INTRODUCTION

Activated carbon has important adsorbent, peripheral, thermal, mechanical and electrical features for new practices of activated carbon [1]. It is a highly ambidextrous substance with wide surface field, wide interior surface field, pore volume, well chemical steadiness, diverse oxygen-having functional groups on the surface. Because of its perfect adsorption ability, it is extensively usaged in refinement and distinction in numerous industrial operation inclusive medical usage, gas storage, contaminant and smell rustication, gas decomposition and catalysis [2,3]. Activated carbons are manufactured from distinct crude materials of organic origin that are rich in elemental carbon, including biomass waste. The selection of the crude material for the manufacturing of activated carbon is decided by the ultimate designation of the activated carbon as well as the availability and cost of the raw material [4]. Activated carbon can be produced such as agricultural wastes, inclusive a corn crust [5,6], peanut crust [7], cotton handle [8], coal, wood, coconut shell, peat [9], almond crust [10], date palm [11], rice shell [11], hazel nut [12], walnut shells [13]. There are two significant process to manufacturing activated carbon [14]: (a) Carbonization which is pyrolysis of the pioneers in an inert atmosphere. Along this process, the organic materials carbon ingredient increases and the constricted or plugged pores are created. Carbonization warmth of cellulosic biomass is at under than 800 °C and (b) Activation is a method used to further improve pores. There are two types of chemical activation and physical activation. Throughout physical activation, a substance is carbonized in an inert atmosphere and after that activated by vapour or carbonic-acid gas to improve inner porousness. The activation temperature for cellulosic carbon is between 800 and 1000 °C [15,16]. In chemical activation, the raw sample is carbonized by mixing with activation agents like zinc chloride, phosphoric acid, sodium hydroxide and potassium hydroxide [17,18]. Chemical activation generally gets location at lesser temperature and smaller process time than physical activation [15,19]. The energy spending and the cost are also decreased by inferior temperatures [15,16]. In addition to, chemical activation keeps bigger surface field and more elevated carbon efficiency than physical activation [19]. Usually, the activation temperature, activation time, impregnation ratio, carbonization conditions, heating conditions, nitrogen flow rate and activation agents are the most important factors impressing the surface fields and pore volumes of the produced activated carbon [20,21]. In this study, activated carbon is produced from Eriobotrya japonica seed by chemical activation with ZnCl₂. The characterization of the produced activated carbons are determined by SEM, BET, FT-IR, TGA and XRD analyzes.

EXPERIMENTAL

*Corresponding author: E-mail: fevzi@kmu.edu.tr

This study reports the utilization of Eriobotrya japonica seed in activated carbon arranging by chemical activation with ZnCl₂ agent and the effect of carbonization situations surface properties. The influences of diverse activation temperatures (500, 600, 700 °C) on the pore-structure properties of activated carbons were discussed by using N₂ adsorption/desorption isotherms, scanning electron microscopy, thermogravimetric analysis, Fourier transform infrared spectroscopy and X-ray diffraction. The conclusions indicated that the carbonization temperature have favourable effect on Brunauer-Emmett-Teller (BET) surface field, aggregate pore volume. The highest Brunauer-Emmett-Teller surface area of activated carbon produced at 700 °C was 1079 m²/g, total pore volume was 0.52 cm³/g, carbonization time was 2 h and impregnation rate was 1/1. Eriobotrya japonica activated carbon (EJAC)-ZnCl₂ can be considered as an ideal low-expense and eco-friendly adsorbent.

Keywords: Eriobotrya japonica seed, Chemical activation, Activated carbon, Characterization.
spring and summer in Karaman Turkey. The *Eriobotrya japonica* seed was separated from the fruit and dried by warming at 80 °C during night. After that, the crude material was milled in grinder and screened to 20-200 mesh grain dimension by standard sieves. The obtained samples were stored in sealed containers until the experimentation.

All reagents (analytical grade AgNO₃, HCl, ZnCl₂ and H₂O) were bought from Merck. pH studies were done usage a pH meter (Model Cyberscan 510). The FT-IR spectra for functional group determination were recorded in KBr discs on a spectrophotometer (Model Perkin Elmer-400 Series). A scanning electron microscopy (SEM, HITACHI S-520) was utilizationed for scanning the adsorbent surface. The outer and micropore surface fields, sum pore, micropore capacity and pore size distribution were obtained with the Brunauer–Emmett–Teller (BET) method (Micromeritics ASAP 2020). Powder X-ray diffraction (XRD) models of the examples were registered at room temperature on the Bruker D8 Advance diffractometer handling CuKα radiation. The thermal evolution of the precursor and ZnCl₂-impregnated samples were studied by the thermogravimetric analysis (TGA-50 analyzer) from room temperature until 1000 °C at a ratio of 10 °C min⁻¹ with a N₂ flow rate of 100 mL min⁻¹. The total weight losses of the examples were detected from the corresponding TGA curves.

