

## Original Research Article

# Effect of Cultural Conditions on Biosurfactant production by *Candida* sp. isolated from the sap of *Elaeis guineensis*

### ABSTRACT

**Aims:** This study is aimed at determining the effect of cultural condition on biosurfactant production by *Candida* sp. isolates from saps of *Elaeis guineensis*

**Methodology:** Chemical analysis of the sap was carried out. Yeast isolates from the sap were screened for biosurfactant production based on emulsification index ( $E_{24}$ ), emulsification assay, haemolytic assay, oil displacement test, CTAB and tilted glass slide ability. The best biosurfactant-producing yeast isolate was identified based on its phenotypic, microscopic, and biochemical characteristics. The emulsification capacity of the produced biosurfactant on selected oils was studied. Optimum cultural and nutritional requirements (temperature, pH, inoculum concentration, nitrogen sources and carbon sources) for biosurfactant production by the isolate were determined.

**Results:** The characteristics of the sap from *Elaeis guineensis* were reducing sugar ( $0.51 \pm 0.03$  mg/ml), alcohol ( $14.04 \pm 0.15$  %), specific gravity ( $0.827 \pm 0.024$ ), and pH ( $5.68 \pm 0.03$ ). The crude biosurfactant produced displaced a thin film of crude oil on petri dish by 55 mm, and revealed high emulsification index ( $E_{24}$ ) of 52.5% using Olive oil as substrate compared to  $E_{24}$  of 60.6% by sodium dodecyl sulphate (SDS). Based on colonial, microscopic, and biochemical characteristics, the isolate SA2 was identified as *Candida* sp. The crude biosurfactant showed varying capacity in emulsifying the different oils that were examined. Optimization data revealed maximum biosurfactant production after 7 days of incubation, inoculum concentration of 10%, at temperature of 20 °C, pH of 2 with cassava peel as substrate.

**Conclusion:** The study has demonstrated the capacity of *Candida* sp. from the sap of *Elaeis guineensis* to produce biosurfactant utilizing cassava peel as substrate. The use of cassava peel, which represents a low-cost substrate, is important in reducing the cost of biosurfactant production. Moreover, using yeasts from *Elaeis guineensis* make the production process ecologically friendly.

**Keywords:** Biosurfactant, *Candida* sp., optimization, *Elaeis guineensis*

### 1. INTRODUCTION

Biosurfactants are green extracellular molecules synthesized by microorganism such as bacteria, yeasts, and fungi. They are amphipathic in nature comprising hydrophilic and hydrophobic moieties that form partitions between oil/water or air/water interfaces. According to Satpute *et al.* [2], this inherent amphipathic property, increases the solubility of hydrophilic molecules, hence reducing both surface and interfacial tensions at air/water interface.

Recent preference for biosurfactants over chemically synthesized surfactant is due to its higher biodegradability, environmental friendliness, and ability to withstand extreme high temperature, salinity and pH, ease of production from renewable agro-wastes, active and

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22 non-toxic nature, multi-functionality, and specificity in terms of its industrial applications [3].  
23 These surface active molecules are classified as glycolipids (rhamnolipids, sophorolipids and  
24 trehalolipids) [4]; fatty acids and phospholipids [5]; polymeric biosurfactants (emulsan,  
25 alasan) [6-7], and particulate biosurfactants [1], based on their chemical structure and  
26 microbial origin.

27 Microorganisms that produce biosurfactants are naturally abundant; they are found in  
28 ecological places like land (polluted soil, sediment, sludge), water bodies (fresh water,  
29 ground water, marine water), and also in some extreme environments (e.g., oil reservoirs),  
30 where they can flourish in wide range of temperatures, pH values, and salinity [8]. In the past  
31 decades, yeast has proven their proficiency for production of biosurfactant, despite having  
32 been isolated from different sources as recorded by some researchers. This is majorly  
33 attributed to their importance in food and pharmaceutical industries on the basis of “generally  
34 regarded as safe” (GRAS) status and, also ability to produce biosurfactant in a larger  
35 quantities than bacteria [1]. The following *Candida* species: *Candida tropicalis* [9]; *Candida*  
36 *albicans* [10]; *Candida antarctica* [11]; *Candida bombicola* [12]; *Candida* sp. SY16 [13];  
37 *Candida sphaerica* UCP0995 [14]; *Candida utilis* [15]; *Candida glabrata* [16], *Candida*  
38 *guilliermondii* [17] are known biosurfactant producers.

39 The type, quality, and quantity of biosurfactant depend on production process conditions  
40 such as pH, temperature, agitation, aeration, inoculum concentration, nature of substrates,  
41 carbon sources and nitrogen sources [18]. Since environmental factors may significantly  
42 affect the yield and characteristics of the produced biosurfactant, it is therefore essential to  
43 optimize the process conditions in order to achieve high yield.

