Research article

Examination of gas and solid products during the preparation of activated carbon using phosphoric acid


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A B S T R A C T

Activating agents play significant roles in the preparation of activated carbon (AC) from biomasses and their wastes, which are widely used in AC production. Application methods are also important for the production process. Products give remarkable ideas regarding the method and heat treatment process. The activated carbon was produced from waste tea in accordance with either the conventional method or microwave energy pre-treated method using phosphoric acid (H₃PO₄) as activating agent. The yields of the activated carbons were 51.8% for conventional method and 46.0% for microwave pretreated method. The acid suppressed the formation of tar and promoted the amount of solid and aromatic structure accordance to sp² hybridisation. Additionally, the waste tea was directly carbonised (without H₃PO₄) and the yield was 36.3%. Major gas (H₂, CH₄, C₂H₆, C₂H₄, CO₂ and CO) products obtained during heat treatment process in a conventional furnace were examined in terms of quantity and quality. The solid products were characterised in terms of surface area, pore size and surface properties. The result of gas analysis showed that phosphoric acid affected formation of activated carbon mechanism and significant reactions occurred during microwave pretreatment process.

1. Introduction

Lignocellulosic biomasses and their wastes are low cost potential raw materials to produce high surface area activated carbon. The cellulose, hemicellulose and lignin contents control the porosity development and surface properties of the activated carbon. The activated carbon is widely used in different fields due to its composition and renewable nature, and plays a significant role in adsorption technologies, battery electrodes, capacitors, and gas storage (Dias et al., 2007; Peláez-Cid et al., 2016; Nabais et al., 2013; Gokce and Aktas, 2014; Inal et al., 2015). A variety of lignocellulosic materials such as waste tea (Yagmur et al., 2008; Gurten et al., 2012; Fadhil et al., 2012), corn cob (Song et al., 2013), rice husk (Somasundaram et al., 2013), sugar cane bagasse (Rufford et al., 2010), palm shell (Arami-Niya et al., 2012), lignin (Gao et al., 2013), and so on have been used as precursors to produce the activated carbon. The activated carbon is prepared by physical activation in an oxidizing atmosphere (such as CO₂, H₂O, O₂) or by chemical activation with chemical reagents such as KOH (Wu et al., 2011; González-García et al., 2013), NaOH (Foo and Hameed, 2012; Islam et al., 2017), ZnCl₂ (Zyoud et al., 2015), H₃PO₄ (Jagtoyen and Derbyshire, 1998; Yagmur et al., 2008; Gokce and Aktas, 2014) and K₂CO₃ (Gurten et al., 2012) under different conditions. The activation temperature depends on the type of activating agent, for example, 700–900°C for KOH, NaOH and K₂CO₃ (Gao et al., 2013; Foo and Hameed, 2012; Gurten et al., 2012), 500–700°C for ZnCl₂ (Molina-Sabio and Rodriguez-Reinoso, 2004), and 400–500°C for H₃PO₄ (Jagtoyen and Derbyshire, 1998; Yagmur et al., 2008).

H₃PO₄ is widely used for the production of activated carbon from lignocellulosic materials (Jagtoyen and Derbyshire, 1998; Yagmur et al., 2008; Gokce and Aktas, 2014; Hared et al., 2007; Dobele et al., 1999; Zuo et al., 2009). Phosphoric acid integrates in the interior of cell walls of the lignocellulosic materials during heating process. Eventually the biomass is converted into a solid product with a highly developed pore structure. The amount of solid product increases due to the crosslinking reactions (above 300°C) as results of hydrolysis and de-hydration by H₃PO₄ (Jagtoyen and Derbyshire, 1998; Solum et al., 1995; Zuo et al., 2010). Jagtoyen and Derbyshire (1998) stated that phosphoric acid has two functions in the process. The first is to promote bond cleavage reactions and the creation of crosslinks by means of processes, for example cyclisation and condensation. The second is to combine with organic species to form phosphate and polyphosphate bridges that connect and crosslink biopolymer fragments. Zuo et al.
reported that the presence of phosphoric acid noticeably depressed the evolution of CO, CO$_2$, and CH$_4$ and promoted H$_2$.

