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Biochemical Composition of Soursop Fruit, *Annona muricata* L., as Affected by Two Harvest Seasons

C. O. Omoifo

Department of Crop Science, Ambrose Alli University, Ekpoma, Nigeria.

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ABSTRACT

Soursop, Annona muricata L., is a crop with promising economic value. The effects of season of harvesting and state of hardness on biochemical composition of the fruit juice were investigated. Analysis of variance of mean determinants of treatments, season versus hardness, was done and means separated by the LSD, p<.01. Value of pH for class 1 (hard-but-ripe) and class 11 (ripeand-soft) were 3.6 and 4.6, respectively. High significance occurred for sugar content, 17.0 19.0% and 20.0 23.0% for wet and dry seasons harvests, respectively, identified as glucose, fructose and maltose. The organic acids were citric and malic. Titratable acidity, as citric acid, also significantly different, were 156.85 438.40mg/100ml and 297.60 842.24mg/100ml. Similar significance was reflected for other parameters including volatile acidity, as acetic acid, 6.0 39.0,g/100ml and 168.0 108.0mg/100ml, fixed acidity, 150.80 399.63mg/100ml and 129.6 735.53mg/100ml, ascorbic acid, 108.75 - 196.13mg/100ml and 166.21 204.01mg/100ml, respectively for wet season and dry season harvests. Percent protein was 1.19-1.71% and 0.50-1.40% for the wet and dry seasons, respectively. Soft fruits had alcohol content of 1.2% and 0.9% for wet and dry season harvests, respectively. Seasonal changes, rather than storage conditions, were thought to have influenced parameters determined.

Key words: Chemical composition, dry season harvest, hardness, seasonal changes, soursop juice, Wet season harvest.

INTRODUCTION

Soursop fruit, *Annona muricata* L., a native of tropical America and West Indies (Dalziel, 1948; Harrison *et al.*, 1969; ANON, 1975) hardly grows in temperate countries (Janick *et al.*, 1969; Purseglove, 1974). It was introduced to China, Australia and Africa (ANON, 1975) and thrives in the rain forest zones of West Africa (Glendhill, 1972; Adams, 1972). Observation showed that it grows throughout Southern Nigeria. The fruit, which has a pleasant aroma, is taken as dessert or made into a refreshing beverage (Dalziel, 1948; Sanchez - Nieva *et al.* 1970; ANON, 1975).

It can be processed without losing its aromatic flavor (Sanchez - Nieva *et al.* 1970; ANON, 1975). Benero *et al.* (1971) found that lye peeling of the fruit discolored pulp and impaired juice quality, but handpeeling proved adequate.

Pulping was done with a 0.060- inch screen and a finisher with a 0.020-inch screen, which had higher extraction performance (Sanchez-Nieva *et al.*, 1970). Frozen pulp without addition of sugar could be used in the manufacture of nectar, soft drinks, ice cream and similar products and is served as nectar base when sugar is added.

Addition of ascorbic acid improved the puree quality. Benero *et al.* (1974) prepared soft drinks using soursop, 15° Brix and a blend of soursop tamarind, 15° and 17° Brix, respectively and sensory evaluation showed that they were acceptable for one year and ten months, respectively.

Sanchez Nieva *et al.* (1970) found that soursop pulp contained total solids, 16.6 - 18.6%; total sugar (as invert sugar), 11.6 - 12.5%; reducing sugars, 9.5 - 11.7%; acidity (as

anhydrous citric acid), 0.89 0 - 096%; soluble solid, 15.8 - 17.4%; and pH 3.61 - 3.65. However, Chan and Lee (1975), using a gas chromatograph with flame ionization detector reported a lower total value for sugars (10.58%) detected as glucose, fructose and sucrose. In one study, Enweani et al., (2004) reported that total carbohydrate was 12.52g/l of juice content. The study referred showed quality differences between ripe and unripe fruits of soursop and sweetsop, respectively. However, it does not appear that a study showing the effect of seasonal changes on quality parameters of the soursop juice has been conducted. This paper is on the biochemical analysis of soursop fruit harvested at two different seasons of the year.

