Effects of Nitrogen Fertilization on Subtropical Peach Fruit Quality: Organic Acids, Phytochemical Content, and Total Antioxidant Capacity

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ABSTRACT. Producing temperate-zone fruit crops in subtropical environments requires alterations in fertilizer application and rates. Nitrogen (N) is a critical mineral nutrient required in high amounts by the tree; however, it is often over- or under-applied for optimal fruit quality and can affect the phytochemical composition of fruits. The effects of different N fertilizer rates and harvest date on total phenolic content, total flavonoid content, total anthocyanins, total antioxidant capacity, total soluble solids, titratable acidity, and organic acids (citric and malic acid) of two subtropical peach (Prunus persica) cultivars, TropicBeauty and UFSharp, were investigated. N rate did not affect total soluble solids in ‘TropicBeauty’, although total soluble solids decreased as N rate increased in ‘UFSharp’. Titratable acidity and organic acid content was significantly higher in ‘UFSharp’ as compared with ‘TropicBeauty’, although there was no effect of N rate on titratable acidity. An overall increase in phenolic content, flavonoid content, anthocyanins, and antioxidant capacity were observed with decreasing N rates in both subtropical peach cultivars. A stronger genotype × N treatment interaction was observed for ‘TropicBeauty’ for phenolic content, flavonoid content, and antioxidant capacity than for ‘UFSharp’. In ‘TropicBeauty’, among the treatments with no N and highest N, an almost 100% increase in phenolic content, 200% increase in flavonoid content, 50% increase in anthocyanin content, and 80% increase in antioxidant activity was observed. A positive correlation among phenolic content, flavonoid content, and antioxidant capacity was observed in both ‘TropicBeauty’ and ‘UFSharp’. Late harvest date decreased phenolic content in ‘TropicBeauty’, ranging from 6% to 32% among different N treatments. Late harvest increased anthocyanin content as compared with fruit that were harvested on early dates. The results suggest that subtropical peach phytochemical composition can be affected by different cultivars and tree age, and can be manipulated with cultural practices like N fertilization and harvest time to produce fruit with altered or desired nutritional composition for consumers.

As consumers learn about the health benefits of fruits (Prior and Cao, 2000), they are demanding fruit with higher antioxidant and phytochemical capacities, as clearly demonstrated by the popularity of fruits (Gilbert et al., 2014; Olmstead et al., 2015). Increased consumption of fruits has been associated with potential protection against age-related diseases (Ames et al., 1993). A clear inverse relationship between the consumption of fruits and incidences of cardiovascular and cerebrovascular degenerative proliferative diseases, oxidative stress, and mortality has been largely supported by a number of epidemiological studies (Ames et al., 1993; Ko et al., 2005; Sun et al., 2002). In a consumer survey conducted by researchers at the University of Florida, 39% of the consumers identified health benefits as the most influential factors for southern highbush blueberry (Vaccinium darrowii) fruit consumption; whereas, 61% of the consumers suggested the flavor was more critical for southern highbush blueberry consumption (Gilbert et al., 2014), indicating a high number of consumers are paying attention to health benefits of fruits over other attributes and often base their decision of consumption on the health benefits offered.

Fruits are excellent functional foods as they are high in antioxidant capacity and phytochemicals (Scalzo et al., 2005; Tomas-Barberan and Robins, 1997). Phytochemicals are naturally occurring substances that play an important role in the visual appearance and flavor. Phytochemicals comprise numerous compounds such as polyphenolics, carotenoids, alkaloids, etc., and are well known for imparting antioxidant activity. Antioxidants act as scavenging agents for harmful free radicals, which have been implicated in the aging process and many degenerative diseases (Harman, 1981; Rice-Evans et al., 1996). Polyphenolic compounds are secondary plant metabolites, and their reox properties allow them to act as reducing agents, hydrogen donors, and singlet oxygen quenchers (Fukumoto and Mazza, 2000; Prior et al., 2005; Rice-Evans et al., 1996). One of the largest classes of polyphenolic compounds in fruits and vegetables is flavonoids. Flavonoids are very well characterized...
for their free radical scavenging activity. Flavonoid radicals have lower reduction potential than in alkyl peroxyl radicals, and the superoxide radical, meaning flavonoids have the potential to inactivate these oxy species and prevent the deleterious consequence of their reactions (Jovanovic et al., 1992; Rice-Evans et al., 1996; Sogawa et al., 1993). Anthocyanins, a flavonoid, are natural, water soluble pigments found in flower, fruit, and leaves. Anthocyanins contain potent antioxidant properties modulated by their different hydroxylation and glycosylation (Rice-Evans et al., 1996). In addition to their health benefits, flavonoids, and other phenolic compounds are important for normal plant growth development, fruit and flower color development, and defense against ultraviolet light, pathogens, insect herbivory, and other biotic and abiotic stresses (Benoit and Berry, 1997; Berhow and Vaughn, 1999; Brignola et al., 1998; Murphy et al., 2000; Ryan et al., 2002).

Subtropical peaches are greatly appreciated for their great taste, flavor, and aroma. Ideal subtropical peach fruit quality and high nutritive value drives consumer satisfaction measured by initial and repeat consumer purchases (Colaric et al., 2005; Delgado et al., 2013; Hudina and Stampar, 2000; Olmstead et al., 2015). In addition to excellent fruit taste, subtropical peaches are also a good source of vitamin C and have a fair amount of antioxidant activity (Cantin et al., 2009). Olmstead et al. (2015) conducted an IdeaMap (Moskowitz Jacobs, White Plains, NY) consumer survey to identify “ideal” subtropical peach attributes that contribute to repeated consumer purchase. According to the survey, vitamin C scored high after flavor and texture, suggesting consumers prefer high nutrient value in subtropical peaches. In subtropical peaches, the primary source of antioxidant capacity is polyphenolic compounds (Gil et al., 2002). The major anthocyanins in subtropical peach are cyanidin-3-glucoside and cyanidin-3-rutinoside (Tomas-Barberan et al., 2001).

