

Isolation, Identification and Production of Biobutanol by Different *Clostridium* species Isolated from Soil Using Waste Paper and Sugar Cane Molasses.

ABSTRACT

Butanol as fuel component has some advantages compared to ethanol. The Study was aimed at isolation of different types of *Clostridium* species from the soil and screening for their capacity for production of bio-butanol. Soil samples were collected from three different areas in keffi. *Clostridium* species were isolated using Reinforce Clostridial media and identified using standard microbiological methods, mutation was induced in the *Clostridium* isolates and 16s rRNA identification techniques was used to identify the isolates. Quantitative screening of the *Clostridium* species isolates capacity in bio-butanol was carried out using sugar cane molasses and waste paper substrate. The result using 16s rRNA identification the *clostridium* isolated from old barracks was *C. acetobutylicum* the one isolated from Amgwan kwara was *C. perfringenes* and *C. difficile* was isolated from Amgwan Jaba. Screening for their bio-butanol producing capacity, it was observed that *C. acetobutylicum* produces the highest before exposing to UV light to induce mutation with 9.61 ± 0.94 mg/l using sugar cane molasses and 4.89 ± 0.19 mg/l using waste paper followed by *C. perfringenes* 6.24 ± 0.61 mg/l using molasses and 3.21 ± 0.11 mg/l using waste paper. After exposing to UV light to induce mutation *C. acetobutylicum* produced 18.03 ± 0.17 mg/l using molasses and 8.63 ± 0.06 mg/l using waste paper substrate follow by *C. perfringenes* 14.19 ± 0.11 mg/l and 5.99 ± 0.31 mg/l. From this study it was observed that *C. acetobutylicum* has the capacity to produce bio-butanol and also inducing mutation will enhance the capacity of *Clostridium* species in the production of bio-butanol.

Key: *Clostridium*, Bio-butanol, Molasses, Substrate

1. Introduction

The rate of energy demands combined with little resources of petroleum and environmental condition have raised a new interest in production of renewable energy from biomass, such as agricultural crops residuals, lignocellulosic waste and from industrial waste. Apart from ethanol, butanol is superior in energy content, has lower volatility and is also less corrosive to existing infrastructure [1]. There are three main areas remaining to be sorted out if bio-butanol is to become a major counterpart in the bioenergy market. This includes optimizing

34 feedstock utilization, reaching theoretical maximum yields of butanol and minimizing energy
35 consumption during separation and purification [2].

36 One important driver for a bio-based economy is the exploitation of the bio-refinery concept
37 [3] where maximum value can be derived from the biomass through the generation of
38 multiple products and the effective use of process integration. Chemical pulp mills are
39 current examples of bio-refineries that can convert lignocellulosic biomass into energy, pulp,
40 cellulose derivatives, tall oil, etc. One strategy for mills to counteract competition from
41 tropical countries using fast-growing raw materials is to further expand the product portfolio
42 into additional value-added products. Cellulose is the primary fraction utilized for chemical
43 pulp production, while lignin, hemicellulose and extractives could be considered as by-
44 products in the process. Currently lignin and a portion of the hemicellulose, and the pulping
45 inorganics end up in the black liquor which is concentrated and burnt in a recovery boiler to
46 recover the chemicals and to supply energy for a large fraction of the mills process steam
47 requirements. Considering the fact that lignin has double the heating value as
48 polysaccharides, the hemicellulose fraction solubilized during alkaline pulping represents an
49 underutilized resource in many mills.

50 One option to enhance the value of the hemicellulose is to recover this fraction prior pulping
51 by using hot-water extraction [4, 5]. However, despite high yields of xylene, the removal of
52 hemicellulose from birch wood chips prior Kraft cooking has a negative impact of some pulp
53 properties affecting the quality of the paper. If a decrease in pulp strength properties cannot
54 be accepted, another option is to recover the xylene fraction from the black liquor.
55 Acidification of alkaline pulping liquors precipitates lignin as well as any hemicellulose
56 present in the liquor [6]. Technologies for recovery of lignin from Kraft liquors via CO₂
57 acidification has been the subject of pilot and demonstration-scale processes [7,8] with
58 proposed process of generating fuels, materials, and chemicals from the lignin as well as the
59 opportunity for “de-bottlenecking” capacity-limited recovery-boilers. Hemicelluloses are
60 typically degraded to hydroxyl- acids during Kraft pulping [9], although oligomeric xylene
61 from hardwoods may be more resistant to alkaline pulping than the glucomannans which are
62 the predominant softwood hemicelluloses due to the protection of glucuronic acid
63 substitutions against end-wise alkaline degradation, and as such, these xylenes may offer an
64 opportunity for recovery and utilization as a feedstock for bioconversion. In the present
65 study, the aim was to produce bio-butanol using waste paper by two clostridium species
66 isolated in soil of Keffi.

