

**FLOCCULATING PROPERTIES OF A BIOFLOCCULANT PURIFIED FROM
BACILLUS SUBTILIS ISOLATED FROM THE STREAM SEDIMENTS OF
ONYEARUGBULEM MARKET, AKURE, NIGERIA.**

Abstract

Soil samples (sediments of stream, its bank and abattoir soil) were collected from Onyearugbulem market abattoir, Akure, Ondo State, Nigeria. Bacteria were isolated from the above soil samples by dilution and pour plate methods. Screening for best bioflocculating bacteria was also performed. Effects of metal ions (such as Mg^{2+} , Ca^{2+} and Al^{3+}), temperature and pH were also determined on flocculating activities of bioflocculant. Six bacterial isolates producing flocculating substances were isolated and the isolate with the best flocculating property was selected. The identified bioflocculant producing bacteria are *Bacillus anthracis*, *B. subtilis*, *B. thuringiensis*, *B. cereus*, *Streptomyces griseus* and *S. somaliensis*. The best bioflocculant producing bacterium was *Bacillus subtilis* and the flocculating activity of its bioflocculant was stimulated in the presence of Mg^{2+} , Ca^{2+} and Al^{3+} . This bioflocculant was thermostable and retained more than 80% of its flocculating activity after being heated at 100°C for 25 minutes. It had the highest flocculating activity of 85% at pH 6 with optimum bioflocculant dosage of 0.8 mL. Bioflocculant exhibited good bioflocculating properties.

Key Words

Bioflocculant, Onyearugbulem stream, pour plate, metal ion, flocculating property, thermostable.

22 Introduction

23 Flocculation is a form of chemical reaction that involves the addition of clarifying agents
24 such as Iron (II) sulphate, Aluminium sulphate, and Iron (II) chloride in water treatment which
25 results in the formation of colloids (Vasantharaj *et al.*, 2013). It can also be described as a
26 physical and chemical process used for the removal of the visible sediments and material from
27 water which makes it a colloidal solution. Flocculation can be carried out through agitation or by
28 the addition of flocculating agents (Moghimpour *et al.*, 2014). Bioflocculants are
29 microorganism-produced special natural inorganic macromolecule substances that can flocculate
30 suspended solids, cells, colloidal solids etc (Zaki *et al.*, 2011). Several biopolymer flocculating
31 microorganisms have been screened and isolated from activated sludge, waste water, and soil
32 (Zaki *et al.*, 2011). Species of microorganisms that have bioflocculant producing characteristics
33 include bacteria (such as *Bacteroidites*, *Bacillus* sp., *Bacillus muscilaginosus*, *Bacillus subtilis*)
34 fungi, actinomyces and algae (*Chlamydomonas reinhardtii*, *Chlorella minutissima* *Arcobacter*,
35 *Cellulosimicrobium cellulans*, *Aeromobacter xyloisidans*,) (Okaiyeto *et al.*, 2013). Bioflocculants
36 stand out among others as they have the advantage of innocuousness, biocompatibility,
37 biodegradability and environmental friendliness, unlike organic and inorganic flocculants which
38 are toxic and whose degradation intermediates are difficult to remove from the environment
39 (Okaiyeto *et al.*, 2015). Besides, organic flocculants such as polyacrylamide and polyethylene
40 imine derivatives have been involved in adverse human health effects. (Nwodo *et al.*, 2014).
41 Conversely, the enormous advantages associated with bioflocculants motivate its consideration
42 as an alternative, hence the vast interest in the scientific and industrial community worldwide
43 (Nwodo *et al.*, 2012).

44

45 **Materials and Methods**

46 **Description of study area**

47 Onyearugbulem abattoir was selected as the study area because of its location in the large
48 expanse of built up area comprising of low medium and high income earners with residential
49 buildings in the north by office complexes and west and east by private schools and shops. The
50 abattoir is about 50 meters off the express (Ilesha-Owo) and covers about 1000m² land mass.

51 **Period of study**

52 This research was carried out from October 2016 to September 2017. The first set of soil
53 and water samples were collected in October 2016 which were immediately analysed. Samples
54 were thereafter collected as required based on failure in experimental set up.

55 **Sample Collection**

56 Soil sample from the Onyearugbulem abattoir slaughtering site and stream bank, was collected
57 with the aid of an auger. The soil was dug in a V-shape to a depth of about 0-5cm, thin slices of
58 the soil was removed from the sides and transferred into a clean container. With the aid of an
59 auger, composite sediment was taken upstream where it was maximally free from contaminants
60 (Nontembiso *et al*, 2011). Well water was collected with the aid of a sterilized fetching bucket
61 into a clean 50 liter container. Stream water was collected at three different points (upstream,
62 mid-stream and downstream) together with a clean bowl into a clean 50 L container. Abattoir
63 waste effluent was collected directly from abattoir drainage into a clean 50 L container.

64 **Physicochemical Properties of the Soil and Water Samples**

65 The physicochemical properties (pH, exchangeable magnesium and calcium component, particle
66 size analysis, nitrogen, phosphorus, carbon, sodium, potassium and organic content
67 determination) of the soil samples used for the isolation of biofloculant producing bacteria were
68 determined according to Association of Analytical Chemists (A. O. A. C, 1990). Collected water
69 samples were subjected to chemical analysis such as Dissolved oxygen, pH, Electrical
70 conductivity, Total Dissolved Solids, Chloride content, Nitrate, Phosphate, Magnesium content
71 and total hardness before their respective treatment.

72 **Determination of the pH of Soil Samples**

73 Twenty grams of each soil sample was weighed and put in a 100 ml beaker. Twenty milliliters of
74 distilled water was added to the sample. The suspension was left for 2 minutes, with occasional
75 stirring using glass a rod by hand in order to enable it reach equilibrium. The pH of the
76 suspension was determined using a pH meter (AOAC, 1990).

77 **Determination of exchangeable magnesium and calcium component of soil sample using** 78 **EDTA titration method**

79 One litre standard flask was filled to the half mark with distilled water and 60 ml
80 concentrated acetic acid and 70 ml ammonia solutions were added. The mixture was shaken
81 together and was made up to 1 liter mark with distilled water. This mixture was left to settle
82 overnight. A 10g quantity of soil sample was weighed into beaker and 100 ml ammonium acetate
83 was added and the mixture was stirred and allowed to stand for 1 hour. The mixture was then
84 filtered using whatman filter paper (pore size 2.5 μ m). The filtrate was collected and bottled. A
85 50 ml burette, which previously had been washed and dried was filled to the level mark with
86 0.01M EDTA solutions. The filled burette was placed vertically on a retort stand. 10 ml of the

87 prepared soil sample solution was then pipette and transferred into 250 ml conical flask. Five
88 drops of 2% KCN was then added. A 7 ml volume of concentrated ammonium solution was
89 added. Three drops of Eriochrome Black T indicator was then added and a wine red colour was
90 obtained. The titration was repeated and the mean values were calculated. To obtain Ca^{2+} alone,
91 10 sml of the sample was pipette into a 250 ml conical flask. Five drops of 2% KCN, 5 drops of
92 5% hydroxyl ammonium chloride and 5 ml 20% KOH solution were added respectively. A pinch
93 of calcium indicator was added to the mixture. The resulting solution was then titrated with
94 0.01M EDTA solution. The titration was repeated and the mean value was found. To determine
95 Mg^{2+} ions in the solution, the value obtained for Ca^{2+} was subtracted from the total value
96 obtained for $\text{Ca}^{2+}/\text{Mg}^{2+}$ (AOAC, 1990).