Production of subtropical peach cultivars requires alterations in cultural practices as the active growth period is considerably longer than in temperate production areas. It has been suggested that these practices may influence overall fruit quality and polyphenolic compound concentrations in fruits. Overall fruit quality, distribution and composition of phenolic phytochemicals are affected by maturity, cultivars, horticultural practices, fertilization, geographic origin, growing season, postharvest storage conditions, and processing procedures (Burda et al., 1990; De Freitas and Glories, 1999; Donovan et al., 1998; Kalt et al., 1999; Kim et al., 2001; Lee and Jaworski, 1987; Spanos and Wrolstad, 1990). Phytochemical composition is strongly influenced by genotypes; different cultivars can produce different concentrations of total phenolic compounds as well as specific phenolic compounds (Atkinson et al., 2006; Buendia et al., 2010). Cantin et al. (2009) reported that the phytochemical profiles of subtropical peach genotypes varied depending on yellow/white flesh color, and antioxidant capacity was positively correlated to total phenolic content. N fertilization has been reported to affect phenolic content in several different fruits and vegetables such as Chinese cabbage [Brassica chinensis (Zhu et al., 2009)], broccoli [Brassica oleracea var. italica (Jones et al., 2007)], apple [Malus domestica (Reay et al., 1998)], basil [Ocimum basilicum (Nguyen and Niemeyer, 2008)], lettuce [Lactuca sativa (Coria-Cayupan et al., 2009)], tomato [Solanum lycopersicum (Bénard et al., 2009)], olive [Olea europaea (Fernandez-Escobar et al., 2006)], and strawberry [Fragaria ananassa (Anttonen et al., 2006)]. In all these examples, an increase in phenolic content has been observed with application of reduced N rates. High N rates in olives and grapes (Vitis vinifera) have reduced polyphenol concentration, including anthocyanins, and resulted in decline of overall oil and wine quality, respectively (Fernandez-Escobar et al., 2006; Hilbert et al., 2003; Keller and Hrazdina, 1998; Malusà et al., 2004; Schreiner et al., 2013; Wade et al., 2004). N rate can also affect fruit quality parameters such as total soluble solids [TSS (sweetness indicator)] and titratable acidity [TA (acidity indicator)]. Sweetness, acidity, and astringency are key traits that are positively correlated with overall liking for subtropical peach fruit (Olmstead et al., 2015; Predieri et al., 2006). The effect of N on TSS and TA has been reported in a number of fruit crops such as grape (Bavaresco et al., 2001), apple (Nava et al., 2008), and tomato (Simonne et al., 2007). Another factor that can influence the fruit quality and phytochemical content of fresh produce is the fruit developmental stage and harvest time. Fruit ripening stage can strongly influence the content of TSS, TA, and phenolic compounds as well as total antioxidant activity of fruit extracts (Guerra and Casquero, 2008; Tulipani et al., 2008).

The production of temperate zone fruit crops in subtropical environments has increased significantly in the last 30 years. Interest continues to grow because of two factors: 1) early flowering and fruit set result in growers being able to harvest fruit earlier in the domestic market window (Morgan and Olmstead, 2013), giving higher economic returns, and 2) breeding advances have resulted in availability of low-chill accumulating plants that provide alternatives for growers seeking to diversify (Byrne and Bacon, 1999). Low-chill cultivars of apple (Hough and Bonnetti, 1988), southern highbush blueberry, plum (Prunus domestica), and subtropical peach (Lyrene, 2005; Sherman and Lyrene, 1985) have been developed by several breeding programs and are in commercial production (Byrne and Bacon, 1999). Currently, we lack in understanding of the effect of preharvest production practices on the fruit quality of subtropical peaches. As the production of subtropical peaches continues to increase in southeast United States, there is a desperate need to ensure that the production practices such as fertilization and harvest time are optimized to yield high quality, nutritious fruit, in addition to optimum production. Therefore, in the present study, we focused on the impact of different rates of N fertilization on fruit quality including organic acid, TSS, TA, total phenolic content (phenolic content), total flavonoid content (flavonoid content), total anthocyanin content (anthocyanin content), and total antioxidant capacity (antioxidant capacity) of subtropical peaches. We hypothesize that increasing N rate will lead to lower phytochemical content, whereas later harvest dates will favor higher phytochemical content. The two subtropical peach cultivars used in this study are TropicBeauty (melting texture) and UFSSharp (nonmelting texture). ‘Tropic Beauty’ is one of the most popular low-chill subtropical peach cultivars; the medium-sized, semifreestone fruit have yellow, melting flesh and develop 70% blush over a yellow ground color, and have low firmness. ‘UFSSharp’ fruit develop 60% red blush over a deep yellow to orange ground color with excellent fruit size, shape, and firmness.

Materials and Methods

Chemicals. All chemicals were of analytical grade. 3,4,5-trihydroxybenzoic acid (gallic acid), acetic acid (CH₃COOH),
aluminum chloride (AlCl₃), catechin, Folin-Ciocalteu’s phenol reagent, ferrous sulphate heptahydrate [TPTZ [(FeSO₄·7H₂O), 2,4,6-tris (2-pyridyl)-s-triazine]], ferric chloride hexahydrate (FeCl₃·6H₂O), hydrochloric acid (HCl), potassium chloride (KCl), sodium carbonate (Na₂CO₃), sodium hydroxide (NaOH), sodium acetate (CH₃COONa), and sodium nitrate (NaNO₂) were purchased from Sigma-Aldrich (St. Louis, MO).