As reported above many studies are available in the existing literature relating to the preparation of activated carbon from lignocellulosic biomasses in the presence of phosphoric acid. In the majority of these studies, attention has been focused on the solids and their characteristics. The literature investigating both the gas and solid products at the same time during heat treatment process (Zuo et al., 2012) is very limited. Research investigating the relationship between the gas evolution and the solid product characteristics would provide significant knowledge regarding the mechanism of the process.

Microwave heating method has been recently used for the preparation of the activated carbon as well as the conventional heating processes (Yagmur et al., 2008, 2013; Yagmur, 2012; Foo and Hameed, 2012). Our previous studies (Yagmur et al., 2008) showed that microwave pretreatment enhanced surface properties (particularly total surface area increased nearly 10–24%) of the activated carbon compared to the conventional method. In this research, experiments were performed to prepare activated carbons from factory waste tea according to the conventional and microwave pretreatment methods using phosphoric acid.

The objective of the study is to examine the gas and solid products and to establish the relationship between the gas and solid products. It is also to show the effects of microwave pretreatment in the presence of phosphoric acid on the amounts and properties of gas and solid produced.

2. Experimental

2.1. Materials and material fabrication

Factory waste tea (WT, particle size: less than 500 μm) was chosen as a raw material to prepare activated carbon in the presence of phosphoric acid (Merck, 85%) which was used as an activating agent. The waste tea was collected from a tea plant located in Eastern Black Sea region in Turkey.

Three sets of experiments were performed. The first involved direct carbonisation of the raw material (WTR). The second used a conventional method where the mixture of the raw material and phosphoric acid was carbonised-activated (WTC). Unlike our previous works (Yagmur et al., 2008), a temperature programmable rotary tube furnace was used to be able to collect properly gaseous product. The third case utilised a microwave pretreatment process (WTMW). In this third case, the acid–raw material mixture was pretreated first in a domestic microwave oven and then the mixture was carbonised-activated in the same furnace as specified above. The whole process is schematically presented in Fig. 1.

Heat treatment of the samples was conducted in a quartz tube reactor placed in the furnace. Pure nitrogen gas was provided as an inert gas atmosphere and the flow rate of nitrogen was 150 ml/min. For each sample (the raw material, mixture of raw material + H$_3$PO$_4$) was heated to temperature of 450 °C at the constant heating rate (10 °C/min). The solids in the reactor were held at 450 °C for 1 h. Gaseous material was collected in Tedlar sample bags during the heat treatment process at different temperature intervals (Table 1).

The raw material (waste tea, WTR) was directly carbonised in the furnace without using any activating agent. Activated carbon was produced in two ways: conventional (WTC) and microwave pretreatment (WTMW) methods. In the conventional method (WTC), the waste tea was mixed with H$_3$PO$_4$ (85%, wt) at the acid/waste tea ratio of 2/1 (wt/wt). The mixture was loaded to the rotary tube furnace for the heat treatment process. In the microwave pretreatment method (WTMW) the mixture was directly subjected to microwave radiation for 30 s before loading the furnace (Yagmur et al., 2008). A domestic microwave oven (Vestel, MDG-620) was used for the pretreatment process. The frequency and the input power of the microwave oven were 2.45 GHz and 900 W, respectively. The microwave pretreated sample was placed in the rotary tube furnace for further activation/ carbonisation process as described for the conventional method (see Fig. 1). Following treatment in the furnace in the presence of phosphoric acid, the sample was removed and mixed with hot distilled water (500 ml, 80 °C) and held in the water for 12 h and then washed and filtered. The solid fraction was dried in an oven at 110 °C overnight. The sample was then turned in a mill (IKA, M20 Universal Mill) at 20 000 rpm for 3 min, in order to liberate trapped acid. The milled sample was re-washed with distilled water until the filtrate became neutral and the solid fraction was dried for analyses.

