Canopy Light Effects in Multiple Training Systems on Yield, Soluble Solids, Acidity, Phenol and Flavonoid Concentration of ‘Frontenac’ Grapes

Christina M. Bavougian¹,⁴, Paul E. Read¹, Vicki L. Schlegel², and Kathryn J. Hanford³

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SUMMARY. Phenolic compounds contribute greatly to the sensory attributes of wine and have a wide range of human health benefits as well. In this study, four trellis/training systems were evaluated for effects on fruit-zone light environment, fruit chemical composition (including phenol and flavonoid concentrations), and yield of ‘Frontenac’ grapes (*Vitis* sp. MN 1047) grown in southeastern Nebraska over two growing seasons. Photosynthetically active radiance (PAR) was measured above the canopy and within the fruiting zone at berry set, veraison, and harvest. Point quadrat canopy analysis was performed at veraison. Both bound and free (un-bound) flavonoid and total phenolic contents were determined for the skins and seeds of fruit samples in 2008. At all sampling dates in 2008, vines grown on Geneva double curtain (GDC) and high cordon (HC) had higher midday percentage PAR transmittances than vines grown on Smart-Dyson (SD) and vertical shoot positioned (VSP) training systems. In 2009, transmittance relationships between trellises were not consistent throughout the season. In both years, leaf layer number (LLN) was lower for GDC and HC than for SD and VSP. Flavonoid and total phenol concentrations of the bound seed and bound skin extracts did not differ among trellises. Within the free extracts, VSP had higher total phenol concentration than SD (GDC and HC were intermediate) and there were no differences in flavonoid concentration. In 2008, GDC had higher pH than other trellises and higher soluble solids than SD and VSP; titratable acidity (TA) was lower in GDC and HC than in SD and VSP. In 2009, SD and VSP had the highest soluble solids concentrations; HC had lower pH than SD and VSP and there were no differences in TA. Results were incongruent regarding light environment effects on fruit chemical composition.

Phenolic compounds include a diverse group of substances that perform a variety of functions in vascular plants. Phenolics are commonly divided into flavonoid (tannins, anthocyanins, flavan-3-ols, and flavonols) and nonflavonoid (hydroxycinnamates and stilbenes) compounds (Cheynier et al., 1999). Flavonoids are differentiated from nonflavonoid phenolics by their carbon skeleton (Kennedy et al., 2006). Most phenolics are purported to protect plants from ultraviolet radiation damage; their positive effects on human health. The capacity to neutralize free radicals (reactive molecules that cause mutations, heart disease, skin problems, and aging), also called antioxidant activity (AOA), is common to phenolics as a group (Jacopini et al., 2008). The AOA of many polyphenols is much greater than that of the essential dietary vitamins, and grapes and wine contain some of the highest phenol concentrations in the human diet (Jacopini et al., 2008). In addition to their antioxidant properties, phenolics act as anti-inflammatory agents to improve human health. Polyphenols in wine reduce the risk of cardiovascular disease (Fang et al., 2008) in two ways: by inhibiting the aggregation of some help defend against bacterial and fungal infection (Downey et al., 2006). Anthocyanins and other pigments attract animals that perform such services as pollination and seed dispersal (Downey et al., 2006). Tannins and flavan-3-ols discourage herbivory because of their astringent and/or bitter flavors (Santos-Buelga and Scalbert, 2000).

In addition to their many functions within plants, polyphenols contribute greatly to the sensory attributes of wine; increased concentration has generally been associated with high quality (Cheynier et al., 2006; Reynolds et al., 1995; Ristic et al., 2007). According to Downey et al. (2006), wine flavonoid content is mostly determined by grape composition and only partly by winemaking procedures; however, Holt et al. (2008a, 2008b) demonstrated that increased total phenol, tannin, and anthocyanin concentrations in must did not have an effect on wine quality scores.

A majority of the research on phenolic compounds has focused on their positive effects on human health. The capacity to neutralize free radicals (reactive molecules that cause mutations, heart disease, skin problems, and aging), also called antioxidant activity (AOA), is common to phenolics as a group (Jacopini et al., 2008). The AOA of many polyphenols is much greater than that of the essential dietary vitamins, and grapes and wine contain some of the highest phenol concentrations in the human diet (Jacopini et al., 2008). In addition to their antioxidant properties, phenolics act as anti-inflammatory agents to improve human health. Polyphenols in wine reduce the risk of cardiovascular disease (Fang et al., 2008) in two ways: by inhibiting the aggregation of...
platelets and by protecting low-density lipoprotein (LDL) cholesterol against oxidation (Santos-Buelga and Scalbert, 2000). Phenolics also protect against lung cancer, act as general antitu- moral and anticarcinogenic agents, reduce systolic blood pressure, and lower plasma cholesterol levels (Fang et al., 2008; Santos-Buelga and Scalbert, 2000).

