Characterization Of Activated Carbon Prepared From Coconut Shell Using Various Reagents For A Low Cost Water-Filter

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Abstract—The chemical and physical conditions used in the synthesis of activated carbon, an adsorbent used in water filtration, influence the specific surface area and porosity. In this paper, coconut shells were the raw material and activation reagents such as Hydrochloric acid, Zinc Chloride and Phosphoric Acid were used. A single and double burn procedure was carried out for each method. The activated carbon obtained was compared in terms of absorption capacity, bulk density, BET surface area and pore volume. The best results were yielded for activated carbon prepared with phosphoric acid using the single burn technique.

Keywords: Activated carbon, Adsorption, Hydrochloric acid, Phosphoric acid, Zinc chloride

I. INTRODUCTION

Activated carbon has been known as one of the highly reliable adsorbents for removal of contaminants from liquids due to its properties such as a large specific surface area and large pore volume that can provide high adsorption capacity, porous interiors and excellent mechanical properties. Adsorption on activated carbon is reliable because of its simple operation, and easy handling without any hazardous conditions.

The most common forerunners for the preparation of activated carbon are organic substances rich in carbon. These include agricultural wastes such as Tea waste, Apple waste, Banana peels, Coconut shells, Bagasse and Sawdust and fossils such as bituminous coal and by destructive distillation of bones.

In this study, coconut shell was used for production of activated carbon and the different methods of activation were studied to find the best activating reagent, since coconut shells are a waste material available at cheap rates in India and are abundant in nature.

Activated carbon is produced either by physical or chemical activation. Physical activation involves carbonization followed by steam or carbon dioxide activation whereas chemical activation involves carbonization followed by chemical activation by reagents such as H₃PO₄, ZnCl₂ and HCl.

In our study, chemical activation method using single burn and double burn techniques was used and the methods were compared.

II. PARAMETERS FOR ACTIVATION

A. Chemical impregnation:

The coconut shell was broken down into fine bits by a hammer and it was ensured that all crushed particles obtained were of a size range of 3-5mm[³]. These particles were then placed in crucibles, each crucible having around 50 grams of the crushed shells. The particles were further grounded using a household mixer grinder to a size of around 1mm.

For the Double-Burn technique, carbonization in the furnace was carried out at a temperature of 450°C[²] and then impregnation was done followed by a second carbonization, while for the Single-Burn technique chemical impregnation was carried out first and then the samples were carbonized.

Different activation agents were used for impregnation, which include Hydrochloric acid, Zinc chloride and Phosphoric acid, all of them at 25% concentration. The crucibles were kept at room temperature in a dry and dark place for a period of 24 hours to facilitate impregnation of the chemicals into the shells. The chemical agents are dehydrating agents that penetrate deep into the structure of the carbon rich shells forming pores within them. Pores affect the specific surface area as larger number of tiny pores will result in a larger specific surface area.
B. Activation Duration:

The activation duration for the coconut shells have a pronounced effect on the formation of pores and hence, development of maximum specific surface area. The duration should be adequate to allow for the evaporation of all moisture and volatile matter in the raw shells.

In the Single-Burn technique, the samples were carbonized over a period of 30 minutes in an inert furnace at a temperature of 450°C [2]. Inert conditions were maintained by supply of N₂ gas during the activation [1]. In the Double-Burn technique, the coconut shells were once burnt at 450°C for a period of 20 minutes followed by chemical impregnation using 25% HCl, ZnCl₂ and 25% H₃PO₄ for 24 hours and a second-burn stage for another 20 minutes was carried out at 450°C. It was observed that smoke containing volatile matter stopped evolving after 10-12 minutes of placing the samples in the furnace for the first time, and thus it might be inferred that the minimum time duration required for activation ranges between 15 to 20 minutes.

Out of these six experiments, highest BET specific surface area and pore volume was observed for the H₃PO₄ impregnated Single-Burn technique.

III. METHODS USED

A. Filtration and pH adjustment:

The samples prepared by both the Single-Burn and Double-Burn techniques were filtered using distilled water, 8-10 times through ordinary filter paper and the filtrate pH was checked after every two filtrations until the pH of the filtrate varied between 6.5 to 6.8 [2]. The initial filtrate had a pH in the range 3.9 to 4.7. The samples that did not reach a pH in the specified range were treated with 1% w/v Sodium bicarbonate solution [3] and filtered with the solution and again with distilled water until the pH was adjusted. The conductance of the filtrate too was checked and was found to be lower than that of tap water, which clearly indicated a low concentration of ions.

