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Evolution of cherries texture in brine: Impact of harvest conditions during long-time storage

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Abstract

Texture is a primary quality attribute of brined sweet cherries (Prunus avium L.) and its preservation is a major objective for candying industry. In order to identify the harvest factors influencing textural changes during long period brine storage, different itineraries were applied: harvest at two different maturity stages, treatment or not with ethephon, manual or mechanical harvest, removal or not of peduncles. The cherries were immersed in brine and examined over a 12-months period for firmness, calcium and total soluble solids diffusion and cytohistological remodelling. Mechanical harvesting, harvest at late maturity stage and storage with peduncle decreased firmness while ethephon treatment had no effect. However, only presence or absence of peduncles influenced salt and total soluble solids diffusion, suggesting that peduncle removal promotes osmotic exchanges.

Brine storage led to a texture gain in the first two months in most cases compared to fresh cherries, as confirmed by a beneficial reshuffle at cytohistological level. This explains why it can allow storage of cherries for candying over the whole duration between two harvest seasons.

Keywords:
Prunus avium L.
Firmness
Candied cherries
Brine
Cytohistology
Cell wall

1. Introduction

Sweet cherries (Prunus avium L. cv. Napoléon) produced in Vaucluse (South France) are mainly intended for processing to glacé cherries, represented 10,000 tons of final product, with 70% destined for more than 60 countries. They must be harvested within a short time (3–4 weeks) and are then stored in brine for candying all along the year. The narrow window of harvesting prompted growers to adopt mechanical methods based on the use of ethephon (2-chloroethyl phosphonic acid) and harvesting at an earlier stage. The ethylene released by Ethephon induces a decrease of the pedicel-fruit retention force in sweet cherry (Smith & Whiting, 2007). This is mediated by its binding to receptor sites that initiates a signal transduction leading to acceleration of cell-wall degrading enzymes and cell death in the abscission zone (Estornell, Agusti, Merelo, Talon, & Tadeo, 2013; Hall, Shakeel, & Schaller, 2007). In the case of cherries, approximately 80% fruits fall without peduncle. However, this harvesting method is not a rule and manually harvested cherries still represent 20% of total quantity delivered to industry (Ulrich Fleury, personal communication).

Between harvest and candying, cherry fruits are stored in brine containing sulfur dioxide and calcium salts. Acidity and sulphites protect fruit from microorganism proliferation due to SO2 anti-septic effects (Dupuy, 1959; Julien, 1972). Calcium salts act on cherries firmness, as also noted to improve texture for pickled cucumbers (Hudson & Buescher, 1985). Sulfur dioxide and calcium are directly related to brined cherry quality. Improper use of these chemicals may result in cherries that are soft, poorly bleached, or spoiled due to fermentation (Payne, Beavers, Cain, & Station, 1969). Atkinson and Strachan (1962) pointed out other advantages of sulfur dioxide as a preservative. It is inexpensive and provides a quick method of handling large quantities of fruit. The
product may be placed in bulk containers for shipment, and the preservative is effective for a sufficient period of time to allow for storage and remanufacture. In addition, most of the sulfur dioxide can be removed easily and inexpensively before candying, so that this allergen is absent (Aptunion, internal analysis) in the glacé cherries. Moreover, storing for a long period without loss of fruit quality is one of industries objectives.

Firmness loss during fruit development is associated with cell-wall polysaccharide turnover (Brummell & Harpster, 2001). Cherry cell walls are modified in harvesting, whether after chemical ethephon treatment (Batisse, Coulomb, Coulomb, and Buret (1998)) or natural ripening-over ripening mechanisms (Batisse, Filiskycaon, and Buret (1994)) leading to softness. Some of the genotypic variations in fruit firmness have been linked with differences in the patterns of cell-wall disassembly. For example, Cheol, Toivonen, Wiersma, and Kappel (2002) reported that soft varieties present lower total cell wall contents in fresh fruit. Taillan, Ambid, Pech, and Raynal (1992) demonstrated that pectic fractions were the main cell wall component modified during cherries storage in aluminium or calcium brines; fruits stored in calcium brine present a better firmness than in aluminium brine leading industry to continually improve brine formulation to increase the quality of stored fruits.

