Comparative Study on the Adsorption Capacity of Activated Carbon Prepared from Lapsi Seed Stone and Betel nut using Phosphoric Acid

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Abstract: This paper presents the comparative study on the adsorption capacity of activated carbons prepared from Lapsi (Choerospondias axillaris) seed stone and Betel (Areca catechu) nut. Activated carbons (ACs) were prepared from Lapsi seed stone (LSS) and Betel Nut (BN) by chemical activation with \( \text{H}_3\text{PO}_4 \) (in the ratio of 1:1 by weight) at 400°C. The pore structure of activated carbons was determined by iodine number and methylene blue number. Surface morphology of ACs was studied by scanning electron microscopy (SEM). Surface functional groups were analyzed by Fourier Transform Infra Red Spectroscopy (FTIR). As indicated by TGA analysis, the appropriate temperature required for carbonization was 400 °C. Betel nut AC showed high iodine number and methylene number of 888 mg/gm and 369 mg/gm respectively. SEM micrographs of Betel nut AC show the presence of well developed pores on its surface. FTIR result indicated that both ACs contain −OH, >C=O groups as oxygen containing surface functional groups. Based on the result, the AC prepared from betel nut by activation with \( \text{H}_3\text{PO}_4 \) is comparable with commercial activated carbon and could be used as potential adsorbent for removal of pollutants from water and waste water.

Keywords: Lapsi seed stone, Betel nut, phosphoric acid activated carbon

1. Introduction

Activated carbon (AC) is a predominantly amorphous solid carbon material with highly surface area, microporous structure and high degree of surface functional groups. These unique characteristic properties make AC a versatile adsorbent for the removal of organic and inorganic pollutants from water and waste water. Adsorption capacity of AC is highly influenced by the preparation conditions such as activation condition, activating agent and nature of precursor [11].

Activated carbon can be generally prepared by two different methods (i) Physical activation and (ii) chemical activation. Physical activation involves carbonization of precursor followed by the activation of the carbonized product at high temperatures in presence of oxidizing gases such as CO\(_2\) and steam. In chemical activation, the precursor is carbonized at low temperatures in presence of chemical agent under inert atmosphere. Chemical activation has been reported as more
advantageous over physical activation because it requires relatively lower temperature and shorter time for activating the material. In addition, very high surface area AC can be obtained. The most commonly used chemical activating agents are H$_3$PO$_4$, ZnCl$_2$, and KOH. Among these, H$_3$PO$_4$ is widely used for activation of lignocellulosic materials.

Activated carbons have been commonly prepared from variety of carbonaceous materials such as wood, coconut shell, coal. The demand for AC has been increased due to the increased utilization of the carbon in pollution control. As a result, cost of AC is also growing depending on the application. Hence, there is a need for the sorting out new precursors for the preparation of AC which should be cost effective in comparison to the commercially available AC. Recently, use of agricultural wastes as an alternate precursor for the preparation of AC has notably increased as it is widely available at low cost. Presently, the use of coconut shell [9], Bamboo [1], olive oil stone [5], peach stone [10] etc has been reported in literature as a precursor for AC production.

In present study, Lapsi (Choerospondias axillaris) seed stone and Betel (Areca catechu) nut were selected as AC precursors because of their wide availability in Nepal. Moreover, little work has been reported on the production of AC using Lapsi seed stone and Betel nut. So Lapsi seed stone and betel nut is of special interest towards preparation of AC. This study focuses on to make comparison in the adsorption capacity of the AC from Lapsi seed stone and Betel nut by H$_3$PO$_4$ activation.

2. Experimental

2.1. Materials

The Lapsi and Betel nut were collected from vegetable market at Kalimati, Kathmandu. Lapsi seed stone and Betel nut were first washed with tap water then with distilled water, dried in an oven at 110°C for 12 hours. The dried materials were crushed with mortar and electric grinder. The crushed particles are sieved into the fraction of size 312 µm. Chemicals used for this analysis were all analytical grade purity and the solutions were prepared in distilled water. The nitrogen gas is of ultra high pure (UHP) grade.