**PLANT MATERIAL.** Experiment 1 investigated the effects of N fertilization rate on fruit quality of subtropical peach cultivars TropicBeauty and UFSharp. Experiment 2 investigated the effects of harvest date (early, mid, and late harvest) on the fruit quality in ‘TropicBeauty’. Experiment 1 used 4-year-old subtropical peach trees of ‘TropicBeauty’ and ‘UFSharp’ grafted on ‘Flordaguard’ rootstock planted at 7.6 m between rows and 4.8 m between trees grown at the University of Florida Plant Science Research and Education Unit, Citra, FL (UF PSREU). The experimental design was a split plot design (N = 4), where two sets of five trees were regarded as one replicate. Five rates of N treatment (0, 45, 90, 179, and 269 kg N ha⁻¹ per year) were in the form of ammonium nitrate [NH₄NO₃ (Liberty Acres Fertilizer, Darlington, SC)] were applied annually from the time of planting (2010). The fertilizer was applied through the irrigation system and was divided into 12 applications per year. The trees were pruned, irrigated, and fruit thinned according to standard industry practices to minimize the environmental component (Olmstead et al., 2013). Fruit from both cultivars, TropicBeauty and UFSharp, were harvested on 11 June 2014. All of the fruit meeting commercially accepted standards for ground color and fruit size (Crisosto and Valero, 2008) were harvested by trained harvesters. A subsample (∼10 pieces of fruit) of harvested fruit was immediately pitted, chopped with skin, frozen in liquid nitrogen, and stored at −80°C until samples were processed for further analysis.

Experiment 2 used 7-year-old ‘TropicBeauty’ trees grafted to ‘Flordaguard’ rootstock planted 7.6 m between rows and 4.8 m between trees and grown at the UF PSREU. Treatments were replicated in a completely randomized design (N = 3), where two trees were regarded as one replicate (number of replicates were reduced in this experiment due to limitation in available trees). Four rates of NH₄NO₃ (45, 90, 179, and 269 kg·ha⁻¹ per year) were applied annually, plus a no N control (0 kg·ha⁻¹ per year). These treatments were initiated in 2011 when the trees were 4 years old; before initiation of N treatments, all of the trees were on the same fertilization program (90 kg·ha⁻¹ per year N). Trees were pruned, irrigated, and fruit thinned according to standard industry practices to minimize the environmental component (Olmstead et al., 2013). All of the fruit meeting commercially accepted standards for ground color and fruit size (Crisosto and Valero, 2008) were harvested by trained harvesters. Harvests occurred on 22 May 2014, 2 June 2014, and 12 June 2014, representing early, mid, and late harvests, respectively (commercial harvest date for ‘Tropic Beauty’ was in the same date range). A subsample of harvested fruit was immediately pitted, chopped with skin, and frozen in liquid nitrogen and stored at −80°C until samples were processed for further analysis.

**FRUIT QUALITY ANALYSIS.** TSS and TA were measured for a subset of chopped fruit. One hundred (±0.5) grams of chopped fruit were homogenized and centrifuged at 5000 g for 10 min (Sorvall Legend XTR Refrigerated; Thermo Fisher Scientific, Waltham, MA). The supernatant of homogenate was used for measuring TSS with a refractometer (Pocket PAL-1; Atago USA, Bellevue, WA) and TA with a titrator (808 Titrand; Metrohm, Riverview, FL). Titratable acidity was determined by titrating 6 mL of juice with 0.1 N NaOH to a pH endpoint of 8.2. The results were expressed as percent malic acid. The ripening index (RI) was calculated as the TSS/TA ratio.

Citric and malic acid concentrations were determined by using the enzymatic ultraviolet method. Quantification was done using kits for citric acid and L-malic (catalog no. 10139076035 and no. 10139068035; R-Biopharm, Darmstadt, Germany). Briefly, the frozen fruit material was homogenized to a fine powder without letting it thaw and then 1 g fresh weight (FW) of fine powder was mixed with 1 mL deionized water at room temperature. Clear supernatant was removed from each sample after centrifugation and citric and L-malic acid quantification was done as per manufacturer’s instructions. Absorbance was measured at 340 nm on a spectrophotometer (Eppich Microplate; Biotek, Winookski, VT). Results were reported as milligram per milliliter of subtropical peach juice.

**PHYSIOCHEMICAL ANALYSIS.** Hydrochloric acid extraction was used for quantification of phenolic content, flavonoid content, anthocyanin content, and antioxidant activity as described by Cantin et al. (2009). Briefly, the frozen fruit material was homogenized to fine powder without letting it thaw and then 5 g of fine powder was mixed with 15 mL of extraction solution, consisting of 0.5 N HCl in methanol/Milli-Q water (80% v/v; Millipore Sigma, Billerica, MA). The mixture was incubated overnight at 4°C and then centrifuged for 20 min (20,000 g) at 4°C. The supernatant was recovered and used for further phytochemical analysis.

Phenolic content of extracts was determined by the Folin–Ciocalteu assay as described by Vashisth et al. (2011). Briefly, 0.5 mL of extract was pipetted into a test tube containing 8 mL of deionized water. Then, 0.5 mL of Folin–Ciocalteu’s phenol reagent (Millipore Sigma) was added and followed by 1 mL of a saturated Na₂CO₃ solution. The contents in each test tube were vortexed for 10 s and left to stand at room temperature for 1 h to allow for maximum color development. Three hundred microliter aliquots were transferred from each test tube into a 96-well microtitre plate and absorbance readings were taken at 750 nm using a spectrophotometer (Synergy HTX Multi-mode Reader; Biotek). The standard curve was constructed with varying concentrations from a stock solution by dissolving 80 mg (470.2 µmol) of gallic acid in 25 mL of methanol and then diluting it with deionized water to 1 L. The total phenolic contents of samples were expressed as gallic acid equivalents [GAE (milligram per 100 g FW)].

