Optimization of the Activated Carbon Preparation from Avocado Seeds, using the Response Surface Methodology

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Avocado seeds based activated carbon was prepared using chemical activation method which consisted of potassium hydroxide treatment. The main factors influencing the preparation of activated carbons at the calcination temperature, the concentration of the activating agent and the duration of calcination were investigated. One used as a mathematical model the response surface methodology to correlate the response. The significant factors identified by the analysis of variance (ANOVA) through the t-test, the Pareto diagram and the diagrams of surfaces. The optimum avocado seeds based activated carbon was obtained by using calcination temperature of 450°C, concentration of activating agent of 0.3 mol/L and time of calcination of 3.0h, which resulted to an avocado seed based activated carbon iodine number remove of 1142.1 mg/g and yield of 75.09 %, by mass. The best activated carbon obtained under the previous conditions and the raw biomass was characterized by Fourier transform infrared and Scanning Electronic Microscope.

Keywords: activated carbon, avocado seeds, central composite design, optimization

In recent years, solid waste management ranked equal to water and air pollution as the most intricate environmental turmoil in Cameroon. With circular economy legislation being introduced in Cameroon, desired direction of solid waste management was oriented towards waste minimization, resource and materials reuse. All strategies adopted were aimed to decrease quantity of solid wastes needing disposal and create available products. Cameroon as one of the largest agricultural country, its vast agricultural wastes were produced annually. However, these wastes were underutilized and often burned in open field, thus causing heavily environmental pollution. The recovery of waste is of particular interest to the extent that it allows the disposal of waste by obtaining quality products economically[1]. Accordingly, the use of agricultural wastes for fabrication of value-added materials seems very attractive and promising from environmentally and economically viable view.

Activated carbon (AC), as a well-known multi-porous material, was widely used in various fields, such as separation and concentration of useful or harmful components from mixed liquids or gases, and catalyst support [2-7]. However, the manufacture remove costs of commercial AC are in fact rather high [8, 9]. As such, it is urgent to produce AC from cheaper, renewable and readily available materials. Recently, some agricultural wastes, such as rambutan peel [2], oil palm fiber [3], bamboo waste [4], cotton stalks [5], mango-steen peel [10] and artichoke leaves [11] were commonly used to prepare AC.

In the literature, physical activation and chemical activation are the two methods used to produce activated carbons [6, 12, and 13]. Based on previous works on these methods of preparation, chemical activation has been chosen as the method to prepare activated carbon in this work. Its advantages are simplicity, shorter production time, lower activation temperature, good development of the porous structure and higher yield [14-15]. In the chemical activation process, both carbonization and activation steps precedes simultaneously by carrying out thermal decomposition of the raw material impregnated with an appropriate chemical activating agent, such as phosphoric acid in an inert atmosphere [17,18]. These impregnating materials are used as dehydrating agents and oxidants that influence pyrolytic decomposition and inhibit the formation of tar, thus enhancing the yield of activated carbon [15, 19]. The quality of activated carbon obtained depends mainly on the precursor-material used and the preparation conditions [20, 21].

To investigate the effect of some factors on the production of activated carbons, a number of the factors are kept fixed at a certain level, while varying one other so as to determine the best condition for this parameter. Thedisadvantage of this method is that, there is a lack of research on the interactive effects of these factors studied, and that, there is a large number of experiments required, which consequently requires more time [22,23]. The evaluation of the interactions between these factors is essential in determining the characteristics of the prepared activated carbons.

The response surface methodology made it possible to study several factors simultaneously (temperature of pyrolysis, concentration of activating agent and the duration of the stay in the furnace) intervening in the preparation of the activated carbon by chemical process of activation. The aim of the study is of double interest. First of all, it is a question of optimizing the factors influencing the method of preparation of the activated carbon by chemical activation with the Potassium hydroxide. In the second place, to develop the local biomasses by using them like precursors of the activated carbon, considering the annual production of co-products agricultural like the lawyer cores, remove cocopods and the peanut shell in the under-area Central Africa is about thousands of tonnes.

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Experimental part
Preparation of activated carbon
The avocado seeds were used as raw materials to produce activated carbons by chemical activation with Potassium Hydroxide followed by carbonization. The seeds were obtained from the local Menoua Subdivision, in the West Region, in Cameroon. Raw materials were initially washed twice with pure water to remove dust and subsequently dried at 105°C for 24 h to remove moisture content. Then, dried raw material was ground and sieved to size of 1-2 mm. Afterwards, 10 g sieved particles were selected to mix with Potassium Hydroxide pellets with one impregnation ratio (IR) 1: 10 and (50 ml) deionised water was added to dissolve all Potassium Hydroxide pellets with different concentrations.