2.2. Analyses

2.2.1. Gas analysis

The gases collected in Tedlar sample bags were analysed in a gas chromatograph (Micro GC T3000, SRA Instruments) equipped with a dual-column (MS 5A and PPU) and micro thermal conductivity detectors. The SOPRANE software package was used to identify the collected data by quantifying the peak areas. The collection temperatures of the gas samples are tabulated in Table 1. Gas analysis details and micro-GC conditions (Table S1) are reported in the Supplementary Information.

2.2.2. Solid analysis

The carbon, hydrogen, nitrogen and sulphur contents of the raw materials and the solids were analysed using a LECO CHNS 932 Elemental Analyser. The oxygen content was calculated by difference. The BET surface area and pore size distributions (the Non Local Density Functional Theory, NLDFT, method) of the samples were determined using a Quantachrome NOVA 2200 series volumetric gas adsorption instrument (Evans and Tarazona, 1989). The analysis details are presented in the Supplementary Information.

2.2.3. FTIR analysis

FTIR (Fourier Transform Infrared Spectroscopy) analyses of the solids were performed using a Shimadzu FTIR - 8040 spectro-photometer. The spectra were completed in the range of 400–4000 cm$^{-1}$. Thin KBr pellets have been prepared with the mass ratio of 300:1 (KBr:Sample) for the samples.

2.2.4. XPS analysis

X-ray photoelectron spectroscopy (XPS) measurements were conducted using a Thermo Scientific, K-Alpha, x-ray photoelectron spectrophotometer with Al Kα x-ray source (1486.6 eV). The binding energy scales of the high resolution spectra were calibrated by shifting Cls peak position to 284.8 eV. Peak deconvolution was carried out using Gaussian-Lorentzian curves. The survey scans were obtained using the pass energy of 200 eV and a step size of 1 eV. High resolution scans were performed using a pass energy of 30 eV and a step size of 0.1 eV.

3. Results and discussion

3.1. Analysis of gases

Thermal decomposition of lignocellulosic materials (hemicellulose, cellulose and lignin) occurs at two different temperature steps: high (> 300 °C) and lower (< 300 °C) temperatures. Major noncondensable gases such as CO, CO$_2$, H$_2$, and CH$_4$ are released during the carbonisation process of these materials (Ronse et al., 2015; Li et al., 2015; Ferrera-Lorenzo et al., 2014a). However, it is well known and commonly reported that phosphoric acid changes the depolymerisation and dehydration mechanisms during the heat treatment process (Jagtoyen and Derbyshire, 1993, 1998; Dobele et al., 1999). According to Jagtoyen and Derbyshire (1998), phosphate esters form on cellulose side chains at < 450 °C in the presence of phosphoric acid. The acid is eliminated at higher temperature (> 450 °C). In this process,
aromaticity and loss of aliphatic, carboxyl, and carbonyl groups in- 
crease (Solum et al., 1995).

Fig. 2 represents the amount of hydrogen evolved during heat 
treatment process for the experiments. The raw material (WTR) without 
H₃PO₄, conventional (WTC) and microwave pretreated (WTMW) 
methods in the presence of the acid produced similar amounts of H₂. 
The amount of hydrogen was insignifi- 
cant up to 400–420 °C, however it 
increased sharply at 450 °C, which was the 
final temperature of the 
process, and reached to a maximum when the material was held in the 
furnace for 30 min at the same 
final temperature. The amount of gas 
sharply declined after 30 min holding time at 450 °C. The presence of 
phosphoric acid did not significantly change the amount of hydrogen 
released.