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2.0 MATERIAL AND METHODS

2.1 Study Area

The study was carried out in Keffi, Nasarawa State, Nigeria. Keffi is approximately 68km away from the Federal Capital Territory (FCT), Abuja and 128km away from Lafia, the capital town of Nasarawa State. Keffi is located at longitude 8°5'E along the Greenwich Meridian and at the equator and situated on longitude 850m above sea level [10].

2.2 Sample Collection

Three (3) soil samples were randomly collected (at the topsoil 5mm) from three different locations of Keffi metropolis such as Angwan Jaba, Old Barracks and Angwan kwara using a clean spoons and stored using disposable screw bottle and transported immediately to the Microbiology Laboratory, Nasarawa State University, Keffi for isolation.

2.2.1 Isolation of *Clostridium* species

The isolation of *Clostridium* species from three different areas in Keffi was carried out using spread plate method. One (1.0g) of the soil sample was suspended in 9.0ml of sterile distilled water containing tween 80 and 7 folds dilutions were made and 0.1ml of the aliquot was spread on petri dishes of freshly prepared Reinforce Clostridial media containing yeast extract 3.0; lab-lemco powder 10.0; peptone 10.0; soluble starch 1.0; glucose 5.0; cysteine hydrochloride 0.5; sodium chloride 5.0, sodium acetate 3.0 and agar agar 1.0. Incubated in anaerobic jar at 37°C for 48 hours.

2.1.3. Identification of *Clostridium* species

Cultural and morphological identifications of *Clostridium* species were carried out in accordance with Bergey's Manual of Determinative Bacteriology [11] such as swarming test, gram staining test, endospore Staining egg yolk agar test, nagler test, lipase test Indole test, catalase test and urease test.

2.1.4 Hank's Buffer/UV light treatment of *clostridium* isolates

Induction of mutation in *Clostridium* isolates was carried out by the modification of a method described by [12]. *Clostridia* isolates were grown on Reinforce Clostridial medium maintained in Hank's buffer (137mM NaCl, 5.4mM KCl, 4.4mM KHIO₄, 0.33mM Na₂HPO₄, 1.3mM CaCl₂, 0.81mM MgSO₄, 4.2mM NaHCO₃, 1g/L glucose, pH 7.4) and were subsequently exposed for 1 hour to ultra violet light rays from 2 white fluorescent tubes (TFC FL-20 SD/18W Day Light). Isolates were exposed at a distance of 15cm. isolates were added to a mixture of 75% methanol and 25% acetic acid for 1 hour. The isolates were subsequently incubated in 70% ethanol containing 0.07N NaOH overnight at 4 °C. Isolates were analyzed using 16S rRNA gene sequencing for identification.

2.1.5 Extraction of bacterial DNA for 16S rRNA gene sequencing

Eighty (80 µl) of NaOH (0.05M) was added to 20 µl of *Clostridium* isolates, were suspended in distilled water and the mixture was incubated at temperature of 60°C for 45 minutes, 6 µl of Tris/HCl (pH 7.0) were addition to achieve a final pH of 8.0. The mixture was diluted by ×100 and 5 µl of the diluted extract was used for the polymerase chain reaction (PCR).

