Production of active coal from pyrolyzed wood wastes
by alkaline activation of KOH

Oleg Kuzmin¹, Julia Tamarkina², Tetiana Shendrik², Valentyna Zubkova³, Olga Koval¹, Tetiana Roman¹

¹ – National University of Food Technologies, Kyiv, Ukraine
² – L.M. Litvinenko Institute of Physical–Organic Chemistry and Coal Chemistry of the National Academy of Science of Ukraine, Kyiv, Ukraine
³ – Jan Kochanowski University in Kielce, Kielce, Poland

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Abstract

Introduction. The purpose of this publication is to evaluate an alternative renewable raw materials obtained from the food industry wastes (pyrolyzed wood wastes – PWW) as precursors for production of active coal (AC) used in a process of water purification in alcoholic beverages’ production.

Materials and methods. PWW of meat processing industry. Chemical activation of PWW by alkaline activation of KOH. Method of adsorption–desorption of nitrogen to determine a porous structure at 77 K; mesopores’ distribution by size and by mesopores’ volume – BJH-method; micropores’ division by size – QSDFT-method; volume of micropores – Dubinin-Radushkevich method; subnanopores’ volume – QSDFT-method.

Results and discussion. The microporous structure has the following characteristics: pores’ diameters are in the range of \( D_{mi}=0,61–2,5 \) nm, most represented pores’ diameters are 0,61; 1,19; 1,54 nm; volume of micropores \( V_{mi}=0,11–0,30 \) \( cm^3/g \); pores’ surface area \( S_{mi}=407–852 \) \( m^2/g \); pores’ differential volume \( dV_{mi}/dD=(0,023–1,400)\times10^{-2} \) \( cm^3/g \); pores’ differential area \( dS_{mi}/dD=(0,18–45,60) \) \( m^2/g \). There are 70,31% of micropores in a total pores’ volume. Dominant contribution of micropores into specific surface of the pores shows a proportional dependence between pores’ volume and surface area of pores. It is also confirmed by the linear dependence between the pores’ differential volume and the differential area. The smallest pores – subnanopores with \( D<1 \) nm were defined at the micropores structure. Subnanopores’ diameters are in the range of \( D_{inn}=0,61–1,00 \) nm. Subnanopores’ volume varies in the range of \( V_{inn}=0,11–0,25 \) \( cm^3/g \). Pores’ surface area is \( S_{inn}=407–783 \) \( m^2/g \); pores’ differential volume: \( dV_{inn}/dD=(11,3–140,0)\times10^{-4} \) \( cm^3/g \); differential area is \( dS_{inn}/dD=(2,33–45,60) \) \( m^2/g \). The subnanopores’ portion at the micropores’ volume is 84,12%. The share of subnanopores’ at the total pores’ volume is 59,15%. It can be argued that the alkaline activation of KOH leads to a development of subnanopores in the porous structure of the adsorbent. The cited data shows that the proposed method allows to obtain AC with an output ratio of 70,4%. The obtained AC has a developed specific surface of \( S_{BET}=777 \) \( m^2/g \) and porosity. Total pores’ volume is \( V_z=0,421 \) \( cm^3/g \).

Conclusion. An energy–saving method is proposed for the production of high porous AC from the secondary «renewable» resources – PWW. It is advised to use it in alcoholic beverages production.
Introduction

A search for economically feasible ways of obtaining cheap sorbent materials in a purification of contaminated environment remains an urgent problem for all the countries around the world. Carbon adsorbents occupy a significant place among such materials. This kind of raw material has a gigantic range of precursors (natural coal, peat, wood, carbonaceous wastes of various origins, etc.) [1–12]. Therefore, there is an urgent need to obtain AC from an alternative material. The search for these materials could involve existing technologies of food industry. Wastes from these industries can be used to produce the adsorbents [4-8, 11-26].

There are many ways of receiving AC (Kumar, Jena, 2017; Yorgun, Yildiz, 2015; Kucherenko et al, 2010; Lillo-Ródenas, 2003) [4, 11, 27, 28], (Pat. 61059 Ukraine): grinding carbon-containing material with (1–2)·10^{-3} m, mixing it with KOH in solid form in a weight ratio 1:1, carbonizing and activating at heatstroke mode, cleaning with water and drying. This method (Pat. 61059 Ukraine) has its disadvantages: raw materials grinding has a high energy consumption; the small size of raw materials’ fractions – it became charcoaled after carbonization and activation and evaporates with a gaseous components; high temperature carbonization and activation of AC; activation in a heatstroke mode causes tearing of the structure and reduction of AC shares; low rate of AC release.