Fig. 3 shows the amounts of methane in the gas samples at different 
temperature ranges. Direct carbonisation of the raw material (WTR) 
produced significant amount of CH₄ compared to the samples WTC and 
WTMW. CH₄ is released mainly from the cracking of methoxyl groups 
and the C–C bond cleavage in aliphatic chains at relatively lower 
temperatures (Ferdous et al., 2001). Lower amounts of CH₄ produced 
from WTC and WTMW are attributed to presence of the acid which 
possibly changes decomposition mechanism of biomass. The gas re- 
leased with temperature increased sharply reaching a maximum at 
450 °C and then sharply decreased with time at the constant tempera- 
ture (450 °C). Microwave pretreated samples produced less methane 
compared to the WTC. The reason is that some of methane released 
during microwave pretreatment process (30 s). A similar trend is ob- 
served for ethane that is shown in Fig. 4. This supports that aliphatic 
structure significantly decomposes due to cleavage of aliphatic side 
chains in the presence of H₃PO₄ during heating process (Jagtoyen and 
Derbyshire, 1998).

The situation is slightly different for ethylene as shown in Fig. 5. The 
maximum amounts of ethylene were produced at lower temperatures 
than 450 °C for the activated carbons prepared in the presence of 
H₃PO₄. Phosphoric acid in the mixture shifted the maxima to a lower 
temperature. Due to microwave pretreatment, less ethylene formed

### Table 1

Data for gas samples collected.

<table>
<thead>
<tr>
<th>Sample no</th>
<th>Temperature (°C)</th>
<th>Collecting time, (min.)</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>180–200</td>
<td>16–18</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>250–270</td>
<td>23–25</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>300–320</td>
<td>28–30</td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>350–370</td>
<td>33–35</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>400–420</td>
<td>38–40</td>
<td></td>
</tr>
<tr>
<td>6</td>
<td>450 (1)</td>
<td>43–45</td>
<td>1st min. at 450 °C</td>
</tr>
<tr>
<td>7</td>
<td>450 (30)</td>
<td>73–75</td>
<td>30th min. at 450 °C</td>
</tr>
<tr>
<td>8</td>
<td>450 (60)</td>
<td>103–105</td>
<td>60th min. at 450 °C</td>
</tr>
</tbody>
</table>

Ambient temperature: 20 °C.

Fig. 1. Schematic of the processes used to produce gases and solids.

Fig. 2. Hydrogen contents of the gas fractions.

Fig. 3. Amounts of methane in the gas samples at different temperature ranges.

Fig. 4. Ethane contents of the gas samples at different temperature ranges.

Fig. 5. Ethylene contents of the gas samples at different temperature ranges.
Fig. 3. Methane contents of the gas fractions.

Fig. 4. Ethane contents of the gas fractions.

Fig. 5. Ethylene contents of the gas fractions.

Fig. 6. Carbon dioxide contents of the gas fractions.

Fig. 7. Carbon monoxide contents of the gas fractions.

Fig. 3 and 4 show the amounts of CO2 and CO released during the heat treatment processes, respectively. The release of CO2 and CO starts at a lower temperature range between 180 and 200 °C and is almost completed at 450 °C after 30 min. The maxima of the gases shift to lower temperatures for the samples treated (WTC, WTMW) in the presence of H3PO4. Majority of the gases were released at higher temperature for the raw material (WTR). Therefore, its maxima moved to the right, which corresponds to a higher temperature when compared to the samples treated in the presence of acid. CO2 and CO were most likely generated from the degradation of hemicellulose and the pyrolysis of cellulose (aromatic and aliphatic carboxyl groups), respectively (Lv and Wu, 2012). It was also reported by Yang et al. (2007) that the presence of C=O groups in hemicellulose caused formation of CO2.

The presence of H3PO4 in the mixture significantly affected the reaction mechanism. CO2, CO, and ethylene gases that were released at lower temperature compared to the experiment performed using WTR (in the absence of acid). In general, the amounts of gases which were produced in the presence of the acid were lower than that of the gases obtained from the raw material. In brief, phosphoric acid catalysed the thermal decomposition of the lignocellulosic raw material and hydrolysed glycosidic linkages in lignocellulosic and depressed the evolution of the gases. It is also reported that the acid cleaved aryl ether bond in lignin in the raw material (Jagtoyen and Derbyshire, 1993, 1998; Dobele et al., 1999, 2001; Zuo et al., 2012; Nowakowski et al., 2008). The gases are produced from carboxyl, carbonyl and ether groups in the structure as a result of the cracking and reforming reactions (Wang et al., 2009; Liu et al., 2008).

