English and Black Walnut Phenolic Antioxidant Activity in Vitro and Following Human Nut Consumption

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ABSTRACT

Background: Walnut consumption may reduce the risk of cardiovascular disease by providing antioxidant protection to low density lipoproteins (LDL). Aim: This study compared the phenolic profile and antioxidant activity of English versus black walnuts. Methods: Nuts were extracted in methanol or acetone prior to analysis with HPLC/LC-MS-MS for phenolic identification and quantitation. The ability to prevent oxidation of LDL was examined in vitro using walnut extracts and ex vivo after walnut consumption for 28 days. Results: Flavonoids identified/quantified with HPLC/LC-MS-MS included the phenolic acids 5-caffeoylquinic acid, 3-caffeoylquinic acid (black walnut only), 4-caffeoylquinic acid, and the flavonol glycosides quercetin-3-rutinoside, quercetin-3-galactoside, quercetin-3-pentoside, quercetin-3-arabinoside, quercetin-3-rhamnoside, and the aglycone quercetin (English walnut only). Total phenolic yield of acetone extracts was 166.1 and 24.2 µg/g for English and black walnut respectively, and yield for methanol extracts was 147.6 and 4.1 µg/g for English and black walnut respectively. In vitro LDL oxidation by Cu²⁺ with English walnut extracts significantly extended oxidation lag-time (A₂₃₄) in a dose dependent manner at 1.0 and 0.1 µg/ml and reduced TBARS formation (1.0 µg/ml). Black walnut extracts reduced TBARS significantly but had no effect on A₂₃₄. Human consumption of English or black walnuts (30 g nuts/day) for 28 days resulted in no differences in LDL antioxidant capacity (A₂₃₄) between groups or within groups. Conclusions: This study suggests that the English walnuts have a phenolic profile and in vitro antioxidant capacity that is better than black walnuts, but that walnut consumption for 28 days does not improve LDL resistance to oxidation.

Keywords: Walnut, Lipoprotein, Oxidation, Antioxidant, Phenolic, Flavonol

1. Introduction

Nut consumption has been demonstrated to be protective against cardiovascular disease, an effect attributable to their phenolic [1], antioxidant [2] and polyunsaturated fat [3]. Walnuts are an important dietary item. In 2008, global total walnut production was estimated to be in excess of 2.1 million metric tons [4]. English walnuts (Juglans regia) and the increasingly popular, black walnuts (Juglans nigra) are currently part of the American diet, with black walnuts having a slightly bitter taste relative to English walnuts. Black walnut trees are native to the United States and grow east of the Rocky Mountains.
source of antioxidants [1,9,13]. Walnut consumption has been shown to be associated with improvements in blood lipid profile [14], possibly by causing changes in HMG-CoA activity [15], an observation in agreement with our previous study [16]. Walnuts have also been shown to improve endothelium mediated vasodilation in dyslipidemic subjects following intake of a high fat meal [16,17] and type 2 diabetics [18].

No direct comparisons between English and black walnut phenolic profiles, and their relative in vitro or ex vivo antioxidant activities, have been performed. The present study compared the flavonol-glycoside profiles of English and black walnuts with high-pressure liquid chromatography and liquid chromatography-mass spectrometry (LC-MS-MS) for quantification, and identification. The relative capacities of English and black walnuts to inhibit LDL oxidation were examined using walnut phenolic extracts prepared with acetone or methanol in vitro. Human LDL antioxidant capacity was also examined ex vivo following 28 days of walnut consumption. These comparisons sought to determine if black walnuts would provide comparable health benefits to that of English walnuts. The black walnut, being adapted to more severe temperate climate, would provide an alternative walnut to the intensively cultivated English walnut.

2. Methods

Human Studies: This study was approved by the Institutional Review Boards of the University of Wisconsin-La Crosse and Winona State University. All participants read and signed an informed consent document prior to volunteering to participate in this study.

Walnut source and preparation: Black walnuts were obtained through donations from Hammons Products Company (Stockton, MO), Michael Winfrey (Ettrick, WI), and Harry Lundstrom (Unionville, VA). English walnuts were purchased from the People’s Food Coop (La Crosse, WI). Walnuts (30 g portions) consisting of nut flesh with attached pellicle were frozen until phenolic extraction or human consumption.