2.1.6 PCR, gel electrophoresis, and 16S rRNA gene sequencing

Treated distilled water DNase I and PCR master mix (containing dNTPs, PCR buffer, and Taq polymerase) were used in all PCR reactions by adding 1 U of DNase I (Pharmacia, Uppsala, Sweden) to 40 µl of distilled water or PCR master mix, the mixture was incubated at 25°C for 15 minutes, and later at 95°C for 10 minutes to inactivate the DNase I. The clostridial isolates, DNA extracts and control was amplified with 0.5µM primers (27F, 5'-TGGCGAACGGGTGAGTAA-3' and 1525R, 5'-AGGCCCGGGAACGTATTCAC-3'; Inqaba Biotechnical Industries (Pty) UK). The PCR mixture (50 µl) contained *Clostridium* isolates DNA, PCR buffer (10mM Tris/HCl (pH 8.3), 50mM KCl, 2mM MgCl₂, and 0.01% gelatin), 200 µM of each dNTP, and 1.0 U Taq polymerase (Boehringer Mannheim, Mannheim, Germany). The mixtures were amplified by 40 cycles of 94°C for one minute, 55°C for one minute, and 72°C for two minutes, with a final extension at 72°C for 10 minutes in an automated 0.5 ml GeneAmp PCR system 9700 (Applied Biosystems, Foster City, California, USA). DNase I treated distilled water was used as the negative control. A 10 µl aliquot of each amplified product was electrophoresed in 1.0% (wt/vol) agarose gel, with a molecular size marker (λ DNA Avall digest; Boehringer Mannheim) in parallel. Electrophoresis in Tris/borate/EDTA buffer was performed at 100 V for 1.5 hours. The gel was stained with ethidium bromide (0.5 µg/ml) for 15 minutes, rinsed, and photographed under ultraviolet light illumination.

The PCR products were gel purified using the techniques developed by the National Centre for Bioinformatics, Islamabad, Pakistan. Both strands of the PCR products were sequenced twice with an ABI 377 automated sequencer according to the manufacturers' instructions (Inqaba Biotechnical Industries (Pty) Ltd UK) using the PCR primers (27F and 1525R).

2.1.7 The pre-treatment of waste papers

The pre-treatment of waste papers was carried out as described by [13]. The waste papers were thoroughly washed in water to remove dust. Waste paper substrates were pre-treated with 3% NaOH for 1 hour. Then, they were washed and neutralized with 0.1N HCl at pH 5. The pre-treated waste papers were dried at 45 °C in an oven after neutralization.

2.1.7.1 Acid hydrolysis

Acid hydrolysis was done to break down cellulose into glucose units. It was carried out by soaking 10g of paper in different amounts of sulfuric acid 5%: 100, 200 and 300 ml of water for 2 hours. The mixture was autoclave at 121°C. After hydrolysis, the samples were filtered and centrifuged to obtain the hydrolysate product. The pH of the obtained product was adjusted to 5.0 by adding sodium hydroxide solution 5M

2.1.8 Preparation of Molasses

The preparation of molasses from sugarcane was carried out as described by [14, 15]. The sugarcane was stripped off leaves. Its juice was extracted by cutting, crushing and mashing. The juice was boiled over a Bunsen burner to concentrate it, promoting sugar crystallization. The product of this first boiling called first syrup was used as a component of the fermentation medium.

2.1.9 Quantitative Analysis of Reducing Sugar Present in Fermentation medium

The quantitative analysis was carried out using 3, 5–dinitrosalicylic acid. The concentration of the sugar present in the samples was determine by adding 1cm³ of 3, 5-dinitrosalicylic acid to 1cm³ of each of the samples and boiled for 5 minutes and 10cm³distille water was added. The absorbance of each of sample was determined at 540nm using JENWAY 6400 spectrophotometer. Thus, the concentration values were extrapolated from the glucose standard curve

2.1.10 Preparation of inoculum

The seed culture was carried out as described by [16]. The cultivated organism two colonies was inoculated into 10 ml of Reinforce Clostridial broth and incubated in an anaerobic jar at 37 °C for 6 hours.

3.0 Production of Bio-butanol

3.1 Fermentation

The batch fermentation was carried out as earlier described by [17] with modification. The 100 ml from the starter culture was inoculated in 900 ml of fermentation media (enzymatically hydrolyzed waste papers and molasses) containing 25g of sugar. The fermentation media were incubated under strict anaerobic and sterile conditions at 35°C for 96 hours.

3.1.2 Extraction and purification of bio-butanol from fermentation broth

Extraction and purification of solvents from the fermentation broth were carried out using a modification of the method described by [13]. In this method, *Clostridium* species were used in the fermenter. The extracting solvent was an alkylate which was substantially free from impurities and had a boiling range of 118°C. The apparatus for carrying out the process included a standard fermenter receiving fermentation broth including a carbon source and microorganisms at a feed stream which are agitated and maintained at a desired

fermentation temperature. The butanol-containing broth was drained through a first stream discharging into the upper end of an extraction column where it flowed downwardly through the extractor. A second stream released a light alkylate into the bottom of the column for upward countercurrent flow relative to the downward flow of the broth. The light alkylate was fed from an alkylate source stream, separated in a splitter with the light end boiling in the range of 1 °C. The butanol containing alkylate extract was discharged through a stream at the top of the extractor to flow through a dryer (a desiccator), in order to purify it and remove any excess water content. The extraction was carried out at a steady state with a solvent feed ratio ranging from 5–20 ml. The solvent was removed through the column at a rate of 200 – 450 ml per minute.