The most promising raw material for AC is PWW. PWW is formed by pyrolysis of wood chips (Kuzmin, Shendrik, 2016) [10]. In a proposed method grinding materials are not required as AC wood chips’ size is 1×b×h=(6×12×3)·10^{-3} m; AC fractional increases up to 3,6·10^{-3}＞d≥1,0·10^{-3} m; temperature reduction of charcoal’s carbonization and activation within T=773–973 К; absence of activation heat stroke due to carbonization at non-isothermal heating and isothermal heating at activation; yield increases of AC ratio.

KOH is one of the promising activating agents (Kucherenko V.A. et al, 2010; M.A. Lillo-Ródenas et al, 2007) [27, 28]. It is added to the brown coal. It can withstand up to full impregnation, allowing alkali to interact with organic and mineral components, with the formation of water-soluble substances washed with AC.

Thus, the use of KOH allows to receive AC with a formed pores’ space. Variation of mass part (MP) of activating agent in relation to PWW can affect the surface pores’ factor, yield ratio of AC and volume of wastewater [29-37].

A blend of raw material/agent during carbonization and activation undergoing non-isothermal heating up to an activation temperature during the subsequent isothermal aging. At the same time low molecular parts of thermal distraction of organic matter of PWW and products of PWW’s chemical reactions with alkali are formatted in a PWW’s volume. It’s outflow from the PWW’s volume creates a spatial framework within PWW. It leads to a formation of micropores and subnanopor and, consequently, increases the pores’ specific surface area and total volume. This improves adsorption characteristics of AC (Shendrik et al, 2003; Kucherenko et al, 2010; Zubkova, 2011) [27, 29, 30]. AC’s fractional composition is determined by MP’s residue on sieves with holes with a diameter of 3,6 mm, 1,0 mm and on the pallet itself.

It has been shown that PWW can be an alternative carbon-containing raw material for AC
The aim of this work is to evaluate an alternative renewable raw materials from the food industry wastes (pyrolyzed wood wastes – PWW) as precursors for production of active coal (AC) which can be used for water purification in alcoholic beverages’ production.

**Materials and methods**

Conditions for AC production are presented at Table 1.

### Table 1

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Characteristic</th>
<th>Experimental data</th>
<th>Rationed data</th>
</tr>
</thead>
<tbody>
<tr>
<td>$T_1$</td>
<td>Drying temperature in the open air, K</td>
<td>295</td>
<td>293–298</td>
</tr>
<tr>
<td>$W_1$</td>
<td>Relative humidity, %</td>
<td>74</td>
<td>67–82</td>
</tr>
<tr>
<td>$v_1$</td>
<td>Air traffic speed, m/s</td>
<td>1.5</td>
<td>1–2</td>
</tr>
<tr>
<td>$\tau_1$</td>
<td>PWW’s outdoor drying time, s</td>
<td>336·60$^2$</td>
<td>(336–504)·60$^2$</td>
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<tr>
<td>$T_2$</td>
<td>Drying temperature in the drying cabinet, K</td>
<td>373</td>
<td>373–383</td>
</tr>
<tr>
<td>$W_2$</td>
<td>MP of moisture of PWW, %</td>
<td>6,58</td>
<td>4–8</td>
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<tr>
<td></td>
<td>MP of KOH in aqueous solution, %</td>
<td>50</td>
<td>30–70</td>
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<tr>
<td></td>
<td>Mass ratio (MR) of PWW/alkali, kg/kg</td>
<td>1:1</td>
<td>1:0.5–1:1</td>
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<tr>
<td>$\tau_2$</td>
<td>Time withstand of PWW with alkali, s</td>
<td>24·60$^2$</td>
<td>(18–24)·60$^2$</td>
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<td>$T_3$</td>
<td>PWW alkali holding temperature, K</td>
<td>294</td>
<td>291–295</td>
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<tr>
<td>$T_4$</td>
<td>Drying temperature, K</td>
<td>381</td>
<td>373–383</td>
</tr>
<tr>
<td>$W_3$</td>
<td>MP of moisture of PWW, %</td>
<td>6,78</td>
<td>4–8</td>
</tr>
<tr>
<td>$Q_1$</td>
<td>Volumetric flow of argon, m$^3$/s</td>
<td>5,6·10$^{-7}$</td>
<td>≤5,6·10$^{-7}$</td>
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<td>Non-isothermal heating, deg./s</td>
<td>0,07</td>
<td>≤0,07</td>
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<td>MP of sulfuric acid, %</td>
<td>96</td>
<td>96</td>
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<tr>
<td>$T_5$</td>
<td>Activation temperature, K</td>
<td>1073</td>
<td>873–1073</td>
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<td>$\tau_3$</td>
<td>Time of activation, s</td>
<td>1·60$^2$</td>
<td>1·60$^2$</td>
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<td>Non-isothermal cooling, deg./s</td>
<td>0,1</td>
<td>≤0,1</td>
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<tr>
<td>$T_6$</td>
<td>Final temperature after cooling of AC, K</td>
<td>323</td>
<td>≤323</td>
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<td>$\tau_4$</td>
<td>Time cleaning of AC from activating agent, s</td>
<td>600</td>
<td>300–600</td>
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<td>$T_7$</td>
<td>Drying temperature in the drying cabinet, K</td>
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<td>373–383</td>
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<td>$W_4$</td>
<td>MP moisture AC, %</td>
<td>5,84</td>
<td>4–8</td>
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<td>$Y_1$</td>
<td>AC yield ratio, %</td>
<td>70,4</td>
<td>70–80</td>
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<td>MP of remnant of AC (%) in the sieve with holes, m:</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>$d&lt;3,6·10^{-3}$</td>
<td>0,2</td>
<td>MP≤2,5</td>
</tr>
<tr>
<td></td>
<td>$3,6·10^{-3}&lt;d&lt;1,0·10^{-3}$</td>
<td>87,6</td>
<td>MP≥95,5</td>
</tr>
<tr>
<td></td>
<td>$d&lt;1,0·10^{-3}$</td>
<td>12,2</td>
<td>MP≤2,0</td>
</tr>
</tbody>
</table>

