1

2345678

9

Original research Article

POTENTIAL APPLICATION OF OYSTER SHELL AS AN ADSORBENT IN VEGETABLE OIL REFINING

ABSTRACT

Aim: Generation of filter materials from the shell of invertebrates for the removal of hazardous components from fluid is receiving low attention, both in industry and in academia. However, the shell of invertebrate like oyster has a very high potential in removing color pigments and other impurities in water and vegetable oils. In other to utilize the adsorption potential and to control the disposal effects of oyster shell on the environment, this work studied its effectiveness as adsorbent in vegetable oil refining.

15 Methodology: Both activated and un-activated oyster shells were produced and each was separately used as adsorbent in the bleaching process of Palm Kernel Oil (PKO). The micro-structural images of the adsorbents produced were studied using SEM. The physicochemical properties of PKO before and after the bleaching process were monitored and compared.

Results: SEM images showed a clear difference on visual observation with higher degree of porosity in carbonated oyster shell. The physicochemical properties after adsorption bleaching showed that the carbonated oyster shell has more adsorptive power in its pure and natural form than when chemically activated with H₂SO₄. A maximum of 96.97 and 94.44 percentage color reductions for un-activated and activated oyster shells were achieved respectively. The performance evaluation of the adsorbents was further substantiated by using Freudlich adsorptions isotherms which gives the K and n (constants for a given adsorbate and adsorbent at a particular temperature) to be 0.397 &

0.567 and 12.373 & -0.459 respectively for un-activated and activated adsorbents. Reduction were also observed in
 the percentage free fatty acid and iodine value after the adsorption process with both adsorbents but more reduction
 were noted in the PKO refined with un-activated oyster shell.

28 Conclusion: Oyster shell can be effectively used as adsorbent in vegetable oil refining. These results confirmed the effectiveness of the oyster shell in color and particulate removal.

Key Words: Activated oyster shell, Un-activated oyster shell, Oyster Shell, Palm Kernel oil, Adsorption Isotherms.

33 1. INTRODUCTION34

Oyster shell contributes adversely to the environmental management in the coastal regions. Especially places where fishery appears to be a highly profitable business and oyster as dominant products of shellfish farm. However, such industry has a potential of serious problems about disposal of oyster shell waste and hence constitute a nuisance to the environment. Gil-Lim [1] analyzed the chemical composition of oyster shell and came up with a report that more than 90% of the shell is CaCO₃ (95.99%) while others are clay minerals like SiO₂, (0.69%) Al₂O₃ (0.42%), MgO, (0.65%) P₂O₅, (0.20%) Na₂O, (0.98%) SrO (0.33%) and SO₃ (0.72%) present in minute quantities. The presence of clay minerals, particularly SiO₂ and Al₂O₃, contributes to the high adsorptive property of oyster shell [2].

Vegetable oil is widely used as edible oil. It is used industrially as base stock in the production of lubricants and grease. Other non edible uses include manufacture of candles, paints and resins. In most industrial applications most especially oleochemical industries, Refined, Bleached and Deodorized (RBD) oil are used. Hence, remover of pigments, and other various trace constituents of the oil is one of the major steps in the vegetable oil refinery and it is best perform by adsorption process.

48

Adsorption is a separation process in which a solute in a liquid is deposited at surface of a solid termed the adsorbent [3]. It could be physical or chemical depending mainly on the operating temperature and the interaction between the solute and the solid. At ordinary temperature, physical adsorption predominates and results from intermolecular forces while at elevated temperatures, chemical bond are either broken or created due to the increase in activation energy and form the bases for chemical adsorption commonly referred to chemisorptions [4]. Adsorption is indicative in most natural physical, biological, and chemical systems, and is widely used in industrial applications

55 [5].

56

Adsorption is usually described through isotherms, that is, functions which connect the amount of adsorbate on the
adsorbent, with its pressure (if gas) or concentration (if liquid). Various isotherms developed includes; Langmuir,
Freudlich, Multisite, Lattice gas and Bet Isotherms [6].