MATERIALS AND METHODS

Collection of samples

Fruits were collected from Scot's farm, Ubiaja in Edo State, Nigeria. Selective plucking was done from treetops and fruits were immediately separated into two categories: ripe-andsoft and, hardbutripe fruits known by the yellowish taint of the green oblong-shaped berry (Fig.1). All collections were done within one week in order to avoid physiological variation. The dry season collection was made in January, while the wet season collection was done in June of the same year.



Fig. 1: A ripe soursop (Annona muricata L.) fruit.

Labeling and storage of samples, juice preparation

Wet and dry season harvests (S and Z) were labeled as SS_1 or SS_2 and Z_1 or Z_2 for hard-but-ripe or ripe-and-soft fruit samples. Dry season samples were stored in a deep - freezer at 18° C for 6 months. The fruits were peeled, cored and seeds separated from pulp (Fig. 2); pooled puree thus obtained for each sample was analyzed. The pH of extracts were taken at 20° C using a pH meter, model 7020 (Electronic Industries Ltd) equiped with a glass electrode. Except otherwise stated, all determinations were done in triplicate.



Fig. 2: A sectioned fruit of soursop showing core, pulp and seeds. C, core; P, pulp: S, seeds.

Determination of sugar content

Standard D-glucose solutions were prepared and developed using the anthrone method (AOAC, 1970). Optical density (O.D.) of the solutions was determined at 625nm (Sharma *et al.*, 1984). A standard curve was constructed by plotting O.D. readings against concentration gradient. A 0.05ml of each juice sample was similarly treated. The concentration of sugar in the juice was found by extrapolating from the standard curve.

Determination of titratable acidity

A 20ml of centrifuged (at 3000g for 5min)

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soursop extract was transferred to a 250ml flask and 10 drops of phenolphthalein were added. A white tile was placed under the flask and sample was titrated against 0.1N NaOH from a burette until a definite pink end point was attained. Titratable acidity (as citric acid) per 100ml juice was obtained according to AOAC (1970).

Determination of volatile and fixed acidity

Volatile and fixed acidity of fruit extracts were determined by the standard methods (AOAC, 1970).

Determination of alcohol content by volume

This was done according to the description by AOAC (1970).

Determination of ascorbic acid content

Soursop extract was added to 200ml freshly prepared 0.6%TCA + 0.005M Na₂ EDTA in a 500ml beaker and mixed thoroughly. A 20ml of this solution was transferred to 100ml volumetric flask and made up to mark with 0.3% TCA + 0.0025M Na₂EDTA and mixed thoroughly. This was filtered through fluted Whatman No. 42 filter paper into a 150ml beaker to remove suspended solids. The filtrate was subjected to further analysis following the description in methods of vitamin assays (A.O.V.C., 1966).

Determination of protein content

Digestion of sample was as described by AOAC (1970) but with slight changes as in the modification of 11TA (1979). A 5ml sample was measured into 500ml round bottom flask into which 5ml conc. H₂O₂ was added in a fume chamber and mixed. The mixture was placed on a heating rack with temperature raised in steps until 50°C when it boiled. Further measures of H₂O₂ were added to ensure complete digestion. A blank with distilled water was similarly treated.

Distillation of ammoniacal nitrogen was done following the method described in AOAC (1970). Crude protein was calculated thus,

nitrogen = $\underline{\text{sample titre-blank titre x 0.56}}$ Weight of sample

% protein = $\frac{\text{sample titre-blank titre } \times 0.56 \times 6.25}{\text{Weight of sample}}$

Chromatographic analysis of sugar

This was done using a one-dimensional descending paper chromatography. Drops of samples and reference standards were spotted 3cm apart on Whatman No. 1 filter paper (40 x 45cm) with 0.1cm glass tube and quickly dried under a stream of hot air. The reference sugar solution contained 1% of each of arabinose, fructose, glucose, galactose, glycerol, lactose, maltose, raffinose, sucrose and xylose in distilled water. The chromatograph was equilibrated overnight in a chromatank (82.5 x 23 x 82.5cm internal dimensions, Shandon Sci. Co., London) with 23ml of solvent system, n-butanol-acetic acid-water (4:1:1). Subsequently, the chromatogram was run for 18h, dried in air and developed in silver nitrate solution according to the method of Trevelyan et al. (1950). Darkbrown spots against a white background showed the presence of sugars. The strip was washed in 5% sodium thiosulphate and thereafter in running water for 1h. Sugars in samples were identified by comparing Rf ' values of spots with that of reference sugars run simultaneously, where Rf' was given by the ratio, distance traveled by substance /distance traveled by solvent front.

