Bioactive compounds and quality characteristics of five apples cultivars

Compostos bioativos e características de qualidade de cinco cultivares de maçãs

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ABSTRACT

The aim of this study was to evaluate bioactive compounds in five apple cultivars and to analyze correlation of their quality characteristics with concentration of bioactive compounds. Phenolic compounds measurements were made in a spectrophotometer compared to a standard curve of gallic acid and expressed as gallic acid equivalent (GAE) per 100g of dry weight. Sugar and organics acids in five cultivars were quantified using high-performance liquid chromatograph (HPLC). Antioxidant activities were evaluated using three complementary tests: 2,2-diphenyl-1-picrylhydrazyl (DPPH) scavenging activity, ferric reducing antioxidant power (FRAP), and 2,2'-azino-bis (3-ethylbenzthiazoline-6-sulfonic acid) (ABTS). 'GoldRush' had the highest total phenolic compared to the other four cultivars. Additionally, 'GoldRush' had slightly higher DPPH activity followed by 'Crimson Crisp' and 'Wine Crisp'. 'GoldRush' and 'Crimson Crisp' cultivars also have higher antioxidant capacity based on the ABTS and FRAP methods. The antioxidant capacity was significantly correlated with total polyphenols present in the different cultivars, while organic acids and fruit color showed slightly significant correlation to total phenols.

Key words: Mallus domestica, organic acids, sugar, antioxidant capacity, HPLC/DAD.

INTRODUCTION

Degenerative diseases such as cancers, Alzheimer’s, and cardiovascular disease pose grave threats to public health. These illnesses can require expensive, lengthy treatment which may still fail to restore patient health. Thus, preventative measures are our best and first defense against degenerative disease. Consumption of foods rich in bioactive compounds have been found to reduce some these diseases (TONG et al., 2011). This is due to the food’s antioxidants including vitamins C and E, carotenoids, and polyphenols. Antioxidants prevent the occurrence of free radical damage in the cell (BAGETTI et al., 2011).
Recent work has shown that there are considerable variations of bioactive compounds in apples (DROGOUDI et al., 2008; WOJDYLO et al., 2008). The main factor is related to genetic variability among different cultivars. The such as, skin color, the environmental conditions during production, such as, temperature, sunlight, and irrigation, also influence (DROGOUDI et al., 2008). WOJDYLO et al. (2008) evaluated old and new apple cultivars produced in Poland. They observed significantly higher concentration of bioactive compounds in newly developed cultivars. This suggests that antioxidant content is a trait which can, and should be manipulated by breeders.

Given the ability of functional foods to reduce degenerative disease, it would be valuable to determine which easily visible characteristics are indicators of their antioxidant capacity. For example, previous work has shown a correlation between the pomegranate’s nutritional value and its readily apparent physical properties of color and size (DROGOUDI et al., 2005). There are countless apple cultivars available, and new cultivars are being continuously developed and introduced commercially. To help meet the needs of public health, we evaluated bioactivity of four newly released apple cultivars developed by the Purdue, Rutgers, and Illinois (PRI) breeding program. We then compared it with the older, widely consumed cultivar ‘Golden Delicious’. Finally, we analyzed the relationship between biochemical properties and readily apparent fruit characteristics.

MATERIALS AND METHODS

Fruit from six-year old ‘Golden Delicious’, ‘GoldRush’, ‘Wine Crisp’, ‘Pixie Crunch’, and ‘Crimson Crisp’ trees grafted on ‘Budagovsky 9’ were harvested at optimum commercial maturity. Trees are grown at the fruit farmer of the University of Illinois, Champaign-Urbana. Average size fruits free of any visual damage were harvested. Fruits were kept stored at 0°C for one week then evaluated after seven days at room temperature.

Ten fruits of four replicates were used to determine each parameter. After measurement of quality indices, fruits were subsampled by taking wedges of about 700g per replicate of each cultivar and stored at -18°C overnight prior to freeze-drying.

