Lutein Content in Sweetpotato Leaves

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Abstract. Fresh leaves of 6 sweetpotato (Ipomoea batatas [L. – Lam.] genotypes, ‘Beauregard’, ‘Bienville’, 1.99–35, 1.00–8, 1.01–145, and 1.01–29, were characterized for lutein. Lutein is a carotenoid capable of delaying blindness-related macular degeneration. The content of lutein in sweetpotato ranged from 0.38 to 0.58 mg g⁻¹ fresh weight. Beta-carotene separated from lutein on high-pressure liquid chromatograms and when spiked in pure lutein extract did not interfere with lutein separation. High-resolution electrospray ionization mass spectrometric analysis was used to confirm the presence of lutein in sweetpotato leaves. Stems were also characterized and found not to contain lutein. Our results showed that sweetpotato leaves are an excellent source of dietary lutein and surpass levels found in leafy crucifers. Leaves of sweetpotato and a related species are human food in some countries and may be a major source of lutein for commercial purposes.

Lutein and xanthophylls, 3,3′-dihydroxy-α-carotene and 3,3′-dihydroxy-β-carotene, have been identified and recognized by various interdisciplinary studies as the major dietary carotenoids present in the human retina and capable of delaying the onset of macular degeneration (AREDS Research Group, 2001; Bernstein, 2002; Bernstein et al., 2004). Age-related macular degeneration (AMD) is a major cause of blindness in the Western world. Risk factors include age, family history, exposure to sunlight, complications of diabetes, and high cholesterol levels. There are therapies, but no curative treatments; most patients with AMD progress to legal blindness. Because standard therapies are limited, there is recognition in the scientific community that nutrition with antioxidants and carotenoids is a meaningful preventive approach against the onset or progression of AMD (Blasdale and Bernstein, 2005; Johnson, 2002; Ribaya-Merchao and Blumberg, 2004; Thurmann et al., 2005).

Lutein is not synthesized de novo and must be ingested from outside sources. Dark green vegetables have been identified as major sources of lutein for human consumption. Major edible sources of lutein, by decreasing order, are kale, spinach, lettuce, and broccoli (Johnson, 2002).

Pratt (1999) indicated that early introduction of lutein in the diet may reduce the severity of AMD. The Age-Related Eye Disease Study Research Group (AREDS Research Group, 2001) found that intake of antioxidants combined with zinc may help reduce the odds of developing advanced AMD. Seddon et al. (1994) suggested that ingestion of 6 mg lutein per day could reduce the risk of macular degeneration by as much as 43%. This concentration would require daily ingestion of approximately 2 lbs. of corn, 16 lbs. of tomatoes, two salad bowls of spinach, or one bowl of kale. As a consequence, identifying edible sources of lutein, fortifying foods with lutein, and enhancing lutein bioavailability is important, appealing to health-conscious consumers, and timely (AREDS Research Group, 2001; Chitchumroonchokchai et al., 2004; Jones et al., 2005).

Sweetpotato leaves have no current economic value in the United States, although a report indicates that leaves are potential sources of xanthophylls and protein (Walter et al., 1978). No data have been presented to date documenting the range of xanthophylls found in various sweetpotato varieties (Woolfe, 1992). In 1999 to 2001, the United States produced approximately $214 million worth of sweetpotato of which only the roots were estimated in September 2003 for ‘Beauregard’ and ‘Bienville’ grown in production fields at the Sweet Potato Research Station, Chase, Louisiana. Three random m² areas of canopy were selected and harvested. Leaves were separated from the vines and both components weighed. These samples were subsequently dried at 85 °C for 48 h before a dry weight determination. Plants were spaced 0.3 m apart and rows were spaced 1 m apart.