97 Calculation:

98 % Calcium (Ca) = % Magnesium (Mg)

$$\frac{\text{Ca}^{2+}}{\text{Mg}^{2+}} - \text{Ca}^{2+} = \text{Mg}^{2+}$$

99 T = Titre value obtained from EDTA titration

100 M = Molarity of acid used

101 V_1 = Total volume of initial extracting solution

102 V_2 = Volume of extracted solution used

103 W = Weight of soil sample

104 40 = Atomic weight of calcium

105 24 = Atomic weight of magnesium

106 **Analysis of Particle size of Soil Samples**

107 This was done using the hydrometer method. Those soil particles (Coarse fragment) that did not
108 pass through the 2mm sieve were weighed and reported as a percentage of the whole weight. 50g
109 of the fine earth fraction (greater than 2mm fraction) were put in a beaker and 100ml of 5%
110 dispersing agent, sodium hexametaphosphate, added. The suspension was mixed with a stirring
111 rod and allowed to soak for 30 minutes before transferring it to the bottle cup. The bottle cup
112 was attached to the stirrer and stirred for three minutes to ensure breakage of soil aggregates. The
113 suspension was poured into cylinder and made up to mark, stirred and both hydrometer and
114 thermometer were inserted at specified time intervals to take readings (40 seconds for silt and
115 clay reading and end of two hour for clay).

116 The formula below was used in their calculation

117 a. For an increase in temperature above 20°C

$$\frac{X + 9 (Y \times 0.36)}{W}$$

118 b. For a decrease in temperature

$$\frac{X + 9 (Y \times 0.36)}{W}$$

120 Where X = hydrometer reading at specified time

121 Y= differences between hydrometer calibrated temperature and the
122 temperature of the solution at specified time.

123 W = weight of the fine earth fraction used.

124 Percent silt was derived by subtracting the calculated percent clay from that of silt and clay
125 subtracting percent silt and clay from 100 gives the percent sand. From the texture triangle
126 diagram, percent clay, silt and sand were used to draw lines parallel to the bottom, left side and
127 right side of the triangle respectively. The area in which these lines intersect gives the class name
128 or texture of the soil. Where the intersecting lines fall on the line between two textual names.
129 The name of the finer fraction was used. The suspension was decanted and the process repeated
130 until the supernatant became clear. The sand fraction was transferred quantitatively into a small
131 beaker and dried in oven at a temperature of 105°C; it was cooled in a dessicator after which the
132 sand fraction was weighed. It was passed through 0.2mm- 0.02mm sieve and the coarse fraction
133 remaining on the sieve was also weighted. The total sand weight minus the coarse and fraction
134 weight gives the fine sand fraction weight. They were expressed in percentage.

135 Percent clay: (% clay) =

$$\frac{h_x \times 100}{w}$$

136 Where h_x is the hygrometer reading at 6 hrs 52 mins and w is the weight of sample.

137 Percent silt: (% silt) =

$$\frac{h_y \times 100}{w} - C(\%)$$

138 Where C (%) is percentage Clay, h_y is the hygrometer reading at 40 sec and w is the weight of
139 sample.

140 Percent sand: (% sand) = $100\% - S(\%) - C(\%)$

141 Where S (%) is percent silt and C (%) is percent Clay.

142

143 **Determination of available phosphorus of soil samples**

144 Air-dried soil sample (5g) was weighed into a beaker and 35 ml of phosphorus extracting
145 solution, NH_4Cl was measured and added to the content of the beaker. The mixture was well
146 stirred for 5 minutes before filtered using Whatman filter paper No 1 of which 4 ml of the filtrate
147 was pipetted into a test tube and 4 ml of ascorbic acid was also added. The resulting mixture was
148 allowed to stand for 30minutes on a test tube rack for colour development. The color developed
149 was blue and the procedure was repeated for the other samples. The standard was also prepared
150 by measuring 0.5 ml of 100ppm phosphorus standards and adding 4 ml of indicator M and R
151 solution. Twenty five millilitres of distilled water was added. The solution was transferred into
152 another test tube. A blank was prepared by measuring 4 ml of the ascorbic acid reagent and 25
153 ml of distilled water into another test tube. The available soil phosphorus absorbance was read at
154 660nm wavelength using the corning colorimeter model 253 (AOAC, 1990).

155 **Determination of total nitrogen content of soil samples**

156 A gram of each of the soil samples, which previously has been ground and sieved in a
157 2mm wire mesh was weighed and transferred into a 500 ml micro-Kjeldahl flask and 20 ml
158 concentrated H_2SO_4 was added, the mixture was swirled for a few minutes and was allowed to
159 stand for about 30 minutes. A 20 ml concentrated copper oxide catalyst was added to the
160 mixture. The flask was then transferred to a mechanical heating mantle. The heater was placed in
161 the fume cupboard connected to the electrical outlet socket and was switched. The flask was then
162 left to boil for about 5 hours in the fume cupboard. After the digest has been observed to be clear
163 of H_2SO_4 fumes in the flask, the heater was then switched off. The micro-Kjedahl flask was then

164 removed from the heater and allowed to cool. The digest was then decanted into another flask.
165 100 ml of distilled was then added to the content of the flask. The micro-Kjedahl flask was then
166 attached to the distillation apparatus. 10 ml of 40% NaOH solution was added through the funnel
167 stop cork of the distillation apparatus. A 50 ml boric acid with indicator solution was transferred
168 into 25 ml conical flask. The flask was then placed under the condenser of the distillation
169 apparatus. The tip of the condenser was positioned such that it was about 4mm above the surface
170 of the boric (H_3BO_3) solution in the conical flask. The digest was then distilled by allowing hot
171 steam pot to pass from the steam pot into the digest in the micro-Kjedahl flask, thereby causing
172 the digest to boil and distill over into boric acid. After about 150 ml of the distillate had been
173 collected in boric acid, the distillation was stopped. The distillate was then titrated with 0.5 M
174 standard hydrogen chloride. The colour change at the end point was from green to pink (AOAC,
175 1990).

$$176 \quad \% \text{ Nitrogen} = \frac{M \times T \times 0.014 \times V_1 \times 100}{W \quad V_2}$$

177

178 M = Molarity of acid used

179 T = Titre volume

180 V_1 = Volume of digest

181 V_2 = Volume of digest used

182 W = Weight of sample

183 0.014 = Multiplication factor (i.e. milligram equivalent of nitrogen in ammonia)

184

185

186 **Determination of organic carbon content of soil samples**

187 Five grammes of sample was placed in a ceramic crucible and then heated at 350°C overnight.

188 The sample was then cooled in a desiccator and weighed.

189
$$\text{Organic matter content} = \frac{\text{Initial-Final Sample Weight}}{\text{Initial Sample Weight}} \times 100$$

190 All weights were corrected for moisture water content prior to organic matter content
191 calculation. To convert the organic matter to total organic carbon content, a conversion factor of
192 1.724 was used based on the assumption that organic matter contains 58% organic carbon
193 (AOAC, 1990).