Chromatographic analysis of organic acids

The method of Kunkee (1968) was employed. The solvent system was n-butanol-distilled water-formic acid- bromocresol green (100: 100: 10.7:15). The solvent mixture was shaken for 20min in a separation funnel and allowed to settle. The aqueous layer was let out and the organic layer used. Spotting, equilibration and development of chromatogram were done as

described using acetic, fumaric, lactic, citric, glutaric, malic, oxalic, succinic, and tartaric acids as reference standards. The chromatogram was run overnight, and then air-dried at room temperature. Spots appeared with yellow color against a blue-green background. Identification was done as described.

RESULTS

The Rf values of sugar and organic acids in soursop juice, measured against their standards in each case, were shown in Tables 1 and 2.

Table 1: Rf 'values of spots of standard sugars and soursop juice extracts.

Sugar	Rf(x100) in n butanol - acetate - water							
	n butanc	or - acetate	- water					
Arabinose	26.68							
Fructose	26.20^{a}							
Glucose	12.99ª							
Galactose	18.02							
Glycerol	53.13							
Lactose	6.49							
Maltose	7.93°							
Raffinose	3.85							
Sucrose	13.76							
Xylose	30.53							
S1 ^b	7.29	12.60	23.95					
S2	7.28	12.81	26.00					
Z 1	7.28	12.50	26.10					
Z2	7.30	12.70	25.90					

^a Sugar Rf' values corresponding with fruit sample Rf's. ^b S1, S2, Z1, Z2 are wet and dry season soursop juice samples.

Both classes of fruit hardness had a range of pH 3.60-4.40. Means of parameters determined for juice were shown in Table 3. There was significant difference, p<. 01, between the levels of sugar determined for class 1 and 11 hardness, with higher values for the softened fruits. This means that as the fruit softened higher levels of sugar accumulated in the juice. This trend was

Also reflected in juice contents of ascorbic acid,

titratable acidity and fixed acidity. The sugar content was 17.03 -23.00%. This was chromatographically detected as glucose, fructose and maltose (Fig.3). Chromatographic analysis of organic acids showed the presence of citric and malic acids (Fig. 4). The Rf' values of sample sugars and organic acids were lower than those of standards. Total titratable acidity was 156.85 - 842.24mg/100 1. Volatile acidity, measured as acetic acid had a range from 6.0 168.00mg/100ml, while fixed acidity as expected, was much higher (129.67 -735.53mg/100ml). The ascorbic acid content was high, 108.75 -204.01mg/100ml, but the protein content did not exceed 1.71%. Specific gravity bottle determination of alcoholic content was done for ripe-and-soft fruits only. This was found to be 0.9 and 1.2% for dry season and wet season harvests, respectively.

Table 2: Rf' values of spots of organic acid standards and soursop juice.

Acid	Rf (x100) in n-butanol- formate-water					
	1	omate-wate				
cetic acid	90.60					
Hutaric acid	92.60					
Citric acid	63.06 ^a					
umaric acid	95.67					
Malic acid	70.00^{a}					
actic acid	86.77					
Hyoxillic acid	56.40					
Oxalic acid	37.50					
uccinic acid	86.56					
1 ^b	62.50	70.62				
52	62.50	69.70				
21	62.50	70.00				
22	6.250	69.88				

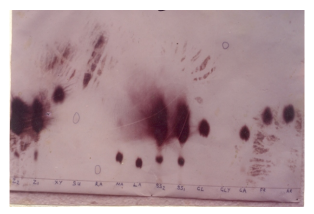
^a Acid Rf' values corresponding with sample Rf's.

^b S1, S2, Z1 and Z2 are wet and dry season soursop juice samples.