Walnut phenolic extraction: English and black walnut flesh with the kernel and attached pellicle was ground into a course powder with a mortar and pestle. A 5 g portion of the powder was extracted twice using either 80% acetone or 80% methanol in water, followed by centrifugation and supernatant concentration using a rotary evaporator (45°C under reduced pressure). Flavonols were isolated from the 80% acetone and 80% methanol extracts with Sep Pack C18 cartridges (5 g, Waters Inc., Milford, MA) preconditioned by sequential treatment with 20 mL of methanol, 20 mL of H2O, respectively, under vacuum. The sample solutions were then distributed onto cartridges (0.5 mL per cartridge). Flavonols were removed from cartridges by elution with 20 mL of methanol followed by water. The eluted methanol solution was concentrated and diluted with HPLC mobile phase-B (2 mL) for HPLC and LC-MS-MS analysis as described below. The acetone and methanol extracts were then diluted in phosphate buffered saline for evaluation of their antioxidant capacity as described later.

Walnut Phenolic isolation, identification and quantification: Extract constituents were preparatively isolated using HPLC (Dionex Inc., Sunnyvale, CA) equipped with a UV-Vis PDA 100 Photodiode Array Detector, AS 50 autosampler and GP 40 gradient pump. Mobile phase: Eluent A: 90% water + 10% methanol + 0.001% formic acid; Eluent B: 20% water + 20% methanol + 60 % ACN + 0.001% formic acid. Flow rate: 0.25 mL·min⁻¹ and injection volume: 5 µL for LC-MS-MS. Column: Pursuit C18 (3 µm particle size; 150 mm length × 3.0 mm ID; Varian Inc, Palo Alto, CA).

Flavonols were identified and quantified by high performance liquid chromatography performed on an API 3000 triple quadrupole mass spectrometer (PE Sciex, Concord, Ontario, Canada). All the analyses were performed using the Electrospray chemical ionization source (ESI) in negative ion mode with the following settings: ion spray voltage –4200 V, nebulizer gas (N2) 7 (arbitrary units), curtain gas (N2) 12 (arbitrary units), collision gas (N2) 6 (arbitrary units), declustering potential (DP) between –60 V to –100 V, focusing potential –400 V, entrance potential –10 V, collision energy (CE) –50 V and collision cell exit potential (CXP) –5 V. The drying gas (N2) was heated to 350°C and introduced at a flow rate of 6000 cm³·min⁻¹. Full scan data acquisition was performed scanning from m/z 100 - 1200 in profile mode, using a cycle time of 2 s with a step size of m/z 0.1 and an interscan pause of 2 ms.

Identified flavonols were quantified using an HPLC equipped with a Waters 996 Photodiode Array Detector (PDA) (Waters, Milford, MA). Quantification of all the phenolic acids was based on a standard curve prepared with 5-caffeoylquinic acid. Quercetin-3-rutinoside, quercetin-3-galactoside, quercetin-3-rhamnoside and quercetin were quantified using the corresponding authentic standard for each compound. Other flavonols were quantified based on a standard curve of quercetin-3-rhamnoside. For all the standards and extracts, 40 µL were injected and separated using the aforementioned column and solvent system, but at a flow rate of 1 mL/min. Scanning by the PDA occurred at 325 nm for phenolics and 366 nm for flavonols, while data acquisition and processing were performed by Waters Empower™ Chromatography Software (Waters, Milford, MA).

In vitro evaluation of walnut antioxidant capacity: Venous blood was collected in K₃EDTA from three
healthy adults following a 12 hour abstinence from all food and beverages except water. Plasma LDL was sequentially isolated sequential ultracentrifugation, dialyzed twice in phosphate buffered saline to remove EDTA, and purged with N₂ gas to prevent oxidation prior to use within three weeks of collection. LDL isolates were merged and the LDL protein concentration determined with the BioRad DC Protein Assay (Hercules, CA) for use in antioxidant assessment [19].

English and black walnut extract antioxidant capacities were determined in vitro by inducing the oxidation of 100 μg LDL protein/mL, with 1.00 or 0.10 μg/mL walnut extracts in the presence of 5 μM Cu ++ and measuring conjugated diene formation at 234 nm. The assay was held at a constant temperature of 37˚C with absorption readings were taken every 85 seconds (300 recordings). Each experiment was performed in triplicate and lag time was determined by finding the midpoint of the resultant curves [19].