The mechanisms for the adsorption in oil bleaching industry are different and diverse [2]. Noorlida [7] in his work found that the removal of carotene from palm oil/hexane miscella by activated rice hull was due to chemisorptions. Proctor [8] in other hand concluded that adsorption of luxein from soy/hexane miscella on rice hull ash was a physical process while that on activated clay was also a chemical one.

Adsorbent is an important filter material for the removal of hazardous components in exhaust gases or liquid, for the purification of drinking water, waste water treatment, or for refining purposes in vegetable oil industries. It can be in form of activated carbon, carbon nano-tubes, silica gel or activated clay [9], [10], [11]. Because of their wide range of use the demand on adsorbent is high and will rise in future as a result of environmental compliances in many countries.

71

Little attention has been given to economic value of shells of invertebrates like oyster shell, rather they are disposed off and consequently become nuisance in the environment. This work studied the effectiveness of activated and unactivated oyster shell as adsorbent in vegetable oil refining. This will reduce the amount of solid waste in the environment and consequently produce raw materials for industrial use.

77 2. MATERIALS AND METHOD

78 79 **2.1 Materials**

Oyster shell was obtained from Kuatan, Malaysia while the palm kernel oil was obtained from Nigeria Institute for
Palm Oil research Benin Nigeria. The reagents used include sodium hydroxide (0.5mole), sodium thiosulphate
solution, starch solution, glacial acetic acid (2%), potassium iodide, ethanol, phenolphthalene (indicator) and
hydrochloric acid (0.5mole). All the reagents are of analytical grade.

The major equipment used are: Electric furnace, electric oven, water bath, Atomic Absorption Spectrophotometer,
 Field Emission Scanning Electron Microscope, Colorimeter, Ph meter and mercury thermometer.

88 2.2 Methods 89

90 **2.2.1** Preparation of adsorbent from oyster shell 91

92 The oyster shell was washed thoroughly with water in a beaker and then dried in an oven maintained at 60 °C for 24 93 hours. This was followed by crushing with mortar and pestle. 200 g of crushed oyster shell was made into slurry by 94 soaking in 0.1 mole H₂SO₄ The resulting mixture was heated at 100 °C till the whole water evaporated. Such is the 95 acid activated oyster shell used. The acid activated oyster shell was dried in an electric oven at 60 °C for another 96 6 hrs before carbonation. Dried acid treated ovster shell was carbonized at a temperature of 400 °C in an 97 environmentally controlled furnace for 1-hour and allowed to cool. The carbonized activated oyster shell was washed 98 in a filtration set to remove the residual acid using distilled water. The residue was dried, pulverized and sieved. This 99 is the activated adsorbent produced from oyster shell. For comparison, non activated adsorbent was made by 100 carbonating the pure oyster shell and then pulverized and sieved. 101

102 2.2.2 Scanning Electron Microscope

103 The surface morphology of the raw and modified oyster shell samples were evaluated using Field Emission Scanning 104 Electron Microscope (JOEL JSM-5600). The oyster shell powder was sprinkled on the sample holder and then coated 105 with a thin gold layer prior to examination using sputter coater. The samples were then mounted on the scanning 106 microscope to view their structural images.

- 107 108
- 1092.2.3Degumming of Crude Palm-kernel Oil (CPKO)110

111 250 ml of crude palm kernel oil was measured into a 1 liter beaker and followed by addition of boiled distilled water 112 (300 ml). The mixture was mixed vigorously in the beaker (1liter capacity). The entire mixture was poured in a 113 separating funnel to remove the gums. The process was carried out three times to ensure complete removal of all the 114 hydratable gums [12].

115 116

117

122

123

137 138 139

140 141

142

150

155

2.2.4 Neutralization of palm kernel oil

118 Neutralization of PKO was carried out by addition of 8.0 ml of 0.5mole NaOH solution into 100ml of degummed 119 oil. The mixture was mixed thoroughly in a beaker. 2 g of common salt was added to facilitate the separation of 120 neutral oil and the soap formed. The neutral oil was filtered from the mixture using vacuum filtration set. 121