44 Biosurfactants find application in different areas. In the environment, play vital roles in  
45 bioremediation of polluted soils and refinery wastewater and microbial enhanced oil  
46 recovery; industrially, they have been used in detergent formulation, household cleaning  
47 agent, pesticides and textile production, agriculture, food and pharmaceutical industries [19-  
48 20]. Several biosurfactants exhibits antibacterial, antifungal, antiviral and antitumor (inhibiting  
49 tumor growth and its toxic effects) properties, making them potential alternatives to  
50 conventional therapeutic agents in many biomedical applications [21-22]. This work was  
51 aimed at isolating, screening and optimizing biosurfactant production from *Candida*  
52 *haemulonis* SA2 obtained from the sap of *Elaeis guineensis*. Finally, the ability of the  
53 biosurfactant produced to emulsify different hydrocarbons was evaluated.  
54

## 55 **2. MATERIAL AND METHODS**

### 56 **2.1 Sample Collection**

57 The *Elaeis guineensis* (oil palm) samples used for the yeast isolation were obtained in a  
58 sterile 500 mL sample containers, each from a palm wine taper within 30 to 60 min of  
59 tapping. The samples were aseptically transported to the laboratory in ice packs within two  
60 hours of collection. Sampling was done on two different locations: Bunu, and Kpite  
61 community within Tai Local Government Area (Ogoni land) of Rivers State, Nigeria.  
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63  
64

### 65 **2.2 Physicochemical Analysis of Oil Palm Samples**

66 The physicochemical parameters analysed were pH, temperature, specific gravity, ethanol  
67 content, total dissolved solids at 25 °C, salinity at 25 °C, reducing sugar and conductivity at  
68 25 °C as described by Ukwuru and Awah [23], and titrable acidity Nwachukwu *et al.* [24].

### 69 **2.3 Isolation of Yeasts**

70 For the isolation of hydrocarbon degrading, 2 % (v/v) of crude oil was added to 100 mL of  
71 palm wine in a 250 mL conical flask. The pH of the medium was adjusted to 6. The conical  
72 flasks were then incubated at 28 °C in a shaker incubator with agitation speed of 150 rpm  
73 for 7 days and 14 days, respectively. At each of the days, 1 mL of enriched palm wine was  
74 used for serial dilution according to Nanhini and Josephine [25]; this was followed by  
75 spreading of 0.1 mL from 10<sup>-3</sup>, 10<sup>-5</sup> and 10<sup>-6</sup> dilutions on triplicates potato dextrose agar  
76 (PDA) plates containing 0.05 mg/mL of gentamycin and chloramphenicol (0.1% wt/v) to  
77 inhibit bacterial growth. The plates were incubated at 28 °C for 48 h [26]. The selected  
78 colonies (confirmed to be yeasts using a microscopic examination) were purified by re-  
79 streaking on PDA agar plates. The pure isolates were maintained in PDA agar slants. The  
80 isolates were sub-cultured from the slants for the various experiments conducted in this  
81 work.

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## 82 2.3 Identification of Yeast Isolates

83 The yeast isolates were examined macroscopically on PDA agar plates for the following  
84 features, colony elevation, pigmentation, colony size, nature and shape. For microscopy,  
85 water mount was employed; with a bacteriologic loop, sterile distilled water was placed on a  
86 glass slide and a light emulsion of the yeast made in this drop of water. The glass slide was  
87 covered with a cover slip and examine under X40 objective lens. The reason is because  
88 yeast settles on a slide more quickly in an aqueous medium making it easier to measure  
89 them. The biochemical features examined were urease test, carbohydrates fermentation test  
90 (glucose, galactose, sucrose, maltose, fructose, lactose, raffinose), germ tube, growth at 37  
91 °C, and pellicle formation.

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## 92 2.4 Screening for Biosurfactant Production

93 The yeast isolates were screened for biosurfactant production using the following  
94 techniques: emulsification stability (E<sub>24</sub>) test, emulsification assay, oil displacement, tilted  
95 glass slide and haemolytic assay as described by Nwaguma *et al.* [18]. The selection of the  
96 biosurfactant producer was based on the ability of a given strain to give positive results in all  
97 the screening test procedures.  
98  
99

### 100 2.4.1 Emulsification stability (E-24) test

101 This screening method for biosurfactant-producing microorganisms has been described as  
102 one of the commonest [27]. The method described by Plaza *et al.* [28], was adopted. In  
103 brief, 2 mL of kerosene was added to 3 mL of cell free broth in a test tube and vortexed at  
104 maximum speed for 2 min to homogenize the mixture. After 24h, the emulsification stability  
105 was calculated using the formula below:  
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107  
108

$$109 \text{ E-24} = \frac{\text{total height of the emulsified layer}}{\text{total height of the mixture}} \times 100$$