Flavonoid content was determined using a colorimetric assay (Zhishen et al., 1999). One milliliter of extract was diluted with water (1:2), and 0.3 mL of 5% NaNO₂ was added. After 5 min, 0.3 mL of 10% AlCl₃ was added. After 1 min, 2 mL of 1 N NaOH was added and vortexed. Absorbance was measured at 510 nm. The results were expressed as catechin equivalents [CAE (milligram per 100 g FW)] on the basis of a standard curve using catechin as a standard.

Anthocyanin content was determined by the pH-differential method (Lee et al., 2005). Briefly, 1 mL of extracted solution was dissolved into a 10-mL volumetric flask and used to prepare two dilutions of the sample: one adjusted volume with KCl buffer, pH 1.0, and the other with CH₃COONa buffer, pH 4.5. After solutions were equilibrated, the absorbance of each dilution was measured at 510 and 700 nm against a blank cell filled with distilled water. The absorbance of diluted samples was calculated as follows:
where A is the absorbance of sample at 510 and 700 nm.

The monomeric anthocyanin pigments were calculated by using the formula below:

\[ \text{Monomeric anthocyanin pigment} = \frac{(A \times \text{MW} \times \text{DF} \times 1000)}{(\varepsilon \times 1)} \]

where MW is the molecular weight, DF is the dilution factor, and \( \varepsilon \) is the molar absorptivity, calculate pigment content as cyanidin-3-glucoside, where MW = 449.2 and \( \varepsilon = 26,900 \) and results expressed as cyanidin-3-glucoside equivalents [C3GE (milligram per kilogram FW)].

**Antioxidant Capacity.** Two assays that are commonly used to measure antioxidant capacity are the ORAC assay (oxygen radical absorption capacity) for the hydrogen atom transfer mechanisms and the FRAP assay (ferric reducing antioxidant power) for the single electron transfer mechanisms. The FRAP assay measures the reducing power of samples (Benzie and Szeto, 1999) and, therefore, indicates a better antioxidant capacity of a given product. Total antioxidant activity of prepared extracts was determined by the FRAP assay as described by Pulido et al. (2000) with modifications. The FRAP reagent was prepared fresh by combining 5 mL of a 10-mmol L\(^{-1}\) TPTZ solution in 40 mmol L\(^{-1}\) HCl with 5 mL of 20 mmol L\(^{-1}\) FeCl\(_3\)-6H\(_2\)O and 50 mL of a 0.3 mol L\(^{-1}\) CH\(_3\)COONa buffer at pH 3.6. The FRAP reagent was warmed to 37 °C in a water bath (Isotemp model 2239; Thermo Fisher Scientific) and held at this temperature until used. Aqueous solutions of known Fe (II) (FeSO\(_4\)·7H\(_2\)O) concentrations in the range of 100 to 2400 mmol L\(^{-1}\) were used to construct the standard curve. In a microtitre plate, 1500 \( \mu \)L of FRAP reagent were mixed with 150 \( \mu \)L of deionized water and 50 \( \mu \)L of the test sample or blank (i.e., 1:1 aqueous methanol). Before reading the absorbance, the plate was incubated at 37 °C for 30 min and agitated for 5 s; the absorbance readings of the samples (at 37 °C) were recorded at 595 nm. Antioxidant activity of the prepared extracts was expressed as Ferrous equivalents [Fe\(^{2+}\)E (millimoles per gram FW)].

**Statistical Analysis.** All statistical analyses were performed using two-way analysis of variance (ANOVA) at \( \alpha = 0.05 \) using SigmaPlot (version 11; Systat Software, San Jose, CA). Mean separation among treatments with significant differences was performed using Tukey’s honest significance test (HSD) at \( \alpha = 0.05 \).

**Results**

**EXPERIMENT 1.** The total yield for both cultivars, TropicBeauty and UFSsharp, as response to different N treatment are presented in Table 1. The moderate N treatment resulted in highest yield as compared with 45 and 179 kg ha\(^{-1}\) per year N treatment. The TSS, TA, and RI for both cultivars, TropicBeauty and UFSsharp, are presented in Table 2. The RI for both ‘TropicBeauty’ and ‘UFSsharp’ in all N treatments were not significantly different \( (P > 0.05) \) suggesting that the fruit were at a similar ripening stage across the treatments and cultivars. In ‘TropicBeauty’, N treatments did not affect TSS; however, in ‘UFSsharp’, both low and moderate N supply, 0 and 90 kg ha\(^{-1}\) per year, resulted in higher TSS content, whereas high rates of 179 and 269 kg ha\(^{-1}\) per year resulted in \( \approx 40\% \) lower TSS. TA was significantly higher \( (P < 0.05) \) overall in ‘UFSsharp’ than ‘TropicBeauty’; however, there was no effect of N treatment. The average citric acid concentration in ‘TropicBeauty’ and ‘UFSsharp’ was 1.4 mg mL\(^{-1}\) and 1.6 mg mL\(^{-1}\), respectively. A significant interaction between cultivar and N treatment \( [C \times N \; (P = 0.043)] \) was found for citric acid quantification (Table 3). In ‘UFSsharp’, citric acid concentration was affected by the N treatment with the lowest citric acid concentration found in the absence of N [negative control (0 kg ha\(^{-1}\) per year)]. In ‘TropicBeauty’, citric acid content did not change as a result of N treatment. Malic acid content was significantly lower in ‘TropicBeauty’ than in ‘UFSsharp’ \( (P = 0.004) \). The average malic acid in ‘TropicBeauty’ and ‘UFSsharp’ was 4.1 and 4.7 mg mL\(^{-1}\), respectively (Table 3). Malic acid content was found to be unaffected by N treatment in both the cultivars.