Several decomposition reactions also take place during the microwave pretreatment process in the microwave oven (Ferrera-Lorenzo et al., 2014b). The release of gas product was observed in the fume cupboard in the course of microwave irradiation process.

3.2. Analysis of solids

The BET surface area, pore volume and yield of the solids were reported in Table 2. As reported in the experimental section, the waste tea was directly carbonised (WTR) under nitrogen atmosphere in the absence of activating agent. The BET surface area and pore volume of WTR were 3.2 m2/g and 0.007 cc/g, respectively. The activated carbons
obtained according to conventional (WTC) and microwave pretreatment (WTMW) methods have significantly higher surface areas. As the BET surface area of microwave pretreated sample was 1255 m²/g, the surface area of the activated carbon obtained from conventional method was 1135 m²/g. Similar findings were reported in our previous studies (Yagmur et al., 2008; Gokce and Aktas, 2014). High surface area is attributed to microwave heating mechanism that occurs from the core of material to outer surface (Yagmur et al., 2008). This process most probably forms new pores and therefore generates higher surface area. The pore volumes and yields of these samples were reported in last two columns of Table 2. The yields of solids were higher for the experiments (WTC, WTMW) performed in the presence of phosphoric acid. On the other hand, the yield of solid is substantially low for the sample WTR (pyrolysed without phosphoric acid). The yield of WTR is consistent with TGA analysis of the same waste tea reported in our previous work (Tiryaki et al., 2014). As reported in Section 3.1, the amount of gas product was highest for WTR compared to the samples WTC and WTMW treated in the presence of acid. This is evidence that phosphoric acid suppresses formation of tar and increases the yield of solid due to crosslinking reactions (Zuo et al., 2012). It is better to point out that the amount of gases is higher for WTR than the others. This clearly shows that significant fraction of the raw material is converted into gaseous product in the absence of the acid.

The adsorption/desorption isotherms of the samples, WTC and WTMW, are presented in Fig. 8. The isotherms of the activated carbon samples are similar and have hysteresis, which is indicative of mesoporosity. The isotherms and hysteresis are Type IV and Type H4, respectively, according to the IUPAC classification (Thommes et al., 2015). The volume of gas adsorbed by WTR (direct carbonised) was very low, therefore the isotherms of this sample was not inserted in Fig. 8.

Fig. 9 shows the pore size distributions of the activated carbon samples obtained from NLDFT method. There is no significant difference in the distribution curves. As pore size distribution of the samples is slightly different between 1.1 and 7 nm, their distributions are very similar beyond 7 nm.

Table 3 reports the results of ultimate analyses and ash contents of the heat treated samples. Additional data about elemental composition of microwave treated-30 sample is reported in Table S2. The carbon contents of the samples; WT, WTC and WTMW are shown in the second column of the table. The ash contents of the solid samples are not significantly different; however, their oxygen contents are dissimilar. The sample WTMW has the lowest oxygen content. This is attributed to the microwave pretreatment causing the release of oxygen rich groups as gases during process. This statement is supported by the study carried out by Dominguez et al. (2007). They reported that microwave pyrolysis caused more deoxygenated chars than that of conventional pyrolysis. The nitrogen content is lower for the samples obtained in the presence of acid. Apparently phosphoric acid promotes the decomposition of nitrogen containing groups in the structure and turns into gases (Kumar and Jena, 2016). There is no significant difference in the contents of hydrogen of the activated carbons. However, hydrogen content is markedly lower than that of the raw material. This indicates that most of hydrogen is converted to gas products during heat treatment processes. For example, this statement is consistent with the FTIR results (reported in Section 3.3.), intensities of the band at 2800-3000 cm⁻¹ (aliphatic C–H groups) are significantly weak in the spectra of the activated carbons.