B. Drying and Crushing:

The samples were dried in a closed oven at 100°C [5] for 24 hours after being filtered, to remove all traces of moisture. The particles were then crushed using a mortar and pestle and sieved through a sieve set consisting of 180, 125 and 90 μ mesh size at regular intervals and then shaken in a sieve shaker. This was carried out till the size was below 125 μ so as to facilitate for more surface area with smaller size of particles.

C. Adsorption Test using Methylene Blue solution:

A stock solution of Methylene blue dye of 1000 ppm was prepared by dissolving 1.0g of Methylene blue dye powder in 1 liter of distilled water and stored in a 1 liter standard flask to be used for all tests pertaining to methylene blue adsorption.

The stock solution was used to prepare 100ml standard solutions of 62.5ppm, 125ppm, 187.5ppm and 250ppm in stoppered conical flasks for the single and double burn samples, both methods being used for each of the HCl, ZnCl₂ and H₃PO₄ activated carbon samples. 0.25g Activated carbon was added to each sample so as to allow for adsorption capacities of 25mg/g, 50mg/g, 75mg/g and 100mg/g. The solutions with the activated carbon samples were shaken in a rotary shaker at 150rpm [3] for duration of 24 hours. Care was taken that the entire mass of the carbon sample was in the methylene blue solution and nothing stuck to the glass surface above the level of the methylene blue solution in each flask. A qualitative result of the changes observed in each flask is given in the table below.

| Reagent Used | Concentration of Methylene Blue (ppm) | Observation (change of color of methylene blue) |
|--------------|--------------------------------------|-----------------------------------------------|
| HCl          | 62.5                                 | Significant change                            |
| HCl          | 125                                  | Insignificant change                          |
| HCl          | 187.5                                | No changes observed                           |
| HCl          | 250                                  | No change                                     |
| ZnCl₂        | 62.5                                 | Decolorized                                   |
| ZnCl₂        | 125                                  | Significant change                            |
| ZnCl₂        | 187.5                                | Insignificant change                          |
| ZnCl₂        | 250                                  | No change                                     |
| H₃PO₄        | 62.5                                 | Decolorized                                   |
| H₃PO₄        | 125                                  | Decolorized                                   |
| H₃PO₄        | 187.5                                | Decolorized                                   |
| H₃PO₄        | 250                                  | Decolorized                                   |
### Table II. Changes in color observed for Activated Carbon prepared by Double-burn technique

| Reagent Used | Concentration of Methylene Blue (ppm) | Observation (change of color of methylene blue) |
|--------------|--------------------------------------|-----------------------------------------------|
| HCl          | 62.5                                 | Insignificant change                           |
| HCl          | 125                                  | No change                                      |
| HCl          | 187.5                                | No change                                      |
| HCl          | 250                                  | No change                                      |
| ZnCl₂        | 62.5                                 | Insignificant change                           |
| ZnCl₂        | 125                                  | Insignificant change                           |
| ZnCl₂        | 187.5                                | No change                                      |
| ZnCl₂        | 250                                  | No change                                      |
| H₃PO₄        | 62.5                                 | Almost decolorized                             |
| H₃PO₄        | 125                                  | Slight change                                  |
| H₃PO₄        | 187.5                                | No change                                      |
| H₃PO₄        | 250                                  | No change                                      |

From the observations in Table I and Table II it can be inferred that Single-burn technique is a more convenient technique than the Double-burn technique since the single burn phosphoric acid method decolorized the highest concentration of methylene blue solution.

### D. BET Surface Area and Pore Volume Determination:

Each of the samples obtained for the three reagents were subjected to tests in a BET Surface Area Analyzer (model Smart Sorb 92/93) after all moisture was removed in a regenerator. The results obtained clearly indicate that Single-burn technique was the more reliable technique and H₃PO₄ was the most efficient activation agent in the activation of carbon prepared from coconut shells.