The color of the peel was carried out individually in the equatorial region of the fruit with digital colorimeter Model CR-200 (Konica Minolta, Osaka, Japan). The color values were obtained from the L*, a*, and b* and the hue angle and chroma (C*) used arctan the formulas $(b/a)$ and $(a^2 + b^2)^{1/2}$. Cultivars ‘Wine Crisp’ and ‘Pixie Crunch’ were performed color readings in the region yellow and red.

To measure the TA, pH and SS content was used homogenized juice of all fruits of each repetition. TA values were determined by titrating juice samples with 0.1M NaOH and expressed as g malic acid 100 mL-1 juice. pH values were measured with a pH meter (Fisher Science Education, Malaysia). For measure the SS content was used analog Leica 10430 refractometer (Fisher Scientific, Waltham, MA, USA).

Freeze-drying was done in a VirTis freeze dryer (VirTis, Gardiner, New York, USA) to dry at 40mmHg). Freeze-dried samples were ground in coffee grinder and analyzed for total phenolic, antioxidants capacity, sugar, and organic acids as follows procedures.

For extraction of polyphenols approximately 0.5g of freeze dried tissue samples were combined with 20mL of 70% methanol and homogenized in a Polytron homogenizer (Kinematica Ag Littau, Switzerland) set at a speed of 4 about 1 minute. The homogenate was centrifuged twice for 10 minutes at 4000g. The supernatant was collected and used to determine total polyphenols (TP) using the colorimetric Folin-Ciocalteau assay as described by SINGLETON et al. (1999). Tissue TP measurements were made in a Shimadzu spectrophotometer set at 725 nm compared to a standard curve of gallic acid and expressed as gallic acid equivalent (GAE) per 100g of dry weight.

The antioxidant capacity was determined by the modified 2,2-Diphenyl-1-picrylhydrazyl (DPPH) radical method (BRAND-WILLIAMS et al., 1995) which is based on the quantification of free radical-scavenging activity, with modifications according to RUFINO et al. (2007). One g of freeze dried tissue were in centrifuge tubes and extracted sequentially with 20 ml of methanol/water (50:50, v/v) at room temperature for 1h. The tubes were centrifuged at 25000g for 15 min and the supernatant was recovered. Then 20ml of acetone/water (70:30, v/v) was added to the residue at room temperature, extracted for 60min and centrifuged. Methanol and acetone extracts were combined, made up to 50ml with distilled water and used to determine antioxidant capacity. A methanol solution containing 0.06mM DPPH was prepared. After adjusting the blank with methanol, an aliquot of 100μl of fruit extract was added to 3.9ml of this solution. Based on preliminary study, the times required to obtain DPPH was after two hours of contact of the sample with the reagent. The antioxidant capacity was expressed as mM DPPH per 100g of dry weight.

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Antioxidant capacity was also evaluated by 2,2'-azino-bis (3-ethylbenzthiazoline-6-sulfonic acid) (ABTS) and ferric reducing antioxidant power (FRAP) method according KU et al. (2011). Briefly the ABTS method involved dissolving 7mM ABTS (ammonium salt) in potassium phosphate buffer (pH 7.4) and combined with 2.45mM potassium persulfate. The mixture was stored in the dark for 10min. The dark blue solution was diluted with potassium phosphate buffer (pH 7.4) until the absorbance reached 1.0 ± 0.02 at 734nm using a microplate reader (Biotek Instruments, Winooski, USA). Two hundred µL of the resulting solution was mixed with 10µL of the sample, and 6min later the absorbance was recorded at room temperature. The results were expressed as Trolox equivalent (TE mM g⁻¹ dry weight). The concentration of standard solution ranged from 0.25 to 4mM. All experiments were carried out in triplicate.

The FRAP method involves mixing 200µL of freshly prepared FRAP reagent with 10µL of the previously prepared extract. The FRAP reagent contained 2.5mL of a 10mM L⁻¹ TPTZ solution in 40mM L⁻¹ HCl in distilled water, 2.5mL of 20mM L⁻¹ FeCl₃•6H₂O in distilled water, and 25mL of 0.3M L⁻¹ acetate buffer of pH 5.9. The mixture was stored in the dark for about 1 minute. The homogenate was centrifuged at a 27,000 × g at 5°C for 30min. A 1mL fraction of the supernatant was filtered through a 0.2µm nylon filter and a 5µL fraction was injected into an Hitachi HPLC equipped with a UV detector set at 210nm and a Phenomenix organic acid column (REZEX 10µ 8% Ca, 300 x 7.80mm column (Torrance, CA, USA), equipped with a refractive index detector and a column heater. Glucose, fructose, sucrose, and sorbitol were separated using a REZEX 10 micron 8% Ca, 300 x 7.80mm column (Torrance, CA, USA) heated at 80°C. Glass distilled water was used as a mobile phase at a flow rate of 0.6mL min⁻¹. Sugar concentration was expressed as g 100g⁻¹ dry weight.