However, other factors have been reported to be important modulators for cherry post harvest behaviour. Richardson et al. (1998) reported that some harvest factors are susceptible to affect sweet cherries P. avium cv. Royal Ann texture, such as machine factors (pattern of tree shaking, duration of shakes, etc.), climatic conditions, crop loads, tree sizes or their spacing. Since this early work, no studies have attempted to evaluate the consequences of other harvest itineraries on the evolution of the quality of cherries in brine. Understanding the impact of the harvest conditions on cherry texture evolution during storage is of major importance for the industry. In the present study, evolution of firmness during brine storage was quantified according to the fruits maturity stage, the harvesting method, the ethephon application and the peduncle presence. Cytohistological investigations have been performed to follow the gross structural alterations during storage in brine.

2. Materials and methods

2.1. Plant material

Fruit of sweet cherry (Prunus avium L., cv. Napoléon) for industry were harvested in Provence (Lagnes 43°53’39” N and 5°06’55” W) in June 2013. Table 1 summarizes the different treatments and harvest conditions. Ethephon (PRM12, Bayer S.A.S, Lyon, France) was applied at 0.36 g l⁻¹ on vigorous trees [conditions (1) and (2)], by high humidity and 20 °C (standard treatment). Untreated trees were also included in the study [conditions (3) and (4)]. Cherries were picked at 10 “early” and 20 “late” days after treatment with ethephon corresponding to the beginning and end of the harvest.

The fruits were harvested mechanically [condition (2)] or manually [condition (1)], (3) and (4) then immediately immersed in brine tanks (Type Aptunion, Apt, France) for storage with 50/50 ratio fruit/brine. Fruits (1) and (3) were stored in brine with their peduncle and fruits (2) and (4) without peduncle (Table 1).

Samples were taken at 0, 2, 5, 8, and 15 days and at 1, 2, 3, 4, 5, 6, 7, 8, 10 and 12 months in brine.

2.2. Fruit texture characterization

For each treatment, 62 cherries (randomly chosen) were placed in holes, with the peduncle scar on top (Fig. 1). They were pitted with a texture analyser TA Plus (Ametek, Lloyd Instruments Ltd, Fareham, UK) using an 8 mm diameter probe moving at 20 mm s⁻¹ for a course of 40 mm after first contact. The maximum load (Newton) during pitting, captured by a 250 N load cell, is our criterion to define the texture quality. It was chosen as the texture test as it is representative of the actual process, pitting being a critical point in the processing to glacé cherries, and presented a good correlation with a compression test such as routinely applied to other fruits, e.g. in Grote, Duprat, Loonis, and Pietri (2001); Missang, Maingonnat, Renard, and Audergon (2011).

2.3. Physico-chemical characterizations

The total soluble solids (TSS) or Brix were measured in the brine with an ATAGO PR-1 Digital Refractometer (Atago Co., LTD, Tokyo, Japan). Indeed, brine TSS is complementary to fruit TSS due to the diffusion phenomenon that leads to an equilibrium state.

Calcium concentration in the brine was measured by Atomic absorption spectroscopy on a Varian AA 55 (Agilent technologies, Santa Clara, USA) equipped with an acetylene/nitrous oxide burner.

2.4. Microscopy

Cytohistological analyses were performed using tissue samples excised from median part of fruits going from exocarp to endocarp (Fig. 2). Immediately after excision all specimens were immersed in fixative solution consisting of 10% Acetic acid 10% Formalin and 80% Ethanol. To promote good penetration of the fixative product samples were subjected to vacuum for 20 min. After 48 h fixation at room temperature, the specimens were rinsed in distilled water and stored in 70% ethanol until required. They were then dehydrated in a graded ethanol series (80–100%) and embedded in methacrylate resin (Kit Technovit 7100, Heraeus-Kulzer GmbH, W. Wahib et al. / LWT - Food Science and Technology 75 (2017) 243-250, DOI: 10.1016/j.lwt.2016.08.059

Table 1 : Summary of the different treatments and harvest conditions for the different sweet cherry samples. Ethephon was applied at 0.36 g l⁻¹, mechanical harvest stored in industrial container and manual harvest stored in bucket for practically. Each sample was picked at to maturity stage “early” and “late”.