110 The emulsion formed by the cell-free broth was compared with that formed by 10 % (w/v)  
111 sodium dodecyl sulphate (positive control) and distilled water (negative control), respectively.  
112  
113

### 114 2.4.2 Emulsification assay

115 Three millimetres of supernatant centrifuged at 10000 rpm for 15 min/RT was mixed with 0.5  
116 mL of kerosene. The mixture was vigorously homogenized by vortexing for 2 min, and was  
117 left undisturbed for 1 h to separate the aqueous and the hydrocarbon phases. The  
118

119 spectrophotometry absorbance of the aqueous phase was measured at 600 nm [29]. Un-  
120 inoculated broth was used as blank.

121

### 122 **2.4.3 Oil spreading test**

123

124 This method is rapid and easy to perform, and most reliable in detecting diverse  
125 biosurfactant-producing microorganisms [28, 30]. The method suggested by Morikawa *et al.*  
126 [31] was used. In brief, 20 µL of crude oil was used in making a thin layer onto a petri plates  
127 (100 mm by 15 mm) containing 50 mL of distilled water. 10 µL of cell free broth was  
128 delivered onto the oil coated surface; a clear zone on the surface indicated a positive result.  
129 The diameter of the clear zone was measured and compared with that obtained with SDS.

130

### 131 **2.4.4 Tilted glass slide test**

132

133 This is an effective modified drop collapse method [2]. A sample colony grown on nutrient  
134 agar plates for 24 h was mixed with a drop of 0.85 % NaCl at the edge of the glass slide.  
135 According to Satpute *et al.* [2], collapsing down of droplet when tilted indicated biosurfactant  
136 production.

137

## 138 **2.5 Optimization of Cultural Conditions for Biosurfactant Production**

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140 The effects of different cultural conditions (inoculum concentration, pH, temperature,  
141 nitrogen sources and agro-wastes as carbon sources) on the growth of selected yeast  
142 isolates and the ability of the strain to produce biosurfactant were determined. The inoculum  
143 for the optimization used was standardized using 0.5 McFarland's standard.

144 The optimum incubation time for growth and biosurfactant production by the selected strain  
145 was studied by varying the incubation time (24, 48, 72, 96, 120, 144, and 168 h) of the  
146 culture medium. The culture medium was inoculated with a 24 h culture broth containing a  
147 total viable cell count of  $2.38 \times 10^8$  cfu/ mL of the selected isolates and incubated at 28 °C  
148 for 168 h in a rotary shaker incubator. Biosurfactant production was measured using E-24  
149 while growth was determined using a spectrophotometer. The yeast isolates were incubated  
150 at different temperature (20, 30, 40, 50 and 60 °C) for 168 h, after which the biosurfactant  
151 production and growth of the strain were determined. The inoculum concentration with  
152 different percentage such as 2, 4, 6, 8, and 10 % (v/v) was added into the culture broth,  
153 incubated for 168 h, after which the growth of yeast isolates and the production of  
154 biosurfactant were determined. The optimum pH of 2, 4, 6, 8, and 10 for the growth of the  
155 yeast isolates and biosurfactant produced were determined after incubation for 168 h. The  
156 yeast isolates were incubated using different agro-wastes (cassava peel, soya bran,  
157 sugarcane bagasse, coconut pulp and beans bran) as carbon sources, and their growth and  
158 biosurfactant production estimated after 168h of incubation. Finally, the yeast isolates was  
159 incubated with different nitrogen sources (urea + yeast extract, yeast extract + NaNO<sub>3</sub>,  
160 NH<sub>4</sub>SO<sub>4</sub> + yeast extract, NH<sub>4</sub>NO<sub>3</sub> + yeast extract, and peptone + yeast extract for 168 h),  
161 and the growth of yeast isolates and biosurfactant production determined thereafter.