#### Table III. Surface Area and Pore Volume of Activated Carbon prepared by Single-burn technique

| Activation agent | Surface Area (m²/gm) | Pore Volume (cc/gm) |
|------------------|----------------------|---------------------|
| HCl              | 22.00                | 0.0212              |
| ZnCl₂            | 250.81               | 0.1501              |
| H₃PO₄            | 779.29               | 0.4234              |

#### Table IV. Double-burn technique

| Activation agent | Surface Area (m²/gm) | Pore Volume (cc/gm) |
|------------------|----------------------|---------------------|
| HCl              | 112.61               | 0.0450              |
| ZnCl₂            | 112.04               | 0.0580              |
| H₃PO₄            | 13.55                | 0.0098              |

The determination of the specific surface area and pore volume further signifies without doubt the fact that single-burn phosphoric acid activation is the most reliable technique as adsorption capacity is directly proportional to the pore volume and the specific surface area.

If a line of best fit is plotted for BET Specific Surface area against Pore Volume for all the three activation agents, it turns out to be a straight line for both the Single-Burn and Double-Burn methods. (Figure 1 and 2)
As seen for the two plots (Figure 4 and 5) the higher values of both Pore Volume and Specific Surface Area are obtained for the Single-burn samples of Activated carbon and considering the case of Phosphoric Acid, Single-burn Pore Volume and Specific Surface Area is several times more than that of the Double-burn Phosphoric acid Activated carbon samples.

IV. METHYLENE BLUE VALUE OF THE ACTIVATED CARBON SAMPLES

The color change of the methylene blue solution in the methylene blue adsorption tests gives an indirect indication of the adsorption capacity of the activated carbon samples. The Methylene Blue value is the milligrams of methylene blue adsorbed by each gram of activated carbon. Thus the concentration of the solution after 24 hours of shaking can be used to estimate the range of methylene blue value for each of the samples.

For the Single-burn samples, HCl activated carbon has a methylene blue (M.B) value of less than 25, ZnCl₂ activated carbon has an M.B value in the range of 25-50, while the H₃PO₄ has a M.B value far superior to the others, i.e., well above 100.

For the Double-burn samples, HCl and ZnCl₂ activated carbon have an M.B value far below than 25, and H₃PO₄ activated carbon has a M.B value just around 25.

V. CALCULATION OF METHYLENE BLUE VALUES

The reason for choosing Methylene blue concentrations of 62.5ppm, 125ppm, 187.5ppm and 250ppm and using 100ml of each solution for a fixed mass of 0.25g of Activated carbon can be mathematically demonstrated with the example –

Suppose for a 187.5ppm Methylene blue solution,

\[ 187.5ppm = 187.5mg \text{ in } 1000ml \]

Or,\[ 187.5ppm = 18.75mg \text{ in } 100ml \]

If the 0.25g of mass activated carbon adsorbs the entire colour of the solution, thus 0.25g adsorbs 18.75mg methylene blue. Hence 1g of activated carbon will adsorb \((18.75 \times 4 = 75)\) mg methylene blue, or the adsorption capacity in terms of M.B value will be 75.

Similarly for 62.5ppm, 125ppm and 250ppm Methylene blue concentrations, the corresponding adsorption capacities as M.B value would be 25, 50 and 100.
It was decided to prepare more closely spaced Methylene blue concentrations of solutions once the best activation agent along with the best activation technique (Single-burn or Double-burn) would be discovered and the range of values for its Methylene blue value would be known.

VI. STANDARD CALIBRATION CURVE AND EXACT METHYLENE BLUE VALUE DETERMINATION FOR H₃PO₄ ACTIVATED SAMPLE

Since the Single-burn, H₃PO₄ activated sample has not only the highest BET Specific Surface Area and Pore Volume, but also decolorizes the higher concentrations of methylene blue solutions, quantitative tests were carried out to determine the exact adsorption capacity of the sample in terms of methylene blue value.

Since the Methylene blue value of activated carbon prepared using H₃PO₄ as the activation agent ranges above 100, a 1000ppm stock solution of Methylene blue was used to prepare 7 different stock solutions of concentrations 200ppm, 250ppm, 300ppm, 325ppm, 335ppm, 350ppm and 400ppm. Between 300ppm and 350ppm, 325ppm and 335ppm were selected to determine more accurately the M.B value.

