MICROPROPAGATION OF CASSAVA (*Manihot esculenta*) USING LOCALLY SOURCED MATERIALS AS SUBSTITUTES IN A ROUTINE MEDIUM.

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5 Abstract

In spite of global acceptance and wide use of micropropagation as a method for the production of 6 7 disease free planting material and germplasm conservation, this practice has been slow and nonaffordable in Sub-Saharan Africa. This is due to the high cost and non-availability of tissue 8 9 culture media. Considering the importance of growth factors (micro and macro nutrients) in culture medium, it is inevitable to search for an alternative, cheaper and readily available source 10 11 of these nutrients. This research therefore provided a natural substitute media formation for Cassava nodal culture. Sugar cane juice was substituted for sucrose (SC) in this research work. 12 13 The result showed that the explants survived and produced foliage at 20ml SC and 40ml SC based media. The forest Top Soil (FTS) modified media produced more foliage (7), at 14 15 20ml/200ml than conventional media (5). Trona is a soft and porous salty evaporate deposit occurring in association with Neutron, Halite, Thernadite and other salts. Trona is a mixture of 16 Chlorides, Carbonates, and Sulphate salts of Sodium, Calcium, Potassium, and Magnessium thus 17 serving as a good source for these salts. 0.2g of Trona gave the highest percentage 66% of nodal 18 cutting that developed foliage. In conclusion, there was a positive response observed in the 19 growth of the cassava nodes in the media modified with various natural nutrient sources. The use 20 21 of these natural sources is encouraged because it is less costly and readily available rather than 22 having to (delete) wait for the importation of the costly synthetic culture media.

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24 Keywords: Tissue Culture, Cassava, Micropropagation, Nutrients, Media, Explant.

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26 INTRODUCTION

Cassava is a perennial lowland woody shrub with edible roots, which grows in the tropical and
subtropical areas of the World. All cultivated forms belong to the species *Manihot esculenta*

Crantz; and family Euphorbiacea. It is also called Manioc, Mandioca, Tapioca, Yuca and Sagu in different regions (delete) or countries. Cassava has the ability to grow on marginal lands where cereals and other crops do not grow well, it can tolerate drought and can grow in low nutrient soils. The plant grows very tall, at times reaching a height of about 15 feet, with leaves varying in shapes and size. The edible parts are the tuberous roots and the leaves. The tuber is dark brown in colour and grows up to 2 feet long or more depending on the cultivar and the soil conditions (Schery, 1972).

According to the Food and Agricultural Organization (FAO) estimates, about 172 million tonnes of cassava was produced in year 2000 (FAO; 2002). Africa accounted for 54%, Asia 28% and Latin America and the Caribbean for 18% of the total World production. In 1999 Nigeria produced 33 million tonnes making it the World largest producer.

40 In Africa, cassava provides a basic daily source of dietary energy. Roots are processed into a wide variety of granules, pastes and flours or consumed freshly boiled or raw. In some of the 41 42 cassava growing countries in Africa, the leaves are also consumed as a green vegetable, which 43 provides protein and vitamins A and B. In South East Asia and Latin America, cassava is used as 44 a binding agent in the production of paper and textiles, in North America and Europe, cassava is 45 consumed as Tapioca prepared from cassava Hour (Anon, 2005). Although, cassava is adapted to 46 a wide range of climatic conditions and is tolerant to poor acid soils and drought, several research constraints have been identified in the areas of production processing, and utilization 47 (CIAT, 1989). Pests and diseases, together with poor cultural practices, combine to cause yield 48 49 losses that may be as high as 50% in Africa (Asiedu et al., 1992).

50 Micropropagation techniques have been developed to provide solutions to some of the 51 cassava production constraints. Micropropagation through tissue culture techniques have been 52 used for disease elimination, pest resistance, germplasm exchange, distribution and germplasm conservation (Ng and Hahn, 1985). The medium for micropropagation must contain all 53 54 components necessary to nourish growth of explants to be grown (delete). Though plants do not have the same nutritional requirements, the components of any tissue culture medium must 55 contain the following growth factors: Macro and micro nutrients, carbon source (organics), 56 vitamins, growth regulators, complex organics and inert supports (Gelling agents) (Smith, 1992). 57

Although micropropagation technique has been developed for cassava, the nutrient medium has
utilized both the synthetic and industrially produced components which are beyond the reach and

60 utilization of major stakeholders who are ready to carry out the multiplication of cassava.