The IR was estimated from equation 1:

\[ \text{IR} = \frac{\text{mass of KOH}}{\text{mass of raw material}} \] (1)

The mixture was left overnight at room temperature and then dried at 110°C for 24 h. The dried material was set on a ceramic boat which was then inserted in a stainless tube and pyrolysis in furnace and heating rate of 5°C min\(^{-1}\). Temperatures and times were based on design of RSM (Response Surface Methodology) shown in Table 1. After pyrolysis, furnace was cooled to room temperature. Afterwards, obtained samples were washed with 1 mol/L HCl, and then with distilled water to make effluent pH close to 7. Then, AC samples were dried at 120°C for 24 h, sieved and stored in plastic containers for measurement.

Experimental part
The preparation of activated carbon was optimized using response surface methodology. A central composite design methodology was employed to describe and optimize this preparation process. Three independent variables were used in this study: the temperature, concentration and time of calcination. There are several types of use plans for optimizing a response surface such as: central Composite design, Three-Level Factorial design, Box- Behnken design and Draper-Lin design. In this work, the authors’ choice was focused on the central Composite design of making benefits. The ease of using experimental data, the simultaneous variation between several input factors, possibility of determining the regression coefficient for a second-order equation and the interpolation of the response from the factors [24]. Applying this method, 17 complex experiments were performed, with two repetitions.

\[ X_1, X_2, \text{and } X_3 \text{ represent the quantitative factors that influence carbon preparation. The ranges of three factors to be evaluated were: } 450 \leq X_1, 600 \text{°C; } 0.3 \leq X_2, 0.06 \text{ mol/L; and } 1 \leq X_3, 3 \text{ h.} \]

Response \( Y \) values were yield (\( Y_1 \% \)) and iodine number (\( Y_2 \)) of AC. In the optimization process based on the response surface methodology, obtaining model equations for an input factor response is an important step [24]. The iodine number and yield (\( Y \)) responses are analysed, and one used a developed, model which correlates the response to the three variables, using a polynomial equation given by formula 2 [24]:

\[ Y = I + ax_1 + bx_2 + cx_3 + dx_1^2 + ex_1x_2 + fx_1x_3 + gx_2^2 + hx_2x_3 + ix_3^2 + e \]

(2)

where:
- \( Y \) is the measured response,
- \( I \) a constant,
- \( a, b, \text{and } c \) - linear coefficients,
- \( d, g, \text{and } i \) - quadratic coefficients,
- \( e, f \) and \( h \) - less interaction coefficient,

\( X_1, X_2, X_3, X_1^2, X_2^2, X_3^2 \) - the coded values of the activated carbons preparation variables.

\( e \) - the error.

The experimental data were analysed using a statistical software design expert named StatgraphicPlus 5, for regression analysis to fit the equations developed and also to evaluate the statistical significance of the equations obtained.

Validation of the model
It is important to validate the empirical model obtained and to do that. The exactitude of the model has to be proven by comparing the responses of the experimental values obtained with the values obtained during manipulation with those obtained from the mathematical model, the t-test, and the values of the probability \( p \). All make it possible to determine the significant degree of each coefficient. If the value of the t-test is large and the value of \( p \) low, then the coefficient is more significant [25]. In addition to that, the coefficient of determination \( R^2 \) was evaluated and for an \( R^2 \) value greater than 0.75, one considered the model accepted. For an easy calculation, input factors are usually coded with -1, representing the smallest value, 0 being the centre of the domain and 1 being the largest input value [25].

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Coded Level</th>
<th>Actual Level</th>
</tr>
</thead>
<tbody>
<tr>
<td>Temperature ( X_1 ) (°C)</td>
<td>-1</td>
<td>450.0</td>
</tr>
<tr>
<td>Concentration ( X_2 ) (mol/L)</td>
<td>-1</td>
<td>0.3</td>
</tr>
<tr>
<td>Time ( X_3 ) (h)</td>
<td>-1</td>
<td>1.0</td>
</tr>
</tbody>
</table>

Measurement of iodine number and yield of activated carbon
The determination of yield in this work allows us to estimate the amounts of activated carbon that can be obtained from an initial amount of dry bio-mass. The activated carbon yield \( Y_1 \) was calculated using the formula 3 [28]:

\[ \text{Efficiency} (\%) = \left( \frac{\text{initial mass-final mass}}{\text{initial mass}} \right) \times 100 \] (3)