The light environment within the canopy is the most important factor influencing grapevine yield and quality (Dokoozlian and Kliewer, 1995; Smart and Robinson, 1991). Trellises or training systems determine the spacing and orientation of shoots, thereby controlling light interception in the canopy (Dokoozlian and Kliewer, 1995; Howell et al., 1991). It is generally agreed that shading of berries reduces total phenol concentrations (Cortell and Kennedy, 2006; Morrison and Noble, 1990; Wolf et al., 2003), and that fruit shading reduces wine phenolic content (Joscelyne et al., 2007; Macaulay and Morris, 1993; Ristic et al., 2007), but results have not always been consistent.

As well as affecting the total phenol concentration in grapes, shading also changes the relative abundance of various phenolic components. Shade treatments increased seed tannins (low molecular weight) relative to skin tannins (high molecular weight) (Ristic et al., 2007). When Downey et al. (2004) applied artificial shading to `Shiraz' clusters, flavonol synthesis was decreased although there was no significant effect on anthocyanin or tan- nin concentrations. In another study, sunlight-exposed clusters contained nearly 10 times the amount of flavo- nol of fruit grown in the shade (Spayd et al., 2002). Some authors observed that fruit shading decreased total an- thocyanin concentration and altered its composition (Koyama and Gotou-Yamamoto, 2008; Price et al., 1995; Smart et al., 1988). Artificial shading did not change the amount of anthocy- anins in fruit although it did change the composition; however, wines made from the shaded grapes had lower anthocyanin concentration than wines made from sunlit grapes (Ristic et al., 2007). In a study of the effects of artificial shade and increased sun exposure on wine quality, wines made from shaded fruit contained lower concentrations of anthocyanins and were less astringent than control wines.

Yet anthocyanin concentration and astringency of wines made from experimentally exposed fruit did not differ significantly from control wines (Joscelyne et al., 2007). Morrison and Noble (1990) compared wines made from differently shaded `Cabernet Sauvignon' fruit. Cluster shading de- creased the anthocyanin content of wines, but sensory analysis yielded no corresponding differences in flavor or perceived aromas (Morrison and Noble, 1990). Total phenolic con- tent of `Shiraz' fruit was correlated with the percentage of ambient PAR (wavelength 400–700 nm) available in the fruit zone, but anthocyanin con- centration was not related to PAR (Wolf et al., 2003).

The influence of fruit-zone light environment on phenolic compounds in classical red wine grapes has been extensively studied, but similar re- search has not been performed on cold climate hybrid grapes in general and on `Frontenac' in particular. This is particularly significant because the genetic background of `Frontenac' con- sists largely of *Vitis riparia*, whereas cited studies involve *Vitis vinifera* cul- tivars (Minnesota Agricultural Experi- ment Station, 2008).

Because previous investigations have yielded such varied and complex results, it is problematic to draw con- clusions, especially for a relatively new cultivar. Therefore, this study was designed to determine if chemical composition of `Frontenac' fruit is influ- enced by light intensity within the canopy. In addition to soluble solids, pH, and titratable acidity, we measured total phenolic and flavo- noid concentrations of the fruit to provide a broader perspective.

**Materials and methods**

**Research site and experimental design.** Research for this study was conducted at a commercial vineyard 4 miles north of Wilber, NE (lat. 40°32′N, long. 97°W) during 2008 and 2009. The soil is a Crete series (very deep, moderately well-drained, fine Pachic Argiustoll) silty clay loam with 1% to 3% slope. The vineyard was not irrigated during the study. `Frontenac' vines (*Vitis* sp. MN 1047) planted in 2004–05 were trained to four different trellis styles GDC (horizontally divided bilateral HC system with downward shoot positioning), HC (single bilateral HC system with downward shoot positioning), SD (vertically divided canopy with upward and downward shoot positioning), and VSP (single bilateral low- or midcor- don system with upward shoot positioning). The training systems used in this study are described and illustrated in detail by Smart and Robinson (1991) and by Dami et al. (2005). Rows were oriented north-south with vines 2.4 m apart and rows 3 m apart. The training systems were each ap- plied to an entire 120-m row except for VSP (1.5 rows). The vineyard was subject to standard cultural practices for the area (permanent sod alleyways and bare under-row strips maintained with herbicide) and all vines were spru- pruned. Each vine was assigned a number; sample plants from each treatment were selected using a ran- dom number generator (*n* = 20 for GDC, HC, and SD; *n* = 30 for VSP). The number of sample plants was constrained by the time it took to record light measurements.