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**Table 2**

<table>
<thead>
<tr>
<th>Sample ID Pre-treatment</th>
<th>Heat treatment</th>
<th>Surface area, m²/g</th>
<th>Micropore volume, cc/g</th>
<th>Mesopore volume, cc/g</th>
<th>Total pore volume at P/P₀ = 0.99, cc/g</th>
<th>Yield, %</th>
</tr>
</thead>
<tbody>
<tr>
<td>WTR</td>
<td>Carbons at 450 °C</td>
<td>3.2</td>
<td>--</td>
<td>--</td>
<td>0.007</td>
<td>36.3</td>
</tr>
<tr>
<td>WTC (with H₃PO₄)</td>
<td>Activated and carbonised at 450 °C</td>
<td>1135</td>
<td>0.248</td>
<td>0.719</td>
<td>0.988</td>
<td>51.8</td>
</tr>
<tr>
<td>WTMW (with H₃PO₄)</td>
<td>Activated and carbonised at 450 °C</td>
<td>1255</td>
<td>0.279</td>
<td>0.762</td>
<td>1.067</td>
<td>46.0</td>
</tr>
</tbody>
</table>

* Micropore and mesopore volumes were calculated according to the NLDFT method.

**Table 3**

<table>
<thead>
<tr>
<th>Sample ID Pre-treatment</th>
<th>Heat treatment</th>
<th>Surface area, m²/g</th>
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<td>1.067</td>
<td>46.0</td>
</tr>
</tbody>
</table>

* Sample ID C, % H, % N, % S, % O a, % O/C Ash, %

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**Fig. 8.** Adsorption/desorption isotherms of the activated carbon samples.

**Fig. 9.** Pore size distribution and cumulative pore volumes of the activated carbon samples.
The peaks in the range of 1050–1590 cm$^{-1}$ are associated with aliphatic C=O hydrogen-bonded P=O (Puziy et al., 2005). It is deduced from the FTIR spectra that the surface functionalities are very sensitive to heat treatment processes. The raw material contained several oxygen functional groups, however the majority of these groups were partly or completely removed during the heat treatment processes. The broad band at 3100–3600 cm$^{-1}$ corresponds to O–H stretching vibrations from alcohols, phenols and absorbed water. The stretching vibrations of aliphatic C–H groups (2800–3000 cm$^{-1}$) are observed in the raw material. However, the bands at 1375 cm$^{-1}$ for the raw material and 1450 cm$^{-1}$ for the WTR are associated with aliphatic C–H stretching (Lladó et al., 2015). The peaks in the range of 1050–1740 cm$^{-1}$ for the waste tea (ethers, lactone, carbonyl) were significantly reduced after heat treatment process (Zielke et al., 1996). The new peaks observed in the WTC and WTMW samples in the same region indicate increasing aromativity of the carbon in the addition of phosphoric acid (Sych et al., 2012). The absorption band at 1740 cm$^{-1}$ is ascribed to C=O stretching vibrations of conjugated carbonyl groups (carboxylic acid and lactone groups) (Golce and Aktas, 2014). This peak has a lower intensity for the treated samples and is shifted to a higher wavenumber of 1755 cm$^{-1}$. The peak in the 1560–1680 cm$^{-1}$ region consists of two overlapped absorption bands, non-conjugated carbonyl groups at 1661 cm$^{-1}$ and highly conjugated carbonyls (quinone groups) and/or C=O stretching vibrations in the aromatic ring at 1600 cm$^{-1}$, respectively. After the activation and carbonisation processes, the combined peak in the region of 1560–1680 cm$^{-1}$ is observed in the samples WTC and WTMW as two distinct peaks. The bands associated to non-conjugated carbonyl groups and aromatic ring vibrations have shifted to 1694 cm$^{-1}$ and 1590 cm$^{-1}$, respectively. The peak at 1180 cm$^{-1}$ for phosphoric acid activated carbons can be assigned to the stretching vibration of hydrogen-bonded P=O groups from phosphates or polyphosphates, to the O–C stretching vibration in the P–O–C (aromatic) linkage, and to P–OH (Puziy et al., 2005). It is deduced from the FTIR spectra that most of the oxygen containing functional groups were removed from the lignocellulosic structure as results of heat treatment process.