3.1.3 Separation and quantification of bio-butanol

The separation and quantification of bio-butanol produced by *Clostridium* species were carried out by Gas Chromatography and Mass Spectrometry (GC & MS) as described by [13]. One microliter (1.0 µL) of acidified sample was injected into “Schimadzu GC-14, Gas Chromatograph” equipped with flame-ionization detector.

The column used for the separation of solvent was PEG (2.1m x3.0mm). The temperature programming of the column oven was 60 °C/min. One hundred and twenty degrees centigrade (120 °C), Nitrogen gas (30 mL/minutes) was used as carrier gas. The temperatures of injector and detector were 150 °C and 200 °C respectively. The Peaks were recorded on “SHIMADZU C-R-4_A, Chromatograph”, and were identified by comparison of the retention times with that of standard mixture. The experiment was carried out in duplicates and the means ± standard deviations of the yield of butanol were recorded. The standard calibration curve presented was used in calculating the concentration of bio-butanol produced by the various *Clostridium* isolates.

3. Results

The species of *Clostridium* that were isolated from soil in Keffi were identified using Cultural, Morphology and different biochemical test as given in table 1. The three species of *Clostridium* isolated from different location in Keffi was able to produce bio-butanol after screening for bio-butanol production at pH 5.5, temperature of 37°C for 96hours were *Clostridium* specie isolated from Old barracks produced the highest butanol 9.61± 0.94 mg/l using sugar cane molasses and sugar utilization of 9.61±0.71 g/l and 4.8±0. 19mg/l using waste paper substrate followed by *Clostridium* species isolated from Angwan Jaba 6.21± 0.61 mg/l, 9.05±1.11 g/l of sugar utilization using molasses and 3.21 ± 0.11 mg/l using waste paper substrate *Clostridium* isolated from Angwan kwara 5.48±0.99 mg/l and 10.11± 0.99 g/l of sugar utilization sugar cane molasses and 1.97 ± 0.33 mg/l using waste paper substrate.

207 Mutation was induced in the *Clostridium* isolates and were identify using 16s rRNA as shown
 208 in figure 1, 2 and 3 respectively. bio-butanol produced pH 5.5, temperature of 37°C for
 209 72hours after the induce of mutation as given in table 2 using waste paper as fermentation
 210 substrate shows that *Clostridium* species isolated from old barracks was *C. acetobutylicum*
 211 also produce the highest butanol after mutation with 8.63±0.06 mg/l sugar utilization of
 212 6.21±0.11 g/l followed by *C. perfringenes* isolated from Amgwan kwara 5.99 ± 0.31 mg/l,
 213 sugar utilization of 4.89±0.19 g/l and *C. difficile* isolated from Angwan Jaba 3.01± 0.54 mg/l
 214 and sugar utilization of 3.97±0.33 g/l respectively. Table 3 shows bio-butanol produced using
 215 sugar cane molasses by the *Clostridium* isolates were *C. acetobutylicum* produced highest
 216 16.63±0.06 mg/l and sugar utilization of 16.98±0.51 mg/l followed by *C. perfringenes* 14.19 ±
 217 0.11mg/l sugar utilization of 12.99±0.06 mg/l and the lowest was produced by *C. difficile*
 218 10.01± 0.01mg/l and 9.89±0.19 mg/l utilization of sugar respectively.

219 Figure 1, 2 and 3 shows the phylogenetic tree of the different species of the *Clostridium*
 220 isolated from different location.

221 **Table1. Cultural, Morphology and Biochemical Characteristics Test**

Cultural	Morphology characteristics							Inference
	Gram stain	Endo	Egg yolk test					
		Spore	lipase	NR	LE	cat		
			IND					
			UV	Sg/UV				
Glossy, grey	-	+	-	+	+	-	+	<i>C. difficile</i>
Large smooth	-	+	Nil	Nil	-	-	-	<i>C. perfringenes</i>
Raised	-	+	+		-	-	-	<i>C.</i> <i>acetobutylicum</i>

222 Key: UV fluorescence , NR- No reaction, SW- swarming growth, LE- Lecithinnase. Cat-
 223 catalase, IND- indole.

224 **Table2: Bio-butanol produced by different *Clostridium* isolates before mutation at**
 225 **pH5.5, 37°C after 96hours**