194
$$\text{Organic Carbon (g)} = \frac{\text{Organic Matter (g)}}{1.724}$$

195 **Determination of exchangeable sodium and potassium of the content of soil samples using**

196 **Flask emission Photometry method**

197 Ammonium acetate extracting solution was used for the extraction of Na and K in the soil
198 sample. The procedure was observed for the exchangeable Mg and Ca preparation. A 10g of soil
199 sample was weighed into a baker; 100 ml of the ammonium acetate solution was added. The
200 mixture was then filtered using Whatman filter paper. The filtrate was collected, bottled and
201 labelled. The exchangeable Na and K were determined using the flame photometer according to
202 AOAC, 1990.

203 Calculation:

204
$$\text{Sodium (Na) or Potassium (k)} = \frac{R \times V \times D}{W}$$

205 R = Reading

206 V = Volume of extracting solution used

207 D = Dilution factor

208 W = Weight of soil sample used

209 **Determination of organic matter**

210 1g was weighed and transferred to 250-milliliter conical flask. A 10ml of $K_2Cr_2O_7$ Pottassium
211 hepta dichromate was added and swirled to mix. 20 ml of concentrated sulphuric acid H_2SO_4 was
212 rapidly added, shaken and allowed to stand for 30 minutes. The mixture was diluted with 100 ml
213 of distilled water and five drops of ferroin indicator added, it was then titrated with 0.5N ($FeSO_4$)
214 ferrous sulphate (Carter, 1992).

215 A blank titration was prepared in the same was (without soil) to standardize the dichromate
216 solution. Percent organic matter was subsequently calculated, using the formula below;

217
$$\% \text{ Organic Carbon} = \frac{(B-T) \times M \times 0.003 \times 1.33 \times 100}{W = \text{Weight of sample}}$$

218 Correction Factor (CF) = 1.33

219 M = Molarity of solution x ml of solution used

220 $\% \text{ Organic matter soil} = \% \text{ organic C} \times 1.729$

221 **Determination of pH for water samples**

222 This was determined as determined as described by Ademoroti (1996). The Jenway 3015
223 pH meter was first standardized using standard buffer solutions of pH 4 and pH 9. The pH of the
224 sample water was determined by using pH meter on arrival at the laboratory. The electrode was
225 carefully suspended in the sample and allowed to stand until the reading was steady before the
226 reading was finally recorded.

227 **Determination of electrical conductivity (EC)**

228 The samples were thoroughly mixed together thereafter; an aliquot was taken into the
229 meter sample holder. The sample holder was then properly placed into a colorimeter.
230 Immediately the reading knob was depressed, the reading was taken and recorded (Ademoroti,
231 1996).

232 **Determination of Total Dissolved solids**

233 The sample was first filtered using a whatman filter paper. Fifty millilitres (50 ml) of the
234 filtrate was then transferred into a previously weighed evaporating dish. This was evaporated to
235 dryness on an electric hot plate before drying to constant weight in the oven at 105°C. The
236 weight of the dish was subtracted from the final weight (mg) of the total dissolved solid.

$$237 \quad \text{Total Dissolved solid } \left(\frac{\text{mg}}{\text{L}} \right) = \frac{\text{Total Dissolved Solid (mg)} \times 1000}{\text{filtrate taken (ml)}}$$

239 **Determination of Chloride**

240 The Mohr method as described by AOAC (1990) was used, 100ml of the sample was
241 measured into a conical flask and a pinch of powder CaCO_3 was added. This was following by
242 addition of 2ml of the indicator. The whole mixture was then titrated against standard AgNO_3

243 solution to a permanent reddish-brown precipitate A blank titration was equally carried out by
244 substituting the sample with distilled water.



247 The chloride was expressed as

$$\text{Cl} \left(\frac{\text{mg}}{\text{L}} \right) = \frac{(A - B) \times M \times 70,900}{\text{ml of sample}}$$

248 where A= ml of AgNO₃ for sample

249 B= ml of AgNO₃ for blank

250 M= molarity of AgNO₃

251 **Determination of sulphate**

252 The turbidity method was used by using BaCl₂ as precipitant as described by Ademoroti,
253 (1996). Ten millimeters (10 ml) of the sample was introduced into 25 ml volumetric flask and 10
254 ml of distilled water was added. This was following by addition of 1ml of gelatin- BaCl₂ reagent.
255 The mixture was made up to mark with distilled water. The mixture was allowed to stand for 30
256 minutes before the optical density was determined at 420nm.

257 Calculation

$$\text{SO}_4 \left(\frac{\text{mg}}{\text{L}} \right) = \frac{\text{mass of SO}_4 \text{ from curve} \times 1000 \times D}{\text{ml of sample}}$$

258 where D is the dilution factor

$$D = \frac{\text{total volume of mixture}}{\text{sample volume}}$$

259

260 **Determination of Phosphate**

261 Vanado-Molybdophosphoric acid colorimetric method as described by Ademoroti (1996)
262 was used. Vanado-Molybdo-Phosphoric Acid Colometric Method: Excessive color of sample
263 was removed by shaking 50 ml portions of each with about 0.2g activated carbon type No 33033
264 is an Erlenmeyer flask about 5 minutes. It was then filtered through whatman filter paper.

265 **Standard phosphate solution**

266 0.1295g anhydrous KHPO_4^{3-} was accurately weighed with the aid of an analytical
267 weighing balance. It was then dissolved in 1 litre of distilled water. Serial dilution was thereafter
268 prepared from the standard solution.

269 **Calculation**

$$\frac{\text{mg}}{\text{PO}_4^{3-}} P = \frac{(\text{reading from curve} \times 1000 \times D)}{\text{ml of sample}}$$

270

271 **Determination of Hardness**

272 The total hardness was determined using titration method (Ademoroti, 1996). A 25ml of
273 water sample was diluted to 50ml with distilled water in a conical flask. A millilitre of the buffer
274 10 solution was added and a pinch of solochrome black T indicator and KCN (for masking) were
275 added and titrated against 0.01M EDTA to the final end point which is blue.

$$\text{Total hardness (EDTA) as } \frac{\text{mg}}{\text{L}} \text{ CaCO}_3 = \frac{(V \times A \times 1000)}{\text{ml of sample}}$$

276 Where V = ml titration for the sample

277 A= mg CaCO₃ equivalent to 1ml EDTA titrant

278 **Determination of Alkalinity (Hydroxide, Carbonate and Bicarbonate)**

279 A few drops of phenolphthalein were added to 20ml of water sample in a 50ml conical
280 flask but there was no color change which indicated the absence of hydroxide and carbonate. A 2
281 drops of methyl orange was added to fresh 20ml of water sample in a 50ml conical flask. The
282 colored solution was then titrated against 0.025M H₂SO₄ till the color changed from yellow to
283 pink. Blank titration was also carried out.

$$\text{HCO}_3 \text{ alkalinity} = \frac{V \times A \times 1000}{\text{ml of sample}}$$

284 Where V = methyl orange titration

285 A= concentration of acid

286 **Determination of Metals**

287 The sample for metal analysis was prepared prior determination 5ml of concentrated HNO₃
288 was added to 200ml of water sample in a 250cm³ beaker. The solution was evaporated to near
289 dryness (less than 25 mL). After cooling, the solution was made up 2ml with concentrated HNO₃
290 and transferred into sample bottles prior analysis (Ademoroti, 1996). The heavy metal (Cd, Pb,
291 Cu, Cr, Ni, Zn, Co) were determined with Atomic Absorption Spectrophotometer (AAS) by
292 using appropriate wavelength for each and the alkali metals (Na and K) were determined by

293 using flame photometer. The absorbance and the concentration of the metals were thereby
294 obtained.