The characterization of the activated carbon by the iodine number gives an idea of its micro porosity. The experiment consists in putting the activated carbon in contact with a solution of known concentration of iodine [13]. The activated carbon thanks to its pores fixes the molecules of the iodine (di-iodine) during this contact. To determine this iodine number, one puts in a 100 mL Erlenmeyer flask, weighed using a precision balance, 0.1 g of activated carbon. One added 30 mL of a 0.02N (normal) iodine solution. According to the standards established by the American Society for Testing and Materials (ASTM) [14], the mixture was stirred for 3 h and filtered with Whatman N° 1 filter paper. Afterwards, 10 mL of the filtrate was titrated with 0.005 N sodium thiosulfate solution.

The iodine number \( Q \) is obtained from the relation 4 [14]:

\[ Q = \frac{(C_0 - C_f)V}{M} \] (4)

where:
- \( M \) (g) is the mass of the activated carbon,
- \( V \) (ml) - the volume of the sodium thio-sulphate solution at the equivalence point,
- \( C_0 \) - the initial concentration,
Cț the equilibrium concentration.

Results and discussions

The main constituents of biomass

The purpose of this analysis was to determine the content of the main constituents of biomass. One used different methods.

X-ray fluorescence

X-ray fluorescence spectrometers allow to present trace elements present in biomass. Table 2 gives the expressed composition in mg/kg dry matter.

<table>
<thead>
<tr>
<th>Element</th>
<th>Avocado seeds (mg/kg dry matter)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Zirconium, Zr</td>
<td>7081</td>
</tr>
<tr>
<td>Lead, Pb</td>
<td>1396</td>
</tr>
<tr>
<td>Zinc, Zn</td>
<td>2470</td>
</tr>
<tr>
<td>Copper, Cu</td>
<td>92</td>
</tr>
<tr>
<td>Iron, Fe</td>
<td>763</td>
</tr>
<tr>
<td>Manganese, Mn</td>
<td>-</td>
</tr>
<tr>
<td>Calcium, Ca</td>
<td>-</td>
</tr>
<tr>
<td>Potassium, K</td>
<td>32.3 - 10^3</td>
</tr>
<tr>
<td>Chromium, Cr</td>
<td>-</td>
</tr>
</tbody>
</table>

Table 2 COMPOSITION IN mg/kg DRY MATTER

Table 3 gives the main components of avocado seed, obtained according to standard protocols [26].

Table 3 THE MAIN COMPONENTS OF THE BIOMASS (AVOCADO SEED)

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Content (%) by biomass, dry matter</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total solid</td>
<td>84.28</td>
</tr>
<tr>
<td>Volatile solid</td>
<td>63.22</td>
</tr>
<tr>
<td>Total moisture</td>
<td>15.48</td>
</tr>
</tbody>
</table>

Optimization of activation temperature, time and concentration of agent activating by response surface methodology (RSM)

Statistical experimental design is an efficient way to improve experimental works, has been widely used in chemistry, food and environmental engineering [13-12].

Among these design methods, RSM is considered as a powerful technique for testing multiple process variables and identifying interactions between these variables, and a combination of factors generating an optimal response can be identified by this technique [13]. Besides the quality of the raw materials, external parameters of activation temperature, calcination time and concentration play key roles in AC preparation, accordingly, to determine optimal preparation conditions. RSM experiment was applied and yield (Y1) and iodine number (Y2) were analysed as response values. Each factor has three levels and corresponding 17 experiments were designed. Each experiment was carried out in duplicate and repeated twice and the average of experimental results was used.

Optimization of iodine number and yield of avocado seed

Table 4 shows the complete design matrices together with both the response values obtained from the experimental work. The predicted values of responses were obtained from quadratic model fitting techniques using the software design expert.

The negative difference shows that the experimental value is lower than the real one.

The direct observation of the results allows affirming that the yields vary (55.12 and 77.72 % by mass and 342.63 mg/g and 1203.01 mg/g) for the iodine number. The highest values of the yields are obtained when the temperature (450°C) and the carbonization time (1 h) are low, while the best yields are obtained when the temperature (525°C) and the calcination time (3h) are high. This variation can be explained by the fact that an increase in temperature facilitates the breaking of the bonds of the biomass thus facilitating the release of the volatile matter and consequently causes a gradual decrease in the mass of the compound.