This research work is therefore designed to provide a method of micropropagation whichprovides all the necessary growth factors from natural sources.

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64	MATERIALS AND METHODS			
65	EXTRACTION AND PREPARATIONS OF LOCAL MATERIALS			
66	(i) Cane sugar juice extraction			
67	Fresh sugar cane sticks were purchased from local markets. The stems were cleansed			
68	by scrapping. The bark was removed and the cane was shredded with a grater, the			
69	shredded cane was then squeezed to release the juice.			
70	(ii) Preparation of Trona powder:			
71	Impure form of Trona was procured from the local market and ground to powder with			
72	pestle and mortar.			
73	(iii)Forest top soil preparation:			
74	The soil was collected from a forest plot, soaked in excess water and allowed to settle			
75	for about 12 hours. The water was decanted into a bottle for use.			
76	(iv)Lichen and Moss ash preparation:			
77	The Lichens and Mosses were collected from old citrus trees (age) by scrapping bark			
78	of the trees with a scapel. The majority of the collection was Lichens. These Lichens			
79	were the crustose type and only a few were foliose Lichens. Mosses collected were of			
80	various kinds.			
81	The crypto samples were then placed into three crucibles which were put into an			
82	oven. The oven was allowed to operate at a temperature of 600°C for 7 hours. The ash			
83	obtained after this procedure (crypto ash) was allowed to cool and stored.			

84

85 MEDIA PREPARATION

Specific aliquots i.e. 20ml, 40ml, 60ml and 80ml of the sugar cane juice were used to substitute
sucrose in the standard Murashige & Skoog (MS) basal medium (Table 1).

0.1g, 0.2g, 0.3g and 0.4g of powdered Trona were weighed and introduced directly into the
medium without MS basal medium.

10ml, 20ml, 30ml and 40ml of the forest Top soil (FTS) was used as medium to substitute MS
basal medium (Table 1). 1.3g of stored Lichen and Moss ash was weighed and infused into the
medium preparation.

93 **pH ADJUSTMENT**

The pH of all prepared media was adjusted to 5.7 using 1M NaOH. 0.8g of agar was added to each medium and made up to 200ml.

All the media were heated in a microwave to melt the agar. With the aid of an automatic dispenser, the preparations were poured into test tubes and were placed in an autoclave at 121°C at 15psi for 15 minutes. These were (delete) left on the shelf for about 8 hours to cool.

99 CULTURING OF EXPLANTS:

After taking the necessary precautive measures of disinfecting the work bench, healthy plants (age) were collected. The plantlets were removed from test tubes, the nodes were excised and placed on the medium. The test tube was recapped and sealed with a piece of parafilm.

103 The test tubes were placed on the shelf in the culture room under fluorescent light at 27°C room 104 temperature and exposed to 12 hours of light and 12 hours of darkness.

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107 **RESULTS**

The result showed that the media modified with sugar cane juice had the percentage growth of green leaves as high as 87% on 20ml/200ml while the lowest percentage was 50% on 60ml/200ml which favorably compared with the conventional MS media at 88%.

0.29g of Trona gave the highest percentage of nodal cuttings that developed green leaves and
roots (10%) while 86% and 14% produced only green leaves and roots respectively.

113 On the average, the percentage foliage production was higher in the media modified with 114 20ml/200ml FTS which was even higher than the foliage production on the conventional media.

115 Table 1 Routine Cassava Tissue Culture Medium

Component	Quantity in 1 litre
MS Basal Medium	4.43g
Inositol	100mg
Sugar	30g
NAA	0.01mg
BAP	0.05mg
Agar	4g

116 MS – Murashige and Skoog, NAA- Naphthylacetic acid, BAP – Benzyl Amino Purine

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118 Table 2 Response of cassava Nodes to Media Modified with SC after 4 weeks

