Evaluation of the Influence of Salt Treatment on the Structure of Pyrolyzed Periwinkle Shell

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ABSTRACT: This paper investigates the effect of use zinc chloride in varying impregnation ratios ranging between 0.1 and 1.5 to chemically activate carbonized periwinkle shells. Studies to characterize the activated carbon were conducted at ambient conditions. The experimental results revealed that the impregnation ratio at 1.0 gave the optimum values for parameters such as iodine number and porosity; and minimum values for parameters like pH and moisture content which represent properties for characterizing maximum adsorption. This study successfully demonstrates the potential of ZnCl$_2$ can be used as an activating agent for producing high quality activated carbon which overall has the capacity to impact positively on the quality of the environment.

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Activated carbon is a black solid substance resembling granular or powdered charcoal. It is a processed carbon material with a highly developed porous structure and a large internal specific surface area (Abdullah et al., 2001, Lysenko, 2007). They are widely used as adsorbents and represent extremely versatile adsorbents of industrial significance. They are employed in many applications which concern principally the removal of undesirable species from liquids or gases (Sharma and Janveja, 2008, Boudrahem et al, 2011).

Typically, the preparation of activated carbon can be divided into two processes. First, the physical method consists of the pyrolysis of the precursor material and gasification of the resulting char in steam or carbon dioxide. The formation of the porous structure is achieved by elimination of a large amount of internal carbon mass. High porosity carbons can be obtained only at a high degree of char burn off. For the chemical method, pyrolysis char would be impregnated with some chemical reagents, such as ZnCl$_2$, H$_3$PO$_4$, NaOH, and KOH. Because of the dehydrogenation process, the chemical reagents promote the formation of cross-links, leading to the development of a rigid matrix, less prone to volatile loss and volume contraction upon usage. Major advantages of the chemical activation compared to the physical activation are lower treatment temperatures and shorter treatment times. In addition, activated carbon obtained by chemical activation exhibits a larger surface area and better developed mesoporosity than physical activation (Sricharoenchaikul et al., 2007, Jabit, 2007).

A number of reports abound on the preparation of activated carbon using various agricultural wastes (Martinez et al., 2007, Wang et al., 2010), low-cost biomass materials such as shea nut shell (Itodo and Itodo, 2011), Parthenium biomass (Rajeshwari et al., 2010), rice husk (Wuana et al, 2007, Nasehir et al., 2010, Alokoh and Adebayo, 2007 Goodhead and Dagde, 2011), coconut shell (Wei et al, 2006, Ash et al., 2006, Gimba and Muyiwa, 2008, Rahman et al., 2006), bituminous coal (Cuhadaroglu and Uygun, 2008), wood (Abdullah et al., 2001, Lysenko, 2007, Goodhead and Dagde, 2011 ), sugarcane bagasse (Qureshi et al., 2007), animal horns (Aluyor and Badmus, 2008), oil palm shells (Luo and Guo, 2001, Tan et al, 2008, Hameed et al, 2009), physic nut waste (Sricharoenchaikul et al., 2007) and periwinkle shells (Badmus et al., 2007) to name a few has been widely studied.

Activated carbon from periwinkle shell forms a cheap, effective, renewable and readily available source of activated carbon. Periwinkle shells are animal waste obtained from dumpsites or market places and are of no industrial importance besides use as land refill. They contain carbon which when carbonized and activated can serve as adsorbents which has various important applications in such areas as analytical chemistry, environmental studies, medicine, fuel storage, gas and chemical purification, distilled alcoholic beverage purification.

In this study, preparation of activated carbon from locally sourced aquatic waste, periwinkle shells using ZnCl$_2$ as activating agent is advanced. Other applications of this salt for activation have been with agricultural materials such as corn cob, groundnut shell, coffee residue (Malik et al, 2006, Jambulingam et al., 2007, Oliveira et al, 2009). The use of acids and bases has variously been applied in previous
research on periwinkle shell carbon (Badmus et al., 2007, Jabit, 2007). The implication of these research findings is that there is a dearth of reports on activated carbon prepared from periwinkle shells using ZnCl₂ as activating agent. The focus of this paper therefore is to provide laboratory scale information on the effects of ZnCl₂ on pyrolyzed periwinkle shell with a view to determining the impregnation ratio of ZnCl₂ that would produce periwinkle shell with high sorption capacity.

