VARIABILITY OF SUGARS IN STAPLE-TYPE SWEET POTATO (Ipomoea batatas) CULTIVARS: THE EFFECTS OF HARVEST TIME AND STORAGE

Evelyn Adu-Kwarteng¹, Esther O. Sakyi-Dawson², George S. Ayernor², Van-Den Truong³, Fred F. Shih³, and Kim Daigle⁴

¹Crops Research Institute, Kumasi, Ghana
²Department of Nutrition & Food Science, University of Ghana, Legon, Ghana
³USDA-ARS, SAA Food Science Research Unit, Department of Food Science, North Carolina State University, Raleigh, North Carolina, USA
⁴USDA-ARS-SRRC, New Orleans, Louisiana, USA

Total soluble sugar content and composition was studied by high performance liquid chromatography in four high dry-matter sweet potato cultivars at 3, 4, and 5 months maturity. Total soluble sugar consisted of sucrose, glucose, and fructose, ranging from 4.10–10.82 g/100 g (dry-weight basis). At harvest, there were significant differences in total soluble sugar due to maturity (p < 0.001) and cultivar (p < 0.05). The highest total soluble sugar contents were in 5-month samples at harvest (7.36–10.34 g/100 g) and 4-month samples after short-term storage under tropical ambient conditions (8.66–10.82 g/100 g).

Estimated amylase enzyme activity varied significantly with harvest age (p < 0.05). Although reducing sugar contents were low, fructose levels in 5-month samples increased considerably after storage.

Keywords: Sweet potato (Ipomoea batatas), Soluble sugars, Storage, Harvest timing, Amylase activity.

INTRODUCTION

Sweet potato (Ipomoea batatas L.), a highly productive and nutritious root crop, is grown in more than 100 countries, ranks seventh among the world’s food crops, and ranks fifth in developing countries where it has the potential to play key roles in food security.[1] Few major food crops are so genetically diverse and underexploited as the sweet potato.[2] Although nutritionally superior to most starchy staples, being rich in vitamins, minerals, and dietary fiber; having a low glycemic index;[3] and containing several bioactive compounds with anti-oxidant, radical-scavenging, and other therapeutic and immunity-boosting properties,[4−6] the crop has experienced persistent low utilization for decades.[7] Sugar in sweet potato is a fundamental aspect of its eating quality;[8,9] this has been directly linked with its characteristic flavor and there have been speculations by specialists in the field
that reducing sweetness will lead to increased utilization of sweet potato as a staple.[7,10] Categorization of sweet potato into staple, supplemental staple, and luxury types is based on sugar and dry matter contents;[11] for certain ethnic groups, the preferred types are non-sweet or only slightly sweet being regarded as a staple carbohydrate energy source (or supplemental staple type), while sweeter types are preferred in some Western countries as dessert or sweet vegetable (luxury type). Sweetness in sweet potato is due to the presence of endogenous sugars (sucrose, glucose, and fructose) present at harvest, and additional sugar (maltose) formed through starch hydrolysis by amylase enzymes during cooking. Breeding efforts targeted at obtaining low- or non-sugar sweet potato have, therefore, aimed at controlling genes linked with the formation of endogenous mono- and di-saccharides and those linked with amylase enzyme activity.[10,12] Different sugars at the same concentrations are known to have different perceivable sweetness levels. For example, glucose is twice as sweet as maltose, sucrose is three times sweeter than maltose, and fructose is five times as sweet as maltose.[13] In addition to taste/flavor characteristics, sugars are known to also affect the processing behavior of starchy raw materials, for example, gelatinization temperature, degree of gelatinization, and retrogradation.[14−16] The presence of high levels of reducing sugars may also significantly affect the appearance of some heat-processed products by causing excessive darkening.[17] Sweet potato sugars have been the subject of much interest and several reports in this area have been documented.[18−23] In the search for factors affecting variability of sugar content, various pieces of information have emerged. For example, high N fertilizer was found to increase free sugar content, and free sugar correlated positively with N, Ca, and Mg content.[24] However, it appears that the influence of the age of the crop at the time of harvesting on sugar content and composition has not been studied much. Harvest age in sweet potato is not hard and fast and may vary at the discretion of the farmer depending on target markets, personal economic situation, type of cultivar under cultivation, etc. Early-maturing types may be harvested with good yields from 90 days after planting, although it is known that roots can be maintained in the field for extended periods up to 6 months or more. In many tropical regions, there are no long-term storage practices for sweet potato due to the warm climate and lack of sophisticated facilities, unlike some other countries where storage for up to 8 months or more is possible.[8,25] It is documented that under marketing conditions in tropical developing countries sweet potatoes have a shelf-life of only 1−2 weeks.[26] The objective of this study was to assess free sugar contents in the roots of staple-type high dry-matter sweet potato cultivars at different levels of maturity right after harvest and after 3 weeks storage under ambient conditions. If sugar contents and composition of sweet potato cultivars (and how these change during storage) are significantly affected by harvest age, then timing the harvest appropriately would be a simple and effective means of controlling the eating quality and processing characteristics of such cultivars. This could possibly be applied in targeting desired sugar levels for specific end-uses, for instance, low sugar for staple food uses and high sugar for industrial production of sweeteners from sweet potato.