For the iodine number, the best values are temperature (450°C), concentration of the activating (0.6 mol/L) and the duration of calcination (3h), while low values are obtained at temperature (525°C), concentration (0.4 mol/L) and duration/time (3h). Indeed, one can say that the decrease of the temperature with the increase of the concentration of the activating causes an opening and a widening of the pores and consequently an increase of the adsorption of the molecules of iodine.

<table>
<thead>
<tr>
<th>Run</th>
<th>Factors</th>
<th>Y1 (%)</th>
<th>Y2 (mg/g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>No</td>
<td>X1 (°C)</td>
<td>X2 (mol/L)</td>
<td>X3 (h)</td>
</tr>
<tr>
<td>1</td>
<td>525.0</td>
<td>0.45</td>
<td>3.0</td>
</tr>
<tr>
<td>2</td>
<td>450.0</td>
<td>0.5</td>
<td>2.0</td>
</tr>
<tr>
<td>3</td>
<td>325.0</td>
<td>0.5</td>
<td>2.0</td>
</tr>
<tr>
<td>4</td>
<td>450.0</td>
<td>0.6</td>
<td>1.0</td>
</tr>
<tr>
<td>5</td>
<td>525.0</td>
<td>0.6</td>
<td>2.0</td>
</tr>
<tr>
<td>6</td>
<td>525.0</td>
<td>0.7</td>
<td>2.0</td>
</tr>
<tr>
<td>7</td>
<td>525.0</td>
<td>0.8</td>
<td>2.0</td>
</tr>
<tr>
<td>8</td>
<td>525.0</td>
<td>0.9</td>
<td>2.0</td>
</tr>
<tr>
<td>9</td>
<td>600.0</td>
<td>0.1</td>
<td>1.0</td>
</tr>
<tr>
<td>10</td>
<td>600.0</td>
<td>0.5</td>
<td>2.0</td>
</tr>
<tr>
<td>11</td>
<td>600.0</td>
<td>0.6</td>
<td>3.0</td>
</tr>
<tr>
<td>12</td>
<td>450.0</td>
<td>0.45</td>
<td>2.0</td>
</tr>
<tr>
<td>13</td>
<td>525.0</td>
<td>0.45</td>
<td>2.0</td>
</tr>
<tr>
<td>14</td>
<td>600.0</td>
<td>0.6</td>
<td>1.0</td>
</tr>
<tr>
<td>15</td>
<td>525.0</td>
<td>0.45</td>
<td>2.0</td>
</tr>
<tr>
<td>16</td>
<td>450.0</td>
<td>0.5</td>
<td>1.0</td>
</tr>
<tr>
<td>17</td>
<td>450.0</td>
<td>0.6</td>
<td>3.0</td>
</tr>
</tbody>
</table>

*Exp. Value = experimental value ** Pre. Value = predicted value.

*Exp. Value = experimental value ** Pre. Value = predicted value.

Table 4 EXPERIMENTAL DESIGN MATRIX AND RESULTS FOR PREPARATION OF AVOCADO SEED
Statistical analysis

Results concerning the probabilities of each of the factors and their interactions as a function of each of the responses using the analysis of variance (ANOVA) are shown in Table 5. Factors are more significant when the confidence interval is 95%, that is, a probability less than or equal to 0.05.

Regression coefficients for Y1 and Y2 are shown in Table 5. Thus, all regression equations of iodine number and AC yield can be achieved according to these coefficients. Correlation coefficient of regression models of iodine number and AC yield were 92.97% and 96.36%, which meant a high fitting accuracy. There is only a 0.01% chance in both models occurring due to noise. Moreover, values of Prob>F, less than 0.0500, indicate that model terms are significant. In this table, we observe that: A, B, C, A², AC, BC are significant for AC yield and factors A2, AB, B2 and C2 are significant for the iodine number. From the statistical results obtained, it was shown that the models were adequate to predict the avocado seed based activated carbon iodine number and yield, within the range of variables studied.

Accordingly, it was concluded that activation temperature, calcination time and concentration of agent activate influenced AC preparation significantly.