2.2 Preparation of Activated Carbon

The Lapsi seed stone (LSS) and Betel nut (BN) powder were separately mixed with H$_3$PO$_4$ in the ratio of 1:1 by weight. Then the mixture was heated in hot plate till dry mass was obtained. Then, the samples were oven dried at 110°C for 24 hours. The dry mass was separately transferred to a quartz tube and placed in horizontal tubular furnace (Accumax, India) and carbonized at temperature of 400°C for 3 hours in continuous flow of nitrogen gas at flow rate of 100 mL/minute. The product was cooled to room temperature. Then treated with 0.1 M NaOH and washed with distilled water till free from acid. The resulting carbons were then dried in an electric oven at 110°C for 3 hours. The well dried ACs were sieved into the particle size of 106 µm and used for further investigations.

2.3 Characterization of Precursor

In order to predict thermal behavior of the precursor, TGA analysis of Lapsi seed stone and Betel nut was carried out. Thermo gravimetric analysis (TGA) was carried out by a thermo gravimetric analyzer (EXSTAR 6300 SEIKO TG/DTA, Korea).
2.4 Characterization of Activated Carbon

Characterizations of ACs were performed by determining iodine number, methylene blue number, scanning electron microscopy (SEM) and fourier Transform-Infrared (FTIR) Spectroscopy.

2.4.1 Determination of Iodine Number

Iodine number ($I_N$) indicates the extent of micro pore distribution in the carbon. It is defined as the milligram of iodine adsorbed by 1.0 gm of carbon \[4\]. The iodine number correlates with surface area. The iodine number was determined according to the standard method (ASTM method, 2006). In this method, 100 mg of activated carbon was added to 5 mL of 5 % HCl and then boiled and cooled. To the cooled solution 10 mL of 0.1 N iodine solution was added, shaken for 30 second and filtered. The filtrate was titrated with 0.1 N sodium thiosulphate solution using starch as an indicator. Iodine number is determined by the following equation.

\[
Iodinenumber = \frac{\text{Weight of activated carbon (mg)}}{\text{Weight of activated carbon (gm)}}
\]  

2.4.2 Determination of Methylene blue number

Methylene blue number ($MB_N$) indicates the extent of meso pore distribution in the carbon. It is defined as the maximum amount of methylene blue (MB) dye adsorbed on 1.0 gm of carbon \[4\]. $MB_N$ of the activated carbon was determined according to Method \[8\]. For the determination of methylene blue number, 0.1 gm of AC was added to 100 ml of MB of concentration 100 ppm and then equilibrated for 3 hours at 200 rpm. Then the solution was filtered through filter paper Whatman 41, and the remaining concentration of methylene blue was determined by measuring the absorbance at 664 nm using a UV/Vis spectrophotometer (CECIL-CE-100).

Methylene blue number of the AC is calculated by equation (2).

\[
MB_N (mg/gm) = \frac{(C_o - C_e) \times V}{M}
\]

Where $C_o$ (mg/L) and $C_e$ (mg/L) is the initial and equilibrium concentration of the methylene blue solution, $V$ (L) is the volume of the solution and $M$ (gm) is the mass of the AC.

2.4.3 Fourier Transform-Infrared (FTIR) Spectroscopy

FTIR spectra were recorded on the Thermo Electron Corporation, Nicolet 4700 at room temperature. The % of transmission of samples was recorded over 500–4000 cm$^{-1}$.

2.4.4 Scanning electron microscopy (SEM)

Surface morphology of the AC samples was determined using scanning electron microscope U-8000, Hitachi Co. Ltd. Japan.

3. Results and discussion

3.1 Thermogravimetric Analysis (TGA): TGA graph of Lapsi seed stone and Betel nut is shown in Fig. 1
TGA is used primarily to determine the composition of materials and to predict their thermal stability up to elevated temperatures. As it can be seen in TG curve, thermal decomposition of both precursors Lapsi seed stone and Betel nut proceed through three major steps. The weight of the precursors decreases only slightly until 220 °C. It corresponds to the elimination of absorbed water and then a rapid weight loss is observed up to 380 °C. It might be caused by decomposition of cellulose, hemicellulose. A little loss of mass from 380 to 600 °C corresponds to the decomposition of lignin. It indicates that, thermal decomposition of precursors was completed by 400 °C. Hence 400°C was selected as the appropriate carbonization temperature for the preparation of activated carbons.

