Research Article

Effects of harvest time on functional compounds and fruit antioxidant capacity in ten strawberry cultivars

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Abstract

BACKGROUND: Huelva (Spain) is the main region for strawberry production in Europe. Most fruit production is exported for fresh consumption to European countries, where consumers demand high fruit quality and appreciate its healthy properties. Strawberry intake is a valuable source of antioxidants compounds with important health benefits. The higher the antioxidant capacity of a cultivar, the better the enhancement of human health.

OBJECTIVE: The comparative knowledge of fruit composition on antioxidant compounds and its variation along the cropping season, in ten strawberry cultivars cropped at Huelva.

METHODS: Fruit yield and citric acid, ascorbic acid, total phenolics, anthocyanins content as well as antioxidant capacity of fruits were evaluated in ten strawberry cultivars at three harvesting times during the 2014 field campaign.

RESULTS: Yield and fruit parameters analyzed were strongly influenced by the genotype and by the time of harvesting. Strawberry fruit quality and antioxidant properties were greater when harvested from mid- to late- season and were not associated with higher yields.

CONCLUSIONS: Healthy properties of strawberry fruits depend on cultivar and harvest time. Knowledge of the nutritional properties of these strawberry cultivars might translate into benefits to growers and enhancement of health for consumers.

Keywords: Antioxidant, cropping conditions, harvesting, health, fruit quality, vitamin C, phenolics, yield

1. Introduction

Strawberry is among the most widely consumed fruits in the world. The Huelva region in Spain (south western coast of Spain), is the main strawberry cropping area in Europe [1], and its production, characterised by high quality fruits, is largely exported to most European countries [2]. Freshly consumed fruits produced in Spain belong to different strawberry cultivars (Fragaria × ananassa Duch). In 2014, more than 65% of the cropping area of Huelva was distributed among ‘Splendor’ (26%), ‘Sabrina’ (24%) and ‘Florida-Fortuna’ (16%), followed by ‘Primoris’ (7%), Candonga (6%) and others. Most of these cultivars were developed by different public and private strawberry breeding programs [3] aiming to achieve better agronomic traits (yield and fruit quality) and, recently, to improve healthy features of fruits.

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Fruit healthy benefits have been related to the presence of different compounds with antioxidant properties and the capacity of neutralizing free radical activity [4, 5]. The chemical nature of these compounds is diverse but includes a group of enzymes (superoxide dismutase, catalase, peroxidase, etc.), phenolic acids (catechins, flavonols, anthocyanins, etc.) and vitamins (C, E and A) [6]. In addition of being an enriched source of vitamins and minerals, intake of fruits and vegetables enriched in these bioactive products has been associated with low incidence of proliferative [7, 8] and cardiovascular diseases [9, 10]. More concretely, berries, including strawberry, contain high amounts of flavonoids, phenolic acids and anthocyanins (responsible of red fruit color) [11–18]. The later has been directly associated with a large number of human health benefits ranging from antioxidant potential, anticancer activity, anti-inflammatory and anti-angiogenic properties [19–25].

The spectra of bioactive compounds vary in different berries according to species, cultivation conditions and production area [26]. In strawberry, it has been reported that genotype influences the concentration of these compounds and their relative amounts [13, 26, 27, 28]. These reports are pointing out that synthesis of phytochemicals in fruits and vegetables is the result of genotypic and environmental interactions [29]. In this sense, it is reasonable to hypothesize that changes in environmental conditions (i.e. light, temperature and relative humidity) along the cropping season might also influence the fruit concentration of bioactive compounds at different harvesting periods.

Therefore, assessment of strawberry nutritional quality has to take into account the whole chain of production, including cultivars and cropping conditions (i.e. crop management, cultivation techniques, variability of environmental conditions), which are quite variable among productive areas worldwide. However, little information is available about the content on bioactive compounds of the cultivars currently growing on the main strawberry productive area of Europe. In addition to the organoleptic features, knowledge of the fruit health properties would represent an extra-value for consumers and would increase benefits for growers.

To analyze, the influence of harvesting time on organic acids and antioxidants compounds on different genotypes, we evaluated 10 short-day strawberry cultivars well-adapted to the productive area of Huelva growing under the conventional cropping system.