2.2.5 Bleaching of Palm Kernel Oil

124 35g of the degummed and neutralized PKO was measured into a beaker of 250ml capacity. This was followed by 125 addition of the activated oyster shell (10% weight of oil) into the oil. The mixture was heated in a water bath at 120 126 °C with continuous stirring for 20 minutes. After the adsorption period, the mixture was filtered and the bleached oil 127 collected. The percentage color reduction was calculated by using the Equation 1. This same procedure was adopted 128 when un-activated adsorbent was used. The collected oil was then subjected to various analyses. 129

130 2.2.6 **Determination of Percentage Colour Reduction (%CR)**

131 The percentage colour reduction of the oil after bleaching was determined using colorimeter (Uniscope, SM-3004). A 132 full absorption spectrum was made between 420 nm and 670nm. The result showed that maximum absorption 133 occurred at 530 nm. The absorbance of the samples was determined at the maximum absorbance wavelength 134 (530nm). The samples were diluted in n-hexane in the proportion 0.1g of oil to 2 ml of solvent [13]. The bleaching 135 capacity of the adsorbent was determined using the following relation: 136

$$\% CR = \frac{Ab_o - Ab_x}{Ab_o} * 100 \tag{1}$$

Where Ab_0 and Ab_x represents absorbance of the crude palm kernel oil and bleached palm kernel oil respectively.

2.2.7 **Determination of Free fatty Acid (FFA)**

143 Acid value and free fatty acid content of the oil before and after bleaching was determined using titrimetric 144 method [14].. Oil sample (2.5g) was accurately weighed in a 250 ml conical flask and 50ml of ethanol and 1ml of 145 phenolphthalein was added to it. The mixture was boiled for 2 minutes and titrated while still hot against standard 146 potassium hydroxide solution (0.1M) in a 25 ml burette (graduated in 0.1ml) under rigorous shaking until the 147 persistence of the pink colour for about 15 seconds. The acid value was calculated from the amount of titrant used 148 using the following relation. 149

Acid value =
$$\frac{V \times N \times 56.11}{M}$$

(2)

151 Where, V is the volume of Potassium hydroxide used in ml, N is the exact molarity and M is the mass in g of oil 152 sample.

153 The free fatty acid was inferred from the acid value according to equation 2 154

% Free fatty acid (FFA) = $0.503 \times \text{acid value}$ (3)

156 2.2.8 **Determination of Iodine value** 157

158 The iodine value of the oil was determined using titrimetric method [15]. Oil sample (0.1 g) was weighed into a 300 159 ml stoppered flask and 20ml of carbon tetra chloride was added. The oil was allowed to dissolve. Iodine bromide 160 (25ml) was also added and the stopper was inserted again. The mixture was gently shaken and kept in the dark for 161 1hr. Potassium Iodide (20 ml) and 150ml of water was added to the mixture. The mixture was then titrated against 162 standard sodium thiosulfate (0.1 mole) until the yellow color due to iodine as almost disappeared. A few drops of 163 starch was then added as indicator and the titration was continued with rigorous shaking until the blue colour just 164 disappeared. The iodine value was calculated from the following relation:

165

iodine value = $\frac{(b-a)*0.01269*100}{weight (in g)of substance}$ Where, a is the number of ml of sodium thiosulphate, b is the volume of titrant in ml. 166 (4) 167 168 169

2.2.9 Effectiveness of adsorbents produced using adsorption isotherms

The adsorptive bleaching was carried out using the standard procedure. The mass of adsorbent was varied between 1 to 3% of PKO (w/w). The results were subjected to Freudlich adsorption equation given in equation 2:

$$Log \frac{x}{m} = nlogc + logk \tag{5}$$

177 Where, x, m and c are amount of substance adsorbed, the amount of adsorbent and the concentration of the residual 178 substance respectively.

RESULTS AND DUSCUSSION 3.

182 Microstructure Analysis of the Oyster Shell Sample 3.1

183 Figure 1 (a), (b) and (c) show the microstructural images of the raw oyster shell, carbonated or un-activated oyster 184 shell and acid activated oyster shell. It is clearly shown in Figure 1 (a) that the oyster shell in its pure natural form is 185 non-porous materials and hence not effective for adsorption bleaching. Compared Figure 1 (a) with (b) and (c), the 186 porosity is obvious on visual observation but more pronounced in 1(b) than (c). This suggests that carbonated oyster 187 shell may have a better adsorption capacity compared to other two.