3.2 Iodine Number and Methylene Blue Number

Iodine and methylene blue adsorption is considered a simple and quick test for evaluating the porous structure of micro and mesoporous carbons. Iodine has a small molecular size and can readily penetrate deep micropores of the AC. So iodine number gives approximate measure of the micropore content of the carbon. Likewise, methylene blue number indicates the mesopore distribution in the carbon and indicates the ability of AC to adsorb medium size molecules. $I_N$ and $MB_N$ of $\text{H}_3\text{PO}_4$ activated LSS carbon and BN carbon is shown in Fig. 1.
The iodine number and methylene blue number of the $\text{H}_3\text{PO}_4$ activated BN carbon was slightly higher than that of the LSS carbon. The values are comparable to commercial AC. It indicated that, $\text{H}_3\text{PO}_4$ activated LSS carbon is highly microporous and mesoporous in nature. Thus, botanical texture of the precursor affects the pore structure of $\text{H}_3\text{PO}_4$ activated LSS carbon significantly which directly links to its adsorption property.

### 3.3 Scanning electron microscopy (SEM) image

Scanning Electron Microscopy (SEM) was used to observe the surface morphology of AC. SEM images of the $\text{H}_3\text{PO}_4$ activated LSS carbon and BN carbon is shown in Fig. 2.

![SEM images of the $\text{H}_3\text{PO}_4$ activated LSS carbon and BN carbon](image)

SEM images show that, both ACs contain numerous pores and crevices over their surface. $\text{H}_3\text{PO}_4$ activated LSS carbon contained open heterogeneous macropores on the external surface. The external surface of $\text{H}_3\text{PO}_4$ activated BN carbon also shows full of pores of different shape and
sizes. It seems that the cavities on the surfaces of carbons resulted from the evaporation of the activating agent (which in this case is phosphoric acid) during carbonization, leaving the space previously occupied by the activating agent. H₃PO₄ functions in two ways during activation. As an acidic catalyst, it promotes bond cleavage reactions and forms crosslink in cellulose chains. Then, it combines with cellulose chains to form phosphate linkages, such as phosphate and polyphosphate esters, that can serve to connect and crosslink biopolymer fragments. H₃PO₄ activated BN carbon showed more pores and cavities on their external surface. Hence, it suggested that, surface morphology of the H₃PO₄ activated LSS carbon and BN carbon is influenced by botanical texture of the precursor material.

3.4 Fourier Transform Infrared (FTIR) Spectra

The adsorption capacity of AC depends upon porosity as well as the chemical reactivity of functional groups at the surface. Various functional groups present on the AC surface are responsible for preferential adsorption for different molecule. FTIR spectra of H₃PO₄ activated LSS carbon and BN carbon is shown in Fig. 3.

![Fig. 4: FTIR of H₃PO₄ activated LSS carbon and H₃PO₄ activated BN carbon](image)

FTIR spectra of H₃PO₄ activated LSS carbon and BN carbons were quite similar. Only slight differences on the intensity of the bands were detected. A broad band at around 3400 cm⁻¹ indicated the presence of the –OH group of phenol. The band located at around 1700 cm⁻¹ indicates C–O stretching of carboxyl or carbonyl groups. The band around 1591 cm⁻¹ and 1416 cm⁻¹ is indicated to C=C stretching vibration in aromatic skeleton generally found in carbonaceous material, such as activated carbon [3]. Both ACs contain oxygenated functional groups such as hydroxyl, carbonyl. It shows that, the nature of precursor has no significant effect on the surface functional groups of H₃PO₄ activated LSS and BN carbon.

4. Conclusion

Adsorption properties of the ACs prepared from Lapsi seed stone and Betel nut by chemical activation with H₃PO₄ was compared. Iodine and methylene blue adsorption indicates that H₃PO₄ activated Betel nut carbon is highly micro and mesoporous with high adsorption capacity. As
shown by SEM micrograph, \( H_3PO_4 \) activated BN carbons shows well developed pore structure on its external surface. FTIR results indicate that both ACs contain oxygenated functional groups such as hydroxyl, carbonyl. The result of adsorption properties indicated that, \( H_3PO_4 \) activated Betel nut carbon can thus be applied for the removal inorganic and organic compounds from water and effluents by adsorption process.

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