Water Relation Parameters of the Cortex Tissue of Fresh ‘Fuji’ Apple Fruit Determined by Centrifugation
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Additional index words. Malus ×domestica, apple fruit, pressure–volume curve, Höfler diagram, centrifugal force, water potential

Abstract. This experiment was carried out to obtain a pressure–volume (P–V) curve and Höfler diagram of the cortex tissue of fresh ‘Fuji’ apple fruit (Malus ×domestica Borkh.) with a novel centrifuge method. Based on the P–V curve and Höfler diagram, several water relation parameters of fruit cortex were determined and the interrelationship of these parameters was established. Turgor loss point (TLP) occurred at –1.74 MPa and 73.7% of relative water content (RWC). At full hydration, osmotic potential (φw) was –1.30 MPa and symplastic water accounted for 86.8% of RWC. Bulk elastic modulus decreased linearly by 28% as pressure potential declined from 1.30 MPa at full hydration to zero at the TLP. This centrifuge technique can provide a simple and efficient way to determine water relation parameters of fleshy fruits.

The P–V curve has been used to estimate symplastic water volume, water potential, turgor loss point, and elastic modulus of plant cells, tissue, and organs (Holbrook and Sinclair, 1992; Richter, 1978; Scholander et al., 1965; Tyree and Hammel, 1972). It is considered to be genetically determined (Lenz et al., 2006) and reflects drought-response properties of plant species (Dickio et al., 2003; Joly and Zaer, 1987; Saito and Terashima, 2004). From analysis of the P–V curve, a Höfler diagram can be derived to display the components of water potential as a function of relative water content of living plant materials (Joly and Zaer, 1987; Nobel, 2005; Richter, 1978). Traditionally, a pressure chamber has been used to obtain the P–V curve and Höfler diagram of leaves (Dickio et al., 2003; Jones and Higgs, 1979; Ruiz-Sánchez et al., 1993; Saito and Terashima, 2004) and stems (Joly and Zaer, 1987; Tyree and Hammel, 1972). However, little information is available regarding the P–V curve and Höfler diagram of fruit because of the difficulty in measuring fruit water relation parameters by the pressure chamber technique (Gelly et al., 2004).

It is generally accepted that fruit water status is closely related to physiological processes, which control fruit growth and development and plays an important role in fruit quality and responses to environmental stresses (Behboudian et al., 1994; Mills et al., 1996, 1997; Mpelasoka et al., 2001; Steudle and Wiencek, 1985; Wada et al., 2008; Yamada et al., 2005). Nevertheless, monitoring fruit water status remains a challenge using a pressure chamber technique (Gelly et al., 2004) or pressure probe (Steudle and Wiencek, 1985). The psychrometer technique used to measure fruit water parameters is time-consuming to reach equilibrium and prone to produce errors (Behboudian et al., 1994; Gelly et al., 2004; Mills et al., 1996, 1997; Mpelasoka et al., 2001; Steudle and Wiencek, 1985; Wada et al., 2008; Yamada et al., 2005). Therefore, the run time of centrifugation was set as follows: 60 min for RCF = 1,020 g·cm⁻², 120 min for RCF = 10,208 g·cm⁻², and 180 min for RCF = 15,271 g·cm⁻² (Fig. 1). Therefore, the time of centrifugation was set as follows: 60 min for RCF = 1,020 g·cm⁻², 120 min for RCF = 10,208 g·cm⁻², and 180 min for RCF = 15,271 g·cm⁻², or less. Average distribution of centrifuge-created water potential (ψ) along the fruit sample of thickness (L, cm) subjected to an angular velocity was calculated according to Reatto et al. (2008) as follows:

\[ \psi = \frac{k \omega^2 L}{6 g} (L - 3r) \]  

where \( k \) is a constant (9.807 MPa·cm⁻¹), \( g \) is the acceleration of gravity (981 cm·s⁻²), \( \omega \) is angular velocity (rad/s), and \( r \) is radius of centrifuge rotor (cm).
solids concentration (SSC) of apple fruit juice expressed at full hydration or extracted during centrifugation were measured with a psychrometer (Wescor Inc., Logan, UT) and a hand refractometer (Chengdu Optics Inc., Sichuan, China), respectively. After being subjected to each level of centrifugation for a predetermined time, the total volume of water extracted was measured. The centrifugation process was repeated four times and four samples were centrifuged at one time. Altogether, data of 16 samples were pooled to calculate the mean and SE for each point. The negative reciprocal of average water potential was plotted against the volume of water extracted to obtain a P-V curve. TLP on the P-V curve was determined using the PVA method given by Schulte and Hinckley (1985).