MATERIALS AND METHODS

Materials: Carbonized periwinkle shell samples were obtained from previous study of Owabor et al., (2010).

Activation: 100 g of the carbonized sample was measured and placed in a plastic container which was then mixed with Zinc chloride pellets at an impregnation ratio (IR) ranging from 0.1 to 1.5. The IR is calculated as shown below:

\[ \text{IR} = \frac{W_{\text{ZnCl₂}}}{W_{\text{Sample}}} \times 100\% \]

Where \( W_{\text{ZnCl₂}} \) is the weight of ZnCl₂ and \( W_{\text{Sample}} \) is the weight of sample

Distilled water was then added to dissolve all the ZnCl₂ pellets. The mixture was placed in a water bath and made to boil for 1 hour to allow impregnation after which it was cooled to room temperature. After cooling, the slurry was immersed in 1M HCl solution for 24 hours to leach out the excess ZnCl₂ on the activated carbon particles. It was then washed with hot distilled water until the pH of the supernatant was between 6.5 to 7. The slurry was thereafter filtered. The filtered sample was subsequently dried in an oven at 100 °C until a constant mass was obtained. The dry activated carbon sample was stored in airtight plastic containers (Rajeshwari, 2010, Cuhadaroglu and Uygun, 2008, Abdullah et al., 2001).

Porosity: 10 g of AC sample was measured using an electron weighing balance into a graduated cylinder and was then tapped until there was no change in volume. The weighed sample in the measuring cylinder was then immersed into water in a beaker and brought to boil. After the pore has been displaced, the sample was superficially dried and weighed again. The weight divided by the density of water gives the pore volume porosity was therefore calculated from pore volume by dividing the pore volume of the particle with the total volume of the particle (Aloko and Adebayo, 2007).

\[ \text{Porosity} = \frac{W_s - W_d}{P_{V_s}} \times \frac{1}{V_p} \]

Where \( W_d \) = Weight of dried sample; \( W_s \) = Weight of superficially dried wet sample; \( P_{V_s} \) = density of water; \( V_p \) = total volume of particles

Iodine Number: 2 drops of starch solution was added to 10 ml of 0.1N iodine solution in a conical flask. It was observed that the pale yellow of the iodine solution turned blue. The resulting solution was titrated with 0.05N sodium thiosulphate until it became colourless. The burette reading corresponded to blank reading (B). 0.2 g of activated carbon was accurately weighed and introduced into a completely dry conical flask. 40 ml of 0.1N iodine solution was added. The flask shaken for 4 minutes and the contents were filtered. The filtrate was collected in a dry flask after which 10 ml of the filtrate was titrated against standard sodium thiosulphate solution using starch as indicator (Abdullah et al, 2001, Lori et al, 2007). The new burette readings corresponded to (A).

\[ \text{Iodine Value} = C \times Cf \times \frac{mB/g}{\text{wt of carbon}} \times 20 \]

Where \( Cf = \text{conversion factor} \); \( N = \text{normality of iodine} \);

Carbon Yield: The weight of dry sample was measured before and after activation using the electric weighing machine using the procedure of Nasehir et al., (2010), Aloko and Adebayo, (2007). The % carbon yield was calculated as:

\[ \%	ext{carbon yield} = \frac{W_a}{W_b} \times 100\% \]

Where \( W_a = \text{weight after activation} \); \( W_b = \text{weight before activation} \)

Ash content: 2.0 g of the activated carbon was heated at 500 °C for 4 hours, cooled in a dessicator and reweighed (w) following the method described by Ekpete and Horsfall, (2011), Abdullah et al., (2001), Aloko and Adebayo, (2007).

\[ \%	ext{ash content} = \frac{2a - w}{2a} \times 100\% \]

pH: 2.0 g of the activated carbon sample was weighed and transferred into a beaker. 100 ml of

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distilled water was measured, added to the beaker and the contents stirred for one hour. The samples were allowed to stabilize before the pH was measured using a pH meter (Ekpete and Horsfall, 2011, Wuana et al., 2007).