<table>
<thead>
<tr>
<th>Sample</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ethephon</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Peduncle</td>
<td>+</td>
<td>-</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>Container (m³)</td>
<td>0.025</td>
<td>3</td>
<td>0.025</td>
<td>0.025</td>
</tr>
</tbody>
</table>

Fig. 1. Sweet cherry disposition and orientation for firmness measurements.
Wehrheim, Germany). Sections (3 µm thickness) were serially cut using a retraction microtome (Supercut 2065; Reichert-Young, Wien, Austria), and collected on microscope slides. They were stained to visualise polysaccharides and proteins using periodic acid-Schiff’s reagent and naphthol blue-black procedures, respectively (El Maataoui & Pichot, 1999). Observations were performed using Leica DMR photomicroscope equipped for brightfield, darkfield, phase contrast and UV illuminations. Images were captured using Leica DFC 300 FX digital camera and processed using LAS software (Leica, Wetzlar, Germany). At least 6 fruits were analysed per treatment.

2.5. Statistics

All statistics were performed using R i386 3.1.2 software (R Foundation for Statistical Computing, Vienna, Austria). Prior to analysis data were tested using Shapiro-Wilk normality test to verify the distribution. Mean and Sd were calculated and one-way analysis of variance (ANOVA) and the Tukey post-hoc test were applied to verify the significance of differences between firmness, TSS and calcium levels in fruits from different harvest conditions.

3. Results and discussion

3.1. General evolution of texture during storage

During storage in brine, evolution of cherry texture was characterized by a first stage of rapid increase in firmness, up to about 10 days. This was followed by a second stage (10 days–2 months) of texture stabilisation, and between 2 and 12 months texture remained constant or showed limited decreasing trend (Fig. 3C and D). Storage in brine thus increased fruit firmness with a maximum reached in about 2 months. Thereafter comparison will concern only D0, D8, M2 and M12 conservation stages as those present...
characteristics points during cherries evolution in brine. This can be compared to the short shelf life of fresh cherries, with both texture loss and material degradation. For example, Wei, Qi, Guan, and Zhu (2011) maintained firmness for 30 days after 1-methylcyclopropene (1-MCP) treatment combined with 0 °C storage and 8 days for untreated sweet cherry ‘Summit’. Preservation effects in sulfur dioxide brine are mainly due to the inhibition of deleterious enzymatic activity thanks to brine and by the calcium complexation in different fruit tissues. Conservation by brine-process confers a stable firmness to pickled capers (Capparis spp.) in brine (Ozcan & Akgul, 1999) and allows improving texture in first stage and stabilization for some months in pickled cucumbers (Bell & Etchells, 1961; Hudson & Buescher, 1985).

3.2. Maturity stage and harvest method

The cherries picked at different maturity stages had different initial firmness: fresh “early” cherries presented a median firmness of 25 N at harvest, and “late” fruit 18 N (Fig. 3A). They reached 35 N for “early” and 25 N for “late” after 2 months and firmness then remained constant over one year storage (Fig. 3C). Indeed maturation is characterized by loss of firmness, as has been reported by Batisse et al. (1994) and Filslycaco and Buret (1990) for sweet cherries including ‘Napoléon’ cultivar. The initial loss observed here was of 38% of the firmness, with a mean difference of 6–7 N that persisted throughout brine storage but the gap becomes 19% of the firmness after 12 months.

Harvesting modalities had an impact on initial texture (25 N for manual, 18 N for mechanical) (Fig. 3B), at 2 months (35 N for manual, 24 N for mechanical), but more important also on texture evolution after 1 year storage in brine (Fig. 3D). Mechanically harvested cherries showed a trend for texture loss reaching 19 N after 12 months, while the texture of manually harvested cherries remained stable at about 34 N after one year. The difference in firmness thus increased to reach 15 N, which was significant and corresponded to the quality differential observed industrially, with soft fruit and increased loss for mechanically harvested cherries after prolonged storage. However they remained above the limit of processability, which has been assessed at 17 N. The mechanical harvest, in spite of its great advantages in cost and speed, was deleterious to fruit quality, probably by increasing mechanical damage and decreasing fruit firmness, as also observed by Timm and Guyer (1998) for tart cherries. Storing in brine over a long period conferred a texture gain, improving effectively the fruit quality for a manual harvest whereas the gain was minimal after a mechanical harvest with fruits that remained above the limit of processability.

