Microwave Assisted Synthesis and Characterizations of Decorated Activated Carbon

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A novel intermittent microwave assisted method was developed for surface impregnation and electrical properties improvement of decorated activated carbon (DAC), from commercially available activated carbon (AC) and Vulcan carbon (VC). These materials were characterized by UV-visible spectroscopy, Raman spectroscopy, X-ray diffraction (XRD), Brunauer–Emmett–Teller (BET), scanning electron microscopy (SEM) and electrical conductivity. The UV absorption peak in carbon materials is related to electronic transitions between the bonding and antibonding π orbital. Raman spectroscopy and XRD results showed that the higher disorder of amorphous carbon phase arose in DAC, when compared to typical AC. Results from BET analysis had relatively high surface area of 1141 m²/g and total pore size volume of 0.524 m³/g for DAC. SEM analysis revealed that undefined structure of activated carbon material (~1-3 µm in size) breakage into ~ 100 nm in size over this Vulcan carbon material (~60nm in size) was decorated. From the electrical conductivity (σ) study, the value of σ followed the tendency DAC < VC < AC and increased almost with pressure. High surface area and high conductivity decorated carbon nano structures could be used as candidate hosts for metal electrocatalysts for fuel cell applications.

Keywords: Activated carbon; carbon support; microwave assisted synthesis; Nanoparticles;

1. INTRODUCTION

Carbon materials have been widely investigated as functional materials for various applications including electrodes for batteries and fuel cells, adsorbents for water purification, and catalyst supports
It is well known that the size, structure, density and shape of the materials influence their electrical properties and their potential applications [6,7]. Hence, the fabrication of carbon materials of a particular morphology is very important for their various applications. Recently, there has been increasing interest in the preparation and processing of various nanostructured carbon materials proposed for potential applications in supercapacitors [4], field emission [8], catalyst support [9], and solar energy conversion [10].

Graphitic carbon materials with a high surface area have attracted attention due to their potential applications. Traditional methods for the synthesis of graphitic carbon materials commonly need additional high thermal treatment [11, 12]. However, a large reduction of the surface area and pore volume of the carbon materials usually results after treatment at high temperatures. Microwave assisted heating offers the apparent advantage of reducing treatment time, which may represent a reduction in the energy consumption. It has been reported that microwave energy was derived from electrical energy with a conversion efficiency of approximately 50% for 2450 MHz and 85% for 915 MHz [13]. Microwave assisted heating caused a reduction in micropore volume and micropore sizes. Also, the microwave treatment is a very effective method for modifying the surface chemistry of the activated carbon fibers [14,15].

In the present work, we have designed a simple and efficient microwave assisted method for surface impregnation of Vulcan XC72R carbon on activated carbon materials leading to high surface area. Also achieved the morphological, electrical and structural properties of the decorated carbon material were characterized. To the best of our knowledge this is the first time that microwave was exploited for surface impregnation of carbon materials. These modified materials could be used as a supporting material for metal electrocatalysts, adsorbents, and electrodes in microbial fuel cells.

2. EXPERIMENTAL PROCEDURE

2.1 Microwave assisted decoration of activated carbon

The activated carbon (AC) (Sigma Aldrich) and Vulcan XC 72R carbon (VC) were dispersed in water in the ratio of 1:1 and allowed ultra sonication for 60 min. Then the carbon mixture was centrifuged at 8 000 rpm for 10 min and the precipitated powder was collected. Afterward, ethylene glycol was added to the carbon mixture for further microwave irradiation. A microwave irradiation programme was used for 20 seconds ON and 20 seconds OFF, repeating the programme 8 times. Finally the modified carbon materials (AVC) were collected and washed with water several times to eliminates impurities. The resulting powder was dried overnight, kept in desiccators and further used for physical characterization.

2.2 Physical characterization of carbon materials

The absorption spectra were obtained using a UV-vis spectrophotometer (Varian Inc. Cary 100/300, model 100/300). The Raman spectroscopy was performed with a Horiba HR800 model
operating at 514.5nm excitation. X-ray patterns of the decorated carbon powder were obtained in a Diffract Bruker AXS, D8 Advanced Plus diffractometer operated at 30 mA and 35 kV using a monochromatic CuKα radiation (λ = 0.15443 nm). A 2θ scanning range from 10° to 90° was explored at a step scan rate of 0.02° min⁻¹. Topography analyses were carried out using field emission scanning electron microscopy (GEMINI-Carl Zeiss). An Autochem II 2920 (Micrometrics) instrument was used to measure the nitrogen adsorption isotherms at -196 °C for determining the total pore volume and the specific surface area through the Brunauer–Emmett–Teller (BET) equation [16].