**Bulk Density:** 10 g of sample was weighed and transferred to a measuring cylinder. Shaking or tapping of the cylinder was avoided. The surface was leveled off with a spatula. The cylinder was then tapped until a constant volume was observed (Aloko et al., 2007) and the bulk volume determined as:

\[
\text{Bulk Density} = \frac{\text{mass of sample (g)}}{\text{volume of sample (cm}^3)}
\]

**Moisture Content:** 2.0 g of the dried sample was weighed and placed in a washed, dried and weighed crucible. The crucibles were placed in an oven and dried at 105 °C to constant weight for 4 h (Ekpete and Horsfall, 2011, Aloko and Adebayo, 2007). The percentage moisture content (%MC) was computed as follows:

\[
\%MC = \frac{\text{loss in weight on drying}}{\text{initial sample weight}} \times 100
\]

**Mean Particle Size:** The weight average mean particle size was determined using the procedure adopted from Richardson et al., (2003), McCabe et al., (2005). 20 g of each sample was measured using the electric weighing machine and placed in test sieves, which were arranged according to various sizes of the screens. The amount retained on each screen was collected and weighed; this was done for each of the screens up to the final screen.

\[
\bar{D}_w = \sum_{i=1}^{n} x_i D_{pi}
\]

\(\bar{D}_w = \text{mean particle size}\)

\(D_{pi} = \text{the mean size of each screen used in the sizing}\)

\(x_i = \text{the weight fraction of the particle retained on the screen}\)

**RESULTS AND DISCUSSION**

**Iodine Number:** The amount of iodine adsorbed increased rapidly with increasing ZnCl₂ impregnation and attained an optimum of 104.95mg/g at an impregnation ratio of 1.0 after which a reduction was observed as shown in Fig 1 below. The increase in the iodine number was due to increase in pore formation and hence adsorptive capacity of the activated carbon as impregnation increased. The observed drop in iodine number drop may be due to abundant increase in the pore volume which eventually resulted to internal collapse of the pores. This result was found to be consistent with the report of Viswanathan et al., (2009).

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**Fig 1:** Variation of ZnCl₂ impregnation ratio with iodine number

**Ash Content:** The influence of impregnation ratio on the % ash content of PSC shown in Figure 2 indicate that as the impregnation ratio increases, the ash content increased and reached an optimum point of 4.72% at impregnation ratio of 1.0 after which a decline was observed. The increase in ash content can be adduced to an increased formation of insoluble inorganic compounds with increase in IR. While the drop observed can be linked to a low char burn off at the respective IR.

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<th>%ash content</th>
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**Fig 2:** Effect of ZnCl₂ impregnation ratio on ash content of PSC

**pH:** The pH decreased and attained a constant value as the impregnation ratio increased as shown in Figure 3. Again this can be attributed to the incidence of the metal salt hydrolyzing in the presence of natural alkalinity to form metal hydroxides thereby reducing the alkalinity of the carbonized periwinkle shells with impregnation. The constant pH observed between impregnation ratios of 1.0 and 1.5 might be due to the activating agent possessing similar hydrolyzing ability at these ratios. There is an increase in activated carbon adsorption property as the impregnation ratio increases because the

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percentage of adsorption decreases with increase in pH.

**Bulk Density:** The variation of the bulk density with the experimental ZnCl$_2$ impregnation ratio used in this study showed that bulk density decreased with increasing impregnation ratio until a constant value was obtained as shown in Fig 4. The observed decrease in the bulk density may not be unrelated to the increase in pore volume resulting from increasing activation. The constant value observed between impregnation ratios 1.0 and 1.5 indicates that increasing impregnation no longer affected the pore size development due to a fully developed pore structure.

**Percentage Carbon Yield:** The result obtained showed that there was a weight loss when the carbonized samples were activated. The weight loss was due to devolatilization of the precursors as a result of intense dehydration and elimination reactions. This was primarily to increase the pore development and create new pores. The carbon yield increased as the concentration of ZnCl$_2$ impregnating solution increased as presented in Figure 5 and approached an optimum point at 20 % impregnation ratio of ZnCl$_2$ with 95.25 % yield. Experimental result revealed that a further increase in ZnCl$_2$ loading led to a decrease in yield. This development may be attributed to the inhibition of tar formation and increased pyrolytic decomposition as the impregnation ratio increases. This observation is affirmed by the result of Rajeshwari et al., (2010).