2. Material and Methods

2.1. Plant material and experimental design

Ten short-day strawberry cultivars, ‘Antilla’, ‘Sabrina’, Candonga®, ‘Fontanilla’, ‘Florida-Florida’ (‘F. Fortuna’), ‘Liberty’, ‘Primoris’, ‘Rabida’, ‘Sahara’ and ‘Splendor’, were planted in mid-October 2013 at the IFAPA experimental station “El Cebollar” (Huelva, Spain). All cultivars are well-adapted to Huelva agroclimatic conditions [30] and were grown following conventional cropping-practices. Planting was done in a double row mulched raised beds (35 cm high and 50 cm wide) of a sandy soil with 5.8% clay, 5% silt and 89.2% sand, 0.09% organic matter, previously biosolarized [31]. Fruit set takes place from January (mid-winter) to end of May (late spring). Polyethylene-covered tunnel structures (macrotunnel; [32]) were installed in mid-November and removed at the end of the cropping season.

In order to evaluate effects of cultivar and harvest time on the nutraceutic features of the fruits, a field experiment was setup in a complete randomized block design with three replicate plots per cultivar and 50 plants per plot spaced at 25 × 25 cm. Throughout the crop season (January-May 2014), all mature fruits per plot were harvested once to twice a week. Total yield (g plant⁻¹) for each cultivar was calculated for the whole season and in three harvesting periods: ‘extra-early’ (from January to February), ‘early’ (March) and ‘late’ (April to May). According to these harvesting periods, fruit quality analyses were done on three sampling dates: 19th February, 21st March and 9th April.

2.2. Sample preparation for fruit quality determination

At each harvest date, ~250 g of mature strawberry fruits per plot (8–10 fruits) were taken and homogenized with a blender immediately after harvesting. Pulp samples were stored at ~20°C until processed accordingly to the following assays at the laboratory.
2.3. Titratable acidity and ascorbic acid content

For titratable acidity (TA) determinations, the pulp was filtrated and diluted with distilled water (1g: 100 mL). Titration to pH 8.1 (end point the third pK value of citric acid; AOAC 22.058) with 0.01M NaOH was done at room temperature with Titroline Easy (Schott Instruments® GmbH) portable pH meter. Total acidity was expressed as grams of citric acid per 100 grams of fresh weight (FW) [33].

For ascorbic acid quantification (AA), pulp samples were diluted with distilled water (1g: 10 mL) and homogenized. Reagent test strips were used with the reflectometer set of Merck Co (Merck Rqflex 10). Results were expressed as milligrams of ascorbic acid per 100 grams of FW.

2.4. Total phenolics and anthocyanins content

Two grams of pulp samples were diluted with 10 mL of extraction solvent: methanol (99.9%) and CH$_3$OH (0.1%), stored at 4ºC during 24 h, and centrifuged at 10000 rpm for 15 min at 4ºC. The supernatant was diluted in extraction solvent (2:1) and stored at −20ºC until analyzed.

Total phenolic content in extracts were determined by the Folin–Ciocalteu method [34, 35] modified by Tulipani et al. [36]. Appropriately diluted extracts (2 ml) were mixed with 0.2 ml Folin-Ciocalteu reagent and 0.4 ml sodium carbonate (35% w/v) was added. After 1 h at room temperature and darkness, absorbance at 725 nm was measured in a spectrophotometer. Gallic acid (Sigma) was used as standard, and results were expressed as milligrams of gallic acid equivalents (GAE) per 100 grams of FW.

Total anthocyanin content was measured with the pH differential absorbance method [37, 38]. The pH values of the diluted pulp juice were 1.0 (in 0.025 M potassium chloride buffer) and 4.5 (in 0.4 M sodium acetate buffer). Absorbance at 510 and 700 nm was measured in a UV–VIS spectrophotometer. The absorbance values of the diluted samples (A) were calculated as follows: $A = [(A_{510} - A_{700}) pH_{1.0} - (A_{510} - A_{700}) pH_{4.5}]$. The total anthocyanin pigment was expressed as milligrams of pelargonidin-3-glucoside (Pg-glc) equivalent per 100 grams of FW. It was calculated as the product of A, the molecular weight of Pg-glc, and the dilution factor divided by the molar extinction coefficient of Pg-glc [39]. Total Anthocyanins = $A \times MW \times df \times 1000/\varepsilon$.

2.5. Antioxidant capacity: ABTS assay

To determine antioxidant capacity, 2.8 g pulps were added to 10 ml methanol (60%) and homogenized. Afterwards, sample was centrifuged at 3000 rpm for 15 min at 4ºC. The supernatant was stored at −20ºC until analyzed.