Analytical procedure

Freshly harvested, petiole-free leaf tissue was frozen at −18 °C. Frozen leaf tissue was ground in a mortar and pestle and 2 g were then extracted with 80 mL of acetone in screw-capped containers at room temperature in the dark overnight. The acetone extracts were filtered through Whatman No. 4 filter paper (Whatman Inc., Florham Park, N.J.) and evaporated in an exhaust hood overnight. The residue remaining on the filter paper was extracted again in 40 mL of acetone overnight and added to the 80 mL after filtering. The residuals were saponified with 15 mL of 10% potassium hydroxide with gentle shaking overnight. The unsaponifiable compounds were extracted with an ether-hexane (1:1) mixture. The solvent mixture was evaporated in an exhaust hood overnight. Samples were safeguarded throughout extraction from light to minimize degradation. The residue was dissolved in 20 mL of a methyl tert-butyl ether:methanol (5:95) mixture and filtered through a 0.20-μm filter (Nalgene Pune, Inc., Rochester, N.Y.). Twenty microliters were injected into an YMC C18 column S 3 μ, 4.6 × 250-mm high-performance liquid chromatography (HPLC) column (Waters, Milford, Mass.). The flow rate on a Waters 600E using a Waters 717 plus autosampler and Waters 486 Turnable Absorbance detector system (Waters) was 1 mL/min, separation was isocratic using methyl tert-butyl ether:methanol (5:95) as the separation mixture, and total separation time was 35 minutes. These procedures were adapted from Lee et al. (2001).

Standard lutein (Sigma Chemical Co., St. Louis, Mo.) was spiked into a sweetpotato carotenoid extract at concentrations ranging from 0 to 200 ppm and the mixture was separated under conditions described previously. A serial dilution of the lutein standard was used for routine quantification of constituent sweetpotato leaf lutein. Similarly, a β-carotene carotene standard (Sigma Chemical Co.) was spiked into a sweetpotato carotenoid extract to verify that the elution time differed from lutein. Finally, FloraGlo

Materials and Methods

Materials

Sweetpotato leaves from each of the following varieties: ‘Beauregard’, ‘Bienville’, 1.99–35, 1.00–8, 1.01–145, and 1.01–29 were harvested in mid-July 2004 from field-grown plants at the Burden Research Station, Baton Rouge, La. The experimental design was a completely randomized design with four replications of each cultivar. The plots consisted of 30-m long rows with 1.2 m between rows and each row contained 20 plants spaced 0.3 m apart. Commercial sweetpotato cultural practices were followed as recommended (Boudreaux, 1994). Mostly, the upper three to four open leaves from the apical meristem were harvested from each of three to four randomly selected plants in each replication and bulked. Leaf and vine mass were estimated in September 2003 for ‘Beauregard’ and ‘Bienville’ grown in production fields with the Sweet Potato Research Station, Chase, Louisiana. Three random m² areas of canopy were selected and harvested. Leaves were separated from the vines and both components weighed. These samples were subsequently dried at 85 °C for 48 h before a dry weight determination. Plants were spaced 0.3 m apart and rows were spaced 1 m apart.

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lutien (Kemin Industries, Inc., Des Moines, Iowa) from marigold (Tagetes erecta L.) was analyzed and found to be consistent with profiles obtained from standard lutein and our sweetpotato samples (data not shown).

High-resolution electrospray ionization mass spectrometric analysis of sweetpotato xanthophylls was performed on an ABI QSTAR QTOF (Applied Biosystems, Foster City, CA). The samples were dissolved in 1 methanol:1 water containing 1% acetic acid and introduced through a syringe pump at a flow rate of 7 μL/min at room temperature. Ion extraction was performed at voltage maintained at 4900 V. The mass spectrometry data were acquired in the positive mode for all xanthophylls.