162

## 163 **2.6 Biosurfactant Production**

164

165 The optimized parameters were used in setting up the biosurfactant production media. The  
166 production was carried out in a 500 mL Erlenmeyer flask containing 300 mL of the  
167 production media with the following ingredients: KH<sub>2</sub>PO<sub>4</sub>, 0.03g; MgSO<sub>4</sub>, 0.03g; NaNO<sub>3</sub>, 0.3g;  
168 yeast extract, 0.1g, 4% of olive oil as carbon source. The conical flasks were then incubated  
169 at 28 °C under 180 rpm for 7 days

170

## 171 **2.7 Application of the Biosurfactant on Hydrocarbon Emulsification**

172 The biosurfactant produced was applied on different oils (soya oil, red oil, olive oil, coconut  
173 oil, orange oil, and castor oil) and the ability to emulsify these oils determined using E-24  
174 Index.  
175

## 176 2.8 Statistical Analysis

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178 The results were compared by one-way analysis of variance (one-way ANOVA) and multiple  
179 range tests to find the differences between the measurement means at 5 % (0.05)  
180 significance level using IBM® and SPSS® Statistics Version 20.0 (Gally and Alder, US) [32].  
181  
182

## 183 3. RESULTS AND DISCUSSION

### 184 3.1 Sample Source

185 According to Olowonibi [1]. (2017), palm wine are naturally synthesized milky alcoholic juice  
186 from the saps of *Elaeis guineensis* (oil palm), proven to be highly nutritious, which support  
187 the growth of yeast species. Figure 1, shows the picture of milky coloured palm wine sap  
188 from oil palm.  
189  
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191  
192 **Figure 1: Sap of *Elaeis guineensis***

### 193 3.2 Physicochemical Analysis of Palm Wine Sap

194  
195 The physicochemical characteristics of the palm wine are presented in Table 1. The palm  
196 wine sap had a temperature of  $17.1^{\circ}\text{C} \pm 1.27$  and a pH value of  $5.68 \pm 0.03$  at the point of  
197 collection. The pH value decreased to 3.8.6 after 6 h interval. The specific gravity,  
198 conductivity, salinity and total dissolved solids values @  $25^{\circ}\text{C}$  were  $0.827 \text{ kgm}^{-3}$ , 2.67, 1.4  
199 % and 1355, respectively.  
200  
201  
202

203 **Table1. Physicochemical properties of the sap of *Elaeis guineensis***

Parameters	Palm wine Saps of Oil palm
Temperature ( $^{\circ}\text{C}$ )	$17.1 \pm 1.27$
pH	$5.68 \pm 0.03$
pH (after 6 h interval)	$3.86 \pm 0.1$
Alcohol content (%)	$14.04 \pm 0.15$
Alcohol content (after 6 h interval, %)	$15.74 \pm 0.27$
Reducing sugar (mg/ml)	$0.51 \pm 0.03$
Reducing sugar after 6h interval (mg/ml)	$0.50 \pm 0.02$

Specific gravity (kgm <sup>-3</sup> )	0.827 ± 0.024	205
Titrateable acidity	2.3 mL of NaOH	
Conductivity (at 25 °C) (µS/cm)	2.67 ± 0.33	
Total dissolved solid (TDS) @ 25 °C (mg/L)	1355 ± 28.28	
Salinity (at 25 °C)(‰)	1.4 ± 0.56	

### 3.3. Selection and Identification of Biosurfactant-producing Yeast Isolates

Out of the five (5) yeast isolates screened, two (2) isolates were selected as biosurfactant producers based on their ability to give positive results to all the screening methods employed. From the two biosurfactant-producing yeasts, the best isolate SA2 was chosen (Table 2). The distribution of yeast isolates within the different palm wine saps of *Elaeis guineensis* are shown in (Table 3). The cultural and colonial characteristics of the best biosurfactant-producing yeast isolate are shown in Figure 2 and Table 4. Table 5 presents the biochemical characteristics of the biosurfactant-producing yeast isolate. Microscopically, using wet mount, budding yeast-like cells which are ovoid in shape were seen.

**Table 2. Screening results of the selected yeast isolates**

Isolate code	source	Emulsification index (E <sub>24</sub> )% (using kerosene)	Oil spreading (using crude oil)(mm <sup>2</sup> )	Haemolytic assay (mm)	Tilted glass slide test	Emulsification assay (OD <sub>600</sub> nm)
SA5	OP	61.3 ± 6.36	37 ± 5.66	γ	+	2.156 ± 0.06
*SA2	OP	62.5 ± 7.78	55 ± 7.07	γ	+	1.977 ± 0.023
SA7	OP	12.9 ± 2.69	7 ± 2.83	γ	-	0.244 ± 0.010
SA3	OP	-	36 ± 8.46	γ	-	0.256 ± 0.024
SA8	OP	45.2 ± 5.94	-	γ	-	2.314 ± 0.154

Legend: OP = oil palm; γ = gamma haemolysis; + = positive test; - = negative test; \*=isolate showing positive results in all the screening methods; and OD =optical density

### 3.4 Count of the Yeast Isolates within the Sap of *Elaeis guineensis*

The result obtained from the sap of *Elaeis guineensis* revealed count of 2.38x10<sup>8</sup>.