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Table 3: Chemical attributes assessed in juice of soursop fruit harvested in two different seasons

Fruit properties		Sugar %		Ascorbic acid		Protein %		Titratable		Volatile acidity		Fixed acidity		Alcohol content	
				mg/100ml				acidity		mg/100ml		mg/100ml		%	
					mg/100ml										
H ardness	Class	Wet	Dry	Wet	Dry	Wet	Dry	Wet	Dry	Wet	Dry	Wet	Dry	Wet	Dry
		season	season	season	season	season	season	season	season	season	season	season	season	season	season
	1	17.0	20.0	108.3	116.16	1.16	0.5	156.85	297.57	6.0	168.03	150.85	129.54	ND	ND
	2	19.3	23.0	196.45	202.89	1.71	1.4033	438.40	842.24	39.01	108.00	399.39	734.24	1.21	0.90
		P<0.01 Lsd: Season, 0.1195; Hardness, 0.1195		P<0.01		P<0.01		P<0.01		P<0.01		P<0.01			
				Lsd: S	Lsd: Season, Lsd: Seas		Lsd: Season, Lsd: Season, 0.0261; 0.3391;		Lsd: Season,		Lsd: Season,				
				, , , , , , , , , , , , , , , , , , ,		0.02			1 91;	0.1501;		0.2693;			
						Hardness,		Hardness,		Hardness,		Hardness,			
						0.0	261	0.3	391	0.1	501	0.20	693		





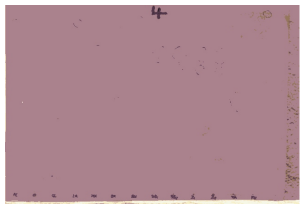


Fig. 4

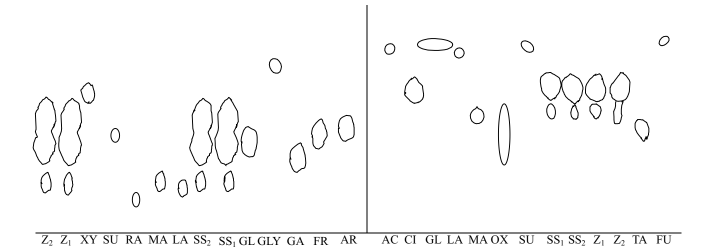


Fig.3: Chromatographic detection of sugars in soursop fruit extract. The solvent system was – butanol: acetic acid: water (4:1:1) Symbols: AR,arabinose; FR, fructose; GA, galactose; GL,glucose;GLY, glycerol; LA, lactose; MA, maltose;RA, raffinose; SU, sucrose; XY, xylose;SS₁, freshly harvested matured and ripe fruits; SS₂, freshly harvested ripe and soft fruit; Z_1 , matured and ripe fruit stored for 6 months; Z_2 , ripe and soft fruit stored for 6 months.

Fig. 4: Chromatographic detection of organic acids in soursop fruit extract. The solvent system was n-butanol: distilled water:formic acid: bromocresol green (100:100:10.7:15). AC, acetic acid; CI, citric acid; FU, fumaric acid; GI, glutaric acid; MA, malicacid; OX, oxalic acid; SU, succinic acid; YA, Tartaric acid; SS₁,SS₂,Z₁ andZ₂ as in fig.3.

Dry season samples had higher values of percent sugar, titratable acidity, volatile acidity, pH and ascorbic acid in comparison with the wet season harvests. But this was not so for fixed acidity and percent protein where the wet season lot had higher values.

Table 3 showed that softened fruits in either seasons contained high percent sugar. The sugar content of juice increased from 17.00% to 19.30% for the wet season harvest, and from 20% to 23% for the dry season harvest. Similar increase was also reflected in the total titratable acidity, where the wet season lot increased from 156.85mg/100ml to 438. 4 mg/100ml. The dry season lot also increased from 297.60mg/100ml to 842.24mg/100ml. But while there was increase in volatile acidity for the wet season crop, 6.0 to 39.0 mg/100ml, the dry season harvest showed a decrease (168 to 108mg/100ml), as the fruit softened.