Isolates	Initial sugar concretion (g/l)	Sugar utilization (g/l)	Bio-butanol produced using waste paper (mg/l)	Bio-butanol produced using sugar cane molasses (mg/l)
<i>Clostridium</i> (AJ)	20	9.05±1.11	3.21±0.11	6.21±0.61
<i>Clostridium</i> (AK)	20	10.11± 0.04	1.97±0.33	5.48±0.99
<i>Clostridium</i> (OB)	20	9.61±0.71	4.89±0.19	9.61± 0.94

226 Key: AJ= Angwan Jaba, AK= Amgwan kwara, OB= Old barracks

227 **Table3: Bio-butanol produced by different *Clostridium* isolates after mutation at**
 228 **pH5.5, 37°C after 96 hours**

Isolates	Initial sugar (g/l)	Sugar utilization (g/l)	Bio-butanol produced using waste paper (mg/l)
<i>C. acetobutylicum</i>	20.1±0.50	6.21±0.11	8.63±0.06
<i>C. difficile</i>	20.1±0.50	3.97±0.33	3.01± 0.54
<i>C. perfringenes</i>	20.1±0.50	4.89±0.19	5.99 ± 0.31

229 **Table 4: Bio-butanol produced by different clostridium isolates after mutation at**
 230 **pH5.5, 37°C after 96 hours**

Isolates	Initial sugar (g/l)	Sugar utilization (g/l)	Bio-butanol produced using sugar cane molasses (mg/l)
<i>C. acetobutylicum</i>	25.01±0.50	16.98±0.51	18.03±0.17
<i>C. difficile</i>	25.01±0.50	9.57±0.93	10.01± 0.01
<i>C. perfringenes</i>	25.01±0.50	12.99±0.06	14.19 ± 0.11

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232

233 **Figure 1: Phylogenetic Tree of the Molecular Characterization of *Clostridium***
 234 ***acetobutylicum* isolated from old Barracks Keffi.**



Figure 2: Phylogenetic Tree of the Molecular Characterization of *Clostridium difficile* isolated from Angwan Jaba in Keffi.



Figure 3: Phylogenetic Tree of the Molecular Characterization of *Clostridium perfringens* isolated from Angwan kwara in Keffi

4. Discussion

Butanol as fuel or blending component has some advantages compared to ethanol. For example a lower vapour pressure and higher energy density. The production of butanol in a microbial fermentation was first reported by Pasteur in 1861. Three different species of clostridium were isolated and identified using 16s rRNA from different location from soil in Keffi, were *C. perfringens*, *C. difficile* and *C. acetobutylicum*. This however, is not in contrary with studies earlier reported by [18] that *Clostridium* species are one of most common bacteria isolated from the soil environment. Bio-butanol production by different species of *Clostridium* isolated in this study is in agreement with other studies earlier described by [19, 20, 21] that *Clostridium* species are some bacteria that has the ability in production of bio-butanol. In the study it was observed that *Clostridium* species isolated from old barracks has the highest accumulation of bio-butanol at 37°C, pH5.5 after incubation of 96 hours. Fermentation parameters and type of fermentation substrate play an important role in production of bio-butanol. It was observed that were was high production of bio-butanol

using sugar cane molasses and glucose which contain noncomplex sugar than using waste paper which is made up of lignin which need to be detoxified to enhance the used of waste paper in the production of bio-butanol to remove the inhibitor that will affect the growth of the *Clostridium* isolates as earlier reported by [22]. In this study it was observed that the sugar cane molasses has higher yield of bio-butanol that when used in a media containing the waste paper, also base on the isolates that yield has the highest accumulation of the bio-butanol it observed that *C. acetobutylicum* has the accumulation of 8.63 ± 0.06 mg/l in use of waste paper as substrate and 18.03 ± 0.17 mg/l on sugar cane molasses as substrate which is agreement with work earlier reported by [18]. Also it was observed that there was increase in the production of biobutanol after the isolate was expose to UV light for mutation which suggested that for economical bio-butanol production using *Clostridium* species, Induction of mutation will enhance the production of bio-butanol industrially

5. Conclusion

Clostridium species isolated from soil environment showed high capacity of production of bio-butanol. It was observed that the use of both molasses and waste paper will be a good source of carbon or media in the production of bio-butanol. Inducing mutation in *Clostridium* species will help to overcome the challenges of using clostridium in industrial scale production of bio-butanol.

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