**Table 3. Colony morphology of biosurfactant-producing yeast isolate**

Isolate code	SA2
Size	medium
Shape	Ovoid
Margin	entire
Elevation	Flat
pigment	- ve
Colour	cream
Texture	Dry
Surface	Flat & smooth
Opacity	Opaque

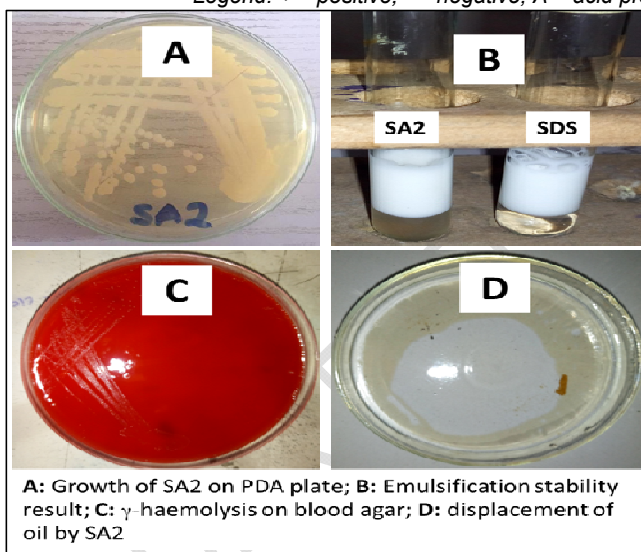
Legend - = negative

**Table 4: Biochemical identification of the biosurfactant-producing yeast isolate**

Isolate code	SA2
Carbohydrate fermentation	
Maltose	+/A
Lactose	-/-
Sucrose	+/-
Glucose	+/A
Galactose	+/A
Fructose	+/A
Raffinose	-/-
Pellicle formation	-
Growth @ 37 ° C	-
Germ tube	-
Microscopy (wet mount)	Ovoid to globose, budding yeast-like cells
Urease test	+
Probable genus	<i>Candida</i>

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Legend: + = positive; - = negative; A = acid production



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232

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Figure 2. Growth and screening characteristics of the biosurfactant-producing isolate.

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### 3.4 Optimization of Cultural Conditions for Improved Biosurfactant Production

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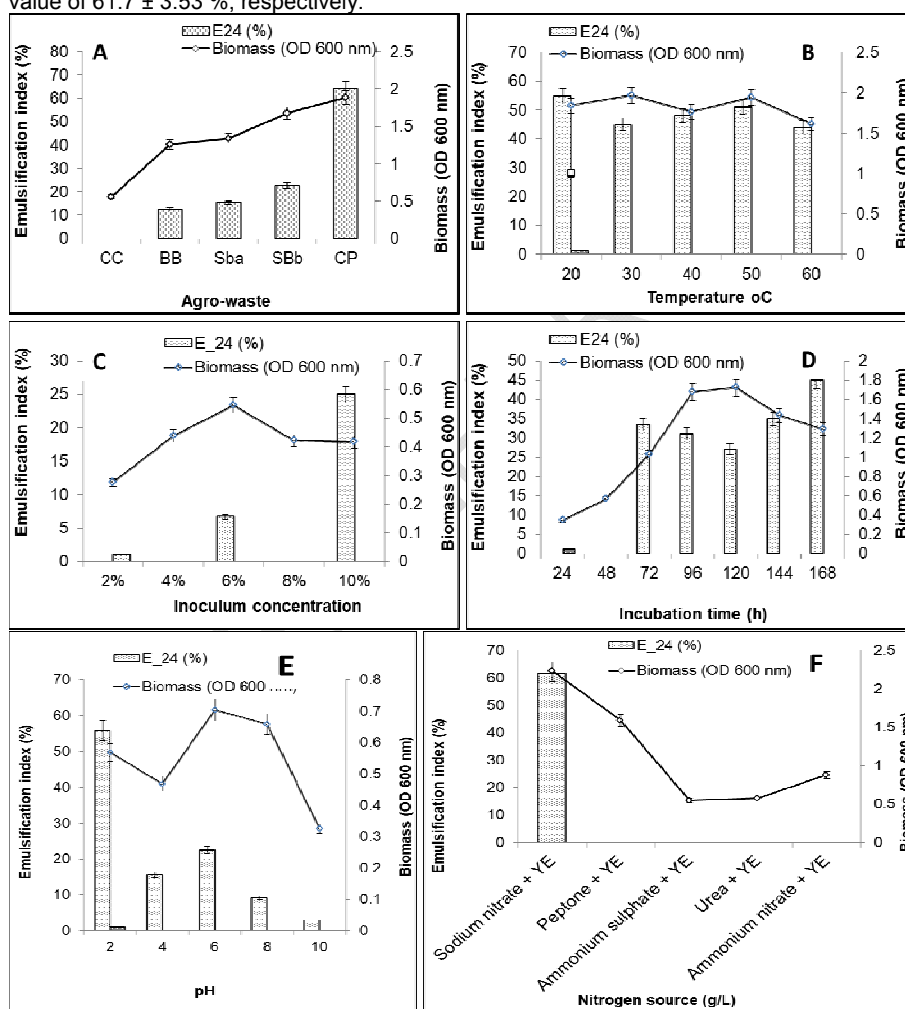
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Based on the analysed results, the optimum incubation time for growth and biosurfactant production were 120 and 168 h with the OD (optical density) reading of  $1.720 \pm 0.009$  and  $E_{24}$  value of  $45 \pm 7.07$  %, respectively. Fig 1A shows the effect of different agro-wastes as carbon sources on growth and biosurfactant production by the yeast isolate; OD reading of  $0.703 \pm 0.01$  and  $E_{24}$  value of  $55.9 \pm 2.82$  % were obtained. Cassava peel was the best carbon source for biomass formation and biosurfactant production, with  $E_{24}$  value of  $64 \pm 1.41$  % and OD reading of  $1.8840 \pm 0.01$ , respectively. The effect of different incubation temperatures on growth and biosurfactant production by the yeast isolate showed the optimum incubation temperatures to be 30 °C and 20 °C for growth and biosurfactant production, respectively (Fig 3B).