295 **Effects of some physicochemical factors on flocculating activity**

296 **Jar Test determination of bioflocculant dosage**

297 0.2 to 1.0 mg/mL of the purified bioflocculant was prepared. Their bioflocculating
298 activity was measured against 4 g/L of Kaolin clay suspension. 3.0 mL of 1% weight per volume
299 of CaCl₂ was added to the different concentrations of the purified bioflocculant and mixed with
300 100 mL of kaolin clay suspension in 500 mL beakers. The solution was rapidly mixed at 160 rpm
301 for 2 min, followed by gradual flocculation at 40 rpm for 2 min and sedimentation for 5 min.
302 After sedimentation, 2 mL was gently withdrawn from the upper clarifying layer in order to
303 measure the flocculating activity Agunbiade *et al.*, (2017).

304 **Effect of cations on flocculating activity**

305 According to Agunbiade *et al.*, 2017 the effect of different cations on bioflocculant
306 production was gotten by using Na⁺, K⁺, Mg²⁺, Mn²⁺, Al³⁺ and Fe³⁺ in the place of CaCl₂ in the
307 production medium.

308 **Effect of pH on flocculating activity**

309 With the use of 0.1 M HCL and NaOH as buffer solutions in adjusting the pH of the
310 production medium, the effect of pH on the flocculating activity of the bioflocculant produced
311 was gotten. pH range of 3-12 was used in this set up. (Elkady *et al.*, 2011).

312

313 **Effect of temperature on flocculating activity**

314 Heat stability was evaluated by incubating the bioflocculant solutions in water bath at a
315 temperature range of 50, 60, 70, 80, 90 and 100°C for 25 minutes. Afterwards, the residual
316 flocculating activity was determined using the protocol of Gong *et al.*, (2008) and Agunbiade *et*
317 *al.*, (2017).

318 **Microbial analysis of water samples**

319 Fifth fold serial dialution was carried out on the collected water samples, using pour plate
320 method to determine microbial load. Biochemical identification methods were employed in the
321 identification of the isolated bacteria. (Omezuruike, 2008).

322 **Results and Discussion**

323 **Physicochemical parameters of soil and water samples**

324 The abattoir slaughtering site was richer in organic content (75%), organic matter (74%),
325 phosphorus (74%), potassium (43%), sodium (62%) and magnesium (42%) than stream bank and
326 sediment. This indicates that the site of slaughter contains higher organic nutrient than the soils
327 of the stream bank and stream sediment. Soil from the stream sediment is sandier than others
328 (Table 1). Abattoir waste water presented high values of the following: sodium (61%), potassium
329 (64%), iron (45%), copper (46%), zinc (52%), pH (38%), conductivity (95%), soluble solids
330 (37%), dissolved solids (91%), Total Dissolved solids(93%), Dissolved oxygen (99%),
331 Biochemical oxygen demand (97%), alkalinity (97%), chloride (97%), chemical oxygen demand,
332 (99%) hardness (81%), sulphate (91%) and phosphate (99%) when placed side by side with well
333 and stream water from the same environment. Well water had more calcium content than abattoir

334 waste water. Also, stream water contained more lead, cadmium and cobalt than the remaining
335 water samples (Table 2).

336 Isolation of bioflocculant producing bacteria

337 *Escherichia coli*, and *Bacillus cereus* were isolated from the three soil samples, ABSS,
338 SBNK and STSD. Among the isolates from abattoir slaughtering site are *Citrobacter freundii*
339 ABSS, *Bacillus subtilis* ABSS and *Monococcus luteus* ABSS which were not isolated from
340 stream bank and sediment. However, *Staphylococcus aureus* was not isolated from abattoir
341 slaughtering site. *Proteus mirabilis* SBNK was found in stream bank but was absent in
342 slaughtering site and stream sediment. The following organisms were isolated from stream
343 sediment but not found in slaughtering site and stream bank. They are *Clostridium botulinum*,
344 *Shigella dysenteriae*, *Streptomyces somaliensis* and *Salmonella typhi* (Table 3).

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354 **Table 1 Physicochemical analysis of soil samples**

355	Soil Sample	ASS	SB	SS
356	pH	5.16±0.06 ^a	5.23±0.06 ^a	5.19±0.06 ^a
357	MC	21.55±0.06 ^a	34.25±0.06 ^b	5.19±0.06 ^c
358	OC	1.55±0.06 ^a	0.17±0.06 ^a	0.36±0.06 ^b
359	OM	2.67±0.06 ^a	0.30±0.06 ^b	0.63±0.06 ^c
360	N	0.40±0.06 ^a	0.08±0.06 ^a	0.06±0.06 ^b
361	P	32.62±0.60 ^a	25.20±0.06 ^b	16.52±0.06 ^c
362	K	1.26±0.06 ^a	0.49±0.06 ^b	0.28±0.06 ^c
363	Na	1.30±0.01 ^a	0.73±0.01 ^b	0.44±0.01 ^c
364	Ca	2.60±0.10 ^a	1.50±0.01 ^b	3.00±0.01 ^c
365	Mg	1.30±0.10 ^a	0.73±0.09 ^a	1.00±0.06 ^b
366	Sand	56.80±0.06 ^a	52.80±0.06 ^b	80.80±0.06 ^c
367	Clay	27.20±0.58 ^a	23.20±0.58 ^b	11.20±0.58 ^c
368	Silt	16.00±0.00 ^a	24.00±0.00 ^b	8.00±0.06 ^c

369 Data are presented as Mean ± S.E (n=3). Values with the same superscript letter(s) along the
 370 same row are not significantly different (P<0.05).