**Preparation of activated carbons:** The preparation of the activated carbon (AC) primarily contains carbonization and activation. The aim of carbonization is to get first pore form for activation and suitable carbon concentrate. The aim of activation is to compose new pores and widen the surface field of activated carbon. The flow scheme for the production of activated carbon is given in Fig. 1.

The *Eriobotrya japonica* powder was impregnated into the zinc chloride with rates 1/1 (10 g/10 g). 150 mL of distilled water was joined to the mixture. The blend was kept to stand at room temperature for 1 day. The mixtures were dehydrated in an oven at 110 °C throughout 24 h. Carbonization of the precursor is applied in a hollow cylindrical rust proof steel reactor of 30 cm in width. The reactor has an inlet and a gas outlet. It is warmed with a tubular oven (Model Protherm) equipped by a heat programmer controller. This system allows the management of the temperature rate, the rise level and the analyses duration of the final temperature. The examples were warmed to last temperatures of 500, 600, 700 °C with 5 °C min⁻¹ warming ratio for residence time of 2 h. Finally, temperature drop until room temperature. The activation is carried out by using nitrogen (9,880 mLmin⁻¹) during the hall experiment (Fig. 2). After cooling to room temperature, the produce activated carbon was bathed with 2 M HCl solution by warming
at about 95 °C for 0.5 h, strained and rinsed with warm pure water till neutral pH, desiccated at 110 °C in a furnace for just about 12 h and weighed to compute the yield.

### RESULTS AND DISCUSSION

**Yield and specific surface BET:** Influences of carbonization temperature on the surface field (S) and yield (%) of activated carbon were assessed below the terms of the impregnation rate of 1/1 (g/g) and carbonization period of 2 h. The carbon yield was computed based on the mass of *Eriobotrya japonica* (EJ) seed on a dry base from the following equality:

\[
\text{Carbon yield (wt %)} = \frac{\text{Weight of activated carbon (g)}}{\text{Weight of EJ seed (g)}} \times 100
\]

The most important study parameters affecting the yield of activated carbon are carbonization temperature, carbonization duration and impregnation rate. As seen from Table-1, with the rising in carbonization temperature, the yield of active carbon increased. This may be connected to the promotion of carbon flaming-off, tar evaporation and the vaporization of biomass would be increased because of rise of temperature [22,23]. The specific surface BET of activated carbons for diverse carbonization temperatures were detected by nitrogen adsorption at 77.3 K. The surface field has changed in direct proportion to the temperature. As viewed from Table-2, adsorption at 77.3 K. The surface field has changed in direct proportion to the temperature. The BET surface field of active carbon was obtained. The curves formed at low pressure show adsorption into the micropores [27]. The increase in the graph indicates the growth of the pores and the improving tendency of complex Type I and Type IV isotherms accordingly to the International Union of Pure and Applied Chemistry (IUPAC) [25].

![Graph showing N2 adsorption and desorption isotherms](image)

Pore magnitude distribution is a very important property of poriferous sorbents because the dissimilarity in the pore magnitude effects the adsorption capability in order to molecules of distinct dimensions and shape. According to the rating IUP activated carbon accepts, sorbent pores are examined in three groups as micropores (< 2 nm), mesopores (2-50 nm) and macropores (> 50 nm) [26]. Fig. 4 shows that large surface area activated carbon was obtained. The curves formed at low pressure show adsorption into the micropores [27]. The increase in the graph indicates the growth of the pores and the improving of mesopores.

![Graph showing pore size distributions](image)

**Adsorption isotherms:** The chemical properties of the adsorbent, the porous structure and the heat adsorption are determined with adsorption isotherms. The porosity of activated carbon substances is generally described by handling N2 adsorption method [24]. Fig. 3 shows the N2 adsorption/desorption isotherms at 77.3 K of the three dissimilar activated carbon examples made from *Eriobotrya japonica* seed. The isotherms created from these works were found to follow the tendency of complex Type I and Type IV isotherms accordingly to the International Union of Pure and Applied Chemistry (IUPAC) [25].

<table>
<thead>
<tr>
<th>Sample</th>
<th>Impregnation rate</th>
<th>Carbonization temperature (°C)</th>
<th>Carbonization time (hold time) (min)</th>
<th>Yield (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>AC1</td>
<td>1/1</td>
<td>500</td>
<td>120</td>
<td>42.58</td>
</tr>
<tr>
<td>AC2</td>
<td>1/1</td>
<td>600</td>
<td>120</td>
<td>43.09</td>
</tr>
<tr>
<td>AC3</td>
<td>1/1</td>
<td>700</td>
<td>120</td>
<td>43.40</td>
</tr>
</tbody>
</table>