The thiobarbituric acid reactive substances (TBARS) assay was performed as a secondary proof that conjugated dienes measured in the Cu++ assay resulted from LDL oxidation. Oxidation was initiated with cupric sulfate (5 μM) and 100 μg LDL protein/mL in PBS incubated at 37˚C for 95 minutes in the presence or absence of 0.1 μg/mL walnut extracts. The TBARS test measured malondialdehyde oxidation products using tetramethoxypropane (TMPP, 8 ng/100-μL incubate) as a standard [20].

**Human walnut consumption and antioxidant capacity:** Ex vivo antioxidant effects of walnuts were observed in 36 subjects randomly assigned to consume either 30g portions of English or black walnuts each day for 28 days, in addition to their usual diet. Fasting venous blood was drawn pre-study (day 0) and day 28. Collected plasma was stored at ~80˚C with 0.6% sucrose for stabilization until simultaneous LDL isolation was performed on a single day using sequential ultracentrifugation for day-0 and day-28 samples as describe previously in this manuscript and used within three days of preparation. The antioxidant capacity of each subjects LDL pre- and post walnut consumption was determined by inducing oxidation of 100 μg/mL LDL in the presence of 10, 4, and 1 μM cupric ions and measuring conjugated diene formation at A₂₃₄ as described previously in this manuscript.

**Statistical analysis:** The data are expressed as means ± standard deviation. ANOVA was used to determine if the variable interaction in the LDL assay and TBARS was significant. Dunnet’s Multiple Comparison procedure was used to compare the extract treatments to the control (LDL with no extract lag-time). Tukey HSD was used to statistically represent the relationship of the extract solvent and concentration used in LDL oxidation. ANOVA and a repeated measures analysis of variance was used to identify significant differences among treatments, time, and interaction between treatment and time. Significant differences among least squares means (P < 0.05) were determined using the Tukey-Kramer adjustment (SAS Inst. Inc., Cary, NC).

### 3. Results

The phenolic acid and flavonol profiles of the identified English and black walnuts were obtained and compounds quantified (Figure 1, Table 1). Phenolics in both 80% acetone and methanol extracts included phenolic acids and flavonoids.

<table>
<thead>
<tr>
<th>Name of compound</th>
<th>( \lambda_{max} ) (nm)</th>
<th>[M-H] fragments in ESI-MS</th>
<th>English 80% acetone (µg/g wet wt)</th>
<th>English 80% methanol (µg/g wet wt)</th>
<th>Black 80% acetone (µg/g wet wt)</th>
<th>Black 80% methanol (µg/g wet wt)</th>
</tr>
</thead>
<tbody>
<tr>
<td>3-caffeoylquinic acid</td>
<td>218, 236, 293, 317</td>
<td>353, 191, 179, 127, 111</td>
<td>ND</td>
<td>ND</td>
<td>0.67</td>
<td>1.05</td>
</tr>
<tr>
<td>5-caffeoylquinic acid</td>
<td>217, 242, 298, 326</td>
<td>353, 191, 179, 127, 111</td>
<td>20.43</td>
<td>17.37</td>
<td>0.19</td>
<td>2.20</td>
</tr>
<tr>
<td>4-caffeoylquinic acid</td>
<td>218, 243, 297, 327</td>
<td>353, 191, 173, 127, 111</td>
<td>2.03</td>
<td>2.93</td>
<td>0.23</td>
<td>ND</td>
</tr>
<tr>
<td>Quercetin-3-rutinoside</td>
<td>200, 251, 351</td>
<td>609, 301</td>
<td>5.37</td>
<td>0.64</td>
<td>0.24</td>
<td>ND</td>
</tr>
<tr>
<td>Quercetin-3-galactoside</td>
<td>203, 254, 351</td>
<td>463, 301</td>
<td>55.63</td>
<td>0.33</td>
<td>1.39</td>
<td>0.05</td>
</tr>
<tr>
<td>Quercetin-3-pentoside</td>
<td>204, 253, 355</td>
<td>433, 301</td>
<td>71.33</td>
<td>78.74</td>
<td>21.14</td>
<td>0.21</td>
</tr>
<tr>
<td>Quercetin-3-arabinoside</td>
<td>204, 253, 355</td>
<td>433, 301</td>
<td>0.45</td>
<td>39.79</td>
<td>0.09</td>
<td>0.44</td>
</tr>
<tr>
<td>Quercetin-3-rhamnoside</td>
<td>204, 253, 355</td>
<td>447, 301</td>
<td>0.67</td>
<td>0.09</td>
<td>0.23</td>
<td>0.14</td>
</tr>
<tr>
<td>Quercetin [b,c]</td>
<td>207, 266, 355</td>
<td>301, 151</td>
<td>10.21</td>
<td>7.70</td>
<td>ND</td>
<td>ND</td>
</tr>
</tbody>
</table>

| Total Phenolics (sum of flavonol constituents; µg/g) | 166.12 | 147.59 | 24.18 | 4.09 |