On completion of the centrifugation process, fruit samples were dried in an oven at 85 °C for 24 h and their weight reached a constant level. $R^*$ of the apple fruit samples after each step of centrifugation was calculated using the equation as given by Kramer (1983):

$$R^* = \frac{W - DW}{SW - DW} \times 100$$  \hspace{1cm} (2)

where $W =$ weight measured after each centrifugation step, $DW =$ weight of the oven-dried fruit sample, and $SW =$ weight of fruit sample at full hydration.

Average pressure potential ($\psi_p$) was calculated as the difference between average water potential ($\psi_w$) and average $\psi_S$ derived from analysis of the P-V curve (Richter, 1978) as follows:

$$\psi_p = \psi_w - \psi_S$$  \hspace{1cm} (3)

The bulk elastic modulus ($\varepsilon$) of apple cortex tissue was calculated according to Schulte and Hinckley (1985) as follows:

$$\varepsilon = \frac{d \psi_p}{d R^*}$$  \hspace{1cm} (4)

**Results**

A progressive increase in the SSC of the extracted juice with maximum relative centrifugal force was detected during centrifugation (Fig. 2). This increase presumably resulted from soluble solids leakage and affected the average water potential. This effect was corrected by adding the lost $\psi_S$ back to each data point of the water potential.

Both P-V curves of apple cortex tissue obtained before and after correction (Fig. 3) followed the typical pattern of P-V curves of other plant organs measured by the pressure chamber technique (Joly and Zaerr, 1987; Jones and Higgs, 1979; Ruiz-Sánchez et al., 1993; Tyree and Hammel, 1972). It was characterized by two parts, the nonlinear phase and the linear phase with a breakpoint between them. This inflection point represents the turgor loss point. Based on the P-V curve after correction (with solid symbols), the estimated TLP occurred at the average water potential of ~1.74 MPa and 73.7% of relative water content. Extrapolation of the linear part of the P-V curve (dashed line in Fig. 3) intersects the y-axis at 0.7663 MPa, representing the turgor loss point.

Fig. 1. Run time of centrifuge rotation versus the accumulative quantity of water extracted from apple cortex samples subjected to maximum relative centrifugal force of 1020 g, 10,208 g, and 15,271 g. Each data point represents the mean ± SE of four measurements.

Fig. 2. Relationship of soluble solids concentration (SSC) of extracted juice with maximum relative centrifugal force (RCF) during centrifugation of apple cortex samples [$y = 1.4166 \ln(x) - 3.3306$, $r^2 = 0.9696$, $P < 0.001$]. Each data point represents the mean ± SE of 16 measurements.

Fig. 3. Relationships between the negative reciprocal of average water potential and the volume of water extracted from apple cortex tissue before and after correction. Solid lines indicate nonlinear parts of the curves in the region of positive pressure potential. The dotted line and dashed line below represent the extrapolated linear parts of the original curve ($y = -0.9691 + 0.7661$, $r^2 = 0.9313$, $P < 0.001$) and the corrected curve ($y = -0.959 + 0.8664$, $r^2 = 0.9122$, respectively). Each original data point ($\Delta$) represents the mean ± SE of 16 measurements.
approaching the TLP. However, the pressure potential curve obtained was not obviously concave toward the x-axis as previously reported for Douglas fir twigs (Joly and Zaerr, 1987) and cyclamen leaves (Richter, 1978).

From full hydration to the TLP, \( \psi_w \) of apple cortex decreased from zero to \(-1.74 \) MPa. Specific water capacitance (change in \( R^* \) per change in \( \psi_w \)) was calculated as 15.1% MPa\(^{-1}\), indicating that when there was a 1-MPa reduction of \( \psi_w \), apple fruit would release 15% of relative water content. Below the TLP, the water potential of flaccid apple cortex likely became identical to the \( \psi_s \).