3.3. Ethephon application

Mechanized harvest is dependent on ethephon application. Ethephon application had a slight impact on initial texture (24 N for untreated, 28 N for treated) (Fig. 4) i.e. on average a gain of about 15%. This effect was at the opposite to what Smith and Whiting (2009) reported, i.e. ethephon application leads to a decrease of firmness by about 20% in both “Chelan” and “Bing” fresh cherries harvested 22 days after ethephon treatments. Initial firmness difference disappeared during brine soaking (31 N at 2 months for both batches) and cherry had almost the same firmness (30.5 N for untreated against 29.5 N for treated) after 12 months. Ethephon application had a positive impact on fresh cherries firmness but brine storage led to texture equalization. Ethephon is known to have different consequences on texture depending on plant species; it decreases firmness in both kiwifruit and muskmelon (Yamaguchi, Hughes, Tyler, Johnson, & May, 1977; Zhang, Li, Liu, Song, & Liu, 2012), does not impact texture in cucumber (Miller &

![Fig. 4.](image) Evolution of cherries firmness in brine with/without ethephon application; D0 — fresh fruit and D008, D060, D365 corresponding to period storage of 8, 60, and 365 days, respectively. Each boxplot represents 248 randomly selected cherries stored in brine. Statistical significance at P ≤ 0.01 represented with different letters after Tukey (HSD) post-hoc test.
Lower, 1972) or induces a better ripening of fruits with uniform colour, pleasant flavour, desirable firmness and acceptable sensory quality in guava (Mahajan, Gagandeep, & Dhatt, 2008).

3.4. Peduncle effect

Removing peduncle before soaking in brine improved texture of fruits. Starting from the same firmness level (23 N), cherries without peduncle presented higher firmness than integrals cherries ones immersed in brine (29 N and 24 N, respectively) after 8 days (Fig. 5). Then, the firmness reached 34 N against 31 N after 2 months storage and between 2 and 12 months texture remained constant with the same gap. This suggests that absence of peduncle leads to fruit with a better texture. The peduncle—abscession zone might be a preferential circuit of exchange between cherries and brine salts. The pedicel of a sweet cherry fruit is a useful indicator of its postharvest freshness (Drake & Elving, 2002; Sekse & Lyngstad, 1993; Wani, Singh, Gul, Wani, & Langowski, 2014). Although studies have been carried out on molecular composition of peduncle (Khalid, Gellert, Szendrei, & Duddeck, 1989) and transpiration mechanism (Athoo, Winkler, & Knoche, 2015) in sweet cherries or abscission and physical zone separation in rabbiteye blueberry (Vashisth & Malladi, 2013), little is known about impact of peduncle removal. Here we show for the first time that peduncle removal had a marked effect on texture of cherry during brine storage leading to improved fruit texture, mostly by increasing the firmness gain between crop days and 2 months.

Table 2 summarizes the variations in firmness in our experimental system. It appears that the best combination was a manual harvesting thus not requiring ethephon, in the early maturation stage and immersing in brine without peduncle. However, this is not practical economically as mechanical harvest is highly effective in terms of cost and rapidity. Storage in brine promotes textural changes in cherries, even mechanically harvested, thus allowing their utilisation for processed candied-fruits until the next crop year.