Both cultivars had significantly higher phenolic content in low N rates, and as the rate of N increased, phenolic content decreased (Fig. 1). An approximate 100% increase in phenolic content occurred in trees not receiving N (0 kg ha\(^{-1}\) per year) as compared with the highest N rate treatment, 269 kg ha\(^{-1}\) per year, in ‘TropicBeauty’. Similarly, in ‘UFSsharp’, the highest phenolic content was observed in fruit from trees receiving 45 kg ha\(^{-1}\) per year as compared with 269 kg ha\(^{-1}\) per year, with a difference of \( \approx 60\% \) between the two rates. The phenolic content was found to be linearly dependent on the rate of N for both ‘TropicBeauty’ \( [P < 0.001, \; R^2 \text{ value } 0.90 \) (Fig. 2)] and ‘UFSsharp’ \( (P = 0.013) \). A significant interaction between cultivar and N treatment was observed \( (C \times N, \; P = 0.012) \), suggesting that the N treatment effect was dependent on the cultivar. For example, phenolic content was greatly reduced with high N rate treatment in ‘TropicBeauty’ as compared with ‘UFSsharp’ (Fig. 1). As expected, flavonoid content in ‘TropicBeauty’ and ‘UFSsharp’ followed a similar trend as phenolic content, with lower N rates resulting in higher flavonoid content (Fig. 3). Phenolic content and flavonoid content were linearly correlated with \( R^2 \) values of 0.82 and 0.73 (data not shown) for ‘TropicBeauty’ and ‘UFSsharp’, respectively. Similar to phenolic content, an increase of more than 200% in flavonoid content was observed in trees receiving no N (0 kg ha\(^{-1}\) per year) compared with 269 kg ha\(^{-1}\) per year in ‘TropicBeauty’. A significant interaction between cultivars and N treatments \( (C \times N, \; P = 0.009) \) was also observed; increasing N rate decreased the flavonoid content greater in ‘TropicBeauty’ than in ‘UFSsharp’. No significant N rate and cultivar interaction was observed for anthocyanin (Fig. 4). Overall, anthocyanin content in ‘TropicBeauty’ was significantly higher \( (P < 0.001) \) than ‘UFSsharp’ irrespective of N treatment.
The overall antioxidant activity was significantly higher in ‘UFSharp’ than in ‘TropicBeauty’ \( P = 0.026 \) (Fig. 5). As the N rate increased, a decrease in antioxidant activity was observed in ‘TropicBeauty’, with a similar trend in ‘UFSharp’. The antioxidant activity followed similar trends to phenolic content and flavonoid content and a significant linear relationship between antioxidant capacity and phenolic content was found in ‘TropicBeauty’ and ‘UFSharp’ \( R^2 = 0.95 \) and 0.52, respectively (data not shown). A significant interaction between cultivars and N treatment \( (C \times N, P = 0.03) \) indicated that cultivar played a critical factor in effect of N treatment. For example, ‘TropicBeauty’ had a more pronounced decrease in antioxidant activity as the N treatment increased in comparison with ‘UFSharp’.

**Table 2.** Total soluble solids (TSS), titratable acidity (TA), and ripening index (RI) for fruit of subtropical peach cultivars, TropicBeauty and UFSharp, grown under five different nitrogen treatments.

| N treatment (kg ha\(^{-1}\) per yr) | TropicBeauty | UFSharp | TropicBeauty | UFSharp | TropicBeauty | UFSharp |
|------------------------------------|--------------|---------|--------------|---------|--------------|---------|
| 0                                  | 12.20 ± 1.80 bA | 19.30 ± 6.85 aAB | 0.55 ± 0.05  | 0.72 ± 0.45  | 21.98 ± 1.46  | 31.77 ± 17.04  |
| 45                                 | 16.43 ± 7.74 aA | 11.88 ± 5.59 aC | 0.55 ± 0.03  | 0.80 ± 0.10  | 30.15 ± 14.51  | 15.73 ± 10.01  |
| 90                                 | 13.00 ± 0.57 bA | 22.65 ± 4.61 aA | 0.57 ± 0.03  | 0.75 ± 0.05  | 22.76 ± 0.52  | 29.92 ± 4.36  |
| 179                                | 11.55 ± 0.79 aA | 14.23 ± 3.45 aBC | 0.52 ± 0.02  | 0.65 ± 0.18  | 22.41 ± 1.96  | 22.14 ± 1.96  |
| 269                                | 12.73 ± 1.16 aA | 12.75 ± 3.95 aC | 0.68 ± 0.16  | 0.72 ± 0.12  | 19.18 ± 2.62  | 17.75 ± 5.38  |

Means followed by different lowercase letters indicate statistically significant differences between cultivars within an N rate \( P = 0.05 \); means followed by uppercase letters indicate statistically significant differences between N rates within a cultivar \( P = 0.05 \); means followed by no letter are not significantly different. Mean separation was performed using Tukey’s honest significance test (HSD) at \( \alpha = 0.05 \).

Table 3. Citric and malic acid levels for fruit of subtropical peach cultivars, TropicBeauty and UFSharp, from trees grown under five different nitrogen treatments.

| N treatment (kg ha\(^{-1}\) per yr) | Citric acid [mean ± sd (mg mL\(^{-1}\))] | Malic acid [mean ± sd (mg mL\(^{-1}\))] |
|------------------------------------|------------------------------------------|-----------------------------------------|
|                                    | TropicBeauty | UFSharp | TropicBeauty | UFSharp | TropicBeauty | UFSharp |
| 0                                  | 1.4 ± 0.24  | 1.02 ± 0.17 B | 4.65 ± 0.38  | 5.32 ± 0.74  |
| 45                                 | 1.6 ± 0.43  | 1.52 ± 0.45 AB | 4.22 ± 0.75  | 5.85 ± 0.55  |
| 90                                 | 1.5 ± 0.14  | 1.77 ± 0.45 A | 3.8 ± 0.35  | 4.62 ± 0.72  |
| 179                                | 1.27 ± 0.25 b | 2.02 ± 0.35 aA | 4.1 ± 1.20  | 3.92 ± 0.75  |
| 269                                | 1.27 ± 0.52  | 1.65 ± 0.25 AB | 3.57 ± 0.56  | 3.87 ± 0.27  |

Means followed by different lowercase letters indicate statistically significant differences between cultivars within an N rate; means followed by uppercase letters indicate statistically significant differences between N rates within a cultivar; means followed by no letter are not significantly different. Mean separation was performed using Tukey’s honest significance test (HSD) at \( \alpha = 0.05 \).