The plot of apple cortex \( \psi_s \) versus \( R^* \) from full hydration to the TLP exhibited a linear relationship. The average osmotic potential of fruit cortex tissue dropped by 34% from \(-1.30 \) MPa at full hydration to \(-1.74 \) MPa at the TLP.

The linear relationship between the bulk elastic modulus (\( E \)) and average pressure potential is shown in Figure 5. The value of \( E \) decreased by 28% from 5.02 MPa to 3.61 MPa as the pressure potential declined from 1.30 MPa at full hydration to zero at the TLP.

**Discussion**

P-V curves have been found for vegetative tissue and organs of higher plants (Holbrook and Sinclair, 1992; Jones and Higgs, 1979; Lenz et al., 2006; Ruiz-Sánchez et al., 1993; Saito and Terashima, 2004). However, to the best of our knowledge, the present study is the first report of the P-V curve for apple fruit. The overall pattern of the fruit P-V curve obtained by the centrifugation method was similar to those of plant leaves and stems obtained with the pressure chamber technique (Dichio et al., 2003; Holbrook and Sinclair, 1992; Lenz et al., 2006; Ruiz-Sánchez et al., 1993), displaying a nonlinear region and then a linear region after considerable water was extracted.

The increase in the SSC of extracted juice suggested that the plasma membrane of apple fruit cells was not totally impermeable to some solutes like in other cases discussed by Tyree and Hammel (1972). Yamaki and Ito (1992) reported that the concentration of total sugars in apoplastic water increased threefold and permeability to sugars across the plasma membrane rose five- to fivefold when they were compared between immature and mature apple fruit. Appoplastic solute accumulation was also observed in grape berries during growth and development (Wada et al., 2008). Thus, the high SSC in juice extracted by centrifugation might be attributed to the sensing stage of fruit used in this study. Nevertheless, the influence of soluble solid leakage on P-V curve could be eliminated if the water potential data were appropriately corrected.

From analysis of the corrected P-V curve, water energy status of fleshy fruit could be studied. The average \( \psi_s \) (\(-1.30 \) MPa) of apple fruit cortex at full hydration was consistent with the values of expressed apple juice measured with a psychrometer (\(-1.28 \) MPa) or calculated (\(-1.24 \) MPa) from the SSC (13.7% Brix) using the van’t Hoff equation (Kramer, 1983). The fruit \( \psi_s \) at full hydration was comparable with that of oak leaves (Saito and Terashima, 2004) but higher than the values reported for leaves of apple, ranging from \(-2.2 \) to \(-2.5 \) MPa (Jones and Higgs, 1979), almond (Ruiz-Sánchez et al., 1993), olive (Dichio et al., 2003), and other woody plant species (Lenz et al., 2006). This parameter, along with the \( \psi_s \) at the TLP, is usually used as an important criterion for determining osmotic adjustment of plant vegetative organs in response to water deficit (Dichio et al., 2003; Joly and Zaerr, 1987; Ruiz-Sánchez et al., 1993; Saito and Terashima, 2004). Osmotic adjustment of apple fruit was observed in the early growth season by measuring the fruit \( \psi_s \) with a hygrometer under different water

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**Fig. 4.** A Höffler diagram derived from analysis of the pressure–volume (P-V) curve in Figure 3 showing relationships of average pressure potential \( \psi_p \) \( (y = -0.0000225x^2 + 0.0458x - 3.4683, r^2 = 0.9913, P < 0.001) \), water potential \( \psi_w \) \( (y = -0.000114x^2 + 0.0485x - 5.951, r^2 = 0.9947, P < 0.001) \), and osmotic potential \( \psi_o \) \( (y = 0.0000225x^2 + 0.0458x - 3.4683, r^2 = 0.9873, P < 0.001) \) with the relative water content of apple cortex tissue.