**Results and Discussion**

**Analysis of lutein in sweetpotato**

The HPLC profiles for commercial standards of lutein and β-carotene were found to be dissimilar. Lutein eluted after 16 min. of fractionation, whereas β-carotene eluted after 3.5 to 4 min. β-carotene spiked into the lutein extract did not affect lutein separation (Fig. 1). Quantification was validated by a serial dilution of the lutein standard across a range of concentrations. Sweetpotato extracts spiked with lutein and β-carotene were quantifiable. The electrospray ionization mass spectrometry of the isolated sweetpotato lutein in the region m/z between 540 to 580 amu (atomic mass units) indicated the presence of lutein at m/z of 568 amu. Peaks at 567 and 569 correspond to a loss or addition of one proton, respectively (Fig. 2). Our results are consistent with previous reports of lutein in ethanol and water (Guarantini et al., 2005). Polyene β-carotene, with an atomic mass of 536, is known to ionize by a radical process with loss of one or two electrons and thus differs greatly in its profile (Guarantini et al., 2005). These preliminary studies support our underlying techniques to assay lutein in sweetpotato leaf extracts.
Lutein content in sweetpotato leaves

The six genotypes ranged from 0.38 for L01–29 to 0.54 mg g⁻¹ fresh weight for ‘Bienville,’ a difference of approximately 30% (Table 1). ‘Beauregard’ and L99–35 had levels comparable to ‘Bienville.’ A preliminary study in 2003 found similar results (data not shown). Extracts of ‘Beauregard’ and ‘Bienville’ sweetpotato vines did not contain any appreciable amount lutein in 2003. These results showed that sweetpotato leaves are among the best sources of lutein among dark green vegetables. Kale has been reported as the edible vegetable with the highest concentration of lutein at 0.38 mg g⁻¹ (Mangels et al., 1993) followed by spinach at 0.12 mg g⁻¹ (Johnson, 2002). Sweetpotato and its relative, Ipomoea aquatica Forsk., are consumed as a leafy vegetable in Southeast Asia and Africa, and this data adds to our understanding of the nutritional value of sweetpotato leaves.

Our data extends that of Walter et al. (1978) who found xanthophylls represented 0.12% to 0.15% of dry sweetpotato leaf protein concentrate. The content of total carotenoids, β-carotene, and α-carotene in sweetpotato leaves were 26.79, 8.99, and 0.99 mg/100 g, respectively, of dry leaf tissues, showing that xanthophylls were the major carotenoid found in leaves (Moshia et al., 1997).

Biomass estimates

Total canopy yield on fresh weight basis for ‘Beauregard’ and ‘Bienville’ were comparable and estimated at approximately 3000 kg ha⁻¹. Leaf yield was estimated at approximately 800 to 910 kg ha⁻¹ for ‘Beauregard’ and ‘Bienville’, respectively. An extrapolation based on 0.40 mg g⁻¹ fresh leaf weight and 800 kg ha⁻¹ suggests a crude yield of 320 g ha⁻¹. A higher leaf yield and concomitant increase in lutein is possible with high rates of fertilizer, which would promote canopy growth. Lutein from marigold flowers is 0.8 mg g⁻¹ and represents a common commercial plant source for lutein. The current market price for raw lutein is $200 per kg of 5% lutein in corn oil from Kemin Foods, Des Moines, Iowa. Extraction protocols will have to be modified to make sweetpotato leaves commercially viable; significantly more solvents are needed to extract lutein from chlorophyll-rich leaves in contrast to chromoplast-rich marigold flower petals.

In conclusion, our study showed that sweetpotato is a rich source of lutein. Its use as a leafy vegetable is novel in most countries and potentially may serve a market niche given its enhanced nutritional status; it is also possible to identify superior leafy vegetable varieties akin to ones used in parts of Africa. The long-term goal is to develop an environmentally sound lutein extraction protocol from sweetpotato leaves.

### Table 1. Lutein concentration in sweetpotato leaves

| Sweetpotato leaf variety | Lutein concentration |
|--------------------------|----------------------|
| Beauregard               | 0.51 ± 0.04          |
| Bienville                | 0.54 ± 0.09          |
| L99–35                   | 0.53 ± 0.16          |
| L00–8                    | 0.44 ± 0.04          |
| L01–14                   | 0.39 ± 0.13          |
| L01–29                   | 0.38 ± 0.10          |

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