The absorbance of the methylene blue solutions were measured in a UV Spectrometer at a set wavelength of 663nm and the calibration curve of absorbance versus concentration of methylene blue solutions was plotted as shown in Figure 5. For the linearization of the standard calibration curve (Figure 5), the equation of the curve is given by:

$$A = 0.1403C + 0.0835$$

Where, $A$ is the absorbance as seen in the UV Spectrometer, and $C$ the concentration of Methylene blue in ppm.

The standard Methylene blue solutions were then taken and 0.2g of Activated carbon prepared by the Single-burn H₃PO₄ activation technique and put into 100ml of solution in conical flasks, of each of the 7 concentrations specified above. The solutions were then stoppered an kept in the rotary shaker for a period of 24 hours.

With the naked eye almost complete decolourization was observed for the 200ppm, 250ppm, and 300ppm Methylene blue solutions. The adsorbed solutions of 7 different concentrations were again subjected to absorbance tests using the UV Spectrometer to determine the exact Methylene blue value. The change in colour of the M.B solutions after being shaken with Activated carbon can be depicted in the absorbance versus concentration curve (Figure 6).
VII. STUDY OF BATCH ADSORPTION TO IDENTIFY ADSORPTION ISOTHERM

The Freundlich Isotherm has a general equation given as:

\[ q_e = K_f C_e^{1/n} \]

where, \( n \) is a Freundlich constant related to adsorption efficiency and energy of adsorption, \( K_f \) is a Freundlich constant measuring adsorption capacity, \( q_e \) is amount of adsorbate adsorbed per unit weight of carbon.

The Langmuir Isotherm\(^7\) has a general equation given as:

\[ \frac{(C_e/q_e)}{(1/Q_o)+(C_e/Q_o)} \]

where \( Q_o \) and \( b \) are Langmuir constants related to adsorption capacity and energy of adsorption respectively\(^7\).

The experiment was carried out using 0.1N oxalic acid as adsorbate\(^8\). 0.1 N NaOH was standardized after titrating it with oxalic acid\(^9\). 5 flasks with 1, 2, 3, 4, 5 grams of the single-burn H₃PO₄ and 150ml of oxalic acid in each was kept in the rotary shaker for 1 hour at 200rpm, the solutions were filtered and 10ml of each solution was titrated against the NaOH.

Table 5. Experimental data to verify the adsorption isotherms

| Amount of AC added (W g) | Titre Value of NaOH (V* ml) | Normality of unadsorbed acid (N) | Initial concentration of oxalic acid (C₀ g/cm³) | Final concentration of oxalic acid (Cₑ g/cm³) | Percentage adsorption | qₑ (g/g) |
|--------------------------|-----------------------------|---------------------------------|-----------------------------------------------|-----------------------------------------------|-----------------------|----------|
| 1                        | 10.4                        | 0.098072                        | 0.0063                                        | 0.006179                                      | 1.928                 | 0.01822  |
| 2                        | 10.2                        | 0.096186                        | 0.0063                                        | 0.006060                                      | 3.814                 | 0.01802  |
| 3                        | 10.0                        | 0.094300                        | 0.0063                                        | 0.005941                                      | 5.700                 | 0.01795  |
| 4                        | 9.8                         | 0.092414                        | 0.0063                                        | 0.005822                                      | 7.586                 | 0.01792  |
| 5                        | 9.6                         | 0.090528                        | 0.0063                                        | 0.005703                                      | 9.472                 | 0.01790  |

Where, Percentage adsorption = \( \frac{(C₀-Cₑ)}{C₀} \times 100 \),
Amount of oxalic acid adsorbed onto unit gram of activated carbon\(^6\), \( qₑ = \frac{(C₀-Cₑ)}{V} \times 100 \)

\( V = (\text{Vol. of Oxalic acid used} / \text{Amount of Activated Charcoal taken}) \) for shaking

The Freundlich constants obtained are \( K_f = 0.0507 \) and \( n = 4.9456 \)

The Langmuir Isotherm constants obtained are \( Q_o = 44.843 \) and \( b = 15.928 \)

![Freundlich Isotherm](image)

\[ y = 0.202x - 1.294 \]
\[ R² = 0.806 \]

Figure 8. Linearized plot of log \( qₑ \) vs log \( Cₑ \) (Freundlich Isotherm)
VIII. ASH CONTENT

Since ash is a factor that undermines the adsorption capacity of activated carbon, the ash content of the Single-burn and Double burn samples exhibiting the highest surface area were found. The HCl activated carbon was chosen for the double-burn technique and the H3PO4 activated carbon sample for the single-burn technique.