188 189

170

171 172

173

174 175 176

179 180

181





Figure 1 (a) Raw Oyster Shell

190 191

192 **Physicochemical Properties of PKO**

193 In order to affirm the observation in Figure 1 above, the oyster shell samples were each used as an adsorbent in PKO 194 refining. The major properties of Oil used to test the effectiveness of the adsorbents produced are percentage color 195 reductions (%CR), acid value (AV), Free Fatty Acid (FFA) and Iodine Value (IV).

196 197 3.2 Color and Percentage Color Reduction

198 Figure 2 shows the graph of mass of adsorbent used relative to the absorbance and percentage color reduction in the 199 PKO.

UNDER PEER REVIEW



Figure 2: Percentage colour reduction (%CR) of activated and unactivated adsorbent

The graph shows that the efficiency of both un-activated and activated oyster shell measured in terms of percentage color removal is higher at 3g (10% mass of the PKO) and this corresponds to the maximum of 96.97% CR and 94.44% respectively at the same quantity of adsorbent. Point "A" in the figure, where the two plots intersect signifies a point where either of the adsorbents gives equal %CR. This implies that above 2.5g (8.33% mass of PKO), un-activated oyster shell has a better adsorption capacity than acid activated one and should therefore recommend for such process for economic reason.

3.3 Physicochemical Properties

Table 1 shows the physicochemical properties of crude PKO and refined PKO using 3g of the adsorbent. Generally, the values of these properties after neutralization with NaOH were 7.74, 3.32% and 16.50 for acid value, percentage free fatty acid and iodine value respectively. It is clearly indicated from the table that most of the properties decreases after treatment with adsorbent.

Table 1: Physicochemical Properties of PKO Before and After Refining

Char	Crude PKO	Refined PKO	Refined PKO	Refined PKO	
		(Crude Oyster Shell)	(Unactivated Oyster Sh	nell) (Activated	Oyster Shell)
Acid Value (mole/Kg) 7.74		6.72	4.40		6.40
%FFA	3.88	3.37	2.21		3.22
Iodine Value	16.50	16.50	13.69		16.49
Color	Light Brown	Light B	rown Light Yel	llow	Light Yellow

3.4 Acid value (AV)

The acid value (AV) is the number of milligrams of alkali required to neutralize 2g of fat or oil. It determines the level of saturation or unsaturation depending on the type of fatty acid present in the PKO. From the table, an appreciable reduction in the AV is observed when both un-activated and activated oyster oil was used in the refining process but un-activated oyster shell is more effective when compared with activated sample with 4.40 and 6.40 mol/Kg AV respectively. This is an indication that apart from color removal, un-activated oyster shell performs better in terms of acid reduction when compared with acid treated sample. This could be traced to the residual acid used for activation that could be added to the free fatty acid in the PKO.

232

233 3.5 Iodine value (IV)

A higher IV usually indicates problems with fractionation process and the yield of palm kernel stearing. The result as indicated in the Table 1 above shows a sharp reduction (from 16.50 to 13.69) in the IV when un-activated oyster shell was used but no significant effect was observed when activated shell was employed in the process. This therefore justified the claim that carbonized and unactivated oyster shell has a better adsorptive power than acid treated one.

240 3.6 Performance Evaluation of the Adsorbents Using Freudlich Adsorption Isotherms

241 Freudlich equation is useful in color measurement as long as the units of measurement are additives and proportional 242 to the actual concentration of coloring materials in the oil [2], [3]. Figure 3 compared the Freudlich adsorption 243 constant (n and k) for both the activated and un-activated oyster shells. The n and k values as determined from the 244 slope and the intercept of the graph respectively for un-activated was 0.3975 and 0.567 while for activated was 12.37 245 and -0.469 With high n-value (0.3975) in activated oyster shell, the un-activated shell proved to be more effective 246 than activated one. This is because n-value determines the degree of decolorization within which the adsorbent 247 exhibit its greatest relative effect. If "n" is high, the adsorbent is relatively inefficient as an agent for effecting high 248 degree of decolorization, but for low n-value (<< 1), reverse is the case [16]. However, k value is just a general 249 measure of the activity of the adsorbent, it's neither denote high or low efficiency but rather it shows the rate of 250 adsorption in each of the adsorbents [8].