Mathematical modelling of the responses

Avocado seeds based activated carbon yield (Y1) and iodine number (Y2) in terms of coded factors, are represented by relations (5) and (6), respectively:

\[
Y_1 = 277.726 + 1.40821A - 197.82B + 
+ 11.0465C - 0.00125778A^2 + 0.0644AB - 0.036933AC + 
+ 126.44B^2 + 13.75BC + 1.61C^2
\]  

(5)

\[
Y_2 = 12286.1 - 35.9853A - 9831.44B - 222.146C + 
+ 0.0318408A^2 + 7.65678AB - 0.602683AC + 7153.54B^2 - 
- 143.975BC + 159.255C^2
\]  

(6)

where:

Y1 and Y2 are yield and iodine number respectively; A—Temperature, B—Concentration, C—Time.

The negative signs before the coefficients of independent and interaction factors indicate that they decrease the responses. Figure 1 shows the main effect plots of the factors on the responses.

It can be observed that the iodine number increases as the concentration and time increases and decreases with

<table>
<thead>
<tr>
<th>Source</th>
<th>Df</th>
<th>Y1 (Yield)</th>
<th>Y2 (Iodine number)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>SS</td>
<td>F-value</td>
</tr>
<tr>
<td>A</td>
<td>1</td>
<td>149.073</td>
<td>31.63</td>
</tr>
<tr>
<td>B</td>
<td>1</td>
<td>34.7071</td>
<td>7.36</td>
</tr>
<tr>
<td>C</td>
<td>1</td>
<td>183.527</td>
<td>38.94</td>
</tr>
<tr>
<td>AA</td>
<td>1</td>
<td>127.492</td>
<td>26.02</td>
</tr>
<tr>
<td>AB</td>
<td>1</td>
<td>7.1442</td>
<td>1.52</td>
</tr>
<tr>
<td>AC</td>
<td>1</td>
<td>61.3832</td>
<td>11.02</td>
</tr>
<tr>
<td>BB</td>
<td>1</td>
<td>20.2931</td>
<td>4.50</td>
</tr>
<tr>
<td>BC</td>
<td>1</td>
<td>34.0313</td>
<td>7.22</td>
</tr>
<tr>
<td>CC</td>
<td>1</td>
<td>6.4985</td>
<td>1.38</td>
</tr>
</tbody>
</table>

Table 5
ANALYSIS OF VARIANCE (ANOVA) AS RESPONSE SURFACE QUADRATIC MODEL FOR AC YIELD AND IODINE NUMBER

R² = 96.36 %; R² adjusted = 91.67 %; R² = 92.9742 %; R² adjusted = 83.941%
SS = Sum of Squares; * significant value; F-value = Fisher value; P-value = probability value.
Df = degree of freedom

Fig. 1. Main effect plots of the factors on the responses
temperature. Concerning the effect of temperature on the yield, we can observe that increasing the temperature increases the yield. Also, it results that by increasing the time, the yield increases.

**Optimize response**

The optimization method makes the location of an extreme point on the surface[24] possible. Based on achieving relative maximum iodine number and AC yield, optimal preparation conditions of activation temperature, activation time and concentration of agent activate were calculated by design-Expert software version 5. Under conditions of 450°C, 3 h and 0.3 mol/L, iodine number reached the relatively highest values of 1092.18mg/g and AC yield 82.59% for 538.78°C, 3 h and 0.6 mol/L respectively. Furthermore, to test accuracy of the value achieved by above model, optimized experimental conditions for AC preparation were carried out.

**Response surface**

The fitted response surface plot was generated by statistically significant above model design Expert program to understand the interaction of the parameters required for optimum iodine number and AC yield. At each point of the domain corresponds a response. The value that a response can take within the field of study can be predicted. The diagrams of the response surfaces thus offer the possibility to see the evolution of the different responses. One can thus note the synergistic effect of all the factors on the different responses.

**Iodine number**

Figure 2 (a and b) shows the three-dimensional response surfaces which were constructed to show the interaction effects. Figure 2 c shows the Pareto Chart for the iodine number. These diagrams allow identifying the combination of input variable parameters that jointly optimize a single response or set of responses. They also allow evaluating the impact of the activating agent, the calcination temperature and the residence time in the furnace on the iodine number and AC yield.

As it can be seen from figure 2 (a, b and c), that the iodine number increased with increased activation temperature, and decreases with increased concentration of the activating agent. The values have an effect on the porosity of the activated carbon [10]. The synergistic effect of the various factors on the response is thus verified.

**Activated carbon yield**

Figure 3 shows the three-dimensional response surface plot of AC yield; (a) effect of concentration and activation temperature, time = 3h, (b) effect of concentration and activation time, T=535.74°C.