ND: Not detected; Based on full scan and product ion scan; Based on Singh et al. 2009.
Figure 1. Representative HPLC chromatographs of English and black walnuts extracted with 80% acetone and 80% methanol demonstrate that English cultivars contain a greater profile of phenolic compounds.
5-caffeoylquinic acid, 3-caffeoylquinic acid (black walnut only), 4-caffeoylquinic acid, and the flavonols quercetin-3-rutinoside, quercetin-3-galactoside, quercetin-3-pentoside, quercetin-3-arabinoside, quercetin-3-rhamnoside, and the aglycone quercetin (English walnut only). Total flavonol yield for the acetone extract was 166.1 and 24.2 μg/g for English and black walnut, respectively, and total yield for the methanol extract was 147.6 and 4.1 μg/g for English and black walnut, respectively.

In vitro LDL oxidation induced by Cu++ (lag-time at A_{234}) was significantly inhibited by English walnut extract relative to control LDL, which was oxidized in the absence of acetone or methanol English walnut extracts. The 1.0 μg/mL dose was significantly better than the 0.1 μg/mL dose of English walnut extract (Figure 2 and 3) for inhibition of oxidation. In contrast, black walnut extracts provided no significant increase in oxidative lag-time and therefore no apparent antioxidant protection of LDL.

In vitro LDL malondialdehyde production was measured with a TBARS assay and supports the results of the oxidative lag-time assay. After 95 minutes of oxidative incubation, TBARS (8 ng TMPP/100-μL incubate) were 0.76 ± 0.01 for control LDL oxidized in the absence of walnut extracts. English walnut acetone and methanol extract TBARS were 0.03 ± 0.001 and 0.003 ± 0.001 (significant vs control LDL; p < 0.001), indicative of antioxidant activity. For black walnut acetone and methanol extracts TBARS were 0.63 ± 0.001 and 0.63 ± 0.02 (significant verses control LDL; p < 0.001). The English walnut extracts provided significantly lower TBARS and antioxidant capacity before and after consumption of 30 g English and black walnuts (control) and in the presence of English or black walnut extracts. English walnut extracts greatly increased lag-time and yielded resistance to oxidation.

Ex Vivo effects of walnut consumption on human LDL antioxidant capacity before and after consumption of 30 g walnut flesh/day for 28 days was evaluated following oxidation with 10, 4, and 1 μM Cu++. There was a slight increase in LDL oxidative lag-time following oxidation by 10 μM Cu++, however, no statistically significant changes in LDL oxidation were observed pre- to post-walnut consumption within or between treatment groups. Percent change in LDL lag-time after 28 days of English walnut consumption and oxidation by 10, 4 and 1 μM cupric sulfate was −0.09 ± 0.08, 0.13 ± 0.08, and −0.09 ± 0.08, respectively. Percent change in LDL lag-time after 28 days of black walnut consumption and oxidation by 10, 4 and 1 μM cupric sulfate was −0.11 ± 0.08, −0.03 ± 0.08, and −0.03 ± 0.08, respectively.

4. Discussion
This is the first study to compare the identity and quantity of the wide range of phenolic acids and flavonol glycosides in English and black walnuts (Table 1). English walnut are known to contain non-flavonoid phenolics including ellagic acid, valoneic acid dilactone and pedunculagin and have been reported to contain a single 433 MW flavonol pentoside [6-11]. In the present study the total English walnut flavonol glycoside yields for 80% acetone and 80% methanol were 166 and 148 μg/g wet weight. In contrast, black walnuts contained a reduced amount of total flavonol-glycosides, with 24 and 4 μg/g with acetone and methanol, respectively. The flavonol glycoside profile of black walnut flesh had not been previously studied prior to this investigation.