252253

253

234

255 256

Figure 3: Freundlich Adsorption Isotherms

4. Conclusion

Adsorbents have been developed from oyster shell using acid activation and carbonization in the presence of Nitrogen as a purging gas. SEM images showed a clear distinction in three samples on visual observation. The effectiveness of the adsorbents produced was tested on PKO and the un-activated oyster shell has better adsorption capacity compared to acid treated oyster shell and crude oyster shell. This result was substantiated by using Freundlich adsorption isotherms which gave better adsorption constant in favor of un-activated oyster shell. Base on these facts, it is therefore concluded that oyster shell (solid waste) can be effectively used as a raw material for the production of adsorbent.

266 **REFERENCES**267

265

- Gil-Lim Y, Byung-Tak K, Baeck-Oon K, Sang-Hun H. Chemical and mechanical characteristics of crushed oyster shell. Waste Management 2003; 23: 825-834
- Salawudeen TO, Dada EO, Alagbe SO. Performance Evaluation of Acid Treatment Clays for Palm
 Oil Bleachin. Journal of Engineering and Applied Sciences. 2007; 2 (11):1677-1680.
- Hui, YH, Yee AH, Nordin MR. Banley Industrial Oil and Fat Products, 5th Edition, John Willey and Sons Inc USA 1996; (4) 192-196.
- 4. Okeke JA. Refining of Sheanut Oil. MSc Thesis, University of Lagos, Lagos, Nigeria. 1990.
- 5. Vanderwiel DP, Pruski M, King TS. A Kinetic Study of the Adsorption and Reaction of Hydrogen on Silical-Supported Ruthenium and Silver-Rithernium Bimetallic Catalysts during the Hydrogenation of Carbon Monoxide. Journal of Catalysis, 1999;188 (1): 186-202.
- Masel RI. Principle of Adsorbtion and Reaction on Solid Surface, John Wiley & Sons, New York. Inc. 1996;108-133.
- 7. Noorlida HMD, Sundram K, Siew WL, Aminah A, Mamot S. TAG composition and solid fat content of palm oil, sunflower oil, and palm kernel olein belends before and after chemical interesterification" Journal of American Oil Society 2002;79 (11), 1137-1144.
- 283 8. Proctor A, Brooks DD. Adsorptive Separation of Oils" John Wiley & Sons, Inc. 2005; Chapter 5.6.
- Salawudeen TO, Arinkoola AO, Jimoh MO, Akinwande BA. Clay characterization and optimisation of bleaching parameters for palm kernel oil using alkaline activated clay. Journal of Mineral and Material Characterisation and Engineering, 2014; 2: 586-597
- 10. James O O, Mesubi MA, Adekola FA, Odebunmi EO, Adekeye JID, Bale RB. Bleaching performance of
 a Nigerian (Yola) bentonite. Latin America Applied Research. 2008; 38(1):45-49.
- Salma O, Badei G, Fakhriya T. Carbonaceous materials from seed hulls for bleaching of vegetable oils.
 Food Research International. 2003; 36:11–17.
- 12. Salawudeen TO, Arinkoola AO, Jimoh MO, Akinwande BA. A suitability assessment of alkaline activated clay for application in vegetable oil refining. International Journal of Engineering and Advanced Technology Studies. 2014; 2(1):1-12.
- Foletto EL, Colazzo GC, Volzone C, Porto LM. Sunflower oil bleaching by adsorption onto acid- activated
 bentonite. Brazialianl Journal of Chemical Engineering. 2010;28:169-174.
- Animal and Vegetable Fats and Oil. Determination of acid value and acidity, ISO Method 660:2009,
 Geneva, Switzerland.
- Animal and Vegetable Fats and Oil. Determination of iodine value, ISO Method 3961:2013, Geneva, Switzerland.
- 30016.Tan IAW, Hameed BH, Ahmad AL. Applicability of the Freudlich and Langmuir Adsorption Isotherms in
the Bleaching of Rubber and Melon Seed Oils" Journal of American Oil Chemists' Society. 2007; 66 (2),
247-252.