Obtained PWW is dried in the open air ($T_1$=293–298 K; $W_1$=67–82%; $v_1$=1–2 m/s) during $\tau_1$=(336–504)·60$^2$ s, followed by more drying at $T_2$=373–383 K up to air–dry state with humidity of $W_2$=4–8%.

*KOH* (MP *KOH* in aqueous solution – 30–70%) used as an activating agent for impregnating
of PWW/alkali in MR 1:0.5–1:1. The received mixture withstands for \( t_f = (18–24) \cdot 60^2 \) s at \( T_f = 291–295 \) K and PWW dried to receive a constant weight of MP moisture \( W_f = 4–8\% \) at \( T_f = 373–383 \) K. Activation carried out in a stream of argon with a volumetric flow of \( Q_1 \leq 5.6 \cdot 10^{-7} \) m\(^3\)/s with drying bubbling after 96\% in sulfuric acid under non-isothermal heating at 0.07 deg./s up to activation temperature \( T_s = 873–1073 \) K and isothermal aging for \( t_f = 1 \cdot 60^2 \) s at the temperature activation and non-isothermal cooling at 0,1 deg./s in a stream of argon up to a temperature \( T_0 = 323 \) K.

The received AC cleaned from activating agent with a usage of water for \( t_f = 300–600 \) s and dried at a temperature \( T_f = 373–383 \) K up to a level of humidity \( W_f = 4–8\% \) with the yield of AC \( Y_f = 70–80\% \), followed by fractioning with the help of MP residue on sieves with holes: \( d \geq 3.6 \cdot 10^{-3} \) m – MP\( \leq 2,5\% \); \( 3.6 \cdot 10^{-3} > d \geq 1.0 \cdot 10^{-3} \) m – MP\( \geq 95,5\% \); \( d < 1.0 \cdot 10^{-3} \) m – MP\( \leq 2,0\% \) with the following selection of working faction on a sieve with holes \( 3.6 \cdot 10^{-3} > d \geq 1.0 \cdot 10^{-3} \) [9].

Figure 1 shows the stages of AC production; Figure 2 – general scheme of AC obtaining as per experimental data at Table 1.