371 **KEY**

372 pH – Hydrogen ion concentration, MC- Moisture Content OC- Organic Content, OM- Organic
 373 Matter, MC- Moisture Content, N- Nitrogen, P- PhosphorusK- PotassiumNa-SodiumCa-
 374 CalciumMg-Magnesium ASS-Abattoir Slaughtering Site, SB- Stream Bank, SS-Stream
 375 Sediments

376

377 **Table 2 Mineral Analyses of selected water samples**

378	Parameters	Well	Stream	Abattoir
379		Water	water	waste water
380	Na	23.80±0.06 ^a	32.90±0.90 ^b	90.50±0.12 ^c
381	K	13.70±0.12 ^a	17.70±0.12 ^b	56.50±0.12 ^c
382	Mg	5.53±0.12 ^a	5.89±0.12 ^b	5.78±0.12 ^c
383	Ca	40.20±0.12 ^a	39.90±0.12 ^b	11.30±0.12 ^c
384	Pb	0.33±0.12 ^a	0.42±0.12 ^b	0.38±0.12 ^c
385	Cd	0.012±0.00 ^a	0.013±0.00 ^b	0.010±0.00 ^b
386	Cr	0.001±0.00 ^a	0.001±0.00	0.001±0.00
387	Fe	0.31±0.00 ^a	0.33±0.00 ^b	0.52±0.00 ^c
388	Cu	0.02±0.00	0.05±0.00	0.06±0.00
389	Ni	0.06±0.00	0.08±0.00	0.07±0.00
390	Co	0.010±0.00	0.012±0.00	0.009±0.00
391	Zn	0.40±0.01 ^a	0.41±0.01 ^a	0.86±0.01 ^b
392	pH	6.49±0.34 ^a	6.90±0.00 ^a	8.30±0.12 ^b
393	Cond	39.80±0.12 ^a	755±0.00 ^b	14480±0.00 ^c
394	SS	1.42±0.12 ^a	1.59±0.12 ^b	1.73±0.12 ^c
395	DS	342±0.00	377±0.00	7225±0.00
396	TDS	179.50±0.12 ^a	378.72±0.88 ^b	7226.73±0.12 ^c
397	DO	5.49±0.12 ^a	5.54±0.1 ^a	910.41±0.12 ^b
398	BOD	1.01±0.01 ^a	1.40±0.12 ^b	82.76±0.01 ^c
399	Alkalinity	139.40±0.12 ^a	320.00±0.00 ^b	13600.00±0.00 ^c
400	Chloride	73.24±0.06 ^a	81.99±0.00 ^b	5466.35±0.12 ^c
401	COD	157.00±0.00	168.00±0.00	51200.00±0.00
402	Hardness	189.20±0.12 ^a	201.60±0.12 ^b	1680.00±0.00 ^c
403	Sulphate	223.89±0.00 ^a	232.01±0.00 ^b	4408.35±0.12 ^c
404	Phosphate	11.98±0.00 ^a	13.68±0.00 ^b	3967.180.00 ^c

405 Data are presented as Mean ± S.E (n=3). Values with the same superscript letter(s) along the same row are not
 406 significantly different (P<0.05).

437 Six bacteria with bioflocculating potentials were isolated. They are *Bacillus cereus*,
438 *Streptomyces somaliensis*, *Streptomyces griseus*, *Bacillus thurigiensis*, *Bacillus subtilis* ABWW
439 and *Bacillus subtilis* STSD. *Bacillus cereus* had its flocculating activity increase progressively
440 with time, the same was recorded for *Streptomyces somaliensis*, *Streptomyces griseus* and
441 *Bacillus thurigiensis*. *Bacillus subtilis* ABWW and *Bacillus subtilis* STSD had their highest
442 flocculating activities at 144hours of production and least flocculating activity at 216 hours.
443 *Bacillus subtilis* STSD had the highest flocculating activity at all times of production. This makes
444 it the bacterium with the best flocculating activity of all the isolates (Figure1).

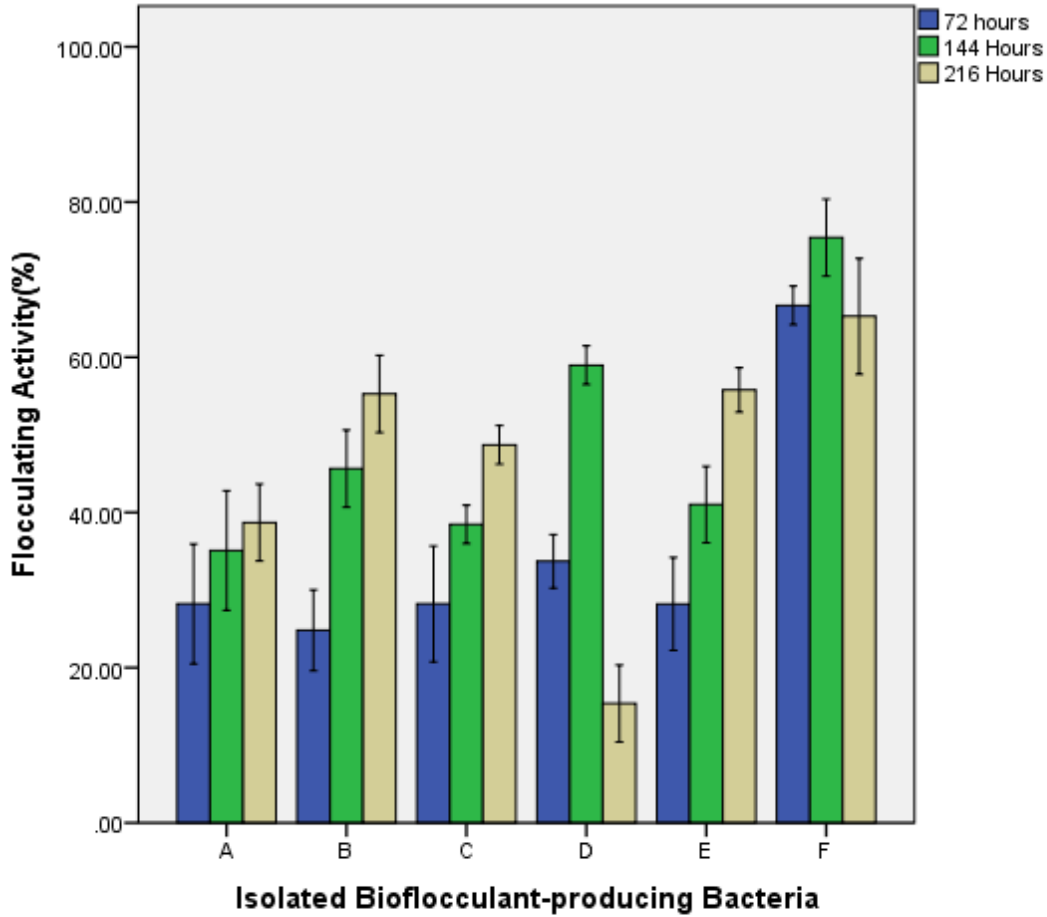
445 Flocculating activity increased progressively with increased dosage till it attained 90% at
446 0.8mg/ml dosage level which is the highest. Thereafter, a progressive decline in the flocculating
447 activities with increased dosage level was recorded (Figure 2).

448 Na^{2+} had the least effect on the flocculating activity of the bioflocculant followed by K^{+} .
449 Mg^{2+} had the best stimulatory effect on the flocculating activity of the bioflocculant produced
450 from *Bacillus subtilis*^b (Figure 3).

451 Least flocculating activity was recorded at pH 4 which progressively increased to a peak
452 of 80% flocculating activity at pH8 and steadily decreased with increased pH (Figure 4).