In ‘TropicBeauty’, with a similar trend in ‘UFSharp’. The antioxidant activity followed similar trends to phenolic content and flavonoid content and a significant linear relationship between antioxidant capacity and phenolic content was found in ‘TropicBeauty’ and ‘UFSharp’ \( R^2 = 0.95 \) and 0.52, respectively (data not shown). A significant interaction between cultivars and N treatment \( (C \times N, P = 0.03) \) indicated that cultivar played a critical factor in effect of N treatment. For example, ‘TropicBeauty’ had a more pronounced decrease in antioxidant activity as the N treatment increased in comparison with ‘UFSharp’.

**Experiment 2.** The average yield in response to different N treatment varied. Moderate N (90 and 179 kg ha\(^{-1}\) per year) treatment resulted in highest yield, whereas 40 and 269 kg ha\(^{-1}\) per year N treatment resulted in lowest yield (data not shown). No significant differences were observed in yield at different harvest times. No significant difference in TSS content as influenced by harvest dates and N treatments were observed \( P > 0.05 \), although an increasing trend in the average TSS on each harvest date was noticed (Table 4). No significant differences were observed in TA among harvest date and N treatment \( P > 0.05 \) (Table 4).
significant decrease \((P < 0.05)\) in TA from early to late harvest was observed. No significant interaction for harvest time and N treatment was found for RI \((P > 0.05)\). As expected, a significant increase in ripening index was observed from early to late harvest \([P < 0.001 \text{ (Table 4)}]\). Similar to TA, average citric acid content was observed to be significantly different \([P < 0.001 \text{ (Table 5)}]\) at each harvest date, although no interaction was observed between harvest date and N treatment \((P > 0.05)\). No effects in malic acid content were observed because of harvest time or N treatment \([P > 0.05 \text{ (Table 5)}]\).

A significant interaction between harvest date and N treatment for phenolic content \([H \times N, P < 0.001 \text{ (Table 6)}]\) was observed. Total phenolic content among all three harvest dates and N treatments was found to be significantly different \((P < 0.001)\) with an exception of treatment 179 kg-ha\(^{-1}\) per year and an overall trend of reduced phenolic content with later harvests. Total flavonoid content followed a similar trend as phenolic content. A significant correlation between flavonoid content and phenolic content was observed for all three harvest dates and N treatments (data not shown). A significant interaction among harvest date and N treatment for flavonoid content was observed \([P < 0.001 \text{ (Table 6)}]\). Overall, flavonoid content decreased with advancement in harvest date. Significant differences among treatments in each harvest date were found, but overall no uniform trend was observed among different N rates. A significant increase in anthocyanin content was found with later harvest date, in addition to a significant interaction among harvest and N treatment \([P < 0.001 \text{ (Table 6)}]\). Total anthocyanin content was consistently high for all N treatments at the latest harvest date with a 40\% increase in anthocyanin content as compared with the early harvest date. The effect of different levels of N was not clear on anthocyanin content.

In regard to antioxidant capacity, a significant interaction among harvest and N treatment was observed \([P = 0.042 \text{ (Table 6)}]\). Total antioxidant capacity in each harvest date was observed to be affected by the N treatment. Total antioxidant capacity was found to be the highest in early harvest, followed by mid and late harvests \(\text{(Table 6)}\). A significant linear relationship between antioxidant capacity and phenolic content was observed among all three harvest dates, \(R = 0.82 (P < 0.001), 0.66 (P = 0.006), \) and \(0.62 (P = 0.014)\) for early, mid, and late harvest, respectively.

Interestingly, in mid and late harvest dates for phenolic content, flavonoid content, and anthocyanin content, all harvest
dates of antioxidant capacity of 0 kg·ha⁻¹ per year and 269 kg·ha⁻¹ per year N treatment were ranked among the highest, whereas most of the observations of 179 kg·ha⁻¹ per year N treatment displayed the lowest values for phenolic content, flavonoid content, anthocyanin content, and antioxidant capacity. Therefore, no clear effect of N treatments on phytochemical content and antioxidant activity was clear in Expt. 2. Overall, late harvest resulted in lowest phenolic content, flavonoid content, and antioxidant capacity and highest anthocyanin content.

**Discussion**

Overall, subtropical peach trees responded to different N treatments; subtropical peach fruit quality and phytochemical composition was positively influenced with lower N rates. Average TA and organic acid content was higher in ‘UFSharp’ than ‘TropicBeauty’, although no effect of N treatment was found on TA or organic acid content for both cultivars. Inherent differences among ‘TropicBeauty’ and ‘UFSharp’ were evident suggesting that every genotype responds differently to abiotic conditions, and there is a possible genotype and mineral nutrient interaction which results in a differential response of two cultivars toward the same treatments. In both experiments, TSS in ‘TropicBeauty’ was unaffected by N treatments, whereas an overall inverse relationship was observed between TSS and N treatments. Similar to our results, an increase in TSS in tomato fruit with decreasing N rate has been reported previously (Simonne et al., 2007), and an increase of 5% to 17% in TSS with a concomitant decrease in TA was observed when N supply was reduced to half of the amount (Bénard et al., 2009). In addition, an increase in individual sugars (glucose, sucrose, and fructose) with a decrease in acids (citrate, isocitrate, fumarate, and malate) was found in tomato leaves when grown in N-deficient medium (Urbanczyk-Wochniak and Fernie, 2005).

Harvest date had little influence on subtropical peach fruit quality, increase in TSS content; a decreasing trend in TA was observed with advancement of harvest date. RI index for late harvest was observed to be significantly higher, suggesting progression of ripening of the fruit at late harvest time point.