**Moisture Content:** As observed in Figure 6, the moisture content decreased successively for ZnCl$_2$ impregnation ratio ranging from 0.1 to 1.0 and thereafter became constant between impregnation ratios 1.0 and 1.5. This reason for the decrease could be deduced from the fact that dehydration rate of the activated carbon increased with impregnation ratio. The constant values obtained showed that a further increase in impregnation had little or no effect on the moisture content of the activated carbon produced.

**Porosity:** The porosity of the activated PSC increased with increasing impregnation ratio and peaked at an optimum porosity of 0.003947 obtained at impregnation ratios 1.0 and 1.2. This occurrence was as a result of the elimination of large amount of internal carbon mass as explained by Sricharoenchaikul et al., (2007). The increased porosity suggests that pores were created by the reactant as a result of the spaces left by zinc chloride

![Fig 3: Effect of ZnCl$_2$ impregnation ratio on pH of PSC](image)

![Fig 4: Effect of ZnCl$_2$ impregnation ratio on bulk density of PSC](image)

![Fig 5: Effect of ZnCl$_2$ impregnation ratio on carbon yield of PSC](image)

![Fig 6: Effect of ZnCl$_2$ impregnation ratio on moisture content of PSC](image)

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after washing (Marsh and Rodriguez-Reinoso, 2006). The constant value obtained between IR 0.2 and 0.3 and 1.0 and 1.2 may have arisen due to pseudo saturation and full saturation of the pores with increasing IR respectively. The decrease observed between impregnation ratios 1.2 and 1.5 maybe due to a breakdown in the physical structure of the precursor caused by excessive pore pathways leading to shrinkage in particle size with increasing concentration of activating agent.

Fig 7: Effect of ZnCl2 impregnation ratio on porosity of PSC

Bulk Volume: The observed increase in the bulk volume may be due to increase in pore formation and hence bulk volume. The constant value observed between impregnation ratios 1.0 and 1.5 indicates that increasing impregnation no longer affected the pore size development due to a fully developed pore structure.

Fig 8: Effect of ZnCl2 impregnation ratio on bulk volume of PSC

Mean Particle Size: The finer the particle sizes of an activated carbon, the better the access to the surface area and the faster the rate of adsorption. Small particles have the fastest rate of attracting and attaching dissolved molecules to their surface (Marsh and Rodriguez-Reinoso, 2006). The observed decrease from 0.482138mm to 0.420263mm as the zinc chloride impregnation ratio increased from 0.1 to 1.0 may again be due to a breakdown in the physical structure of the precursor caused by excessive pore pathways leading to shrinkage in particle size with impregnation. However, a slight increase was observed between impregnation ratios 1.0 and 1.5. The observed increase may be attributed to an increase in the internal pore size and development of pore pathways leading to an increase in the overall particle size with increased impregnation. The result from this study affirms that the best particle size which would ensure increased sorption capacity of the activated PSC was obtained at impregnation ratio 1.0.

Fig 9: Effect of ZnCl2 impregnation ratio on mean particle size of PSC

Generally, the effectiveness of activated carbon as an adsorbent which is attributable to its unique properties including large surface area, high degree of surface reactivity, universal adsorption effect, and favourable pore size cannot be overemphasized. Porosity, iodine number, mean particle size and pH evaluated in this study that directly influence the suitability of an activated carbon sample for adsorption as these properties reflect the nature of internal pore development and adsorptive capacity (a measure of surface area) of the activated carbon.

Conclusion: Preparation of activated carbon from pyrolyzed periwinkle shell waste was performed in a laboratory scale facility. The data obtained indicated that activated carbon with favorable physiochemical properties may be produced from the methods advanced in this study. The effects of impregnation on the physiochemical properties of periwinkle under study affirm that the optimum impregnation of zinc chloride was obtained at IR 1.0. This impregnation ratio gave peak values for porosity and iodine number and minimal values for pH and mean particle...
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size which are the properties used for assessing the suitability of activated carbon for purposes of adsorption.

REFERENCES


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