Total antioxidant capacity was evaluated according to TEAC (Trolox Equivalent Antioxidant Capacity, [40]) assay. This assay employ ABTS molecules (a chromogen and colorless substance that changes into its colored monocatonic radical form (ABTS) by an oxidative agent). The absorption peak of ABTS is at 734 nm. Addition of antioxidants reduces ABTS into its colorless form. Therefore, the extent of decolorization, as percentage of inhibition of ABTS, is determined as a function of concentration and calculated relative to the reactivity of Trolox. Antioxidant activity is expressed as μmol of Trolox equivalents (TE) per gram of fresh weight (FW).

2.6. Statistical analysis

Data were subjected to analysis of variance (randomized complete block design, ANOVA) for means comparison, and differences between mean values were compared by Tukey honest significant difference (HSD) with the analytical software STATISTIX 9.0 (Analytical Software, Florida, USA) and reported as means ± standard error of the mean. Correlations between the levels of the different antioxidants were calculated by Pearson correlation.
3. Results and discussion

3.1. Cultivars effect on fruit quality and yield

The influence of cultivar on organic acids (citric and ascorbic acids) was significant (Table 1). Among the 10 cultivars studied, ‘Sabrina’, Candonga® and ‘Sahara’ showed the highest values of citric acid (more than 0.8 g TA/100 g FW) whilst ‘Antilla’ had the lowest (0.65 g TA/100 g FW). Thus, the amount of citric acid depended on the cultivars as it was previously described by Sturm et al. [41]. Besides its importance in flavor, citric acid is also of great interest because promotes the antioxidant action and seem to contribute to the antioxidant activity [42]. In this sense, ‘Sabrina’, Candonga® and ‘Sahara’ would be more appropriate in terms of health, whereas ‘Antilla’ would be better for taste, since a low amount of acid is associated with a higher sugar/acid ratio [43].

Regarding to ascorbic acid content, the highest value was shown by ‘Liberty’ (55.29 mg AA/100 g FW; 37.7% more ascorbic acid than the lowest one ‘Rabida’) followed by ‘Sahara’ and Candonga® (Table 1). Differences in ascorbic acid among strawberry cultivars were also observed by several authors [36, 44, 45] working with cultivars adapted to Sweden, Italy, and Belgium; in general all seem to give ascorbic acid values in the same range. Likewise, ascorbic acid is an important antioxidant, involved in human health [11, 46] and, in this sense, the consumption of strawberry cultivars with high content in ascorbic acid (i.e. ‘Liberty’. Table 1) would lead to an improvement in health. Specifically this acid has repeatedly been associated with lowered risk of developing several diseases, as cancer [47].

Great variability existed among the examined strawberry fruits regarding to their phenolic compounds (Table 1). ‘Primoris’ and Candonga® showed significantly higher contents than the others cultivars (up to 17%). The results found in this work are comparable and in accordance with those observed by Pincemail et al. [45] and Buendia et al. [48] who found important differences in the content of phenolic compounds among various cultivars of strawberry.

Differences among cultivars were also observed for anthocyanin content (Table 1). In this sense, the highest amounts of anthocyanin were shown by ‘Florida-Fortuna’ and ‘Sabrina’, which showed up to 49% more anthocyanins than ‘Antilla’ (the cultivar with the lowest value). High variability in anthocyanin content among strawberry cultivars was previously described by Tulipani et al. [36]. These type of cultivars are of great interest in a healthy diet, due to anthocyanins may play an important role in controlling oxidative reactions and exhibit antiinflammatory and anti-angiogenic properties [19–25].

As regards antioxidant capacity, significant differences among the 10 cultivars were shown (Table 1). The highest value (39.1 μmol Teq/g FW) was obtained by the cultivar Candonga®, whilst ‘Primoris’ showed the lowest one. Variations in antioxidant capacity among strawberry cultivars were already observed in several studies [29, 49, 50, 51].

