981) also found that rhododendron flowers were injured by ice inoculation at temperatures just below freezing and suggested that supercooling was a mechanism for avoiding freezing injury. In both species supercooling of the flower persisted from autumn through winter until bud break (Ishikawa and Sakai, 1981; Quamme, 1983).

The LTSEM observations that extracellular ice was absent in the flower, and that flower cells did not appear to be shrunk after freezing slowly to below the LTE temperature followed by quench-freezing in supercooled liquid nitrogen substantiates the hypothesis that the flower freezes intracellularly. Furthermore, the plasma membrane did not appear to detach from the cell wall to allow ice to form between the cell wall and the plasma membrane.

Rewarming the flower to -12C during transfer to the freeze-fracture apparatus after freezing to below the LTE was not thought to affect the presence of ice in the flower tissue because its thawing point was previously determined to occur at -2 to -3C (Quamme, 1983). Extracellular ice was present in tissue subtending the flower that was frozen slowly to below the LTE followed by quench-freezing.

The lack of a distinct exotherm on the time-temperature profile of the inoculated excised flower indicated that inoculation suppressed, if not eliminated, supercooling. It was not determined whether injury to the inoculated bud occurred as a result of intracellular or extracellular freezing. If intracellular freezing occurred, it progressed slowly through the tissue and did not produce a sudden deflection on the time-temperature profile as occurred in the uninoculated flower. Small exotherms produced in the range of injury in some flowers indicated that progressive intracellular freezing probably occurred.

The flower on the flower bud axis from which the scales were removed exhibited a LTE if only the flower bud axis was ice inoculated. If both the flower and the flower bud axis were ice inoculated, no LTE was present. This indicates existence of a barrier to ice propagation into the flower from the flower bud axis. However, the temperature to which the flower supercooled on the flower bud axis without scales was higher (-11C) than that of the flower within the intact flower bud with scales (-18C), which suggests that the scales are required for the barrier to be fully effective. The barrier to the ice propagation appears to be a morphological feature because supercooling was not eliminated in freeze-killed (Quamme, 1978) or heat-treated flower buds (Ashworth, 1984).

The barrier to ice propagation into the flower may involve the basal structure of the flower. The BZ and adjoining vascular...
Fig. 5. (A) A longitudinal section of the flower bud that has been frozen below the low temperature exotherm, incubated at room temperature for 3 days, and cut to show the flower and subtending tissue. The injured flower became brown (dark colored in the micrograph), whereas the scales and flower bud axis remained normal green (light colored in the micrograph). (B) The provascular strands and basal zone are shown in a longitudinal section of the flower base that was stained with toluidine and azure blue. (C) A cross section stained with toluidine and azure blue through the basal zone into the provascular strands shows the absence of intercellular spaces and small cell size of these tissues. (D) The vacuoles of cells in the basal zone appeared small and numerous in a transmission electron micrograph. Cell nuclei appeared to be prominent. (E) The presence of intercellular spaces is seen in the toluidine and azure blue stained cross section of the bud axis below the basal zone of the flower. Large vacuoles were observed in this tissue. (F) A cross section of the flower above the basal zone stained with toluidine and azure blue shows the provascular strands and parenchyma cells inside the strands at the base of the flower. The parenchyma cells inside the strands were observed to be large and contain large vacuoles. Abbreviations: F = flower, Sc = scale, FBA = flower bud axis, PV = provascular strands, BZ = basal zone, V = vacuole, NU = cell nucleus, and IS = intercellular space. (A) Bar = 1 mm. (B, C, E, and F) Bar = 20 µm. (D) Bar = 5 µm.
strands were devoid of intercellular spaces, and had smaller cells with smaller vacuoles and thicker walls than the subtending tissue. The tissue to the exterior of the provascular strands near the base of the flower had similar cell characteristics to that of BZ. In flower buds frozen to just below the LTE, the boundary of injury to the flower appeared to coincide with the BZ.

The presence of the BZ in peach buds may be similar to an anatomical feature found in some conifer buds that supercool. The buds of *Abies* and *Larix* spp. that supercool have a crown of medullary tissue that is absent in other conifers. (Sakai, 1978, 1979b). Removal of the medullary crown tissue from the excised shoot primordium of *Abies homolepis* raised the supercooling point from –25 to –15°C (Sakai, 1979b).

In *Prunus* spp., an abscission zone is present at the base of the pedicel at the point of stem attachment in the fruit (Esau, 1953). The BZ may correspond to the early development of an abscission zone but it was not verified in this study.

The absence of intercellular spaces may be critical to preventing ice propagation into the flower. Acid fuchsin (Quamme, 1978) and azosulfamide (Ashworth, 1982) uptake did not penetrate the flower which indicates a fine pore structure. Water held in fine pores does not freeze as bulk water but tends to have a lower melting point and vapor pressure depending on pore size. The critical pore size for this effect to occur appears to be 6 nm. Cell wall pore size was not determined, but in absence of intercellular spaces, it could be in this size range (Ashworth and Abeles, 1984).

The small cell size, lack of vacuoles, and relatively thick cell wall structure may increase the resistance of the cells at the base of the flower to cell wall deformation that results from extracellular freezing. Rigid cell structure may be critical to restricting extracellular freezing and ice propagation along cell walls.

The presence of a single LTE indicates that ice propagates suddenly throughout the flower when freezing begins. LTSEM conducted in this study confirms the occurrence of intracellular freezing. Previously, flower tissue at –15°C was observed to turn from a translucent, greenish to whitish color when inoculated with ice. The change in flower coloration, which spread from the point of inoculation, was possibly caused by intracellular freezing (Quamme, 1978). Once intracellular freezing begins, it may proceed from cell to cell through the pit structure.
Survival of peach flower buds by freezing avoidance is noteworthy because it is not clear what evolutionary advantage this adaptation would have. Possibly, it is a feature associated with early flower development in the spring, and subsequent early fruit development in summer, (Quamme, 1991) but this is speculative.

Inoculation of the flower with ice on the surface of the cuticle was as effective in preventing supercooling as inoculation of the flower with ice at the cut surface. Therefore, the cuticle and epidermis appeared to be a weak barrier to the ice propagation into the flower from the exterior. The scales enclosing the flower may act to exclude water from the surface of the flower that could inoculate the flower during freezing. Otherwise, freezing rain or dew drops would inoculate the flower.

The overwintering peach flower appears to survive by a mechanism of freezing avoidance. This mechanism is complex and involves a combination of physiological and anatomical features that segregate ice formation in the flower bud and restrict its spread into the flower. In this study, observations indicate that provascular tissue and the BZ at the flower base may be a barrier to ice propagation into the flower and that the first and second scales subtending the flower are important to the full expression of supercooling because they may be sites for sequestering ice.

**Literature Cited**


Fig. 8. The cell cross sectional area (CCSA), cell wall thickness (CWT), ratio of cell wall and thickness to cell diameter (RCC), and % intercellular spaces (IS) measured in longitudinal sections at the periphery of the flower through to the second scale. Measurements were made in the flower and near the base of the first and second scale as indicated in the adjacent diagram. Vertical bars denote ± SE (n = 30).


