Biomass Derived Activated Carbon/Epoxy Composite as Microwave Absorbing Material

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Research Article

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ABSTRACT
Herein, activated carbon/epoxy composite was synthesized, in which waste mango leaves biomass were used as carbon source. To create a sufficient amount of porosity in the biomass, a simple process of carbonization followed by its activation was used. The morphology and porosity of activated carbon material were studied by SEM, TEM images and BET analysis, which confirmed the presence of macro and mesopores in the material. The activated carbon/epoxy composite having 3 mm thickness gave the highest Reflection Loss (RL) of -39.57 dB at 6.48 GHz. At a thickness of just 2 mm the maximum effective absorption bandwidth (RL < -10 dB) of 2.14 GHz was achieved. This excellent microwave absorption performance was attributed to large surface area due to porous and bowl type structures, which resulted in high impedance matching and subsequent attenuation of microwaves by interfacial and dipolar polarization. Present study provides porous activated carbon/epoxy composite that is cost effective, lightweight and has high structural stability for microwave absorption applications.

KEYWORDS: Activated carbon, microwave absorbing material, biomass, composite materials, reflection loss.
1. INTRODUCTION

The rapid advancement in the area of electronics, communications, radar systems as well as increasing dependence on wireless technologies has resulted in an ever increasing electromagnetic pollution [1,2]. Therefore, the development of an efficient Microwave Absorbing Material (MAM) is in high demand. A high performance microwave absorbing material must display the properties like wide Effective Absorption Bandwidth (EAB), thin thickness, lightweight, high stability and low cost. Conventional MAM like ferrites and metal powders face drawbacks like heavyweight, poor chemical stability and require thick coating [1,3]. Recently carbon based MAM like carbon nanotubes [2,4–6], graphene [7–11] and graphene foam [12–14] has attracted attention because of their excellent properties like lightweight, high specific surface area, corrosion resistance and thermal stability [1,15], but due to high processing cost and complicated manufacturing techniques have limited their applications.

Recent studies have shown that biomass based porous carbon has the potential to become an efficient microwave absorbing material [1,3,15–19]. Most of the biomasses naturally possess network of microscale porous vascular bundles for water and ion transport, these built-in micropores contribute as an effective media in the composite, and improves the impedance matching and MW absorbing property of an absorber [1]. The porous materials facilitate multiple reflections of microwaves, which increases its path of propagation, leading to microwave energy loss inside the absorber material [3,20]. The solid-air interface formed due to porosity enhances the interfacial polarization hence attenuating a larger percentage of microwave. The porosity also helps to reduce the permittivity of the material which leads to better impedance matching [21]. Materials showing impedance close to air allow maximum amount of microwave to enter, hence reducing the direct reflection from the surface of the absorber material.

Apart from showing enhanced microwave absorbing properties, porous materials have low density which makes the composite lightweight. Therefore, biomass based porous materials have the potential to become efficient microwave material as they are cheap, lightweight, easy to scale and environment friendly. Our previous work has shown the excellent microwave absorption properties of waste mango leaves based activated carbon [16], it gave an excellent EAB of 5.17 GHz at thickness of just 1.75 mm.

But most of the literature published on biomass based activated carbon for MAM application had used paraffin wax as their matrix. There is hardly any study for activated carbon/epoxy composite as MAM. Therefore, in the present work, the microwave absorption properties of waste mango leaves based activated carbon (AC)/epoxy composite is reported. The addition of epoxy to AC was done to provide structural strength and stability. Epoxy resins as a thermosetting matrix display excellent mechanical properties. The ease of processing, low cost and good chemical resistance of epoxy resins makes it a promising candidate for producing structurally strong composite material [22]. In the present study the AC/epoxy composite sample exhibited a maximum Reflection Loss (RL) of -39.57 dB at a thickness of 3 mm.
2. EXPERIMENTAL SECTION

2.1. Materials
Mango leaves were accumulated from Institute premises, potassium hydroxide (KOH) was purchased from Merck life science Pvt. Ltd., Epoxy resin Huntsman Araldite LY 5052 and Hardener Huntsman LY Aradur 5052 were used as a matrix, and hydrochloric acid 35.4 wt% (HCl) AR grade was purchased from Loba Chemie Pvt. Ltd.