247 The effect of different percentage inoculum concentrations on growth and production of  
 248 biosurfactant showed optimum inoculum concentrations of 6 % and 10 % for growth and  
 249 biosurfactant production (Fig 3C). From the data, the optimum inoculum concentration with  
 250 OD reading of  $0.545 \pm 0.028$  and optimum biosurfactant production with E-24 value of  $25 \pm$   
 251  $1.41$  % were obtained. The result on the effect of incubation time on growth and  
 252 biosurfactant production is presented in Fig 3D. The effect of different pH values on growth  
 253 and biosurfactant production, showed the optimum pH values to be 6 and 2, respectively and  
 254 is presented in Fig 3E. Finally, Fig 3F shows that  $\text{NaNO}_3$  and yeast extract favoured growth  
 255 and biosurfactant production by the yeast isolates with OD value of  $2.286 \pm 0.01$  and  $E_{24}$   
 256 value of  $61.7 \pm 3.53$  %, respectively.



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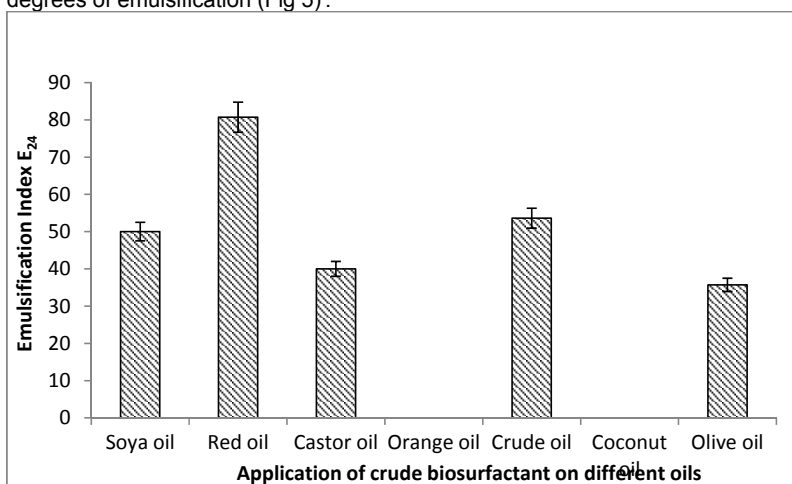
**Figure 3: Effect of different cultural conditions on biomass and biosurfactant production. Legend: A – Effect of different agro-wastes; B - Effect of different**



262 temperature; **C** - Effect of different inoculum concentration; **D** – Effect of different incubation  
263 time; **E** – Effect of different pH; **F** – Effect of different nitrogen sources.

### 264 265 3.6 Application of the Biosurfactant on Oil Emulsification

266  
267 When the crude biosurfactant produced was applied on different oils, it showed varying  
268 degrees of emulsification (Fig 5).