**MATERIALS AND METHODS**

**Sweet Potato Cultivars**

Two relatively new cultivars, Hi-starch and CRI-Otoo, released in 2005 and two old cultivars, Faara and Sauti, all with high dry matter contents (30% or more), planted at the
same location (with 30–34°C day temperatures, 22–25°C night temperatures), and given the same management practices were obtained from the Crops Research Institute, Ghana. They were harvested at 3, 4, and 5 months after planting and processed into flour. At each harvest, some roots were saved, stored at room temperature (25°C ± 4) for 3 weeks, and also processed into flour.

**Flour Preparation**

Representatively sized roots (i.e., small, medium, and large) were selected at each sampling time, peeled, washed, shredded with a hand grater, dried in an air oven at 60°C for 72 h, and milled (using a Cyclotec 1093 sample mill, Foss, Denmark) to pass through a 60–80 mesh screen.

**Sugar Analysis**

**Extraction of alcohol-soluble sugars.** Flour samples (100 mg) were accurately weighed into glass centrifuge tubes (16 × 120 mm; 17 ml capacity). Soluble sugars were extracted by adding 10 ml of 80% aqueous ethanol to the tubes and incubating at 80–85°C for 10 min, with intermittent mixing on a vortex stirrer. The tubes were centrifuged for 10 min at 1000 × g (3000 rpm). The supernatants were carefully poured off into 50-ml beakers; the pellets were re-suspended in another 10 ml of 80% ethanol and the process was repeated. Supernatants were pooled to obtain the total extracts of soluble sugars, quantitatively transferred into 25-ml volumetric flasks, and made up to the volume with 80% ethanol. Duplicate extractions were carried out for all samples.

**Preparation of standards for high performance liquid chromatography (HPLC).**

- **Internal standard solution:** Cellobiose (40 g) was weighed into a 100-ml volumetric flask; 50 ml of distilled water was added and sonicated to dissolve. It was then made up to volume with more distilled water and mixed thoroughly.
- **Standard solution:** Into a 50-ml volumetric flask, 5 mg of glucose (Sigma, St. Louis, MO, USA), 15 mg of fructose (Fisher, Waltham, MA, USA), and 50 mg of sucrose (Fisher) were weighed; 30 ml of distilled water was added, the mixture was sonicated to dissolve, and the contents made up to the volume with distilled water.
- **HPLC standard:** Into a test tube, 9.5 ml of water was carefully measured with a pipette. Then, 250 µl each of internal standard solution and standard solution (described above) was added, vortexed to mix, and filtered through a Dionex OnGuard-H column (Dionex Corporation, Sunnyvale, CA, USA) into an HPLC auto sampler vial. This standard was run at least three times during the course of the analysis.