<table>
<thead>
<tr>
<th>Cultivar</th>
<th>Citric acid (g CA/100 g FW)</th>
<th>Ascorbic acid (mg AA/100 g FW)</th>
<th>Phenolics (mg GAE/100 g FW)</th>
<th>Anthocyanins (mg Pg-glc Eq/100 g FW)</th>
<th>TEAC (μmol Teq/g FW)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Antilla</td>
<td>0.65 ± 0.04 b</td>
<td>36.6 ± 1.3 d</td>
<td>217.6 ± 6.3 e</td>
<td>10.4 ± 0.7 d</td>
<td>35.4 ± 2.3 ab</td>
</tr>
<tr>
<td>Sabrina</td>
<td>0.82 ± 0.02 a</td>
<td>△1.3 ± 1.8 hcd</td>
<td>227.0 ± 6.4 ab</td>
<td>19.5 ± 1.0 ab</td>
<td>37.0 ± 2.4 ab</td>
</tr>
<tr>
<td>Candonga</td>
<td>0.84 ± 0.04 a</td>
<td>43.7 ± 1.4 bc</td>
<td>263.9 ± 5.7 a</td>
<td>17.3 ± 0.6 abc</td>
<td>39.1 ± 1.7 a</td>
</tr>
<tr>
<td>Fontanilla</td>
<td>0.76 ± 0.05 ab</td>
<td>43.5 ± 2.0 hcd</td>
<td>222.2 ± 7.1 b</td>
<td>17.3 ± 0.5 abc</td>
<td>35.0 ± 0.0 ab</td>
</tr>
<tr>
<td>Fortuna</td>
<td>0.73 ± 0.02 abc</td>
<td>38.7 ± 0.5 cde</td>
<td>212.7 ± 5.3 b</td>
<td>21.4 ± 1.0 a</td>
<td>34.3 ± 2.9 ab</td>
</tr>
<tr>
<td>Liberty</td>
<td>0.74 ± 0.02 ab</td>
<td>55.3 ± 0.8 a</td>
<td>232.2 ± 12.8 ab</td>
<td>14.4 ± 0.8 cd</td>
<td>36.1 ± 1.3 ab</td>
</tr>
<tr>
<td>Primoris</td>
<td>0.76 ± 0.03 ab</td>
<td>38.4 ± 1.7 cde</td>
<td>264.0 ± 12.9 a</td>
<td>15.6 ± 1.5 bc</td>
<td>25.5 ± 4.0 b</td>
</tr>
<tr>
<td>Rabida</td>
<td>0.75 ± 0.04 ab</td>
<td>34.4 ± 2.1 e</td>
<td>217.4 ± 13.8 b</td>
<td>15.5 ± 1.6 bc</td>
<td>30.1 ± 3.0 ab</td>
</tr>
<tr>
<td>Sahara</td>
<td>0.81 ± 0.03 a</td>
<td>48.1 ± 1.4 b</td>
<td>216.6 ± 8.2 b</td>
<td>15.8 ± 1.1 bc</td>
<td>35.2 ± 1.5 ab</td>
</tr>
<tr>
<td>Splendor</td>
<td>0.72 ± 0.03 ab</td>
<td>42.9 ± 1.1 bcde</td>
<td>224.8 ± 8.1 ab</td>
<td>18.3 ± 0.6 abc</td>
<td>33.8 ± 2.5 ab</td>
</tr>
</tbody>
</table>
Table 2
Content of citric and ascorbic acid, total phenolics, total anthocyanins and antioxidant capacity (TEAC) in fruits across the ten strawberry cultivars in three harvesting times (19th February, 21st March and 9th April). Data are the mean ± SE (n=30). Means within the same line followed by different letters were significantly different at *p*< 0.05

<table>
<thead>
<tr>
<th>19th February</th>
<th>21st March</th>
<th>9th April</th>
</tr>
</thead>
<tbody>
<tr>
<td>Citric acid (g CA / 100 g FW)</td>
<td>0.70 ± 0.01 c</td>
<td>0.73 ± 0.01 b</td>
</tr>
<tr>
<td>Ascorbic acid (mg AA / 100 g FW)</td>
<td>39.7 ± 1.6 a</td>
<td>44.8 ± 1.1 a</td>
</tr>
<tr>
<td>Phenolics (mg GAE / 100 g FW)</td>
<td>212.2 ± 5.2 c</td>
<td>248.4 ± 6.2 a</td>
</tr>
<tr>
<td>Anthocyanins (mg Pg-glc Eq / 100 g FW)</td>
<td>13.9 ± 0.7 b</td>
<td>17.9 ± 0.7 a</td>
</tr>
<tr>
<td>TEAC (μmol TEq / g FW)</td>
<td>30.1 ± 1.4 b</td>
<td>39.9 ± 0.7 a</td>
</tr>
</tbody>
</table>