269  
270 **Figure 4: Application of crude biosurfactant from isolate SA2 on different oils**

## 271 272 273 4. DISCUSSION

274  
275 This study has demonstrated the ability of *Candida* sp., isolated from oil palm in producing  
276 biosurfactant. Although, there is dearth information available in literature, regarding the  
277 production of biosurfactant by yeasts isolated from oil palm. Konishi *et al.* [33] reported that  
278 biosurfactant-producing yeasts inhabit various vegetables and fruits. Iroha *et al.* [34]  
279 confirmed this by producing glycolipid biosurfactant from cashew fruit bagasse using  
280 *Pseudomonas aeruginosa*. Many researchers have reported that yeasts from different  
281 sources have the potentials of producing biosurfactants. Amaral *et al.* [35], reported that the  
282 majority of microbial biosurfactants are of bacterial origin. However, the pathogenic nature of  
283 this producing organism, has limited the application of these compounds in food and  
284 pharmaceutical industries. The study of biosurfactant by yeast has been of immense  
285 importance, because of 'generally regarded as safe' (GRAS) status that most of the species  
286 present. This GRAS status means that the yeasts do not present pathogenic or toxic  
287 considerations, thus, enhancing the application of their products for industrial usage.

288 The use of efficient screening strategy is the major key to successful discovery of new  
289 biosurfactant producers [36]. The screening methods employed in this study were  
290 haemolytic assay, oil-spreading test, emulsification index (E<sub>24</sub>), emulsification assay, and  
291 tilted glass slide test. These methods have been previously reported for the identification of  
292 biosurfactant-producing microorganisms such as bacteria and yeasts: haemolytic assay [37-  
293 38], oil spreading [31, 30, 39], emulsification index [40-42], emulsification assay [43], tilted  
294 glass slide [44-46]. The yeast isolates screened showed varying results for the different  
295 screening methods employed. The biosurfactant-producing yeasts were selected based on

296 its competence in giving positive results to all the screening methods. According to Satpute  
297 *et al.* [45], the examples of qualitative screening techniques are haemolytic assay and tilted  
298 glass slide test, whereas that of the quantitative screening techniques are emulsification  
299 index and oil spreading test. The screening techniques used in this study, employed both  
300 qualitative and quantitative methods. The use of these techniques is similar to the report of  
301 Satpute *et al.* [45], who used the combination of oil spreading, drop collapse, tilted glass  
302 slide and emulsification index to select biosurfactant producers. Satpute *et al.* [45],  
303 suggested that a single method is not suitable to select all the biosurfactant-producing  
304 microorganisms, and recommended the combination of methods. In addition, Dhimans *et al.*  
305 [47] used different screening methods, such as emulsification index, oil spreading method,  
306 oil displacement assay, surface tension measurement and drop collapse test to detect  
307 biosurfactant production. Ndibe and Usman [46], reported the confirmation of biosurfactant-  
308 production using the following classical techniques: haemolysis test, oil spreading, drop  
309 collapse, and emulsification index test.

310 To develop a process for maximum biosurfactants production is very crucial to optimize the  
311 medium and thus use suitable fermentation conditions. Incubation time has significant  
312 effects on biosurfactant production because microorganisms produce biosurfactant at  
313 different time intervals. This study investigated the effect of incubation time (24, 48, 72, 96,  
314 120, 144 and 168 h) on the ability of the test yeast isolate to grow well (biomass formation),  
315 and produce biosurfactant. The optimum biosurfactant production with  $E_{24}$  value of  $45 \pm$   
316  $7.071$  was observed after 168 h (7 days) of incubation time. However, the optimum growth  
317 ( $1.720 \pm 0.009$ ) was also observed after 120 h (5 days) of incubation time. This is similar to  
318 the result of Cavalero and Cooper, [48] and Felsa *et al.* [49], who obtained maximum  
319 biosurfactant production from *Aspergillus ustus* after 5 days of incubation. Morita *et al.*  
320 (2006) reported that  $16.3 \text{ gL}^{-1}$  of glycolipid biosurfactants was produced by *Pseudozyma*  
321 *antarctica* after seven days of incubation using glycerol as a source of carbon. *Klebsiella*  
322 *pneumoniae* strain IVN 51 isolated from hydrocarbon polluted soil had optimum growth and  
323 biosurfactant production after five and two days of incubation, respectively [18].

324 Microbial processes are temperature dependent and, they usually get affected by change in  
325 temperature. According to Saharan *et al.* [50], most of the biosurfactant productions from  
326 fungi reported so far have been performed in a temperature range of  $25$  to  $30$  °C. It was  
327 observed that the growth of *Candida bombicola* reaches a maximum at temperature of  $30$  °  
328 C, while  $27$  °C was the best temperature for the production of Sophorolipids [51]. This study  
329 is unique, in the sense that the yeast isolate was able to produce biosurfactant at an  
330 optimum temperature of  $20$  °C, with an  $E_{24}$  value of  $54.7 \pm 0.282$  %, and biomass production  
331 with OD value of  $1.965 \pm 0.007$  at optimum temperature of  $30$  °C. Khopade *et al.* [52],  
332 stated that many physiochemical factors such as pH, temperature, growth conditions and  
333 agitation have been shown to strongly influence microbial growth and metabolism. Among  
334 them pH of the production medium has proven to be the key factor for microbial growth.