Organic acid extraction was carried out by weighing a 1.0g of freeze-dried tissue samples. Dried tissue were ground with a mortar and pestle, placed into a 20mL tubes, and combined with 5mL of 0.004N H₂SO₄. The tissue was homogenized in the dark with a Polytron homogenizer set at a speed of 4 about 1 minute. The homogenate was centrifuged at a 27,000 × g at 5°C for 30min. A 1mL fraction of the supernatant was filtered through a 0.2µm nylon filter and a 5µL fraction was injected into an Hitachi HPLC equipped with a UV detector set at 210nm and a Phenomenix organic acid column (REZEX 10µ 8% H₂O in distilled water, and 25mL of 0.3M L⁻¹ acetate buffer of pH 5.9. The results were expressed as Trolox equivalent (TE mM g⁻¹ dry weight). The concentration of standard solution ranged from 0.25 to 2mM. All experiments were carried out in triplicate.

To determination sugars content, 0.2g of freeze-dried tissue samples were placed in thimbles inside 20mL test tubes, and 2mL of 70% ethanol was added to each tube. A 1cm marble was placed on the mouth of each tube to trap the evaporated ethanol. The tubes were heated on a hot plate (Pierce Reactio-Therm III, Rockford, USA) for four hours at 90°C. The ethanol fraction was collected and evaporated under an air stream and the dried residue was brought up 1mL with distilled water. The aqueous extract was filtered through a 0.2µm nylon filter and 60µL sample was injected into a Hitachi HPLC (Tokyo, Japan), equipped with a refractive index detector and a column heater. Glucose, fructose, sucrose, and sorbitol were separated using a REZEX 10 micron 8% Ca, 300 x 7.80mm column (Torrance, CA, USA) heated at 80°C. Glass distilled water was used as a mobile phase at a flow rate of 0.6mL min⁻¹. Sugar concentration was expressed as g 100g⁻¹ dry weight.

Antioxidant capacity required to reduce the original amount of free radicals by 50% (EC₁₀) and values expressed as EC₅₀ = g⁻¹ dry weight. DPPH.

Total phenolic content and antioxidant activity in the five apple cultivars are presented in figure 1. There was a significant variation in total phenolic content among cultivars. The highest total phenolic content (1.90g GAE per 100g⁻¹ dry weight) were found in ‘GoldRush’, followed by ‘Crimson Crisp’, ‘Wine Crisp’, ‘Golden Delicious’, and ‘Pixie Crunch’, which had (1.58; 1.53; 1.42; 1.14g GAE per 100g⁻¹ dry weight) (Figure 1A). The results were similar with previous studies (WOJDYLO et al., 2008; VALAVANIDIS et al., 2009). WOJDYLO et al. (2008) examined old and new apple varieties and reported that total polyphenols concentrations ranged between 0.5 and 2.7g GAE per 100g⁻¹ dry weight. VALAVANIDIS et al. (2009) examined total phenolics content in five apple cultivars (‘Red Delicious Starking’, ‘Golden Delicious’, ‘Granny Smith’, ‘Royal Gala’, and ‘Jona Gold’) and found that whole fruits contained between 0.1-0.2g GAE per 100g⁻¹ dry weight based on pulp weight and peels contained between 0.2-0.4g GAE per 100g⁻¹ dry weight.

Antioxidant capacity differed from five apple cultivars, but different analytical methods gave similar results (Figure 1B, 1C, 1974.

RESULTS AND DISCUSSION

Total phenolic content and antioxidant activity in the five apple cultivars, but different analytical methods gave similar results (Figure 1B, 1C, 1974.