453 The temperature retaining ability of the bioflocculant shown in figure 5 indicated
454 progressive increase in temperature. Highest flocculating activity was 80% at 100°C and least
455 was 75% at 50°C.

456



Error bars: 95% CI

457

458 **Figure 1** Flocculating activities of isolated Bioflocculant producing bacteria

459 **KEY**

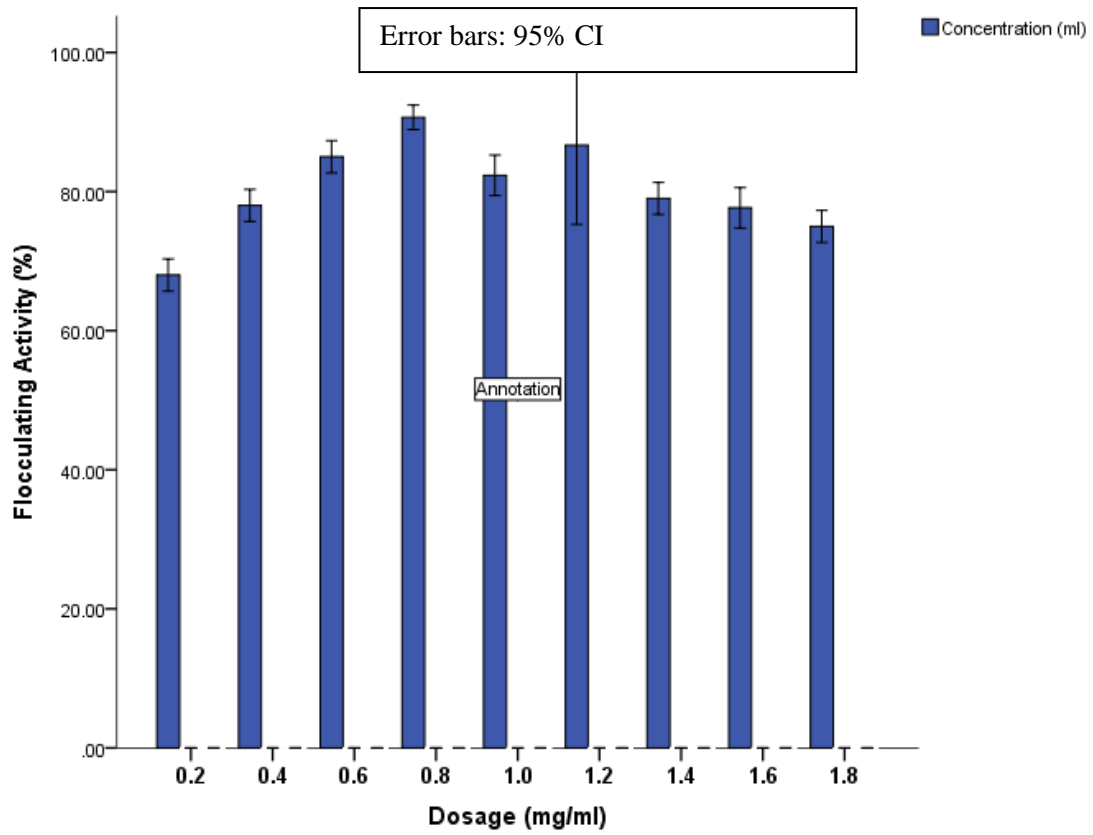
460 **A** *Bacillus cereus* **B** *Streptomyces somaliensis* **C** *Streptomyces griseus*

461 **D** *Bacillus thuringiensis* **E** *Bacillus anthracis* **F** *Bacillus subtilis*

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464 [Type a quote from the document into the text box. You can position the text box
 465 anywhere in the document. Use the wrapping of the pull quote text
 466 box.]



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469 **Figure 2** Effect of Treatment dosage on flocculating activity of the bioflocculant

470 purified from *Bacillus subtilis*

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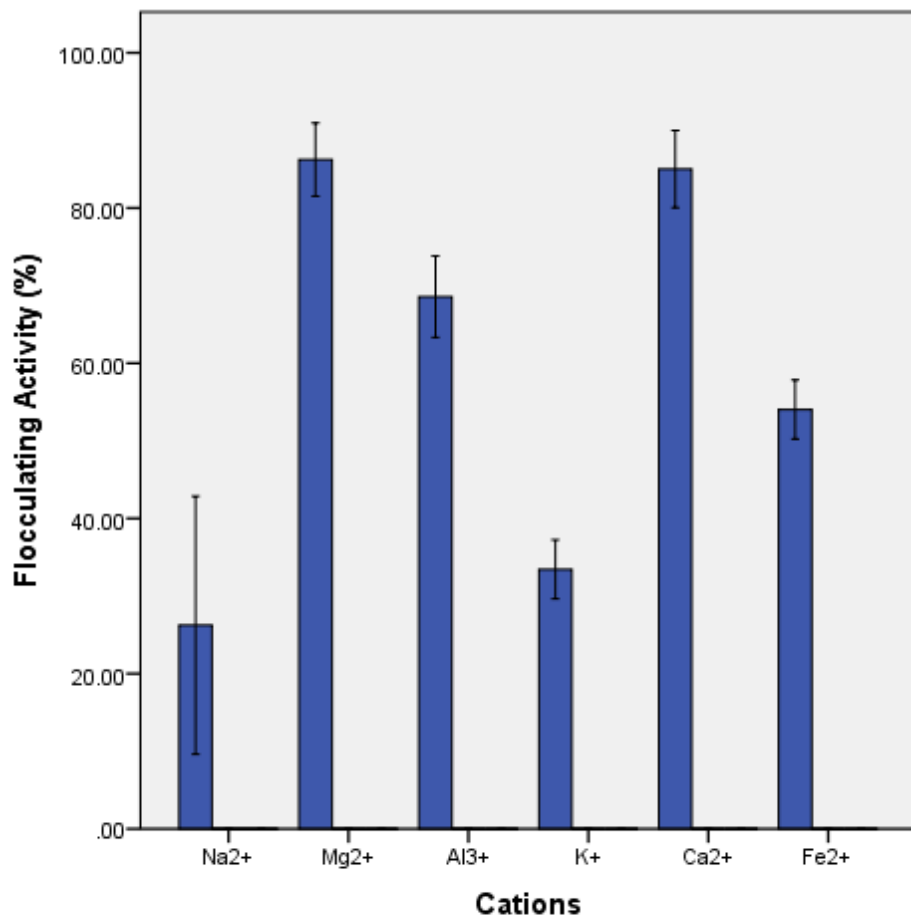
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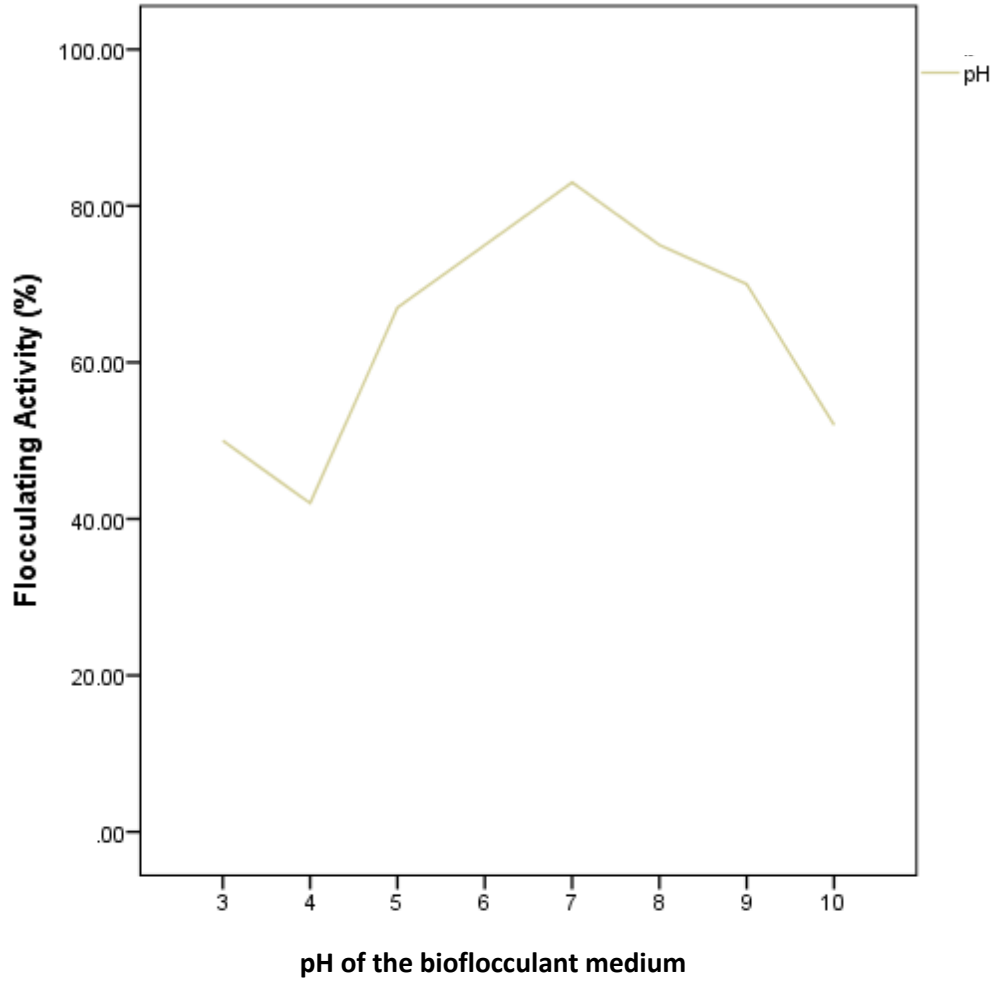