2.2. Synthesis of Activated Carbon
The mango leaves were first washed with tap water and then with distilled water to remove dirt. After washing, the leaves were naturally dried and then grinded into powder. The 15 g of grinded powder was then pyrolyzed at 800 °C for 1 hour in a vacuum tubular furnace with a ramp rate of 5 °C/min in nitrogen atmosphere. The resulting carbonized powder was thoroughly mixed with KOH (w/w 1:3) using mortar and pestle. The mixture was then put into a vacuum tubular furnace at 800 °C for 2 hours in nitrogen atmosphere for activation of carbonized mango leaves. The as prepared sample after activation was then washed with HCl (1:1) solution followed by distilled water using centrifuge, till pH becomes neutral. The sample was then dried in oven and named as Activated Carbon (AC). The activation process using KOH resulted in defects and porosity in AC. The AC and epoxy composite was then prepared by mechanically mixing AC (40 wt %) with epoxy resin (44 wt %) and hardener (16 wt%), the mixture was then pressed into toroidal shape (φ\text{out} = 7.0 mm, φ\text{in} = 3.0 mm) and cured at 100 °C for 4 hours. The resulting final composite was named as Activated Carbon Epoxy (ACE).

2.3. Characterizations
Powder X-ray Diffraction (XRD) of AC was done by Rigaku Miniflex II with Cu Kα radiation (λ=1.54 Å). Raman spectrum was obtained using Renishaw inVia Raman Microscope with laser excitation at 514 nm. Morphology of AC was studies using images of Scanning Electron Microscopy (SEM, JEOL JSM-6390LV) and High Resolution Transmission Electron Microscope (HRTEM, FEI Tecnai G2 20 S-TWIN). Nitrogen adsorption–desorption isotherm was measured on Quantachrome Nova 2200e at 77.3 K. Multiple points Brunauer–Emmett–Teller (BET) and Barrett, Joyner, Halenda (BJH) methods were used to calculate specific surface area and pore size distribution respectively.

2.4. Microwave absorption measurements
The microwave absorption properties of sample ACE were measured at 2 mm, 3 mm and 4 mm thicknesses. The relative complex permittivity (ε_r) and permeability (μ_r) measurement values were obtained by Agilent E8364B PNA series Vector Network Analyzer (VNA) in the frequency range of 2 to18 GHz using the Nicolson-Ross-Weir (NRW) algorithm [17]. The Reflection loss values were obtained using the metal backed values of S_{11}. 
3. RESULTS AND DISCUSSIONS

3.1. XRD and Raman analysis
Fig. 1(a) represents the XRD pattern of the AC. The broad peak at $2\theta = 23.4^\circ$ corresponds to (002) plane and a minor peak at $2\theta = 43.2^\circ$ corresponds to (100) plane. These planes confirms the graphitic carbon nature of AC [3,16].

Raman spectra of AC as shown in Fig. 1(b) displays a typical D band at 1359 cm$^{-1}$ and G band at 1597 cm$^{-1}$, these two bands are the characteristic signature of graphitic materials. The G band is due to the $E_{2g}$ vibrational mode present within aromatic carbon rings and it also represents the degree of graphitization [23], whereas D band arises due to defects and disorder introduced into sp$^2$ carbon. The intensity ratio value of the peaks $I_D/I_G = 0.95$, which is proportional to the degree of graphitization of carbon material [18].

Fig. 1 (a) XRD pattern and (b) Raman spectrum of AC.

3.2. BET and Morphological analysis
Nitrogen adsorption-desorption isotherm of AC has already been discussed in our previous study [16]. In brief the sample gave type IV isotherm suggesting the presence of mesopores in the sample [24,25]. The pore size distribution calculated by BJH with total pore volume was 0.274 cm$^3$/g, it clearly shows that the pore size of all the pores are above 2 nm which confirms the formation of mesoporous in AC. The activation process by KOH resulted in a network of mesopores and macropores, which resulted in a large specific surface area $S_{BET} = 683.146$ m$^2$/g. The morphology of AC was studied by SEM and HRTEM images. Fig. 2(a) clearly shows the presence of porous network formed within the sample due to the activation process, whereas Fig. 2(b) depicts a continuous bowl-shaped morphology which can be a result of partial porosity
induced during activation process. Fig. 2(c,d) depicts the HRTEM images which confirms the presence of mesopores within the sample.

![HRTEM images of AC](image)

**Fig. 2** (a, b) SEM and (c, d) TEM images of AC.

### 3.3. Microwave Absorption Properties

The microwave absorption properties of an absorber are mainly studied by two important parameters that are complex permittivity \( (\varepsilon_r = \varepsilon' - j\varepsilon'') \) and complex permeability \( (\mu_r = \mu' - j\mu'') \). Where the real parts \( (\varepsilon' \text{ and } \mu' \text{)} \) corresponds to energy storage and imaginary parts \( (\varepsilon'' \text{ and } \mu'' \text{)} \) corresponds to energy dissipation [17,21]. The ACE is non-magnetic therefore the values of \( \mu_r \) were taken to be 1, hence the microwave absorption performance shown by ACE was solely because of dielectric part. Fig. 3(a) represents the relation of \( \varepsilon' \text{ and } \varepsilon'' \text{ with frequency.} \)
The $\varepsilon'$ value for the sample at initial frequency range remains nearly constant till 8.6 GHz with minor variations, reaching a maximum value of 17.2 at 6.8 GHz.