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and DPPH, respectively. Cultivars ‘Golden Delicious’ and ‘Pixie Crunch’ showed the significantly higher values than above three apple cultivars which are 1.47 and 1.52 EC_{50} = g dry weight mg^{-1} DPPH, respectively. The values found in this study are similar of the WOJDYLO et al. (2008) that studied the antioxidant capacity from 67 apple varieties.

Antioxidant capacities obtained by the ABTS and FRAP were expressed in TE mM g^{-1} dry weight. The both methods showed similar results among cultivars. The cultivars that had high antioxidant capacity by ABTS method was ‘Gold Rush’ and ‘Crimson Crisp’ (66.6 and 63.5TE mM g^{-1} dry weight) followed by cultivars ‘Wine Crisp’ and ‘Golden Delicious’ with intermediate values (52.40 and 55.37TE mM g^{-1} dry matter) and ‘Pixie Crunch’ (45.80TE mM g^{-1} dry weight) with lower value (Figure 1C). Determined via the FRAP method, the cultivar ‘Gold Rush’ and the ‘Crimson Crisp’ also had higher antioxidant capacity 61.77 and 54.39mM g^{-1} dry weight. The cultivar ‘Pixie Crunch’ had the lowest antioxidant capacity by ABTS and FRAP methods. However, it was not statistically different from the cultivar ‘Golden Delicious’ as measured by the FRAP assay (Figure 1D). Although statistical differences among cultivars vary between the two methods, the assay results were generally similar for different cultivars because these methods are based on electron transfer mechanism. Due to these variations, it is recommended to have at least two antioxidant methods with different modes of actions in study to accurately reflect the antioxidant activity (OZGEN et al., 2006).

The sugar, acid and ratio are closely related with the taste and flavor of the apple fruit. The

Figure 1 - Total phenolic content (expressed mg GAE 100 g^{-1} dry weight) (A), antioxidant capacity (2,2-Diphenyl-1-picrylhydrazyl DPPH, expressed EC_{50} = g dry weight mg^{-1} DPPH) (B), antioxidant capacity – [2,2’-azino-bis (3-ethylbenzthiazoline-6-7 sulfonic acid) ABTS expressed TE mM g^{-1} of dry weight] (C), and antioxidant capacity – (ferric reducing/antioxidant power FRAP, expressed TE mM g^{-1} of dry weight) (D) in tissues of five apple cultivars. Values represented mean ± S.E. Different letters denote significant differences (P≤0.05). TE = trolox equivalent.
fruit acid is beneficial for human body (HECKE et al., 2006). Since apples are recommended for diabetic patients, the knowledge of the sugar content of various apple fruits is of crucial importance today (HECKE et al., 2006). Therefore, it is important to measure sugar and acids from new apple variety for consumers. The sugar content showed significant differences between cultivars (Figure 2). Generally, the cultivar ‘Pixie Crunch’ presented the highest values and ‘Wine Crisp’, the lowest values. Regarding the glucose content of fruit, the cultivar ‘Pixie Crunch’ presented the highest concentration but did not differ significant from cultivars ‘Crimson Crisp’ and ‘GoldRush’ (Figure 2A). The concentration of fructose showed similar behavior among cultivars, the exception was the cultivar ‘Wine Crisp’ that presented the lowest value among them (Figure 2B). Cultivars had sucrose distributions similar to the glucose distribution, highlighted by maximum values in ‘Pixie Crunch’ and the minimum in ‘Wine Crisp’ (Figure 2C), for both sugars. Sorbitol, despite having a much smaller portion compared to the others sugars, followed the same behavior among cultivars, with higher values given by ‘Pixie Crunch’ and the smallest by ‘Wine Crisp’ (Figure 2D). The data presented here show variation between cultivars similar to that found by STURM & STAMPAR et al. (1999) that evaluated seasonal variation of sugars in apple in different growing systems.