English walnut leaves, in comparison to nuts, have been found to contain 3- and 5-caffeoylquinic acids, 3- and 4-p-coumaroylquinic acids, p-coumaric acid, quercetin 3-galactoside, quercetin 3-pentoside derivative, quercetin 3-arabinoside, quercetin 3-xylloside and quercetin 3-rhamnoside with HPLC/DAD in quantities ranging from 0.8 to 21 μg/g dry weight for p-coumaric acid and quercetin-3-galactoside, respectively [12]. Given the difference between the wet and dry weight measures, the quantitative values are consistent to those previously observed [12]. The difference between English and black walnut profiles may be a reflection of the structure of the whole nut. Walnut anatomy consists of the shell, pellicle and nutmeat. The pellicle contains many antioxidant compounds that serve to protect the meat, which is high in unsaturated fat, from oxidation. English walnut nutmeat with pellicle contains 6.5 mmol TAC per serving while the nutmeat alone contains 0.3 mmol TAC per serving [1]. English walnuts are bred to have a thin shell and husk for easier harvesting, making the pellicle more important in protecting the meat from rancidity. Black walnut anatomy consists of a thick husk, liquid layer, and strong shell. The added protective measures of the thick husk, liquid and strong shell may reduce the need for antioxidants in the form of phenolics to protect the nutmeat and germ DNA.

The lower phenolic profile content associated with black walnuts is suggestive of a reduced potential to pro-
Figure 3. English walnut phenolic extracts at 1.0 and 0.1 µg/mL extended the lag-time for LDL oxidation by 5 μM Cu++, while black walnut extracts failed to significantly delay the LDL oxidation lag-time.

vide human health benefits. *In vitro* analysis of English-walnut acetone and methanol extracts demonstrate suppression of cupricion mediated oxidation of LDL at 1.0 and 0.1 µg/mL concentrations (Figure 3). However, black walnut acetone and methanol extracts contained 7- and 36-fold less total flavonol glycosides, respectively, and were found to have non-significant increases in oxidative lag-time. While the TBARS test demonstrated that black walnut extracts have antioxidant activity, the activity and phenolic content was less than that of the English walnut extracts.

In contrast to the robust *in vitro* antioxidant effects found with English walnuts in the present study, walnut consumption did not significantly improve LDL protection *ex vivo*. Several studies have demonstrated that walnut consumption can improve human antioxidant capacity, however, others have failed to demonstrate that human consumption of English walnuts improved antioxidant capacity [17,21,22]. Plant phenolics have been shown to adhere to LDL resulting in improved antioxidant capacities following consumption [23]. However, because the LDL samples were dialyzed prior to oxidation to remove EDTA, it is possible LDL samples were no longer associated with adherent dietary walnut phenolics. The reduced flavonol glycoside content of our black walnuts may also explain the results of our previous investigation which determined that human consumption of English walnuts is superior to black walnuts for improving the blood lipid profile and endothelial function [16].

The present study was designed to obtain preliminary data regarding the phenolic acid and flavonol contents of the walnuts used in antioxidant and human feeding trials. It was not designed to examine the effect of cultural effects nor growing location on walnut phenolic expression or antioxidant activity. In this regard the total leaf flavonol-glycoside content of six English walnut cultivars was found to be fairly uniform with a total phenolic content of (70 ± 3) µg/g wet weight [12]. A similarly tight standard deviation was observed for total phenolics in methanol extracts of English walnut kernel ((494 ± 164) mg/100g of kernel) and pellicle ((13,792 ± 2917) mg/100g pellicle) in ten different English walnut cultivars...
Future studies may wish to examine differences within black walnut cultivars and growing locations. The 7 to 36 fold reductions in flavonol-glycoside content for black walnut suggest the effect would be independent of the specific black cultivar examined.

In the human trial of the present study a major limitation was lack of control over subject intake of other dietary elements that could have affected antioxidant status. While subjects reported no major changes in food or supplement intakes over the course of walnut consumption [16], there was no confirmation of this or attempt to control intake of other foods that may affect LDL oxidation.

5. Conclusions

English walnuts contain a greater content of phenolic acids and flavonols than black walnuts. Walnut flanonols included 5-caffeoylquinic acid, 3-caffeoylquinic acid (black walnut only), 4-caffeoylquinic acid, quercetin-3-rutinoside, quercetin-3-galactoside, quercetin-3-pentoside, quercetin-3-arabinoside, quercetin-3-rhamnoside, and the aglycone quercetin (English walnut only). The greater flavonol content of English may be responsible for its improved ability to protect LDL against oxidation in vitro. A 28 day period of English or black walnut consumption (30 g/day) was not sufficient to alter in vivo LDL antioxidant capacity. Further studies would be needed to confirm these effects across diets and nut sources.

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