### 3.4. FTIR analysis

Fig. 10 shows the FTIR spectra of the raw material, WTR, WTC and WTMW samples. In addition, the FTIR spectra of the raw material, microwave pretreated-30 s and WTMW samples are presented in Fig. S1 to show the effect microwave pretreatment. It is apparent from the spectra that the surface functionalities are very sensitive to heat treatment. The raw material contained several oxygen functional groups, however the majority of these groups were partly or completely removed during the heat treatment processes. The broad band at 3100–3600 cm$^{-1}$ corresponds to O–H stretching vibrations from alcohols, phenols and absorbed water. The stretching vibrations of aliphatic C–H groups (2800–3000 cm$^{-1}$) are observed in the raw material. However, the bands at 1375 cm$^{-1}$ for the raw material and 1450 cm$^{-1}$ for the WTR are associated with aliphatic C–H stretching (Lladó et al., 2015). The peaks in the range of 1050–1740 cm$^{-1}$ for the waste tea (ethers, lactone, carbonyl) were significantly reduced after heat treatment process (Zielke et al., 1996). The new peaks observed in the WTC and WTMW samples in the same region indicate increasing aromativity of the carbon in the addition of phosphoric acid (Sych et al., 2012). The absorption band at 1740 cm$^{-1}$ is ascribed to C=O stretching vibrations of conjugated carbonyl groups (carboxylic acid and lactone groups) (Golce and Aktas, 2014). This peak has a lower intensity for the treated samples and is shifted to a higher wavenumber of 1755 cm$^{-1}$. The peak in the 1560–1680 cm$^{-1}$ region consists of two overlapped absorption bands, non-conjugated carbonyl groups at 1661 cm$^{-1}$ and highly conjugated carbonyls (quinone groups) and/or C=O stretching vibrations in the aromatic ring at 1600 cm$^{-1}$, respectively. After the activation and carbonisation processes, the combined peak in the region of 1560–1680 cm$^{-1}$ is observed in the samples WTC and WTMW as two distinct peaks. The bands associated to non-conjugated carbonyl groups and aromatic ring vibrations have shifted to 1694 cm$^{-1}$ and 1590 cm$^{-1}$, respectively. The peak at 1180 cm$^{-1}$ for phosphoric acid activated carbons can be assigned to the stretching vibration of hydrogen-bonded P=O groups from phosphates or polyphosphates, to the O–C stretching vibration in the P–O–C (aromatic) linkage, and to P–OH (Puziy et al., 2005). It is deduced from the FTIR spectra that most of the oxygen containing functional groups were removed from the lignocellulosic structure as results of heat treatment process.

### 3.5. XPS analyses

XPS analyses were performed to investigate the surface chemical characteristics of the samples (Fig. 11) and to evaluate how the production methods affected the surface functionalities of solid samples. Determination the type of the surface functional groups and the changes in the amount of these groups would be helpful for the examination of the gas product. The surface composition (atomic %) of the activated carbon samples were determined by using the peak area of each elements and the total peak area obtained by survey scan (Figs. S2–S4). According to survey scans, all the samples contain carbon, oxygen, nitrogen and phosphorus and carbon is the major element located on the surface. The oxygen content is considered to be the indicator of the presence of the surface functional groups and higher O/C ratio refers to hydrophilic surface character. The elemental composition and the O/C ratios of the samples obtained from XPS analysis are presented in Table 4.

As given in Table 4, there is no significant difference in the elemental compositions of WTC and WTMW (produced in presence of H$_3$PO$_4$) however the values for WTR are different compared to the WTC and WTMW. The results reported in the same table are different from those obtained by ultimate analysis (Table 3). Different values may be due to analysis methods applied. In the XPS analysis, x-rays can only access to a limited depth of the surface (a few nm), so the XPS analysis results cannot represent the overall chemical composition. Apparently activation in the presence of phosphoric acid causes decreases in the contents of nitrogen and oxygen. This may be ascribed to production of heat sensitive surface functional groups in the presence of H$_3$PO$_4$ during the activation process. Because of the higher heat resistance characteristics of graphitic carbon structure, the removal of nitrogen and oxygen containing surface groups leads to an increase in the carbon content. The phosphorus content of the activated samples in Table 4 is attributed to the phosphate linkages produced by the reaction of H$_3$PO$_4$ with the carbon skeleton or entrapped H$_3$PO$_4$ molecules in the pores. These results are consistent with the FTIR analysis data. The phosphorus content of WTR might be inherent material of the raw waste tea.