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**Figure 1. Stages of receiving AC:**

- a – technological chips of oak large \((6 \times 12 \times 3) \cdot 10^{-3} \) m;
- b – PWW with MP moisture \( W = 43.01\% \);
- c – PWW after drying of moisture MP \( W = 6.58\% \);
- d – AC to fractionation;
- e – AC after fractionation of \( d \geq 3.6 \cdot 10^{-3} \) m;
- f – AC after fractionation of \( 3.6 \cdot 10^{-3} > d \geq 1.0 \cdot 10^{-3} \) m;
- g – AC after fractionation of \( d < 1.0 \cdot 10^{-3} \) m
Figure 2. The general scheme of AC production as per experimental data

PWW dried for $\tau_f=336\cdot60^2$ s outdoors ($T_i=295$ K; $W_f=74\%$; $v_f=1.5$ m/s), followed by drying at the drying cabinet at $T_f=373$ K to air-dry state with MP moisture $- W_f=6.58\%$. Potassium hydroxide with MP KOH $- 50\%$ in an aqueous solution, injected by impregnation of PWW $- KOH$ and kept for $\tau_f=24\cdot60^2$ s at temperature $T_f=294$ K, dried up to a moisture obtained at MP PWW $W_f=6.78\%$ at $T_f=381$ K. The volume of solution has been chosen to create MR $PWW/alkali - 1:1$ kg/kg. Activation was performed in a vertical cylindrical tubular reactor made of steel, with thickness of 3 mm, diameter of cylinder $- 0.15$ m, height $- 0.3$ m.

The reactor was purged with argon volumetric flow of $Q=5.6\cdot10^{-7}$ m$^3$/s, drained bubbling through concentrated sulfuric acid (96%). The heating of reactor’s furnace has been switched on after $0.17\cdot60^2$ s after the start of argon input. The temperature mode of process included a period of non-isothermal heating (0.07 deg./s) up to an activation temperature, isothermal holding at this temperature for $\tau_f=1\cdot60^2$ s and rapid cooling in a...
stream of argon cooled at non-isothermal 0,1 deg./s to $T_0=323$ K. Activation temperature was $T_1=1073$ K when activated via KOH.

Samples of AC activating agent washed with distilled water for $\tau_4=600$ s and dried at $T_7=378$ K to humidity $W_4=5,84\%$ of the yield of AC $Y_1=70,4\%$. Fractionation AC remnant of MP conducted on sieves with holes: $d\geq3,6\cdot10^{-3} \text{– MP}=0,2\%$; $3,6\cdot10^{-3}>d\geq1,0\cdot10^{-3} \text{ – MP}=87,6\%$; $d<1,0\cdot10^{-3}$ (pallet) – MP=12,2\% with the following collection of working fractions on sieves of 3,6 mm and 1,00 mm MP – 87,8\%.

**Results and discussions**

Characteristics of porous structure was determined on a basis of isotherms of an adsorption-desorption of nitrogen at $T=77$ K in the range of relative pressure $P/P_0=0,00–1,00$ (device Quantachrome Autosorb 6B) (Figure 3).

![Figure 3. Isotherms of an adsorption-desorption of nitrogen at AC at T=77 K](image)

The obtained isotherms of type II – according to Brunauer S. et al, 1938 [31] classification, per multimolecular adsorption. Sorption hysteresis loop approaching the point of relative pressure $P/P_0=0,4$, indicating a predominance of micropores of meso- and macropores.

Figures 4–7 shows the distribution of micropores ($QSDFT$–method), the size of the sample and the corresponding volumes accumulated in these pores.

The microporous structure has the following characteristics (Figure 4–5): pore diameters are in the range of $D_{m}=0,61–2,5$ nm, mostly represented by pores with a diameter of 0,61; 1,19; 1,54 nm; volume of micropores – $V_{mi}=0,11–0,30$ cm$^3$/g; pores’ surface area – $S_{mi}=407–852$ m$^2$/g; pores’ differential volume $dV_{mi}/dD=(0,023–1,400)\cdot10^{-2}$ cm$^3$/g; pores’ differential area $dS_{mi}/dD=(0,18–45,60)$ m$^2$/g; micropores are about 70,31\% of the total pore volume.
Figure 4. Distribution of micropores by size of AC sample – (dependence of pores’ volume and pores’ differential volume on pores’ diameter) by QSDFT–method.