2.3 Electrical conductivity

The electrical conductivity (σ) of carbon samples was measured by impedance spectroscopy within the frequency range 10 kHz – 100 mHz, at room temperature. The carbon black samples were placed in a glass tube (inner diameter = 0.5 cm) between two metal pistons, the upper one movable and the lower one fixed. The electrical conductivity was measured at different pressures varied between 10 and 1013 kPa by employing different loadings on the upper piston. Changes in the height caused by loadings were measured with an accuracy of ±0.02 mm. A view of the experimental arrangement is shown in Figure 1.

Figure 1. Experimental arrangement for electrical conductivity measurements.

The carbon samples used were dried at 100°C overnight. 10 mg of each sample were employed for the measurements. A potentiostat/galvanostat, PARSTAT 2273, was used to measure the conductivity of the different samples. 3 min were allowed before any conductivity measurement to
avoid any transient effect of sample relaxation due to compression. The conductivity was calculated using equation 1.

\[ \sigma = \frac{l}{AR} \]  

(1)

Where: \( l \) is the distance between the two pistons (cm), \( R \) is the electrical resistance (Ω), \( A \) the area of the piston surface (cm²). The volume and density of the samples were determined after the measurement of \( l \).

3. RESULTS AND DISCUSSIONS

3.1 Physical characterization of Carbon materials

3.1.1. SEM and BET Surface area analysis

The schematic representation of our synthesis method of DAC is depicted in Figure 2.

**Figure 2.** Schematic representation of the synthesis method for decorated DAC
Intermittent microwave irradiation cracked the large particles of AC in smaller particles (100nm) with high pore volume, where as particle VC was not affected, the latter may due to small size of particle with low pore volume. Microwave may favour the VC particle impregnated over the AC. The carbon materials were further morphologically characterized by scanning electron microscopy analysis, as shown in Fig.3. Activated carbon shows undefined structure ranging from 1 to 3 µm size (Figure 3). Vulcan carbon presented well distributed spherical particles ranging between 50 to 60 nm in size (Figure 3b). However, decorated carbon materials showed the Vulcan carbon impregnated over the AC surfaces and revealed well distributed particles 90 to 100 nm in size (Figure 3c), attributed to breaking of larger sized AC particle in cluster formation along with Vulcan carbon, possible due to the irritation of microwaves.

![Figure 3](image.png)

**Figure 3.** Scanning electron micrograph of carbon materials. Inset: particle size histogram

**Table 1.** Surface area deduced from BET analysis

<table>
<thead>
<tr>
<th>Samples</th>
<th>BET-Surface area- m²/g</th>
<th>Total pore volume- m³/g</th>
</tr>
</thead>
<tbody>
<tr>
<td>VC</td>
<td>261</td>
<td>0.1203</td>
</tr>
<tr>
<td>AC</td>
<td>778</td>
<td>0.3568</td>
</tr>
<tr>
<td>DAC</td>
<td>1141</td>
<td>0.5242</td>
</tr>
</tbody>
</table>
The adsorption rates and capacities of typical carbon materials (AC and VC) increased after several adsorption-microwave regeneration cycles, which were reflected by the BET surface area analysis depicted in Table 1. The values for BET surface area and total pore volume followed the tendency DAC>AC>VC. BET surface area and total pore volume values obtained in the present work were in agreement with those reported previously in literature [14, 17].

3.1.2. UV-Vis Spectroscopy analysis

Figure 4 shows the dependency of the absorption peaks with the wavelength, as well as on the type of carbon materials. The increase in absorption from typical form of carbon materials (AC and VC) to modified carbon material (DAC) was assigned to the increasing surface area of the carbon materials after dispersion, concurrence with our BET surface area result showed in table 1. The increased surface area of the modified carbon material (DAC) exposed to UV radiation, is probably due to the resulting $\pi - \pi^*$ transitions.

Figure 4. UV-vis absorption spectrum of carbon materials.

Hence, by increasing the surface area of the dispersion, a higher absorption in the UV-vis spectra is expected [18]. The UV absorption peak in carbon materials is related to electronic transitions between the bonding and antibonding $\pi$ orbital [19]. The $\pi - \pi^*$ transitions appear in the range of 180–260 nm in all the carbon materials, as reported in ref [20]. Figure 4 shows the UV-vis absorption spectra of the different carbon materials in the range of 200-800 nm. AC and VC showed absorption
maximum peaks ($\lambda_{\text{max}}$) at 258 and 249nm, respectively (Figure 4 curve b and c). The $\lambda_{\text{max}}$ of the DAC materials showed an absorption maximum peak at 247nm (Figure 4 curve a).