**Fig. 5.** Relationship between bulk elastic modulus and average pressure potential of apple cortex tissue. Bulk elastic modulus was calculated using Eq. [4] and data in Figure 4. \( (y = 1.0765x + 3.5572, r^2 = 0.9916, P < 0.001) \).
conditions at predawn, midday, and dusk (Mills et al., 1997). We expect that using the centrifuge method to determine fruit water relation parameters can be of great help in studying osmotic adjustment of fruit.

The average pressure potential (1.3 MPa) of apple fruit cortex at full hydration was similar to the maximum pressure potential (1.2 MPa) of apple cortex cells measured with a pressure probe (Steudle and Wiencke, 1985). Much lower pressure potential of apple fruit, determined before harvest, was reported in the literature (Mills et al., 1996, 1997; Mpelasoka et al., 2001; Yamada et al., 2005). This discrepancy was probably because the fruit samples in those experiments did not reach full hydration. In addition, it could also be attributed to different apple cultivars and measuring techniques used in those studies.

The relative water content of apple cortex at the TLP was lower than the values reported for apple shoots (Diaz-Perez et al., 1995), olive leaves (Dichio et al., 2003), and palm leaves (Holbrook and Sinclair, 1992). Richter (1978) proposed that cells with rigid walls have higher relative water content at the TLP than those with elastic walls. From an anatomical aspect, apple cortex tissue consists primarily of parenchyma cells with large vacuoles and thin cell walls except for a few vascular bundles embedded among the cortex (Esau, 1977). The low $R_*$ of apple cortex tissue at the TLP may result from the thin, elastic walls of apple fruit parenchyma cells.

The corrected value of $\psi_S$ (~1.74 MPa) at the TLP agreed with the $\psi_S$ value (~1.72 MPa) calculated from the SSC of fruit samples using the van’t Hoff relation. Lower $\psi_S$ (more negative values) was reported for ‘Fuji’ apple fruit (Jones and Higgs, 1979) compared with that of apple leaves (Jones and Higgs, 1979) implies that attached fruit may also act as a water storage organ and play an important role in keeping the daily water balance of the tree-like stem of palm reported by Holbrook and Sinclair (1992). This point was supported by the observation of reversal water flow out of attached apple fruit to other parts of tree (Lang, 1990; Tromp, 1984). Our results suggested that if fruit water potential dropped by 1.0 MPa, an apple of 250 g could provide 34 g of water for 28 min of leaf transpiration at a rate of 20 mg m$^{-2}$ s$^{-1}$.

The bulk elastic modulus of apple cortex tissue was comparable with the values of olive leaves estimated by the pressure chamber technique (Dichio et al., 2003). Dependence of the bulk elastic modulus on pressure potential was also observed in apple fruit cells (Steudle and Wiencke, 1985) and olive leaves (Dichio et al., 2003). Steudle and Wiencke (1985) reported that the elastic modulus of apple fruit cells was a function of cell size and larger cells showed a higher elastic modulus. Variation in $S$ values could be the result of the difference in cellular characteristics between plant materials. Steudle and Wiencke (1985) and the experimental methods used as well (Schulte and Hinckley, 1985).

It is well known that elastic modulus is closely related to the rigidity of plant cell walls (Nobles, 2005). Plant cells with rigid walls have a higher elastic modulus than those with elastic walls (Kramer, 1983). A lower elastic modulus of apple probably means higher elasticity of the fruit. An increase in apple tissue elastic modulus at high pressure potential may contribute to the firmness of apple fruit flesh before any water loss during storage.

Psychrometer has been previously used to measure water $w_*$ and to determine pressure potential of pear (Behboudian et al., 1994) and apple (Mills et al., 1996; Mpelasoka et al., 2001; Yamada et al., 2005) fruits. However, this traditional method can only at a given time provide limited snapshot measurements to study growth and development of fruit related to water status. The H"ofler diagram, derived from analysis of the P-V curve, presents the components of fruit water potential in relation to $R_*$. Therefore, dynamic changes in water energy status of attached fruit may be able to be determined in situ by continuously monitoring fruit water content.

In summary, the centrifuge method used here to obtain the P-V curve of apple fruit may be used to determine fruit water status at different developmental stages and under various environmental conditions. This highly efficient and less time-consuming technique allows new to further study the roles that water plays in the growth and development of apple and other fruits in laboratories.

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