DISCUSSION

In this study, ripe-andsoft fruits were analyzed along side hard but ripe fruits. Differences in sugar content may be due to hydrolytic enzyme activities that accompany ripening and softening, which result from solubilization of pectic substances associated with the middle lamellae (Leopold and Kriederman, 1975), when fats and

starch also yield sugars (Beevers, 1961). Similarly, higher values for titratable acidity, volatile acidity, fixed acidity, ascorbic acid and protein in softened fruits may be attributed to effect of hydrolysis of more complex parent compounds. Hulme (1954) showed that protein synthesis accompanied the ripening of apple fruits.

Fruits stored at -18°C for 6months presented higher values, p<.01, for sugar, titratable acidity, volatile acidity, fixed acidity and ascorbic acid. Leopold and Kriederman (1975) noted that plant tissues stored at temperatures below 10°C increase in sugar concentration. It is difficult to attribute the increases in values in the present study to the low storage temperature and, or period, since a portion of the dry season harvest was not analyzed at the time of harvest. However, Sanchez-Nieva et al. (1970) showed that unsweetened soursop puree with or without heattreatment and stored at -10°F for 223-225 days or 396-419 days did not show significant change in total solids, pH and color. In our study, it is tempting not to attribute the differences observed to the influence of season since environmental conditions differed greatly between the two harvesting periods.

The evergreen soursop plant which flowers and fruits almost continuously throughout the year (ANON, 1975), has a tap root system for meeting physiological water requirement, in spite of low relative humidity and high atmospheric temperature that occur in the dry season. Results in this study showed that the dry season harvests had higher values for sugar, titratable acidity, fixed acidity, and ascorbic acid, in comparison with fruits harvested in the wet season, when humidity was relatively high. Seasonal differences may also account for the lower protein value recorded for the dry season harvest. Volumetric measurements showed the presence of alcohol in ripe and soft fruits harvested in both season, with a higher value for the wet season crop. The difference in pH of softened fruits between the two seasons is insignificant to the extent that it is within the acidic range.

Malic and citric acids were detected in the dry-and wet-season crops. Citric acid is a product

of the photosynthetic carbon reduction cycle in C₃ plants (Ting and Osmond, 1973), while malic acid occurs as an intermediate product in C₄ plants, but it is quickly metabolized to other products (Edwards and Haber, 1981). In CAM plants, CO₂ is fixed in malic acid in the dark (Beevers et al., 1966), and eventually released from storage in light for the photosynthetic carbon reduction process (Edwards and Haber, 1981; Leopold and Kriederman, 1975). Since malic and citric acids were the only organic acids detected in soursop juice, one may speculate on the occurrence of CAM process in soursop. It is known that crassulacean acid metabolism is a photosynthetic and biochemical adaptation of CO₂ utilization to water stress (Osmond and Holtum, 1981), a condition that possibly prevails for the crop in the dry season. If this is assumed, then the higher sugar, titratable acidity, fixed acidity and ascorbic acid values detected may not be unexpected. The lower value for volatile acidity in dry season softened fruit cannot be accounted for presently.

The total percent sugar (19-23%) in ripe-and-soft soursop fruits is higher than that presented by other workers, 12% (ANON, 1975); 10.58% (Chan and Lee, 1975); 11.6-12.5% (Sanchez-Nieva *et al.*, 1970). The value determined in this study compares favorably with that for grape (*Vitis* sp.) must (18-25%), the conventional wine Making fruit (Amerine, 1964; Okafor, 1987). Therefore, soursop may serve for wine production.

Sugars previously reported in soursop fruit include invert sugar (Sanchez-Nieva *et al.*, 1970), glucose, fructose and sucrose (ANON, 1975; Chan and Lee, 1975). A new discovery in this study is the detection of maltose, disaccharide and reducing sugar, which occurs naturally in plants (Conn and Stumpf, 1976). This appears to be the first report on the detection of maltose in soursop fruit juice.

Several papers (Sanchez-Nieva et al., 1970;

Benero *et al.*, 1974; ANON, 1975) have reported on the use of soursop puree for the manufacture of nectar, ice cream, soft drinks and other similar

products. The high concentration of sugar makes the juice suitable for alcohol fermentation, without further amelioration, which is done when the sugar content is low (Jarczyk and Wzorek, 1977). It's acid pH and aroma (ANON, 1975), may contribute to a high quality product.

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