**Light measurements.** Photo- synthetically active radiation was measured with a line quantum sensor (LI-191; LI-COR Biosciences, Lincoln, NE) and recorded by a data logger (Polycorder 720 series; Wescon En- vironmental Products, Logan, UT) both above the canopy and within the fruit zones (8 inches above the cordon for SD and VSP, 8 inches below the cordon for GDC and HC) of sample plants. For the GDC vines we took fruit-zone PAR measure- ments in the west canopy, and for SD we used the upper canopy. PAR measurements were performed on three dates during each growing sea- son [25 June, 31 July, 29 Aug. 2008 (day of the year (DOY) 177, 213, 242); 29 June, 8 Aug., 18 Sept. 2009 (DOY 180, 220, 261)]. Sampling dates were chosen to correspond to key phenological stages: berry set, vera- son, and harvest. The dates we chose for sampling were near typical harvest dates for `Frontenac` in this area, although fruit was harvested before the corresponding PAR measure- ments in both years of study. On each sampling date, PAR measurements were obtained for each plant between 1200 and 1400 HR, within 1 h of solar noon. Sample plants were marked so that the instrument could be inserted into the canopy in the same place each time. Percentage PAR transmittances
Table 1. Mean midday fruit-zone percentage photosynthetically active radiation (PAR) transmittances of ‘Frontenac’ grapes grown on four training systems in southeastern Nebraska in 2008 and 2009.\(^a\)

<table>
<thead>
<tr>
<th>Trellis(^a)</th>
<th>2008 Bloom (DOY 177)(^a)</th>
<th>2008 Veraison (DOY 213)</th>
<th>2008 Harvest (DOY 242)</th>
<th>2009 Bloom (DOY 180)</th>
<th>2009 Veraison (DOY 220)</th>
<th>2009 Harvest (DOY 261)</th>
</tr>
</thead>
<tbody>
<tr>
<td>VSP</td>
<td>0.50 a(^a)</td>
<td>0.19 a</td>
<td>0.43 a</td>
<td>0.27 a</td>
<td>0.24 a</td>
<td>0.63 a</td>
</tr>
<tr>
<td>HC</td>
<td>0.48 a</td>
<td>0.23 a</td>
<td>0.36 a</td>
<td>0.25 a</td>
<td>0.22 a</td>
<td>0.52 b</td>
</tr>
<tr>
<td>SD</td>
<td>0.16 b</td>
<td>0.08 b</td>
<td>0.16 b</td>
<td>0.08 b</td>
<td>0.11 b</td>
<td>0.30 c</td>
</tr>
<tr>
<td>VSP</td>
<td>0.21 b</td>
<td>0.06 b</td>
<td>0.21 b</td>
<td>0.11 b</td>
<td>0.20 a</td>
<td>0.33 c</td>
</tr>
</tbody>
</table>

\(^a\)The statistical models included the following fixed effects: year (P < 0.1452), trellis (P < 0.0001), date (P < 0.0001), year by date (P < 0.0001), year by trellis (P = 0.0276), trellis by date by year (P < 0.0001), and a random plant effect. Because there was not a significant trellis by date interaction, the data are averaged over both years.

\(^b\)GDC = Geneva double curtain, HC = high cordon, SD = Smart-Dyson, VSP = vertical shoot positioning.

\(^c\)DOY = Day of the year.

Table 2. Mean yield and leaf layer number (LLN) of ‘Frontenac’ grapes grown on four training systems in southeastern Nebraska in 2008 and 2009.\(^a\)

<table>
<thead>
<tr>
<th>Trellis(^a)</th>
<th>2008 Yield (kg/plant)(^a)</th>
<th>2009 Yield (kg/plant)(^a)</th>
<th>2008–09 LLN (no.)(^a)</th>
</tr>
</thead>
<tbody>
<tr>
<td>VSP</td>
<td>2.36</td>
<td>4.21 a(^a)</td>
<td>1.29 b</td>
</tr>
<tr>
<td>HC</td>
<td>1.10</td>
<td>2.52 c</td>
<td>1.45 b</td>
</tr>
<tr>
<td>SD</td>
<td>1.34</td>
<td>3.73 ab</td>
<td>2.27 a</td>
</tr>
<tr>
<td>VSP</td>
<td>1.34</td>
<td>3.01 bc</td>
<td>2.19 a</td>
</tr>
</tbody>
</table>

\(^a\)For 2009 yield analysis, the statistical models included the following fixed effects: year (P = 0.0006). For LLN analysis, the statistical models included the following fixed effects: (P < 0.0001), year (P = 0.094), year by trellis (P = 0.7533), and a random plant effect. Because there was not a significant year by trellis interaction, the data are averaged over both years.