The concentration of the various acids present in the different cultivars is shown in figure 3. Malic acid was the one with the highest concentration between acids studied (Figure 3A), demonstrating consistency because when evaluating the total acidity of apples, it is represented by the concentration of malic acid (DROGOUDI et al., 2008). The greatest levels of malic acid was found in ‘Gold Rush’ with 272mg 100g⁻¹ of dry weight and the lowest value in ‘Pixie Crunch’, with 192mg 100g⁻¹ of dry weight (Figure 3A).
Similar distributions of malate and oxalate were observed, but malic acid concentrations were a whole order of magnitude greater than oxalic acid concentrations (Figure 3B). Tartaric and ascorbic acids were present in relatively small amounts compared to malic acid. ‘Golden Delicious’ and ‘GoldRush’ had no detectable tartaric acid (Figure 3C). The values of ascorbic acid (Figure 3D) of different cultivars evaluated are lower than found by JAN et al. (2012) who studied different cultivars of apple grown in Peshawar-Pakistan. They reported ‘Red Delicious’ contain 12.5mg of ascorbic acid of 100g of fresh weight. The highest ascorbic acid was observed in ‘Gold Rush’ in this study but less than 4mg in 100g of dry weight. The different ascorbic acid value may be due to the difference from the cultivar, environment, and extraction method. Ascorbic acid is an important quality characteristic of apple fruit because of its essentiality to human health, decreasing many diseases (TONG et al., 2011). SZETO et al. (2002) estimated that 100mg ascorbic acid is contained in one orange, a few florets of raw cauliflower or a handful of uncooked spinach leaves. However, ascorbic concentration of apple is lower than other fruits and vegetables (SZETO et al., 2002). Apples, bananas contain very little ascorbic acid (SZETO et al., 2002).

Although a significant correlation coefficient ($r = 0.97$) was observed between ABTS and FRAP methods, the DPPH method showed relatively smaller correlation negative coefficients with ABTS ($r = -0.78$) and FRAP ($r = -0.85$) methods (Table 1). The significant correlations among the three methods that showed consistent results indicate that antioxidant capacity of the different apples cultivars were reasonably determined.

There are several studies that verify the antioxidant capacity correlate with different quality indices, especially with polyphenols (WOJDYLO et al., 2008). In this study the polyphenolic content showed significant correlation coefficients with the values of DPPH ($r = 0.80$), ABTS ($r = 0.94$), and FRAP ($r = 0.97$) as KHANIZADEH et al. (2007) have reported. In our

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analysis, only malic and tartaric acids had significant correlation (0.53 and 0.52) with antioxidant capacity obtained by DPPH and FRAP methods. DROGOUDI et al. (2008) also found a significant correlation between antioxidant capacity and acids in different cultivars of apple, but the authors obtained with ascorbic acid. In the study with pear, SÁNCHEZ et al. (2003) found weak correlation between antioxidant capacities and organic acids but chlorogenic acid (simple phenolic) showed a slight positive correlation.

The color of the peel of the fruit is generally correlated with antioxidant capacity. When assessing only the peels of the apple, the values for antioxidant activity are greater than whole apple or pulp of apple (KHANIZADEH et al., 2007; WOJDYLO et al., 2008). In the present study, only the characteristic L was significantly correlated with ABTS method (r = 0.45), indicating that ABTS method is sensitive to antioxidant compounds in lightness apple peel. Further correlation was not obtained due to the evaluations are carried out with peel and pulp, causing dilution of antioxidant compounds were present in higher quantities in the peel, since the peel corresponding to approximately 12% mass of apple fruits. The present analysis method using the whole apple fruit can be easily justified because most consumers eat the peel and pulp of apple fruits.

CONCLUSION

Among the apple cultivars we tested, ‘GoldRush’ had the highest total phenolic content. Additionally, ‘GoldRush’ and ‘Crimson Crisp’ have shown greater antioxidant activity than cultivar ‘GoldRush’ in DPPH, ABTS, and FRAP methods. However, ‘Pixie Crunch’, has lower total phenolic and antioxidant ability in ABTS and FRAP, suggesting breeding program may need to focus on total phenolics and antioxidant compounds as well as other phenotype in the future. The antioxidant capacity was significantly correlated with polyphenols in this present study, while the organic acids showed little correlation. The ABTS antioxidant activity significantly correlated with L.

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REFERENCES


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