**Analysis.** Five ml of each extract was placed in a 50-ml beaker and the ethanol was evaporated by leaving uncovered beakers overnight in a working fume hood. Precisely 1 ml of internal standard was added to each beaker and the residue dissolved using swirling and sonication. The solutions were transferred into 2-ml screw-capped vials, capped and kept frozen where necessary till the time of analysis. For analysis, 50 µl of each solution was diluted with distilled water to 2 ml in a small test tube and poured into a syringe fitted with a Dionex OnGuard-H filter. The solution was passed through the filter, the first 1 ml discarded, the remaining 1 ml collected in an HPLC autosampler vial, and analyzed using a Dionex BioLC AD 50 HPLC system (Dionex Co., Sunnyvale, CA, USA). The sample
was injected and eluted through a Carbo PAC PA-1 column (250 × 4.6 mm id) (Dionex Corporation) at 30°C with the mobile phase consisting of 200 mM sodium hydroxide at an isocratic flow rate of 1.0 ml/min. A Dionex PAD (pulse amperometric detector) was used for detection of peaks, and the sugars were identified based on retention times.

\[
\text{Calculation of } \% \text{ sugar} = \left( \frac{A}{B} \right) * \left( \frac{C}{D} \right) * \left( \frac{E}{F} \right),
\]

where

- A = Peak height of sugar;
- B = Peak height of internal standard;
- C = Dilution factor;\(^{[40]}\)
- D = Weight of sample in HPLC vial;
- E = Concentration of sugar in the standard mix;
- F = Standard ratio.

**Flour Pasting Properties**

Pasting properties were determined by a Rapid Visco-Analyzer (RVA) (Newport Scientific Pty. Ltd., Warriewood, NSW, Australia) using 12% (dry basis) flour slurries, both in distilled water and in 0.05 mM AgNO\(_3\), a potent amylase inhibitor.\(^{[27]}\) An estimated index of amylase activity was calculated as (PV2-PV1)/PV1, where PV1 was peak apparent viscosity in amylase-active samples (obtained using distilled water) and PV2 was that in amylase-deactivated samples (obtained using AgNO\(_3\)).\(^{[28]}\) Tests were run in duplicate. Apparent viscosity was expressed in Rapid Visco Units (RVU, where 1 RVU = 12 Centipoise).

**Statistical Analysis**

All experiments were performed in duplicate. Statistical analysis was performed using GenStat Discovery statistical software (Edition 3, VSN International, Hemel Hempstead, UK). Data were analyzed by general linear model (GLM). Differences at \(p < 0.05\), \(p < 0.01\), and \(p < 0.001\) were considered to be significant. Pair-wise comparison of all means was performed using Duncan’s multiple comparison procedure.

**RESULTS AND DISCUSSION**

**Total Soluble Sugar Contents**

Total soluble sugar (TSS) was made up of sucrose, glucose, and fructose and ranged from 4.10–10.82 g/100 g on a dry-weight basis (Table 1). It has been suggested that sugar levels for staple and supplementary staple sweet potato cultivars should be up to 2 and 5%, respectively, while ‘luxury’ types could have variable sugars, generally higher.\(^{[29,30]}\) Sucrose was the major component of TSS in the staple-type, high dry-matter cultivars studied (Fig. 1) and this is in line with previous reports that cultivars with high dry weight tend to have high sucrose and low reducing sugar content (glucose and fructose).\(^{[11,25]}\) Mcharo and LaBonte,\(^{[31]}\) working with 45 sweet potato clones, also confirmed that clones with high sucrose had low levels of glucose and fructose.
Table 1 Total soluble sugar contents (sucrose + glucose + fructose, g/100 g dry matter basis) of four sweet potato cultivars harvested at 3, 4, and 5 months maturity.