Table 3
Pearson correlation coefficient (R²) of the different parameters analyzed (citric acid, antioxidant capacity (TEAC), Anthocyanins, and phenolics compounds) in strawberry

<table>
<thead>
<tr>
<th></th>
<th>TEAC</th>
<th>Anthocyanins</th>
<th>Phenolics</th>
<th>Ascorbic acid</th>
</tr>
</thead>
<tbody>
<tr>
<td>Citric acid</td>
<td>0.1875 ns</td>
<td>0.4668**</td>
<td>0.2740 ns</td>
<td>0.1284 ns</td>
</tr>
<tr>
<td>TEAC</td>
<td>0.3013*</td>
<td>0.3176**</td>
<td>0.3002 ns</td>
<td>0.1335 m</td>
</tr>
<tr>
<td>Phenolics</td>
<td>0.2459 m</td>
<td>0.2516 ms</td>
<td>0.2459 m</td>
<td>0.2459 m</td>
</tr>
</tbody>
</table>

n = 90. ns: not significant, * and **: significant differences at *p* < 0.05 and at *p* < 0.01, respectively.

Differences in the amount of organic acids, phenolic and anthocyanin compounds in the cultivars analyzed, confirm that the genotype largely determines antioxidant capacity in strawberry. Therefore, the knowledge of antioxidant composition of cultivars would favor their consumption and might contribute to improving human health [7, 8, 9, 10]. Also, those cultivars with high antioxidant compounds can be used as parental in breeding programs to obtain new cultivars with increased antioxidant capacity.

A significant correlation (Table 3) was observed between total antioxidant capacity and anthocyanin content (R² = 0.3613, *p* < 0.05, n = 90), ascorbic acid (R² = 0.5044, *p* < 0.001, n = 90), and total phenolic compounds (R² = 0.5176, *p* < 0.001, n = 90); indicating the important contribution of these compounds for the antioxidant capacity on the strawberry cultivars studied. These results are in agreement with Aaby et al. [46], who reported that ascorbic acid was the most important contributor to the antioxidant capacity in strawberry. In contrast, Pinçemail et al. [45] only found those correlations in one out of the twelve cultivars they studied. Previous studies reported that citric acid contributed greatly to antioxidant capacity on strawberry [42], but our results are not consistent with those since no significant correlation between total antioxidant capacity and citric acid was found, despite citric acid was significantly correlated with anthocyanin content (R² = 0.4668, *p* < 0.001, n = 90). Given that anthocyanins are pigments that confer colors from orange to purple, the significant relationship between anthocyanins and antioxidants capacity is pointing out that the more red is the fruit, the greater antioxidant capacity and health benefit.

All study cultivars displayed fruit yields comparable to commercial orchards along the whole season. Total fruit yield was above 900 g/plant but significant differences among cultivars were found (Fig. 1). ‘Sabrina’ and ‘Fontanilla’ displayed the highest values of total production (1240.4 ± 12.6 and 1225.4 ± 13.4 g/plant, respectively), followed by ‘Sahara’, ‘Rabida’ and ‘Florida-Fortuna’ (with a yield decrease of 5.69, 7.29 and 12.05% respect to ‘Sabrina’) being ‘Antilla’ the cultivar with the lowest fruit yield (with a decrease about 30%). ‘Splendor’, ‘Primoris’, ‘Liberty’ and Candonga® displayed intermediate fruit yields. Among the most productive cultivars, ‘Rabida’ and ‘Sahara’, showed also high precocity (189.3 ± 9.6 and 161.7 ± 7.8 g/plant in ‘Extra-early’ period, respectively; Fig. 1) together with ‘Splendor’ and ‘Primoris’ (174.1 ± 14.6 and 148.6 ± 1.8 g/plant in ‘Extra-early’ period, respectively), but the two latter showed lower total yields (1020.6 ± 44.6 and 961.9 ± 30.9 g/plant). These results are indicating that high precocity do not necessarily translate into higher yields at the end of the cropping season and vice versa. Fruit production in ‘Extra-early’ (January to February), ‘Early’ (March) and ‘Late’ (from April to May), was 15 %, 24% and 63%, respectively. This is suggesting that fruit production is associated to the main flowering bloom that takes place in late winter and early spring since averaged ripening from anthesis is about 38 days [52].
3.2. Harvest time influence on fruit quality

Previous studies on strawberry have shown that harvest time affected ascorbic acid, phenolic content and antioxidant capacity of fruits [45]. However, environmental conditions, cropping system (i.e. open field or greenhouse) and harvest dates (i.e. May-November) were quite different.