As shown in Fig. 11, Cls XPS spectra have an asymmetric shape suggested the existence of different carbon species. The deconvolution of the carbon spectra exhibited four peaks for the sample WTR and five peaks for the samples WTC and WTMW at different binding energies (BE). Peak deconvolution of Cls spectra can be used for the identification and the quantitative determination of the carbon containing surface functional groups. Type and the relative content of the surface functionalities obtained by Cls XPS spectra are tabulated in Table S3. The C=O/C–C sp$^2$ peaks observed at 284.6 ± 0.2 eV for all samples were assigned to graphitic carbon structure. Beside C–C sp$^2$ peaks, the activated samples contained C–C sp$^3$ peaks (285.5 ± 0.2) on the solid surfaces. The dominant sp$^2$ hybridisation and relatively mild sp$^3$ hybridisation of carbon atoms correspond to degree of graphitisation or highly graphitic structure having limited defects and disorders (Tai et al., 2006; Aria et al., 2013). According to Cls spectra, the carbonised sample exhibited the lowest C–C sp$^2$ structure (19.16%) but has the highest C–C sp$^3$ structure (59.73%). The graphitic structure contents of WTC and WTMW samples are 60.88% and 64.02%, respectively. The increase in the graphitic carbon content and the formation of the C=O and O=C–O− groups is most probably due to the impregnation of raw material with H$_3$PO$_4$ and its oxidative effect.

The signal obtained at 286.6 ± 0.2 eV refers to the existence of C=O type structures in the form of hydroxyl, phenol, ether or phosphorus complexes. As reported in FTIR analysis, the sample WTR did not contain C=O type functional groups and these functional groups were not detected in XPS analysis as well. The peak at 287.4 ± 0.1 for WTR implies that carbon-oxygen double bonds occurs as quinonic or ketonic groups at the surface. Taking into account of the FTIR results, this peak can be assigned to highly conjugated quinone groups. On the other hand, the signal at 288.9 ± 0.1 eV is associated with carboxyl, ester, anhydride or lactone type surface groups. It is clear that the impregnation of the raw material with H$_3$PO$_4$ inhibits the formation of C=O single bonds to some extent but promotes the development of O=C groups on the surface. The formation of the double bonded carbon-oxygen complexes mainly depends on the interaction between the raw
material and phosphoric acid. The signals observed at 291.2 ± 0.2 eV and 293.2 ± 0.1 are originated from the $\pi-\pi^*$ shake up satellite in aromatic rings for the samples WTR, WTC and WTMW, respectively.

**4. Conclusions**

Substantial changes were observed in the gas and solid products with and without phosphoric acid during heat treatment processes. The decomposition mechanism strongly depends on the presence of phosphoric acid. Phosphoric acid causes a shift in the gas release temperature to a lower level for CO, CO$_2$ and C$_2$H$_4$. The GC analyses showed that the amounts of gases obtained from WTC and WTMW were lower when compared with the sample WTR (without phosphoric acid). The yields of solid products obtained in the presence of the acid are significantly higher for the samples WTC (51.8%) and WTMW (46.0%) than that of the solid produced (36.3% for the sample WTR) in the absence of the acid. High solid yields are evidence that the formation of tar is suppressed by the addition of phosphoric acid.

Although the microwave pretreatment period is short (30 s), several decomposition reactions take place much faster if there is phosphoric acid in the mixture. The majority of oxygen containing functional groups is removed during heat treatment process. Phosphoric acid promotes aromatic structure as a result of sp$^2$ hybridisation.

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Appendix A. Supplementary data

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