A decrease in phenolic content and flavonoid content was observed with an increasing rate of N application, although this effect seemed to be more pronounced in ‘TropicBeauty’ than ‘UFSharp’ suggesting inherent cultivar differences and strong differences among ‘TropicBeauty’ and ‘UFSharp’ were evident suggesting that every genotype responds differently to abiotic conditions, and there is a possible genotype and mineral nutrient interaction which results in a differential response of two cultivars toward the same treatments. In both experiments, TSS in ‘TropicBeauty’ was unaffected by N treatments, whereas an overall increase in TSS with decreasing N rate has been reported previously (Simonne et al., 2007), and an increase of 5% to 17% in TSS with a concomitant decrease in TA was observed when N supply was reduced to half of the amount (Bénard et al., 2009). In addition, an increase in individual sugars (glucose, sucrose, and fructose) with a decrease in acids (citrate, isocitrate, fumarate, and malate) was found in tomato leaves when grown in N-deficient medium (Urbanczyk-Wochniak and Fernie, 2005).

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A decrease in phenolic content and flavonoid content was observed with an increasing rate of N application, although this effect seemed to be more pronounced in ‘TropicBeauty’ than ‘UFSharp’ suggesting inherent cultivar differences and strong
Table 4. Total soluble solids (TSS), titratable acidity (TA) and ripening index (RI) for fruit of subtropical peach cultivar, TropicBeauty, grown under five different nitrogen treatments. Fruit were harvested on 22 May, 2 June, and 12 July 2014, representing early, mid, and late harvests, respectively.

| N treatment (kg ha⁻¹ per yr) | Early          | Mid          | Late          | Early          | Mid          | Late          | Early          | Mid          | Late          |
|-----------------------------|----------------|--------------|---------------|----------------|--------------|---------------|----------------|--------------|---------------|
| 0                           | 10.07 ± 5.49   | 9.1 ± 3.83   | 13.17 ± 3.84  | 1.043 ± 0.07 a | 0.66 ± 0.10 b | 0.50 ± 0.05 b | 9.66 ± 5.10 b   | 13.38 ± 3.76 b | 26.89 ± 10.53 a |
| 45                          | 8.9 ± 1.84     | 12.97 ± 3.75 | 11.03 ± 2.47  | 0.78 ± 0.25 a  | 0.79 ± 0.03 a | 0.62 ± 0.06 b | 12.44 ± 4.86    | 16.34 ± 4.22   | 17.57 ± 2.62   |
| 90                          | 13.3 ± 2.59    | 9.8 ± 2.26   | 8.8 ± 1.01    | 0.88 ± 0.09 a  | 1.10 ± 0.78 a | 0.58 ± 0.02 b | 15.04 ± 2.21    | 11.39 ± 6.02   | 15.11 ± 0.43   |
| 179                         | 10.37 ± 0.40   | 11.27 ± 0.21 | 10.7 ± 0.85   | 0.64 ± 0.05 a  | 0.60 ± 0.01  | 0.54 ± 0.09   | 16.37 ± 1.75 b  | 18.70 ± 0.15 a | 20.01 ± 1.63 a |
| 269                         | 10.93 ± 0.81   | 11.43 ± 0.12 | 14.73 ± 5.95  | 0.93 ± 0.12 a  | 0.67 ± 0.01 b | 0.56 ± 0.05 c | 11.80 ± 0.95 b  | 18.21 ± 0.32 a | 25.98 ± 7.83 a |

Means followed by different lowercase letters indicate statistically significant differences between harvest dates within an N rate; means followed by no letter are not significantly different. Mean separation among treatments with significant differences was performed using Tukey’s honest significance test (HSD) at α = 0.05.

RI = TSS/TA.

Table 5. Citric and malic acid levels for fruit of subtropical peach cultivar, TropicBeauty, from trees grown under five different nitrogen treatments. Fruit were harvested on 22 May, 2 June, and 12 June 2014, representing early, mid, and late harvests, respectively.

| N treatment (kg ha⁻¹ per yr) | Early     | Mid        | Late       | Early     | Mid        | Late       |
|-----------------------------|-----------|------------|------------|-----------|------------|------------|
| 0                           | 4.30 ± 0.76 | 3.03 ± 1.3 | 1.50 ± 0.4 | 4.33 ± 0.68 | 3.76 ± 0.35 | 4.66 ± 0.47 |
| 45                          | 3.45 ± 0.07 | 3.73 ± 1.3 | 2.23 ± 0.40 | 4.53 ± 0.73 | 4.63 ± 0.65 | 4.36 ± 0.75 |
| 90                          | 5.1 ± 0.65  | 3.23 ± 0.5  | 1.66 ± 0.05 | 4.36 ± 0.85 | 4.36 ± 0.15 | 4.26 ± 0.68 |
| 179                         | 3.46 ± 0.94 | 2.33 ± 0.77 | 1.3 ± 0.3   | 3.5 ± 1.2  | 4.13 ± 0.3  | 4.26 ± 0.86 |
| 269                         | 3.9 ± 0.5   | 2.16 ± 0.66 | 1.29 ± 0.61 | 4.36 ± 0.47 | 4 ± 1.2     | 3.7 ± 1.44  |