Fig. 3 Frequency dependence of (a) Relative permittivity, (b) Dielectric loss tangent and attenuation constant for ACE.

But thereafter, $\varepsilon'$ value decreases with increasing frequency. Whereas $\varepsilon''$ value increases with increasing frequency reaching a maximum value of 11.9 at 16.7 GHz and then shows a decreasing trend. The large values of $\varepsilon'$ maybe attributed to the interfacial polarization and dipolar polarization phenomenon [19], the porous structure of AC results in the increase in surface area which facilitates the accumulation of electrons at the pore-solid interfaces [3], resulting in enhanced interfacial polarisation of microwaves [15].

In order to study the materials microwave energy loss ability, dielectric loss tangent [15,26] and attenuation constant [27,28] were calculated using the equation 2 and 3 respectively:

$$\tan \delta = \varepsilon''/\varepsilon' \quad (1)$$

$$\alpha = \frac{\pi f}{c} \left(2 \left(\mu'' \varepsilon'' - \mu' \varepsilon' + \left((\mu''^2 + \mu'^2)(\varepsilon''^2 + \varepsilon'^2)\right)^{1/2}\right)\right)^{1/2} \quad (2)$$

Fig. 3(b) depicts the relation of dielectric loss ($\tan \delta_e$) and attenuation constant ($\alpha$) of ACE with frequency. The value of $\tan \delta_e$ as well as of $\alpha$ increases with increase in frequency, reaching a maximum value of 1.95 and 707 at 18 GHz respectively.

Fig. 4(b) depicts impedance matching (Z) curve for ACE at different thicknesses, which is an important parameter that effects the performance of an absorber [2,17,20], and represented as:

$$Z = |Z_{in}/Z_0| = \sqrt{\mu_r/\varepsilon_r \tanh \left( j \frac{2\pi f d}{c} \sqrt{\mu_r \varepsilon_r} \right) } \quad (3)$$
Where $Z_{in}$ is the input impedance of an absorber, $Z_0$ is the impedance of free space, $c$ is the speed of light, $f$ is the frequency of the electromagnetic waves and $d$ is the thickness of the absorber.

![Fig. 4](image.png)

**Fig. 4** Frequency dependence of (a) Reflection loss and (a) $|Z_{in}/Z_0|$ for ACE.

Closer the value of $Z$ to 1 means better impedance matching, which allows more amount of microwave to enter into an absorber contributing in better microwave absorbing performance. Fig. 4(a) shows the RL curves of ACE sample at different thicknesses. At 2 mm thickness a maximum EAB of 2.14 GHz was achieved. Whereas the highest reflection loss of -39.57 dB at 6.48 GHz was obtained at 3 mm thickness. By comparing Fig. 4(a) and Fig. 4(b) it is clearly evident that the maximum RL for ACE at different thicknesses was achieved when there is maximum impedance matching.

This enhanced performance by ACE can be attributed to high impedance matching allowing more microwaves to enter into absorber and then subsequent attenuation of microwaves by interfacial polarization and dipolar polarization.
4. CONCLUSIONS
Waste biomass (mango leaves) derived Activated Carbon and Epoxy composite (ACE) was synthesized as a microwave absorbing material. The porosity generated by KOH activation process provided large surface area ($S_{BET}$) = 683.146 m$^2$/g, which facilitated high impedance matching and attenuation of microwaves. The highest reflection loss of -39.57 dB was obtained at 3 mm thickness and a maximum EAB of 2.14 GHz was achieved at 2 mm thickness. Thus, activated carbon/epoxy composite is a porous, cost-effective, lightweight composite material and has the potential as microwave absorbing material.

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Figures

Figure 1

(a) XRD pattern and (b) Raman spectrum of AC.
Figure 2

(a, b) SEM and (c, d) TEM images of AC.
Figure 3

Frequency dependence of (a) Relative permittivity, (b) Dielectric loss tangent and attenuation constant for ACE.

Figure 4

(a) Reflection loss for different thicknesses (2 mm, 3 mm, 4 mm).
(b) $|Z_m/Z_0|$ vs. Frequency for different thicknesses (2 mm, 3 mm, 4 mm).
Frequency dependence of (a) Reflection loss and (a) $|Z_{in}/Z_o|$ for ACE.