335 The effect of pH (2, 4, 6, 8, 10) on the microbial growth and biosurfactant production were  
336 investigated. The results showed that maximum biosurfactant production was achieved at  
337 acidic pH of 2, with  $E_{24}$  value of  $55.9 \pm 2.85$  % and the yeast isolates grew best at pH of 6  
338 ( $0.703 \pm 0.009$ ). According to Bednarski *et al.* [53], the acidity of the production medium  
339 was the parameter studied in the synthesis of glycolipids by *Candida antarctica* and *Candida*  
340 *apicola*. When pH is maintained at 5.5, the production of glycolipids reached a maximum.  
341 The synthesis of the biosurfactant decreased without the pH control indicating the  
342 importance of maintaining it throughout the fermentation process. The pH of 6, favours the  
343 growth (biomass formation) and production of biosurfactant by *Pseudomonas aeruginosa*  
344 2297, as reported by Kumar *et al.* [54]. *Candida lipolytica* at pH of 5.0 and *Candida batistae*,  
345 at pH of 6.0 produced maximum biosurfactant [55-56]. Amaral *et al.* [57], confirmed the  
346 production of Yansan, with a stable pH between 3 and 9 from *Yarrow lipolytica*.

347 It is estimated that substrate (carbon source) account for 10 to 30% of the total production  
348 costs of biosurfactant [58]. Thus, to reduce the cost involved in biosurfactant production, it is

349 desirable to use low-cost raw materials like agro-industrial wastes. The effects of agro-  
350 industrial wastes (cassava peel, sugarcane bagasse, soya bran, coconut chaff, and beans  
351 bran) as carbon sources on biosurfactant production and growth of the yeast isolate was  
352 also investigated in this study. The result shows that cassava peel favoured the growth and  
353 production of biosurfactant with OD value of  $1.884 \pm 0.011$  and  $E_{24}$  value of  $64 \pm 1.41 \%$ ,  
354 respectively. According to Nitschke *et al.* [59], microorganisms for biosurfactant productions  
355 can be selected using agro-industrial wastes such as cassava flour waste water. Nigeria has  
356 cassava in abundance, and most of the wastes are discarded. Therefore, finding industrial  
357 use for these wastes will have positive economic benefits.

358 Several nitrate salts such as sodium nitrate, ammonium nitrate, potassium nitrate was used  
359 as nitrogen sources for biosurfactant production. A combination of sodium nitrate and yeast  
360 extract were most influential nitrogen source. The result obtained revealed that these  
361 nitrogen sources favoured the growth of the test isolate with OD value of  $1.884 \pm 0.01$ , and  
362  $E_{24}$  value of  $64 \pm 1.41 \%$  for biosurfactant production. The result is similar to the report of  
363 Abbasi *et al.* [60] that  $\text{NaNO}_3$  (39.3 g) and yeast extract (3.93g) enhanced the optimum  
364 conditions for biosurfactant production by *Pseudomonas aeruginosa* MA01. Silva *et al.* [61],  
365 showed that *P. aeruginosa* 44T1 fail to give good biosurfactant yield with ammonium salts  
366 but instead gave good yield when  $\text{NaNO}_3$ . However, in another study, higher yield of  
367 biosurfactant by *Candida glabrata* UCP 1002 was observed with ammonium nitrate and  
368 yeast extract [62]. The effect of different inoculum concentration on the growth of the test  
369 isolate and for biosurfactant production was carried out. The result shows that inoculum  
370 concentration of 6 % (v/v) ( $E_{24}$  value of  $25 \pm 1.41 \%$ ) and 10 % (v/v) (OD value of  $0.545 \pm$   
371  $0.007$ ), enhanced the biosurfactant production and biomass formation by the test yeast  
372 isolate, respectively.

#### 373 374 **4. CONCLUSION**

375 The results obtained from this study demonstrated the capacity of a yeast isolate from the  
376 sap of *Elaeis guineensis* to produce biosurfactant. The yeast isolate was identified as  
377 *Candida* sp. Production of biosurfactant from ecological safe source has an added  
378 advantage of excluding any risk of toxicity and pathogenic reactions to the environment. The  
379 ability to produce biosurfactant was dependent on the incubation media conditions.  
380 Moreover, the biosurfactant was able to emulsify at varying degrees different hydrocarbons.  
381 Therefore, biosurfactant from *Candida* sp. can be scaled up for industrial production.

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#### 385 386 387 **COMPETING INTERESTS**

388 The authors declare that they have no competing interest.

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