3.5. TSS and calcium evolution in brine

During storage in brine, the sugars and organic acids that constitute most of the TSS of sweet cherries diffuse to the brine by osmotic phenomenon and, inversely, the brine components like calcium diffuse from brine to sweet cherries (Karel, 1973; Taillan, 1991). Accordingly, the measures were performed only in brine, assuming the remainder was in the fruits. The cherries without peduncle released more TSS after 8 and 60 days in brine than those immersed with peduncle. After 1 year storage, TSS in brine was higher for the fruits without peduncle (Fig. 6A). This difference in TSS content of the brine after 1 year (i.e. equilibration complete) starting from identical initial fruit (16.5°B) and brine (same batch) soluble solids might be an artefact from fruit packing in the bins. Indeed, while a 50/50 ratio (weight/weight) was initially aimed for, fruits with peduncle occupied more volume per weight so that more brine had to be added to cover them. Calcium evolution in the brine (Fig. 6B) was faster for cherries without peduncle; after 8 days cherries without peduncle had absorbed about 10% more calcium than cherries with peduncle.

Calcium is well known to enhance texture of fruit and particularly of fruit stored in brine, as has been studied for pickled cucumbers (Howard & Buescher, 1990; McFeeters, 1985). It limits the softening of pickles during heating or storage under different conditions (Tang & McFeeters, 1983). The impact of calcium on texture is due to formation of calcium cross-links between pectin molecules in the cell wall (Van Buren, 1979). Garcia, Brenes, and Garrido (1994) reported that the monovalent sodium ions Na⁺ does not show any action on firmness while calcium Ca²⁺ improves

![Fig. 5. Evolution of cherries firmness in brine with/without peduncle; D0 – fresh fruit and D008, D060, D365 corresponding to period storage of 8, 60, and 365 days respectively. Each boxplot represent a variation of 248 randomly selected cherries stored in brine. Statistical significance at P ≤ 0.01 represented with different letters after Tukey (HSD) post-hoc test.](image-url)
firmness and use of both can lead to competition between them in the cross-linked de-esterified pectin thereby weakening the pectin-calcium-based network.

Further, Taillan et al. (1992) demonstrated that residual pectin methyl esterase (PME) activity is present during brine storage of cherry fruit, and that calcium concentration modulates the stimulation of PME activity. The pedicel-abscission zone allowed an earlier transfer of brine components thus potentially enhancing the PME-calcium interaction and conferring improved texture. In contrast, the removal of Ca\(^{2+}\) bound to middle lamella-cell wall material enhances the rate and magnitude of galacturonan degradation by polygalacturonase (Buescher & Hobson, 1982), and may cause softening. Loss of textural integrity can be controlled by the formation of intermolecular links between Ca\(^{2+}\) and pectin achieved by the addition of calcium salts (Barrett, Garcia, & Wayne, 1998) and/or boosting the endogenous PME activity (Anthon, Blot, & Barrett, 2005). The positive impact of peduncle removal observed in our conditions might thus be due to increased and accelerated calcium availability for pectin cross-linking in the cherry fleshy tissues.

3.6. Microscopy

Cytomorphological observations were performed by light microscopy to better understand the impact of the studied harvest conditions on cherry fleshy tissues during brining. Applied to all conditions, this approach globally revealed that brined fruits display similar tendencies of cell and tissue structural behaviour. These tendencies will be described here taking as example sections of tissue fragments excised from fruits harvested mechanically (without peduncle) at early stage, and brined either for 2 or 12 months. Fig. 7 presents typical micrographs illustrating main changes in comparison with samples from fresh fruits.

In fresh fruits, the peripheral pericarp tissues showed cohesive, normally shaped cells exhibiting intact cell walls (Fig. 7A). In contrast, middle mesocarp and to a greater extent inner mesocarp changed significantly (without peduncle) at early stage, and brined either for 2 or 12 months. Fig. 7 presents typical micrographs illustrating main changes in comparison with samples from fresh fruits.