<table>
<thead>
<tr>
<th>Cultivar</th>
<th>Maturity</th>
<th>At harvest</th>
<th>After storage (3 weeks)</th>
<th>Difference/change</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>3 months</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hi-starch</td>
<td></td>
<td>4.62 [±0.23]a</td>
<td>6.95 [±0.35]ab</td>
<td>2.32 (50.4%)</td>
</tr>
<tr>
<td></td>
<td>4 months</td>
<td>4.46 [±0.03]a</td>
<td>8.66 [±0.70]b</td>
<td>4.19 (94.1%)</td>
</tr>
<tr>
<td></td>
<td>5 months</td>
<td>8.17 [±0.28]b</td>
<td>3.81 [±0.11]a</td>
<td>−4.36 (−53.4%)</td>
</tr>
<tr>
<td>Otoo</td>
<td>3 months</td>
<td>5.17 [±0.11]a</td>
<td>8.90 [±0.47]a</td>
<td>3.72 (72.0%)</td>
</tr>
<tr>
<td></td>
<td>4 months</td>
<td>6.31 [±0.19]a</td>
<td>9.75 [±0.69]a</td>
<td>3.44 (54.6%)</td>
</tr>
<tr>
<td></td>
<td>5 months</td>
<td>10.34 [±0.34]b</td>
<td>7.87 [±0.61]a</td>
<td>−2.47 (−23.9%)</td>
</tr>
<tr>
<td>Sauti</td>
<td>3 months</td>
<td>4.10 [±1.127]a</td>
<td>7.44 [±0.25]b</td>
<td>3.33 (81.3%)</td>
</tr>
<tr>
<td></td>
<td>4 months</td>
<td>4.55 [±0.14]a</td>
<td>10.82 [±0.81]c</td>
<td>6.26 (137.7%)</td>
</tr>
<tr>
<td></td>
<td>5 months</td>
<td>7.36 [±0.88]b</td>
<td>4.51 [±0.06]a</td>
<td>−2.85 (−38.8%)</td>
</tr>
<tr>
<td>Faara</td>
<td>3 months</td>
<td>4.31 [±0.11]a</td>
<td>7.98 [±0.66]a</td>
<td>3.67 (85.4%)</td>
</tr>
<tr>
<td></td>
<td>4 months</td>
<td>5.41 [±0.34]a</td>
<td>8.99 [±0.81]a</td>
<td>3.57 (66.2%)</td>
</tr>
<tr>
<td></td>
<td>5 months</td>
<td>8.82 [±0.68]b</td>
<td>8.64 [±0.49]a</td>
<td>−0.17 (−1.9%)</td>
</tr>
</tbody>
</table>

Numbers in square brackets represent standard deviation of the means. Means followed by a common letter are not significantly different (mean separation by LSD, $p < 0.05$, for main effects of harvest time).

Reducing Sugar Contents

In our cultivars, reducing sugar contents were generally low at all harvest times (glucose was 0.1331–1.006 g/100 g and fructose 0.0665–1.6838 g/100 g dry-weight basis). Wang and others[32] found that starch and sugar (i.e., sucrose, glucose, and fructose) levels correlated with root weight/size during development, with glucose and fructose decreasing gradually, and sucrose and starch contents increasing with the expansion of tuberous roots. This to some extent appears to be in line with our findings, i.e., sucrose contents increasing with increasing maturity (Fig. 1) although their studies were limited to root weights below or up to 200 g, and in our study the effect of root size within each cultivar would not come into play since representative sampling was done before processing.

Effects of Maturity and Storage on Sugars

Fructose levels in samples harvested at 5 months increased considerably after storage (Fig. 1). These observed increases in fructose content may have implications in heat processing due to characteristic darkening of products (for example during frying) and harvest timing could be studied for other cultivars that tend to have high contents of endogenous reducing sugars. TSS contents on the day of harvest were highest at 5 months maturity for all the cultivars, ranging from 7.36 to 10.33 g/100 g (Table 1). After 3 weeks of storage there were large increases in TSS levels in samples harvested at 3 and 4 months, ranging from 50.5 to 137.7% of their original levels (Table 1). On the contrary, in samples harvested at 5 months, TSS contents dropped after storage (Table 1) with the greatest declines occurring in Hi-starch and Sauti (−53.4 and −38.8%, respectively). Since sucrose was the predominant sugar, net changes in TSS showed virtually the same trends as for sucrose. Sucrose contents on the day of harvest were lower at earlier maturity (i.e., 3 and 4 months) and highest at 5 months (Fig. 1). After 3 weeks of storage under ambient tropical
Figure 1 Individual components of soluble sugar (sucrose, glucose, and fructose) in sweet potato varieties at 3, 4, and 5 months maturity both (a) at the time of harvest and (b) after 3 weeks in storage. The error bar in each chart shows the least significant difference (LSD) for effect of harvest time ($p < 0.05$).