**TABLE 2**: Structural characteristics of activated carbon

<table>
<thead>
<tr>
<th>Structural properties</th>
<th>AC1</th>
<th>AC2</th>
<th>AC3</th>
</tr>
</thead>
<tbody>
<tr>
<td>BET surface area, S_(BET) (m²/g)</td>
<td>863.3</td>
<td>ND*</td>
<td>1079</td>
</tr>
<tr>
<td>Micropore area, S_(MIC) (m²/g)</td>
<td>207.0</td>
<td>417.0</td>
<td></td>
</tr>
<tr>
<td>External surface area, S_(EXT) (m²/g)</td>
<td>656.3</td>
<td>662.2</td>
<td></td>
</tr>
<tr>
<td>Total pore volume, V т (cm³/g)</td>
<td>0.44</td>
<td>0.52</td>
<td></td>
</tr>
<tr>
<td>t-plot micropore volume, V_(MIC) (cm³/g)</td>
<td>0.09</td>
<td>0.19</td>
<td></td>
</tr>
<tr>
<td>Average pore size, Dp (nm)</td>
<td>3.31</td>
<td>3.55</td>
<td></td>
</tr>
</tbody>
</table>

*ND = None detected

**SEM analysis:** SEM views of activated carbons produced at distinct temperatures are given in Figs. 5-7.

SEM technique was used to watch the surface physical features of the activated carbon. According to Figs. 5-7: when the SEM images of 1000 and 2000 fold magnifications of each active carbon are taken, activated carbon has a porous structure. The obtained activated carbon displayed a highly porous structure with various sizes of mesopores and macropores. Micro pores were observed at this resolution. Additionally, a three dimensional interconnected and irregular mesopore structure can also be clearly observed. These pores are caused by the
Fig. 5. SEM images of activated carbon produced at 500 °C carbonization temperature

Fig. 6. SEM images of activated carbon produced at 600 °C carbonization temperature

Fig. 7. SEM images of activated carbon produced at 700°C carbonization temperature
The spectra show a number of absorption peaks defining the compound structure of the materials. According to Fig. 8, the existence of the wide and intensive absorption peak at 3320.12 cm\(^{-1}\) shows the O-H stretching oscillation of cellulose, hemicellulose, absorbed water, pectin and lignin. The peak appeared at 3005.2 cm\(^{-1}\) may be assign to the aliphatic saturated C-H tension oscillations of in lignin polysaccharides bearing cellulose and hemicellulose [29]. The peak at 1745.30 cm\(^{-1}\) display the carbonyl (C=O) tension oscillation of the carboxyl classes of pectin, hemicellulose and lignin [30]. The peak at 1634.53 cm\(^{-1}\) may be due to adsorbed water H-O-H bend. The oscillation at 1510.34 cm\(^{-1}\) and 1380.72 cm\(^{-1}\) could be owing to aliphatic and aromatic (C-H) groups in the horizontal oscillation of methyl, methoxy and methylene groups. The band at 1180.13 cm\(^{-1}\) is due to the C-O stretching oscillation of phenol group [31,32]. The band seen nearby 1634.53 cm\(^{-1}\), could be ascribe to (C=C) vibration in aromatic rings [33]. The peaks in the range of 1250-1000 cm\(^{-1}\) may belong to the C-O stretching oscillation of alcohols and carboxylic acids [32]. Many absorption peaks were not observed while the carbonization temperature increased. This is probably due to the evaporation of organic substances at high temperatures.

**Thermal analysis:** The TGA curve showing the % mass change of activated carbon produced by ZnCl\(_2\) activation in Fig. 9. When TGA curve of active carbon produced by ZnCl\(_2\) activation are taken into account, it is seen that the initial mass loss is around 8.29 % at 25-109 °C. This reduction is thought to be caused by the removal of phenolic groups from the environment. The second mass loss is about 4.18 % at 109-445 °C and the third mass loss is around 11.75 % at 445-660 °C. The total mass loss is 2.11 mg (24.22 %).

**XRD analysis:** XRD models of the activated carbon are indicated in Fig. 10. According to Fig. 4, activated carbon sample exhibit with five broad peaks corresponding to 2θ = 38.9°, 44.1°, 64.5°, 77.6° and 80.9°. When the elemental pairings of the peaks are made 2, 3 and 5th peaks are formed C-C bonds, 1th peak is formed Zn-H and 4th peak is formed H-H. The presence of many C-C pairs shows less impurities.

**Conclusion**

In this study, active carbon was produced from *Eriobotrya japonica* seed using chemical activation method with ZnCl\(_2\) agent. Activated carbon was produced at three different carbonization temperatures (500, 600, 700 °C). The highest BET surface field (1079 m\(^2\)/g), micropore surface area (417 m\(^2\)/g) and total micropore volume (0.52 cm\(^3\)/g) were obtained below the following terms: impregnation ratio of 1/1, activation temperature of 700 °C and activation time of 2 h. This suggests that ZnCl\(_2\) is a activating agent in order to producing activated carbon by an improved microporous structure. The results show that the chemical activation with ZnCl\(_2\) from *Eriobotrya japonica* seed is good in terms of specific surface field, pore improvement and the structural order of activated carbon. *Eriobotrya japonica* activated carbon-ZnCl\(_2\) can be contemplated as an excellent low-cost and eco-friendly adsorbent.

**ACKNOWLEDGEMENTS**

The authors are grateful for the financial support of the Scientific Research Project of Karamanoglu Mehmetbey University, Karaman, Turkey (Project Number 01-D-17).

**REFERENCES**