In this work, the three harvesting times tested (19th February, 21st March and 9th April) significantly affected the content of organic acids and antioxidant compounds of fruits (at the same ripening stage) of the 10 cultivars studied (Table 2). Thereby, at first harvest time (19th February) all cultivars displayed the lowest content of organic acids and antioxidant compounds. More concretely, citric acid, ascorbic acid, phenolics acid, anthocyanin and total antioxidant capacity were lower in extra-early period by 12.02%, 8.30%, 11.09%, 22.86% and 17.18% respectively, compared to early and late average value. This is indicating that fruits harvested from the middle to the end of season (early: 21st March and late period: 9th April, Table 2), which coincides with the peak of production in the Huelva area (Fig. 1), would have better healthy properties.

However, significant interaction of cultivars and harvest periods were found ($p < 0.001$) in the analyzed parameters, evidencing that cultivars behave differently across harvest times (Fig. 2). Although in most cultivars there was a tendency to increase citric acid content through the field campaign, in ‘Sabrina’, with intermediate values, it did not change along the crop season (Fig. 2A). Regarding ascorbic acid, all cultivars achieved their highest values in the early harvest with the exception of ‘Fontanilla’, which displayed the highest ascorbic acid content in the extra-early (19th February) period, and ‘Liberty’, in which vitamin C content did not change along the field campaign, being significantly higherst than in any other cultivar (Fig. 2B). The highest ascorbic acid content in extra-early period observed in ‘Fontanilla’ could be of great interest due to its precocity.

Regarding to phenolic compounds, most cultivars followed a similar bell-shaped pattern with the highest values in the early period (21st March). However in ‘Florida-Fortuna’, ‘Sabrina’, ‘Antilla’ and ‘Fontanilla’ values remained almost unchanged (Fig. 2C).

Pattern of variation of anthocyanin content was similar in most cultivars, which showed a marked increase between extra-early and early harvest times, except in ‘Antilla’, ‘Candonga’, and ‘Fontanilla’, in which anthocyanins did not change significantly (Fig. 2D). Regarding to total antioxidant capacity, patterns of variation were quite different among cultivars. ‘Primoris’, ‘Splendor’, ‘Rabida’, and ‘Candonga’ displayed a bell-shaped pattern (i.e. maximum values at early-harvest); ‘Fontanilla’ and ‘Florida-Fortuna’ remained unchanged; ‘Antilla’ increased progressively; and a drop was observed in ‘Liberty’ and ‘Sahara’ in the late harvest (Fig. 2E).
Fig. 2. Fruit content of citric acid (A), ascorbic acid (B), total phenolics (C), anthocyanins (D) and fruit antioxidant capacity (TEAC; E) of the 10 strawberry cultivars during the 2014 cropping-season. Fruits were analyzed at the three harvest times (Extra-early: 19th February; Early: 21st March; Late: 9th April).

Such variability in the accumulation of antioxidant compounds in the fruits, is suggesting that cultivars are differently affected by the changes in the environmental conditions during the cropping season [53]. In this sense, it is known that high temperatures and light intensity influences the synthesis of organic and antioxidant compounds in growing fruits [53–56]. Therefore, selection of cultivars enriched in healthy compounds (i.e. antioxidants) and with high stability across changing environmental conditions of the growing areas might represent an advantage for human health and would translate into benefits for growers.

4. Conclusion

Widely consumed strawberry cultivars produced on the main strawberry production area of Europe, differ largely in the fruit their fruit composition in of organic acid and antioxidant compounds. Differences among cultivars might
not be kept along the cropping season since fruit quality and healthy properties vary at different harvest times. The range of variation depends on the cultivar; therefore harvest time has to be taken into account when comparing nutraceutical properties. Overall, fruit quality is substantially better when fruits are harvested from mid-March to early April in the season. Among cultivars, Candonga®, ‘Sabrina’, ‘Fontanilla’ and ‘Sahara’ have very high content of citrus acid and antioxidant compounds in conjunction with the highest yields. It is remarkable that ‘Fontanilla’ and ‘Sahara’ also displayed precocity, suggesting that their cultivation would increase the economic profit for growers by the early arrival of high quality fruits to the markets.

In addition to giving relevant information about the healthy properties of commercial strawberries, these results have importance for the choice and selection of parental in breeding programs.

Acknowledgments

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