Crucibles were taken and weighed before and after putting in the samples. Then they were placed in a muffle furnace and heated in the open atmosphere at 900°C for a period of 30 minutes for complete combustion of the combustible matter\(^4\). The mass of crucibles were then weighed and the Ash content found out using the relation –

\[
\text{Ash Content}^{[4]} = \left( \frac{(M_{pb} - M_{ec})}{(M_{bb} - M_{ec})} \right) \times 100
\]

where, \(M_{pb}\) – mass of crucible + carbon post burning
\(M_{ec}\) – mass of empty crucible
\(M_{bb}\) – mass of crucible + carbon before burning

The Ash content was found to be 17.3% of the total mass for the Double burn sample and 1.686% of the total mass for the Single burn sample.

IX. BULK DENSITY

The Bulk Density is a significant indicator of the packaging consistency of the powdered sample in terms of weight per volume. The single-burn H3PO4 activated carbon samples were placed till the 5ml mark in a dry 10ml measuring cylinder whose mass was weighed before and after placing the sample and drying it. The activated carbon was then placed in an oven for 60minutes at 100°C and the mass was measured\(^4\). The Bulk Density was calculated as:

\[
\text{Bulk Density}^{[4]} = \frac{(\text{Mass of A.C + Cylinder after drying} - \text{Mass of empty cylinder})}{\text{(Volume occupied)}}
\]

\[
= \frac{(67.47 - 64.529)}{5} = 0.5882 \text{ g/cm}^3
\]

X. RESULTS AND DISCUSSION:

The comparison column chart in Figure10 below can be used to conclude what has been stated above. The original Methylene blue absorbance has been depicted to the left of each concentration abscissa while the Methylene blue absorbance after shaking with Activated carbon is to the right of each concentration.
Thus it can be concluded that amidst the 3 different activating agents tested under two different types of carbonization, the best Activated carbon has been produced using H₃PO₄ as the activating agent in the Single-burn technique.

The Single-burn technique is more viable as it is not only economic and requires lesser expenditure of energy but also allows for less of undesirable ash content, as it has been the case using Double-burn technique. The Single-burn technique using an inert atmosphere allows for larger pores to develop, thus increasing the Pore Volume and Specific Surface Area.

As the absorbance of tap water was around 0.1, it may be safely concluded that the Methylene blue value of the H₃PO₄ activated carbon using the Single-burn carbonization technique lies between 150-155.

Table 6. Summary of parameters and properties for the H₃PO₄ Single-burn activated carbon

| Parameter                                      | Value               |
|------------------------------------------------|---------------------|
| Size of coconut shells post crushing           | 3-5mm               |
| Duration of Chemical Impregnation              | 24 hours            |
| Concentration of acid used                     | 25%                 |
| Activation temperature                         | 450°C               |
| Activation duration                            | 30 minutes          |
| Duration of drying                             | 24 hours            |
| pH of final filtrate                           | 6.8                 |
| Particle size                                  | <125μ                |
| BET Specific Surface Area                      | 779.29 m²/gm        |
| Pore Volume                                    | 0.3968 cm³/gm       |
| Bulk Density                                   | 0.5882 g/cm³        |
| Ash content                                    | 1.686% by mass      |
| Methylene Blue Value                           | 150-155             |

We can thus conclude that among the three activation agents, namely Hydrochloric Acid, Phosphoric Acid and Zinc Chloride, Phosphoric acid is the best one in all the parameters tested for such as Methylene blue value, Specific Surface Area, Pore Volume and Ash Content.

The adsorption follows the Langmuir Isotherm as the correlation coefficient of 0.9838 is close to 1 as compared to 0.8061 for the Freundlich Isotherm. In our experiment the value of n (4.9456), in Freundlich Isotherm being above 1 signifies good adsorption⁸.

In all cases, adjustment of pH after filtration is a necessary factor in order to subject the samples to Methylene blue tests and get positive results because Methylene blue itself being acidic repels H⁺ ions in activated carbon, which occurs due to the chemical impregnation by acid. Hence the pH must be brought to the range of 5.8 to 7.0 either by repeated water washing and filtration or water wash by addition of NaHCO₃ which is alkaline and filtration followed by filtration.
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