\(^b\)GDC = Geneva double curtain, HC = high cordon, SD = Smart-Dyson, VSP = vertical shoot positioning.

\(^c\)2008 yield means not statistically compared because harvest weights for individual vines were not measured that year.

were calculated by dividing the fruit-zone PAR value by the ambient PAR (measured above the canopy).

**Point Quadrat Canopy Analysis.** On 8 Aug. of both years [DOY 221 (2008), DOY 220 (2009)], point quadrat analysis was performed as described by Smart and Robinson (1991). Three insertions were made for each sample plant. Point quadrat data were used to compute LLN.

**Yield and Berry Composition.** Bird-proof netting was used to protect the crop in both years of study. Netting was in place from veraison through harvest. Fruit was harvested on 15 Aug. 2008 (DOY 228) and 19 Aug. 2009 (DOY 231). In 2008, fruit was collected from each treatment and weighed. Average yield per plant was then calculated from those sums (2008 yield data not subject to statistical analysis). In 2009, fruit from each sample plant was harvested separately and weighed. At both years' harvests, randomly selected 30 berry samples were collected from each sample plant, placed in resealable plastic bags and frozen until laboratory analysis. Berry samples were weighed, thawed, wrapped in cheese cloth and then crushed with a mortar and pestle. Juice was reserved for analysis (both years) and fruit solids were returned to their bags and refrozen (2008). Juice pH was measured with a digital pH-meter (Orion model 611; Atago, Bellevue, WA). Titratable acidity (TA) was determined by titration with sodium hydroxide (NaOH), using an automatic TA test (model 620F-1; Kimax, Vineland, NJ). Soluble solids were measured using a digital refractometer (model PR-101; Atago, Bellevue, WA). Titratable acidity (TA) was determined by titration with sodium hydroxide (NaOH), using an automatic TA test (model 620F-1; Kimax, Vineland, NJ). Phenolic analysis was carried out in 2008, but not in 2009 because of personnel limitations.

**Total Phenol Extraction.** Total phenol extraction was performed using the method described by Gorinstein et al. (2010). Berry samples were thawed; skins and seeds were separated from pulp and dried to constant weight in an Isotemp vacuum oven (model 280A, Thermo Fisher Scientific) at 60 °C for ≈24 h. After crushing seeds with a mortar and pestle and skins with a metal spatula, small portions of the samples were placed in 15-mL plastic tubes for extraction (≈0.19 g for seeds and 0.21 g for skins). Free, or unbound, phenols were extracted with 50% methanol:50% water. Two milliliters of solvent was added to each seed sample; skin samples received 3 mL. After shaking for 1 h on a shaker (Labquake; Barnstead Thermolyne, Dubuque, IA), the samples were centrifuged for 15 min at 8000 rpm and supernatants were placed at −20 °C for future use.

**Phenol Content Assay.** Phenolic content was determined using the method described in Singleton and Rossi (1965). To perform the assay, 100 μL of 2 N Folin-Ciocalteu reagent (Sigma-Aldrich, St. Louis, MO), 4.5 mL nanopure water, and 0.3 mL 2% (w/v) sodium carbonate (Na2CO3) were added to the sample extracts, which were diluted as follows: 5 μL of free seed extract, 10 μL of free skin extract, and 20 μL of bound seed extract were combined with nanopure water to a final volume of 100 μL. The bound skin extract was not diluted. The mixture was shaken intermittently for 2 h. The absorbance was measured at 760 nm with a spectrophotometer (model DU800; Beckman Coulter, Brea, CA), and compared with a trans-cinnamic acid standard curve. Results were calculated as concentrations (milligrams per gram dry sample).
FLAVONOID CONTENT ASSAY. Flavonoid concentration was determined using the methods described in Adom and Liu (2002) and Jia et al. (1999). To perform the assay, 0.625 mL nanopure water, 37.5 μL 5% (w/v) sodium nitrite (NaNO₂), 74 μL 10% (w/v) aluminum chloride (AlCl₃), 0.25 mL 4.0 M NaOH, and additional 0.4 mL nanopure water was added to 125 μL of sample extract. Seed extracts were diluted at ratios of 1 part extract in 9 parts nanopure water (free) and 1 part extract in 4 parts water (bound); skin extracts were not diluted. The mixture was vortexed to distribute any precipitate that formed. The absorbance was measured at 510 nm and compared with a (+)-catechin standard curve. Results were calculated as concentrations (milligrams per gram dry sample).