Figure 5. Distribution of micropores by size of AC sample – (dependence of surface area and pores’ differential surface area on pores’ diameter) by QSDFT–method.
Figure 6. Distribution of micropores by size of AC sample – (dependence of pores’ volume on pores’ surface area) by QSDFT–method

Figure 7. Distribution of micropores by size of AC sample – (dependence of pores’ differential volume on pores’ differential surface area) by QSDFT–method
According to distribution of micropores by size, two areas of values can be distinguished: a dynamic range \((D_{mi}=0,5–2,0\ nm)\) of values with several maxima; a stationary range of values \((D_{mi}=2,0–2,5\ nm)\). The main differences are fixed at the dynamic range of values \((D_{mi}=0,5–2,0\ nm)\). We can observe three maximum values: at \(\sim 0,6\ \text{nm}\), weakly expressed at \(\sim 1,2\ \text{nm}\), maximum at \(\sim 1,5\ \text{nm}\). The differential pore volume is in the range of \(dV_{mi}/dD=(1,9–2,1)\cdot10^{-4}\ \text{cm}^3/\text{g}\) at the static range of values \(D_{mi}=2,0–2,5\ \text{nm}\). The differential area is in the range of \(dS_{mi}/dD=(0,18–0,20)\ \text{m}^2/\text{g}\).

The dominant contribution of micropores in the specific surface of the pores shows a proportional relationship between the pores volume and the surface area of pores. This is also confirmed by the linear dependence between the pores’ differential volume and the pores’ differential area (Figure 6–7).

The subnanopores with \(D\leq1\ \text{nm}\) – the smallest pores were considered in the micropores’ structure (Figure 4–5): pores’ diameters are in the range of \(D_{inn}=0,61–1,00\ \text{nm}\); subnanopore’s volume varies in the range of \(V_{inn}=0,11–0,25\ \text{cm}^3/\text{g}\); pores’ surface area: \(S_{inn}=407–783\ \text{m}^2/\text{g}\); pores’ differential volume: \(dV_{inn}/dD=(11,3–140,0)\cdot10^{-4}\ \text{cm}^3/\text{g}\); pores’ differential area: \(dS_{inn}/dD=(2,33–45,60)\ \text{m}^2/\text{g}\). The subnanopore’s portion in the micropores volume is 84,1\%. The share of subnanopore’s in the total pores’ volume is 59,2\%. It can be argued that the alkaline activation of KOH leads to the development of a subnanopore’s in the porous structure of the adsorbent.

Figures 8–13 show the distribution of mesopores (BJH–method) by size in the sample and the corresponding volumes accumulated in these pores.

In Figure 8, the curve of the pores’ size set with its size increasing smoothly, not reaching the plateau, indicating the presence of mesopores with a wide distribution in size.

![Figure 8. Distribution of mesopores by size of AC sample – (dependence of pores’ volume and pores’ differential volume on pores’ diameter) by BJH–method](image-url)
Figure 9. Distribution of mesopores by size of AC sample – (dependence of pores’ volume and pores’ \( dV(\log r) \) on pores’ diameter) by BJH–method.

Figure 10. Distribution of mesopores by size of AC sample – (dependence of pores’ surface area and pores’ differential surface area on pores’ diameter) by BJH–method.
Figure 11. Distribution mesopores the size of AC sample – (dependence of pores’ surface area and pores’ \( dS(\log r) \) on pores’ diameter) by the \( BJH \)-method

\[
y = -7632.9x^2 + 1391.5x - 0.8916 \\
R^2 = 0.99
\]

Figure 12. Distribution of mesopores by size of AC sample – (dependence of pores’ surface area on pores’ volume) by the \( BJH \)-method
Mesoporous structure has the following characteristics: pores’ diameters are in the range of $D_{me}=3,3–50,0$ nm, most represented pores with a diameter of 3,73 nm; mesopore volume varies in the range of $V_{me}=0,010–0,091$ cm$^3$/g; pores’ surface area – $S_{me}=11,7–60,0$ m$^2$/g; pores’ differential volume $dV_{me}/dD=(0,02–4,75)\cdot10^{-4}$ cm$^3$/g; pores’ differential area $dS_{me}/dD=(0,002–0,570)$ m$^2$/g; fraction of mesopores in the total pores’ volume is 2,14–21,6% (Figure 8–13).

Curves of pores’ differential volume and pores’ differential surface area at the interval of $D=15,8–50,0$ nm are in the static area. Maximum location of pores with a smallest diameter is observed at the pores’ differential volume $dV_{me}/dD=6,43\cdot10^{-4}$ cm$^3$/g at the point of 3,73 nm within $D=2,5–15,8$ nm. The biggest number of mesopores are located at the range of $D=2,5–15,8$ nm.