SC Vol. In 200ml	No Growth %	Green Leaf with	Green Leaf (No	Roots No Leaf
		Root %	Root) %	
20ml	8±1.2	13 ± 3.4	57±7.5	22 ± 4.5
40ml	10 ± 2.1	5 ± 2.0	55 ± 7.1	30 ± 5.2
60ml	26 ± 3.0	0	50 ± 6.4	24 ± 4.1
80ml	11± 2.4	0	72 ± 8.3	28 ± 5.5
Control	9± 2.5	33 ± 6.4	30 ± 3.0	28 ± 4.3

119 SC – sugar cane juice; control – Routine MS (Year) Tissue culture medium

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Vol. In 200ml	No Growth %	Green Leaf with	Green Leaf %	Root %
		Roots %		
0.1gTrona	34 ± 3.5	7±2.1	49 ± 5.4	10 ± 2.0
0.2gTrona	10 ± 2.3	10 ± 2.2	66 ± 6.3	14 ± 2.4
0.3gTrona	27 ± 4.5	4 ± 1.1	53 ± 5.2	16 ± 3.1
0.4gTrona	14 ± 2.2	22 ± 4.6	50 ± 6.4	14 ± 2.6
Control	7±1.7	33 ± 5.7	50 ± 5.5	10 ± 2.2

125 Control: Routine MS Tissue culture medium

126 Table 4: Response of Cassava Nodes to media modified with FTS Preparation after 4

127 weeks.

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Media (vol. In	No Growth %?	Green Leaf with	Green Leaf No	Root No Leaves
200ml)		Roots	Root	
10ml FTS	10 ± 2.4	14 ± 3.2	66± 4.5	10± 2.4
20ml FTS	9 ± 2.4	22 ± 4.5	47± 3.3	22± 4.6
30ml FTS	12 ± 3.1	31 ± 4.7	37± 4.1	20 ± 4.2
40ml FTS	10 ± 3.2	28± 3.5	40 <u>+</u> 4.7	22 ± 5.5
Control	10± 2.5	42± 5.6	30± 3.5	18± 3.4

128 Control: Routine MS Tissue culture medium. FTS: Forest Top Soil

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130 Table 5: Response of cassava Nodes to media modified with Lichens and Moses after 4

131 weeks

Media (Vol. In 200ml)	No Growth %	Green Leaf with Root	Green Leaf No Root	Roots No Leaf
1.3g Lichen and	8± 1.3	10± 2.3	64± 6.7	18± 3.7
Moss ash				
Control	10 ± 2.1	42 ± 5.7	50 ± 5.6	8± 2.0

132 Control: Routine MS Tissue Culture Medium

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134 **DISCUSSION**

The cassava explants were observed for survival (days), green leaves formation and rootsformation on each of the modified media.

137 Generally, all the media prepared from locally sourced materials were effective in sustaining the

138 growth and survival of the cassava explants.

139 The sugar cane replaced sucrose as a source of energy required for the heterotrophic nutrition of140 the explants.

141 It is interesting to note that the percentage of plants that survived or produced foliage especially 142 in the 20ml SC - based medium can be equated to that of the control. It showed also that the 143 20ml and 40ml SC based media were the best concentration of SC required for the sustainable 144 growth of cassava.

Fertile top soil (FTS), Trona, Lichen and Moss were used in this study to substitute the MS basal medium containing industrially produced salts. This study has shown that it is possible to use natural and locally available salts in place of the industrially produced salts. This research work agrees with the research works of Santana *et al.*, (2009) and Kwarne *et al.*, (2012) who used different concentrations of locally available fertilizer to micropropagate cassava.

Different kinds of fertilizers at different concentrations were also used by Escobar *et al.*, (2006) to realize a 24.4% cost reduction for the medium prepared. Trona has been established to be a good source of inorganics for many tropical plants in Africa (Esan, 1993).

153 It was observed that most of the explants regenerated roots without addition of Auxins, this is in 154 agreement with Yona et al, (2010) who reported that cassava explants can naturally develop 155 roots without the addition of Auxins. Alfred and Uchenna (2013) also used locally available 156 materials for substrate hardening in the micropropagation of Sweet Potato.

157 CONCLUSION

This research work is an indication that it is possible to formulate nutrient media for sustaining
Cassava growth from cheaper, local and safer materials to promote micropropagation of Cassava
germplasm through tissue culture.

Future prospects should be to increase input in the development of a natural nutrient medium soas to make micropropagation of not only cassava but other crops more affordable.

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