STATISTICAL ANALYSIS. Statistical analyses were performed using the GLIMMIX procedure of SAS® (version 9.3; SAS Institute, Cary, NC). For those variables measured on the same plants once each year (harvest variables and point quadrat data) an initial mixed linear repeated measures model with fixed effects of trellis and year and a trellis by year interaction was used. The residual matrix was tested with both the unstructured and independent covariance structures and the one with the smallest Akaike’s information criteria was used. The PAR measurements were analyzed using an initial mixed linear repeated measures model with the fixed effects of trellis, year, and month, as well as their interactions and a random plant effect. The residual matrix was tested the same as for the other repeated measures models. Total phenol and flavonoid results were examined using a linear model with fixed

Table 3. Mean fruit characteristics of ‘Frontenac’ grown on four training systems in southeastern Nebraska in 2008 and 2009.*

<table>
<thead>
<tr>
<th>Trellis*</th>
<th>2008 Berry wt (g)*</th>
<th>2008 Soluble solids (%)</th>
<th>2008 pH</th>
<th>2008 TA (g L⁻¹)w</th>
<th>2009 Berry wt (g)</th>
<th>2009 Soluble solids (%)</th>
<th>2009 pH</th>
<th>2009 TA (g L⁻¹)w</th>
</tr>
</thead>
<tbody>
<tr>
<td>GDC</td>
<td>1.08 a</td>
<td>22.45 a</td>
<td>3.06 a</td>
<td>17.9 b</td>
<td>0.85 ab</td>
<td>20.35 b</td>
<td>3.10 ab</td>
<td>15.7 a</td>
</tr>
<tr>
<td>HC</td>
<td>1.10 a</td>
<td>21.50 ab</td>
<td>2.95 b</td>
<td>18.6 b</td>
<td>0.82 b</td>
<td>19.74 b</td>
<td>3.05 b</td>
<td>14.6 a</td>
</tr>
<tr>
<td>SD</td>
<td>1.12 a</td>
<td>20.81 bc</td>
<td>2.93 b</td>
<td>20.9 a</td>
<td>0.88 a</td>
<td>21.58 a</td>
<td>3.12 a</td>
<td>15.2 a</td>
</tr>
<tr>
<td>VSP</td>
<td>1.09 a</td>
<td>20.35 c</td>
<td>2.92 b</td>
<td>20.4 a</td>
<td>0.88 a</td>
<td>21.42 a</td>
<td>3.12 a</td>
<td>15.0 a</td>
</tr>
</tbody>
</table>

*The statistical models included the fixed effects of year, trellis, their interactions, and a random plant effect. Statistical probability values for year, trellis, and their interaction for berry weight were <0.0001, 0.1642, and <0.0001; for soluble solids were 0.0539, 0.1755, and <0.0001; for pH were <0.0001, 0.0003, and <0.0001; and for TA were <0.0001, 0.0130, and 0.0033, respectively.

GDC = Geneva double curtain, HC = high cordon, SD = Smart-Dyson, VSP = vertical shoot positioning.

*1 g = 0.0353 oz.

**TA = titratable acidity, 1 g L⁻¹ = 0.1%.

*Means followed by the same letter within a column do not differ at P ≤ 0.05 using Tukey’s adjustment.

Fig. 1. Maximum and minimum daily air temperature and monthly rainfall precipitation near Crete, NE, during the 2008 growing season; (°F − 32) × 1.8 = °C, 1 inch = 25.4 mm.
Results and discussion

Trellis effects on light environment. At all sampling dates in 2008, vines grown on GDC and HC trellises had higher midday percentage PAR transmittances than vines grown on SD and VSP training systems (Table 1). In 2009 the pattern was similar, except GDC had higher transmittance than HC at harvest and VSP had higher transmittance than SD at veraison (Table 1). There was a significant year by trellis by sampling date interaction, as evidenced by the differences in transmittance relationships between the years. In both years, LLN was lower for GDC and HC than for SD and VSP (Table 2). There was not a significant year by trellis interaction for LLN, so the differences presented are averaged over both years. GDC and HC had less dense canopies (higher transmittances and lower LLN) than the other trellises in this study. Open canopies optimize yield and fruit composition; they facilitate pruning, harvesting, and spray penetration. They also tend to have fewer disease problems because of their favorable canopy microclimates (Smart and Robinson, 1991). Light penetration (percentage PAR transmittance) is generally negatively correlated with LLN (Vanden Huevel et al., 2004). In this study, training systems with low LLN values did have correspondingly high transmittances, though the correlation was not statistically significant.