Error bars: 95% CI

477

478 **Figure 3** Effect of cations on flocculating activity of the bioflocculant purified from

479 *Bacillus subtilis*



480

481 **Figure 4** Effect of pH on flocculating activity of the purified bioflocculant from

482 *Bacillus subtilis*

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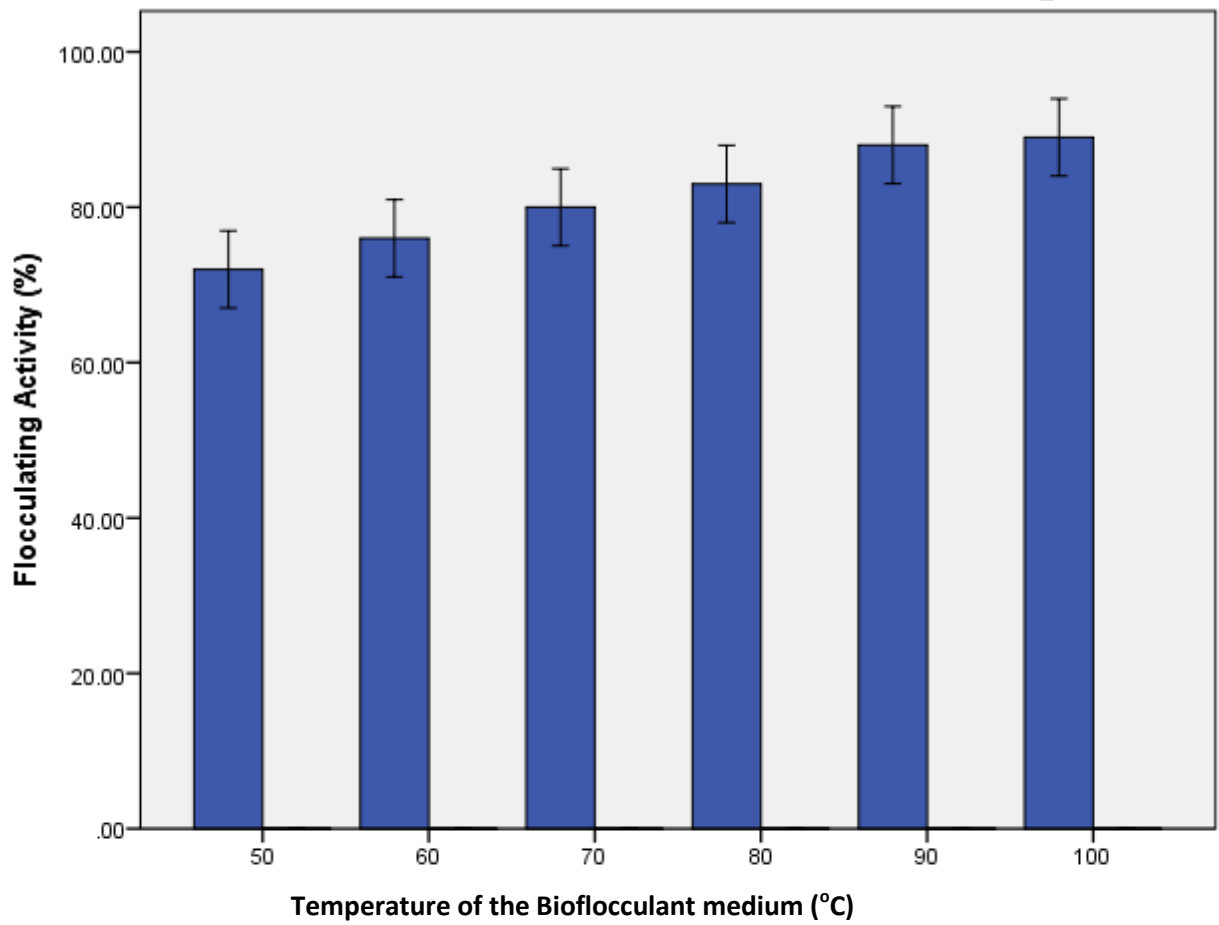
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Error bars: 95% CI

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492 Fig. 5. Graphical representation of temperature retaining ability of the biofloculant during
493 progressive increase in temperature

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498 The increased value of Abattoir Slaughtering Site than Stream Bank and Stream Sediment
499 suggests the impact of abattoir waste on the surrounding soil. Abubakar and Tukur (2014)
500 revealed that the discharge of abattoir effluent to the surrounding soil had significant effect on
501 some soil chemical properties. Going by the findings of Chukwu and Anuchi (2016), the values
502 obtained above are typical for soil samples located within the vicinity of a slaughter house. It
503 can be deduced from the physicochemical parameters that the pH values of the soil samples were
504 below average, indicating that the soil samples were slightly acidic (Chukwu and Anuchi, 2016).
505 There's a possibility of contamination of the soil samples from abattoir activities (Neboh *et al.*,
506 2013).