Table 2

<table>
<thead>
<tr>
<th>Maturity stage</th>
<th>Harvest method</th>
<th>Ethephon</th>
<th>Peduncle</th>
<th>Fresh cherries</th>
<th>15 days in brine</th>
<th>2 months in brine</th>
<th>1 year in brine</th>
</tr>
</thead>
<tbody>
<tr>
<td>Early</td>
<td>Manual</td>
<td>Yes</td>
<td>P+</td>
<td>28 ± 4.9</td>
<td>34 ± 5.1</td>
<td>37 ± 4.8</td>
<td>30 ± 3.4</td>
</tr>
<tr>
<td></td>
<td>Mechanical</td>
<td>Yes</td>
<td>P-</td>
<td>22 ± 4.2</td>
<td>29 ± 5</td>
<td>28 ± 3.8</td>
<td>22 ± 4.2</td>
</tr>
<tr>
<td></td>
<td>Manual</td>
<td>No</td>
<td>P+</td>
<td>24 ± 4.7</td>
<td>32 ± 4.9</td>
<td>36 ± 4.2</td>
<td>33 ± 5</td>
</tr>
<tr>
<td></td>
<td>Manual</td>
<td>No</td>
<td>P-</td>
<td>24 ± 4.7</td>
<td>34 ± 4.8</td>
<td>41 ± 5.1</td>
<td>37 ± 5.4</td>
</tr>
<tr>
<td>Late</td>
<td>Manual</td>
<td>Yes</td>
<td>P+</td>
<td>20 ± 4.5</td>
<td>24 ± 4</td>
<td>25 ± 4.4</td>
<td>25 ± 3.8</td>
</tr>
<tr>
<td></td>
<td>Mechanical</td>
<td>Yes</td>
<td>P-</td>
<td>15 ± 4.3</td>
<td>21 ± 3.8</td>
<td>20 ± 5</td>
<td>17 ± 4.6</td>
</tr>
<tr>
<td></td>
<td>Manual</td>
<td>No</td>
<td>P-</td>
<td>23 ± 4.5</td>
<td>28 ± 4.8</td>
<td>29 ± 3.6</td>
<td>30 ± 4.2</td>
</tr>
</tbody>
</table>

Fig. 6. Evolution of A total soluble solids (°B) and B calcium (g/L) in brine during storage of cherries with (---) and without peduncle (---). Different letters within the means and standard deviation of each date signify statistical separation at P ≤ 0.05.
and 12 months in brine, the peripheral mesocarp cell layers appeared to be structurally similar to those of fresh fruits (compare Fig. 7A, D and G). But surprisingly, in middle and inner mesocarp, cells exhibited regular contour giving them a turgescence-like state (Fig. 7E, F, H and I). This apparent morphological rescue may be interpreted as cell wall stabilising effects of the brine via pectin-calcium interactions. Alonso, Tortosa, Canet, and Rodríguez (2005) described a synergic effect of thermal treatment (70 °C) and immersion in calcium solution to decrease the degree of esterification of water-soluble and EDTA-soluble pectins, thereby favouring the formation of calcium bridges and preventing the depolymerization of pectins in sweet cherry cv Pico Colorado. Other authors (Roy, Jauneau, & Vian, 1994) observed high contents of non-esterified homogalacturonan sequences in the same spaces in accordance with the presence of calcium ions. Calcium pectate plays a role as a last barrier before total cell disaggregation and had an important function in maintaining firmness during brine storage. One-year storage in brine further consolidated the cellular architecture with an intercellular space rich in polysaccharides.

4. Conclusion

In this work, we characterized the different harvest factors involved in the evolution of cherries texture during brine storage. Overall, brine increased texture, thus alleviating slightly the initial damage due to harvest itineraries. Among these factors, ethephon treatment only impacted positively fresh fruit firmness and allowed to maintain acceptable cherries quality for processing over one year.

The harvest modalities that impacted texture of cherries were, in order of importance, harvest type, maturity stage, and presence of peduncle.

However, presence of an abscission zone accelerated osmotic exchange and also improved penetration of brine salts rate, conferring high texture gain with cell wall remodelling. Optimal textures were obtained for fruit harvested manually (to avoid mechanical damage) an early stage (to avoid over-ripening) and stored in brine without peduncle (to promote brine diffusion within fruit tissues). Since mechanical harvesting is unavoidable and fruit growers aim to shorten the harvest period, fruit quality enhancement may be established at post-harvest level. In addition to peduncle and its relevance for calcium diffusion, temperature and/or pressure might modulate residual endogenous PME activity (Taillan et al. 1992). This leads to release a significant proportion of free carboxyl groups from pectin polymers that might then interact with calcium salt to form secondary cross-bridges thus improving cell wall strength, and consequently, a higher quality final product. This will be the object for further work.

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