3.1.3. Raman Spectroscopy and X-ray Diffraction analysis

Figure 5 shows Raman spectra of typical carbon materials (AV and VC) and decorated carbon material (DAC) measured at 514.5nm excitation; the former exhibited two predominant band ranges between 1100-1350 cm$^{-1}$ and 1600-1800 cm$^{-1}$, both related to characteristic feature of carbon associated to the presence of $C-O$ and $C=O$ groups [21]. VC revealed two most intense bands, where G band at 1580 cm$^{-1}$ was attributed to the ordered crystalline graphitic (within the layer) and a band at 1340 cm$^{-1}$ that corresponded to disordered amorphous carbon structure within the carbon sheet termed as D band.

![Raman spectra of carbon materials](image)

**Figure 5.** Raman spectra of carbon materials.

The AC showed a strong peak at 1580 cm$^{-1}$ and a small hump in the D peak position at 1340 cm$^{-1}$, which is clearly an indicative of the more dominant crystalline phase, in agreement with XRD results (see below). In DAC the D band increased in relation to the G band, this is may be due to the
increase of disorder in carbon phase on the modified surfaces and the most prominent band, always observed in graphite G sample, appears at 2600 cm$^{-1}$ in decorated carbon samples [22]. This outcome concurs with the surface modification of crystalline activated carbon materials over amorphous Vulcan carbon and confirm the XRD results. This result is important because the amorphousness/crystalline ratio of graphite directly impacts on the contact properties between the support and metal catalyst (i.e., Pd), which affected the mass-transport properties of the gas diffusion electrode [23, 24].

![XRD pattern of carbon materials.](image)

Figure 6. XRD pattern of carbon materials.

The XRD patterns of typical carbon and decorated carbon materials are depicted in Figure 6. Only two diffraction peaks at 25.4 and 43.2° were observed on the all the carbon materials, corresponding to the <002> and <101> planes of the hexagonal graphite (JCPDS, 75-1621) [25]. AC carbon showed the crystalline phase like sharp peaks. VC carbon exhibited broad peaks, while in the case of the modified carbon material the <101> plane revealed slightly broadening which may be due to the modification of crystalline activated carbon materials over the amorphous Vulcan carbon materials.

3.1.3. Conductivity analysis

Figure 7a shows the variation of the electrical conductivity ($\sigma$) with pressure. In all cases the value of $\sigma$ increases almost linearly with the compression exerted by the loadings. According to the contact theory by Mrozowski and Holm [26, 27], $\sigma$ depends on the distance between particles and on their average size [28, 29]. The three carbon samples (VC, AC and DAC) are constituted by particles, some of these particles can be fused together to form aggregates. When a pressure is applied the
aggregates are forced to be in a more dense packing arrangement, creating closer and new contacts with neighbouring aggregates increasing in this way the conductivity. Even though pressure exerts an important effect on $\sigma$, it is not the only factor that modifies its value. The morphology of the particle has also a great effect on the electrical conductivity [6, 30-31]. In the case of AC, the particles are irregular in shape and larger in comparison with the spherical particles of VC and DAC (Fig. 3). The particle size followed the order VC < DAC < AC while $\sigma$ followed the order DAC < VC < AC i.e., the greater $\sigma$ values of DAC cannot be explained by only taking into account the particle size. Other properties of carbons, such as the change in volume and density due to compression, may affect the value of $\sigma$.

![Figure 7](image)

**Figure 7.** Variation of different properties of three carbon samples: i) VC, ii) AC and iii) AVC.

In Figure 7b, it is observed that the change in volume was significant for the three samples at low pressures ($\leq$80 kPa). Contrary to this, at high pressures ($> 80$ kPa) the change in volume becomes less important (Fig. 7b). The increase of $\sigma$ as the density becomes larger (Fig. 7c); however, the change in density due to compression is clearly different for each sample. The increase in $\sigma$ may be explained taking into account that compression produces closer and new contacts with neighbouring aggregates; this in turn increases the value of $\sigma$, as described by the contact theory. In the present investigation it has been shown that pressure, the particle size, morphology, density and surface chemistry, have a combined effect on $\sigma$. 
4. CONCLUSIONS

A novel method of surface impregnation of decorated AVC materials was developed through intermittent microwave irradiation. Decorated AVC morphologies and textural features exhibited a high surface area and increased pore volume after the microwave irradiation. The AVC revealed a well distributed particle size (~90 to 100 nm), caused by the breakage of larger sized AC particles due to microwave irradiation during synthesis. The XRD patterns shows two diffraction peaks corresponding to the <002> and <101> planes of the hexagonal graphite. Raman analysis showed a predominant peak in G position for AVC sample; this may be due to an increase of disorder in AVC. For all the samples, the value of $\sigma$ increases almost linearly with experimental pressure, $\sigma$ values follows the order AVC < VC < AC. These modified materials could be used as a supporting material for metal electrocatalysts, adsorbents, and electrodes in microbial fuel cells.

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