507 Well water samples consist of expected compositions of minerals for typical well water.
508 This can be attributed to the fact that the well water is not located within the vicinity of the
509 abattoir slaughtering site. This water sample serves as a form of control to other water samples.
510 Mineral composition of the stream water is a little above average and this can be associated with
511 the fact that the utensils, containers and bodies of the slaughter men are washed in the stream
512 where the water sample was collected (Adelegan, 2002). The mineral composition of abattoir
513 waste water was extremely high and can be associated with the deposition of fat contents, animal
514 wastes etc., in the abattoir waste water (Teekenah *et al.*, 2012).

515 During the production of bioflocculant from *Streptomyces griseus*, according to
516 Shimofuruya *et al.*, 1996. The bioflocculant was produced by the bacterium in the death phase of
517 its growth. In this research, the highest flocculating activity was achieved at 216 hours of
518 cultivation indicating the production of more bioflocculant at the death phase. The bioflocculant
519 purified from *Bacillus* sp 1-450 was produced during the log phase as reported by Kumar *et al*
520 (2004). *Bacillus* sp isolated in this research had their bioflocculants produced in less than 72

521 hours of production which increased with time for all at 144 hours but decreased after 216 hours
522 of production in the case of *Bacillus thuringiensis* and *B. subtilis*.

523 Effect of treatment dosage or inoculum size on the flocculating activity of the purified
524 bioflocculant from *Bacillus subtilis* shows that high flocculating activity of 80% and above was
525 achieved with treatment dosages of 0.8, 0.6, 1.0, and 1.2 mg/ml. At dosage 0.2 mg/ml, the
526 flocculating activity was about 70%; which has the lowest flocculating activity. It can be
527 deduced that the bioflocculant gave its best flocculation at different dosage levels of 0.6, 0.8 and
528 1.2. Previous studies have shown that inoculum size play important role in cell growth and
529 bioflocculant production (Okaiyeto *et al.*, 2015). Small size inoculum prolong the lag phase,
530 while large inoculum make niches of strain overlap excessively and consequently inhibit
531 bioflocculant production (Okaiyeto *et al.*, 2016, Zhang *et al.*, 2007). The quantity of the
532 bioflocculant taken at varying quantities per milliliter reflects its bioflocculating ability
533 (Okaiyeto *et al.*, 2016). The biopolymer purified from *Bacillus mojavensis* at a dosage level of
534 5.2g/l attained very fast sedimentation (Elkady *et al.*, 2011). Bioflocculant purified from
535 *Bacillus* sp DP 152 at a dosage of 1mg/l brought about flocculation (Suh *et al.*, 1991).

536 The effect of cations on the flocculating activity of the bioflocculant produced by
537 *Bacillus subtilis* showed that except Na⁺ and K⁺, which drastically reduced the bioflocculating
538 efficiency of *Bacillus subtilis*, virtually all the metal ions stimulated flocculating activity of the
539 bioflocculant to a level above 50%. This result is in tandem with that of Ugbeyen and Okoh
540 (2014), Okaiyeto *et al.*, (2015) where the monovalent ions used completely inhibited the
541 flocculating activity of the bioflocculant used. The variation in the flocculating activity recorded
542 could be as a result of the fact that the bioflocculants being compared (this research, Okaiyeto
543 (2015) and Ugbeyen and Okoh (2014)), were produced from different organisms and this

544 therefore justifies the disparity in the effect Na^+ and K^+ has on the bioflocculating activity.
545 Divalent cations were observed to better stimulate the flocculating activity of the bioflocculant
546 produced by *Bacillus subtilis*. Ugbeyen and Okoh, (2014) had similar finding that divalent
547 cations have good stimulatory effect on bioflocculating activities of bioflocculants. The
548 bioflocculant showed optimum flocculating activity with Mg^{2+} and Ca^{2+} when compared with
549 Al^{3+} and Fe^{3+} . Bioflocculant produced by *Bacillus licheniformis* (CRC 10826) in an aerobic
550 culture medium with citric acid, glutamic acid and glycerol as carbon source had its flocculating
551 activity stimulated by Ca^{2+} , Fe^{3+} and Al^{3+} with a neutral pH (Shih, 2001). *Bacillus* sp Gilbert had
552 its source from Algoa Bay used Mg^{2+} as cation at pH 6.2 (Nontembiso *et al.*, 2011).

553 Studies have shown that the initial pH of the growth medium required for bioflocculant
554 production varies from one microorganism to the other (Li-Fan and Cheng, 2010; Ugbenyen *et*
555 *al.*, 2012). Xia *et al* (2008) revealed that the initial pH of the growth medium affects the electric
556 charge of the cell and the redox reaction which in turn affect the nutrient assimilation and
557 enzymatic reaction. The effect of pH on flocculating activity of purified bioflocculant from
558 *Bacillus subtilis* was assessed at concentration of 0.8mg/ml with the pH of the solution ranging
559 from 3-12. The bioflocculant flocculated a kaolin suspension with over a wide range of pH
560 between 3 and 12 at rates above 50% except at pH 4 which is drastically low and is about 20%.
561 From this study, it was observed that bioflocculant production is possible in almost all the pH
562 conditions except pH 4. Optimum bioflocculant production was observed in pH 6 at 65%
563 followed by pH 9 at 60%. This gives an impression that bioflocculant production from *Bacillus*
564 *subtilis* is possible under weak acidic and alkaline conditions. This finding is similar to that of
565 Liu *et al.* (2010) and Zulkeflee *et al.*(2012). Zheng *et al.*, (2008) and Okaiyeto *et al.*, (2015)
566 revealed that an alkaline pH range of 7-12 was more suitable for bioflocculant production of

567 *Bacillus sp* F19 with its maximum flocculating activity observed at pH 9; bioflocculant
568 production was however inhibited under acidic conditions. The case was however different for
569 bioflocculant produced from *Cobetia sp.*, with its optimum production at pH 6 (Ugbenyen *et al.*,
570 2012).

571 Effect of temperature on the purified bioflocculant from *Bacillus subtilis*, shows that the
572 bioflocculant purified from *Bacillus subtilis* is thermal stable. This was demonstrated with an
573 increase in the flocculating activity of the bioflocculant with increased temperature. It is
574 understood that bioflocculants rich in polysaccharides are more resistant to heat than those that
575 are mainly composed of proteins or have lesser polysaccharide content (Xia *et al.*, 2008;
576 Okaiyeto *et al.*, 2015).

577 **Conclusion**

578 *Bacillus subtilis* isolated from the stream sediments of Onyearugbulem abattoir stream possessed
579 properties capable of forming colloids as a result of flake formation in the macromolecule
580 produced by it. This bioflocculant can be exploited further in water treatment.

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