conditions, however, sucrose contents increased in 3- and 4-month samples but decreased in those harvested at 5 months, while glucose and fructose contents generally increased after storage. A similar observation has been reported\cite{22} in two white-fleshed cultivars during the early stages of a prolonged storage period, where sucrose concentration decreased and alcohol-insoluble solids increased. This was consistent with the belief that sucrose may be used up as a carbon source for postharvest starch synthesis in reserve tissue.\cite{33} Since sucrose is made up of glucose and fructose units, it is suggested that in our 5-month samples, the decrease in sucrose and increase in fructose levels after storage may reflect the uptake of the glucose fraction from sucrose for starch synthesis. Such reduction in sucrose content after storage was only observed in 5-month harvested samples for all four of our cultivars (Fig. 1). Other workers\cite{34,35} had somewhat conflicting results in terms of changes in individual and total sugar concentrations in sweet potato at harvest and during storage.
and this could be due to the fact that maturity at the time of harvest was not taken into account. Lee and Chen,\textsuperscript{[36]} studying sweet potato in short-term storage (7 to 14 days), found TSS to either increase progressively or increase and then decrease depending on cultivar, even though age of the cultivars at the time of harvest was not factored in. The activity of sugar-related enzymes, such as acid invertase and sucrose synthase, in sweet potato have been studied by various workers,\textsuperscript{[37,38]} and sucrose metabolism has been linked with the regulation of processes associated with sink strength and starch accumulation in sweet potato.\textsuperscript{[39,40]} This has direct implications on economic factors, such as dry matter yield, and may explain why, in spite of the quest for low- or non-sugar types, the world’s sweet potato clones were found to be predominantly either high or moderate in terms of sweetness, measured as sucrose equivalents.\textsuperscript{[10]} In freshly harvested roots, differences in TSS due to maturity were highly significant at both $p < 0.01$ and $p < 0.05$, while cultivar differences were only significant at $p < 0.05$ (Table 2). After storage, the influence of maturity was significant but cultivar differences were not ($p < 0.05$).

**Flour Pasting Properties and Amylase Activity**

An index of amylase activity was calculated for each sample using RVA data obtained from both amylase-active and amylase-inhibited slurries. Trends in amylase activity were significantly affected by harvest timing ($p < 0.05$) but not by cultivar (Table 2). The estimated index of amylase enzyme activity right after harvest was highest in 5-month samples, and in 4-month samples after storage (Fig. 2). The relatively low levels of amylase activity in freshly harvested samples at 4 months, a maturity stage that coincides with the highest starch content in the cultivars studied (unpublished data), could be attributed to the action of endogenous inhibitors of amylase. The presence of amylase inhibitors in the development of starchy storage organs is essential for protecting starch granules from breakdown by enzymes, especially during phases of active starch accumulation.\textsuperscript{[41]} The high amylase