Means followed by no letter are not significantly different. Mean separation among treatments with significant differences was performed using Tukey’s honest significance test (HSD) at α = 0.05.
equivalents (C3GE), and total antioxidant capacity (TAC) in ferrous equivalents (Fe$_2^+$E) for fruit of subtropical peach cultivar, TropicBeauty, grown under five different nitrogen

| Treatment | TPC [mg GAE per 100 g FW] | TFC [mg CAE per 100 g FW] | ATC [mg C3GE per kg FW] | TAC [mg Fe$_2^+$E per g FW] |
|-----------|---------------------------|---------------------------|-------------------------|-----------------------------|
| Early     | 45 42.39 ± 2.12 aB        | 40.16 ± 2.94 abAB         | 35.43 ± 2.34 bAB        | 5.62 ± 0.74 abCD            |
| Mid       | 90 42.95 ± 3.50 aB        | 41.34 ± 4.29 aAB          | 29.87 ± 0.27 bC         | 6.85 ± 1.30 aB              |
| Late      | 179 34.30 ± 2.69 aC        | 33.86 ± 2.53 aC           | 32.76 ± 0.99 aBC        | 4.87 ± 0.87 aD              |
| Early     | 269 57.57 ± 6.08 aA        | 44.51 ± 2.98 bA           | 39.83 ± 4.69 bA         | 8.35 ± 0.51 aC              |

Means followed by different lowercase letters indicate statistically significant differences between harvests within an N rate; means followed by uppercase letters indicate statistically significant differences between N rates within a harvest; means followed by no letter are not significantly different. Mean separation was performed using Tukey’s honest significance test (HSD) at α = 0.05.

Anthocyanin content values were well in range of values reported in other subtropical peach studies (Cantin et al., 2009). Anthocyanin content in subtropical peaches is dependent on genetics as well as environmental factors; for example, Reig et al. (2013) reported as much as 11-fold changes in the anthocyanin content observed were due to different nitrogen treatments applied from the time of establishment).

The production of photoprotective pigments such as flavonols and anthocyanins may provide protection against light-induced oxidative damage (Guidi et al., 1998). Flavonols are known to accumulate in the skins of tomato fruit (Stewart et al., 2000) and, therefore, filter out damaging wavelengths of radiation. We also observed a decrease in phenolic content and flavonoid content with advancement in harvesting dates; such a decrease in phenolic content and flavonoid content can be potentially significant because of the changes in weather condition. As the harvest date progressed later in the season, the overcast and rainfall (as suggested by Florida Automated Weather Network, UF) increased which could have contributed to a decrease in phenolic content and flavonoid content. Polyphenolic compounds are reported to vary greatly among cultivars (Aaby et al., 2007; Tulipani et al., 2008), cultural systems, and environmental factors. Flavonoid accumulation can be induced by a number of environmental conditions, including ultraviolet light (Winkel-Shirley, 2002), pathogen attack (Dixon and Paiva, 1995), and nutrient deficiencies (Stewart et al., 2001). The lack of significant trends in phenolic content and flavonoid content observed in Expt. 2 might be due to the tree age; trees used in this experiment were mature, large, and well-established trees and, therefore, may not have been as influenced by different rates of N applied as younger trees (as in Expt. 1 where treatments were applied from the time of establishment). Anthocyanin content values were well in range of values reported in other subtropical peach studies (Cantin et al., 2009). Anthocyanin content in subtropical peaches is dependent on genetics as well as environmental factors; for example, Reig et al. (2013) reported as much as 11-fold changes in the anthocyanin content among different cultivars. Similar results were reported by Cantin et al. (2009) in subtropical peach and nectarine (P. persica) progenies. In Expt. 1, anthocyanin content in 'TropicBeauty' was higher than 'UF Sharp' irrespective of N treatment suggesting cultivar differences. This is also supported by visual fruit observation and description; 'Tropic Beauty' develops about 70% blush at maturity, whereas the 'UF Sharp' develops up to 60% blush (Olmstead et al., 2013).
Medium to low rates of N have been reported to result in higher concentration of anthocyanin in a number of fruit such as subtropical peaches (Cordts et al., 1987), grapes, and strawberries (Yoshida et al., 2003). In this present study, an increase in anthocyanin content was also observed at a later harvest date coinciding with previous studies in southern highbush blueberry (Prior et al., 1998), sour cherry [Prunus cerasus (Poll et al., 2003)], and apple (Reay and Lancaster, 2001). Overall, manipulation of N and harvest date can be used to stimulate increases in polyphenolic compounds that can increase the nutritional value in crops.

Many studies have reported a correlation of antioxidant activity with phenolic, flavonoid, and anthocyanin content in animals and humans. The correlation found between plasma endogenous antioxidant and circulating level of dietary phenolics have been shown in many fruit crops such as strawberry (Azzini et al., 2010) and blackberry [Rubus sp. (Srivastava et al., 2010)]. A significant and positive relationship was observed between antioxidant activity and phenolic content in both cultivars and harvest dates. A reduction in antioxidant capacity was observed with an increasing N rate in ‘TropicBeauty’ and ‘UFSharp’, but no consistent differences in N treatments were observed in different harvests, although antioxidant capacity decreased at later harvest date. Similar data were also reported in subtropical peaches and nectarines (Font i Forcada et al., 2012; Gil et al., 2002), apples (Lata, 2007), sweet cherries [Prunus avium (Serrano et al., 2005)], and plums (Gil et al., 2002).

**Conclusion**

Overall, differences in phenolic content, flavonoid content, anthocyanin content, and antioxidant capacity were observed among different rates of N in subtropical peach cultivars, TropicBeauty (melting flesh) and UFSharp (nonmelting flesh). The differences in phytochemical content and antioxidant activity between early, mid, and late harvest dates were also observed but no consistent trends or patterns were observed in N treatments in Expt. 2. Experiment 1 was conducted in a young orchard with the treatments applied since establishment, which could have resulted in differences among treatments being more pronounced. Experiment 2 was conducted in a mature orchard and treatments were applied for the last 3 years, which could have potentially contributed to such results as before the application of treatments all trees were receiving the same rate of N. Overall, these data suggest that the concentration of health benefiting compounds such as polyphenolics and antioxidants can be altered with cultural practices such as fertilization and harvest maturity. Manipulations to cultural practices such as reduced N fertilization or early harvest can be easily done without any added cost to obtain fruit with higher health benefits, and thus, provide consumers an added advantage with consumption of fresh subtropical peaches.

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