From the observation of these figures and in agreement with the results of values presented in table 5, one can
confirm that the temperature and time impose the greatest effect on AC yield, followed by concentration. The interaction effects between the activation temperature and concentration, as well as activation time and concentration, were considered high.

**Optimum values**

The statistical software was used to obtain the optimum preparation parameters in relation to the responses considered. The Statgraphic Plus5 software considered the unit factor effect, two factors interaction and quadratic factors impact on the resulting models which ordinarily are tedious or impossible with maximum accuracy to execute manually. The optimum values of the responses, as well as the values of the factors that make it possible to obtain these values, are summarized in table 6. These values are the result of the superposition of the different curves of response surfaces. The area of interest is the one found in optimal conditions.

**Characterization of Activated Carbon**

Fourier Transform Infrared Spectroscopy (FTIR)

Fourier transform infrared (FTIR) transmission spectra were obtained to characterize the surface groups on the Avocado seeds and the AC prepared from these precursors. The infrared spectroscopic analysis spectra of the various adsorbents are shown in figures 4 and 5. Examination of these spectra reveals the presence of several absorption bands of valence and deformation vibrations attributable to the different groups existing in these materials. The different vibration bands identified were assigned using the results of the literature on the characterization of lignocellulosic materials [27].

The first band appears at 3330 cm⁻¹ due to the vibration of elongation of O-H functions. The bands of vibration of elongation of the C-H functions are raised between 3000-2840 cm⁻¹, with a larger intensity around 2920 cm⁻¹. One also observes the appearance of a band to 1730 cm⁻¹ corresponding to the vibration of elongation of C=O functions. The band of elongation C=C of the aromatic skeleton of lignin is also visible on these spectra; it appears around 1580 cm⁻¹. In the spectral area 1150 - 1000 cm⁻¹, a band of great intensity (centered at 1030 cm⁻¹) appears, corresponding to the vibration of elongation C-O [27].

**Scanning Electronic Microscope (SEM)**

The images of the surface morphology of the best activated carbon obtained at 450°C, 0.3 mol/L and 3h, as well as the raw biomasses are illustrated in figure 6.

These images show that the active carbon induces a consequent development of the porosity by bursting of the naturally occurring pores in the biomass or induced by activation and pyrolysis. The aim of the SEM examination is to illustrate the porosity, especially that created by activated carbon. A developed porosity makes it possible to increase the specific surface area of the coal and, consequently, the number of active sites on which may be fixed molecules of interest.

<table>
<thead>
<tr>
<th>X1 (°C)</th>
<th>X2 (mol/L)</th>
<th>X3 (h)</th>
<th>Iodine number (mg/g)</th>
<th>Yield (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Predicted</td>
<td>Experimental</td>
<td>Error (%)</td>
<td>Predicted</td>
</tr>
<tr>
<td>450</td>
<td>0.3</td>
<td>3.0</td>
<td>1092.18</td>
<td>1142.10</td>
</tr>
</tbody>
</table>

**Table 6**

MODEL VALIDATION

![Fig.4. FTIR spectra of raw avocado seeds](image)

![Fig.5. FTIR spectra of avocado seeds - based AC](image)

![Fig.6. SEM images for raw biomass(a) and activated carbon (b) (voltage 15 kV and 3000× magnification).](image)
Conclusions

The analysis of the experimental design through the response surface methodology (composite design plant) was used to study the main factors (carbonization temperature, pyrolysis duration and concentration of the activating agent) on the yield and the adsorption properties measured by the iodine number during the preparation of activated carbons. The influences of its various factors on the responses have been modelled by a quadratic model of the second order in satisfactory ways.

The optimum avocado pear based activated carbon was obtained by using activation temperature, activation time and concentration of 450°C, 3.0 h and 0.3mol/L, respectively, thus resulting an iodine number 1142.10 mg/g, and AC yield of 82.51%. The avocado pear, prepared based activated carbon, demonstrated high surface area and well-developed porosity.

Acknowledgements: The authors are grateful for the support of the research work for Mr TagnetenKam Rufs Fregue, through the University Agency of Francophonie (AUF) via the program EUGEN IONESCU 2017/2018, hosted by the Politehnica University of Timisoara (Romania). Also the PhD position of Alin-Cristian Mihiauti, at the same university, is acknowledged.

References
28. FERNANDEZ, E., Etude de la carbonisation et de l'activation de precureurs vegetaux dures et mous, PhDThese de Doctorat, (Study of carbonisation and activation of hard and soft plant precursors, PhDThesis), 2002 (Universite NEUCHATEL)