Trellis effects on yield. This research demonstrated large differences in yield between training systems. GDC had the highest fruit yield of all trellises in 2008, and higher yield than HC and VSP in 2009 (Table 2), which is consistent with previous studies comparing the yield of other cultivars grown on GDC and other horizontally divided canopies to single-canopy controls (Reynolds et al., 1995; Shaulis et al., 1966; Smart et al., 1982). Generally, sunlight penetration and yield are positively correlated because increased shoot exposure improves bud fruitfulness (Perez and Kliewer, 1990; Shaulis et al., 1966; Smart et al., 1982). Because HC had such high transmittance values, one would expect it to yield more than VSP, the only other single-canopy training system; however, HC suffered damage from birds because they were able to access its fruit through the netting. If the crop had been better protected, perhaps HC would have produced a higher yield than VSP. Based on the yields we recorded in 2009, at a vine spacing of 551 plants/acre (2.4 m between vines and 3 m between rows) and price of $1200/t of fruit, the vineyard’s gross income would have been $3062/acre for GDC, $1833/acre for HC, $2713 for SD, and $2189/acre for VSP.

Trellis effects on fruit composition. In 2008, GDC and HC had lower TA than other training systems (Table 3); this is in agreement with the findings of Smart et al. (1988) and Macaulay and Morris (1993) who observed higher TA in shaded treatments. In 2009, the trellises did not differ in TA despite observed differences in percentage PAR transmittance (Table 3). In both years, all of the trellises exceeded the ideal TA concentration, which is
Table 4. Mean total phenol concentrations of skin and seeds of ‘Frontenac’ grapes grown on four training systems in southeastern Nebraska in 2008. *

<table>
<thead>
<tr>
<th>Trellis</th>
<th>Phenol concn of free</th>
<th>Phenol concn of combined</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>extract of skin and</td>
<td>(free + bound) extracts</td>
</tr>
<tr>
<td></td>
<td>seeds (mg g⁻¹ dry wt)*</td>
<td>of skin and seeds (mg g⁻¹ dry wt)</td>
</tr>
<tr>
<td>GDC</td>
<td>60.27 ab</td>
<td>65.38 ab</td>
</tr>
<tr>
<td>HC</td>
<td>61.70 ab</td>
<td>65.65 ab</td>
</tr>
<tr>
<td>SD</td>
<td>51.60 b</td>
<td>55.31 b</td>
</tr>
<tr>
<td>VSP</td>
<td>64.45 a</td>
<td>68.64 a</td>
</tr>
</tbody>
</table>

*Statistical models included fixed effects of sample type (bound seed, bound skin, free seed, free skin), trellis, and their interactions. Statistical probability values for sample type, trellis, and their interaction for free extracts were < 0.0001, 0.0477, and 0.2826 and for combined were < 0.0001, 0.0382, and 0.3220, respectively. Because there were no significant interactions between trellis and sample type, we investigated the main effect of trellis averaged over sample type.

Additional, some phenolic compounds have higher AOA than others (Di Majo et al., 2008; Yildirim et al., 2005). In a study of various red wines made from different grape cultivars, flavonoid concentrations were not significantly different although flavonoid composition varied greatly (Fang et al., 2008).

Conclusions

Although individual phenolic compounds were not identified in this study, it is possible that the composition and antioxidant capacity could have varied among trellis treatments, even though total flavonoid and phenolic concentrations were mostly not significantly different. Therefore, it would be problematic to draw conclusions on the health-promoting properties of the fruit in this study. Future research on grape and wine phenols and/or flavonoids should identify individual compounds where possible, since their AOA—and potential health benefits—are variable.

Of the four training systems evaluated in this research, we feel GDC is an appropriate choice for ‘Frontenac’ grown in the upper midwestern U.S. Although fruit quality results were inconclusive, GDC vines had the highest yield in both years of our study. Vines trained to GDC also had very favorable canopy conditions, as determined both by percentage PAR transmittance measurements and by point quadrat analysis.

Literature cited