\begin{table}[h]
\centering
\caption{ANOVA for cultivar differences and the effect of harvest age on total soluble sugars (TSS) (dry weight basis) and amylase activity index (AAI) of sweet potato flour prepared from roots at harvest and after 3 weeks storage.}
\begin{tabular}{llll}
\hline
 & \textbf{TSS (g/100 g)} & \textbf{AAI} & \textbf{TSS (g/100 g)} & \textbf{AAI} \\
\hline
\textbf{At harvest} & & & \textbf{After 3 weeks storage} & & \\
Differences due to harvest age (average of four cultivars) & & & & \\
3 months & 4.55 (± 0.46)a & 0.46 (± 0.28)a & 7.81 (± 0.83)ab & 1.09 (± 0.67)a \\
4 months & 5.18 (± 0.86)a & 0.37 (± 0.17)a & 9.55 (± 0.95)b & 2.92 (± 1.03)b \\
5 months & 8.67 (± 1.25)b & 0.95 (± 0.42)b & 6.20 (± 2.40)a & 1.08 (± 0.85)a \\
\textbf{F-value} & 83.60*** & 10.13* & 5.64* & 7.89* \\
Differences due to cultivar (average of three harvest ages) & & & & \\
Hi-starch & 5.75 (± 2.09)a & 0.98 (± 0.52)b & 6.47 (± 2.46)a & 1.63 (± 1.68)a \\
Otoo & 7.27 (± 2.71)b & 0.56 (± 0.34)a & 8.83 (± 0.94)a & 1.10 (± 0.75)a \\
Sauti & 5.33 (± 1.76)a & 0.46 (± 0.24)a & 7.58 (± 3.15)a & 1.50 (± 1.04)a \\
Faara & 6.17 (± 2.35)a & 0.38 (± 0.20)a & 8.53 (± 0.51)a & 2.55 (± 1.28)a \\
\textbf{F-value} & 8.79* & 5.67* & 1.71 NS & 1.99 NS \\
\hline
\end{tabular}
\end{table}

Numbers in brackets represent standard deviation of the means. Means followed by a common letter are not significantly different (mean separation by LSD, $p < 0.05$, for main effects of harvest time and cultivar).

***Significant at $p < 0.001$; *significant at $p < 0.05$; NS: not significant.
Figure 2 Estimated amylase activities of sweet potato cultivars harvested at 3, 4, and 5 months maturity using flour from (a) freshly harvested roots and (b) roots stored for 3 weeks. The error bar in each chart shows the least significant difference (LSD) for effect of harvest time ($p < 0.05$).

activity of 5-month samples at harvest and 4-month samples after storage could be due to various regulatory mechanisms including a possible drop in amylase inhibitor action. In the same vein, the reduction in amylase activity after storage observed in Hi-starch and Otoo when harvested at 5 months could be due to increases in amylase inhibitor activity (Fig. 2). The possibility of such postharvest increase in inhibition of amylase enzyme activity may be supported by previously reported evidence of starch synthesis in some reserve tissues during the postharvest phase.\cite{33} Amylase activity in sweet potato storage roots during processing is of much importance since it results in the breakdown of starch through hydrolysis to form more sugar (maltose) during cooking,\cite{35,36,42} giving rise to particular product characteristics. The amylase activity index correlated positively with TSS ($r = 0.60$; Fig. 3). This implies that in the cultivars studied those with relatively higher initial sugar content in the raw/unprocessed state would also have more potential for increases in total sugar levels during cooking based on the higher amylase enzyme activities. TSS correlated negatively with RVA peak apparent viscosity ($r = -0.54$, Fig. 2), reflecting previous reports that higher sugar levels to some extent are linked with less thickening of starchy pastes during the cooking process.\cite{14–16}

CONCLUSION

Maturity was found to significantly affect total soluble sugar content and amylase activity of the cultivars studied, and how these changed during short-term post-harvest storage. This information can serve as a springboard to provide opportunities to possibly control the eating quality of sweet potato through the formulation of recommendations for harvest timing and fresh produce quality of specific target cultivars, hopefully to match specific needs.
Correlation of total soluble sugars in storage roots (uncooked) of four sweet potato cultivars at different maturity stages with: (a) estimated amylase activity ($r = 0.60$) and (b) peak apparent viscosity in rapid visco units (RVU, where 1 RVU = 12 Centipoise) ($r = -0.54$).

ACKNOWLEDGMENT

The authors gratefully acknowledge the Agricultural Sub-Sector Improvement Project (AgSSIP), The World Bank, Ghana, for providing funding for this study.

REFERENCES