The terms of AC and its characteristics are shown at the Table 2. The following characteristics of AC were measured: $Y$ – yield (%); $S_{BET}$ – specific surface area; $V_{S}$ – pores’ total volume; $V_{ma}$ – macropores’ volume; $V_{me}$ – mesopores’ volume; $V_{mi}$ – micropores’ volume; $V_{1nm}$ – subnanopores’ volume; $A_{Phenol}$ – sorption capacity toward phenol; $A_{Pb}$ – sorption capacity toward Plumbum; $A_{MB}$ – sorption capacity toward methylene blue.

Comparison of distribution of porous space according to (Pat. 61059 Ukraine) to the experimental data is presented at the figure 14.

A method that allows the production of AC from PWW, generated after the process of food products’ smoking has been proposed. Moreover, PWW is subsequently heated non-isothermally and chemically activated in the presence of KOH. As a result, AC is produced with a high yield of 70–80%, developed specific surface, porous space and a high sorption capacity. The results of AC production can be adapted for the technology of alcoholic beverages’ production at the expense of 1,0–3,6 mm particles’ fractional composition.
Table 2

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Method of AC production (Pat. 61059 Ukraine)</th>
<th>Method of AC experimental production</th>
</tr>
</thead>
<tbody>
<tr>
<td>Type of raw material</td>
<td>Lignite (brown coal)</td>
<td>PWW</td>
</tr>
<tr>
<td>Activation temperature, K</td>
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<td>1073</td>
</tr>
<tr>
<td>Activating agent</td>
<td>KOH</td>
<td>KOH</td>
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<tr>
<td>State of the activating agent</td>
<td>solid</td>
<td>solution 50%</td>
</tr>
<tr>
<td>MR raw/agent, kg/kg</td>
<td>1:0,5</td>
<td>1:1</td>
</tr>
<tr>
<td>$Y, %$</td>
<td>39,0</td>
<td>70,4</td>
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<tr>
<td>$S_{BET}, m^2/g$</td>
<td>890</td>
<td>777</td>
</tr>
<tr>
<td>$V_{S}, cm^3/g$</td>
<td>0,580</td>
<td>0,421</td>
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<td></td>
<td>100%</td>
<td>100%</td>
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<td>$V_{ma}, cm^3/g$</td>
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<td>1,73%</td>
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<td>$V_{me}, cm^3/g$</td>
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<td></td>
<td>43,10%</td>
<td>21,61%</td>
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<td>$V_{mi}, cm^3/g$</td>
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<td>0,296</td>
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<td>70,31%</td>
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<td>$V_{1nm}, cm^3/g$</td>
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<td>39,66%</td>
<td>59,15%</td>
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<td>$A_{Pb}, mmol/g$</td>
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<tr>
<td>$A_{MB}, mg/g$</td>
<td>92</td>
<td>150</td>
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</table>

Figure 14. Distribution of pores in AC:

\(a\) – according to Pat. 61059 Ukraine; \(b\) – according to experimental data.

The attention is drawn to the volume of pores’ with $D \leq 1$ nm. It accounts to 59,15\% of the total pores’ volume in AC. The obtained data allows us to hope that the studied raw materials can be used for the purification of water-alcohol mixtures.
Conclusions

The data show that the proposed method allows to obtain AC with a high yield of 70.4% compared to the method of obtaining AC from lignite (Pat. 61059 Ukraine) – 39.0%. Experimentally received AC has a lower specific surface $S_{BET}=777 \text{ m}^2/\text{g}$ with respect to AC (Pat. 61059 Ukraine) $S_{BET}=890 \text{ m}^2/\text{g}$ and pores’ space: total pores’ volume $V_\Sigma=0.421 \text{ cm}^3/\text{g}$ to $V_\Sigma=0.580 \text{ m}^3/\text{g}$. Nevertheless, the ratio of micropores in the experimental sample (70.31%), increased in relation to the prototype (55.17%), and the ratio of subnanopores in the experimental sample (59.15%), increased in relation to the prototype (39.66%). The ratio of macropores in the experimental sample (8.08%), increased in relation to the prototype (1.73%). At the same time the ratio of mesopores in the experimental sample (21.61%) reduced relatively to the prototype (43.10%).

It can be concluded that the proposed method of AC production from PWW, produced of smoked foods, with further carbonization at non-isothermal heating and activation at temperature of 873–1073 K in the presence of KOH, leads to sorbents with a high yield of 70–80% and fractional composition of particle size of 1.0–3.6 mm (~ 90%). An energy-saving method is proposed for a production of cheap AC from secondary «renewable» resources – PWW. These AC can be examined for water purification in alcoholic beverages’ production.

References


