Analytical Investigation of Microcrystalline Wood Charcoal Reinforced Polyester Composites Using ED-XRF, FTIR and SEM-EDS Techniques

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Highlights
• This paper focuses on functional and chemical characterization of polymer matrix composites.
• 75µm wood charcoal particles were successfully utilized as reinforcing phase for polyester matrix.
• FTIR study revealed interactions between wood charcoal particles and functional groups in polyester.

Abstract
Various weight fractions (ranging from 0 to 30 wt percent, at 5 wt percent intervals) of microcrystalline wood charcoal powder (75µm) were used to generate polyester-based particle reinforced composites. The developed wood charcoal (WC) particles reinforced polyester matrix composites were successfully characterized quantitatively, qualitatively, and functionally using an energy dispersive x-ray spectrophotometer (EDXRF), scanning electron microscope (SEM) enhanced with ancillary EDS for elemental identification capability and Fourier transform infrared spectrometer (FTIR). High peaks of Fe₂O₃ (3456cps/mA), CaO (4369cps/mA), CuO (2017cps/mA), ZnO (1646cps/mA) and Iron, calcium, copper, zinc were discovered in wood charcoal by EDXRF. The characteristics of reinforced polymer composites were found to be improved by these elements and oxides. The key elements revealed by EDXRF analysis were also validated by EDS elemental mapping. Due to the uniform distribution of the filler particles, SEM images showed that composites with microcrystalline wood charcoal reinforcement had strong interfacial adhesion and interlocking. According to FTIR functional analysis, interactions between the polyester matrix molecules and the microcrystalline wood charcoal fillers resulted in very slight alterations in the frequency bands of functional groups that are typically present in unsaturated polyester resin.

1. INTRODUCTION
Monolithic materials are becoming less prevalent across a wide range of technological sectors as a result of the constantly rising demands of today’s existing and new technologies on high product capabilities, functions, and performance [1]. Several materials can be combined in a natural or synthetic way to create materials with improved qualities and performance. Wood for instance, is a natural composite made up of cellulose (wood fibers) strongly held together by a matrix of lignin. Most structural and functional composite materials used today are man- made and composed of two or more component parts that are macroscopically combined, but does not form soluble mixture [2,3]. In the second constituent, the matrix phase, one constituent known as the reinforcing phase is introduced. Two frequently used approaches for classifying composite materials, the one based on the reinforcing geometrical dimension (which could be particle, flake, or fiber reinforced composites) and the other based on the type of matrix material employed (the matrix could be metallic, ceramic, polymeric or carbon material) [4]. Polymer matrix composites (PMCs), have comparatively high strength, low cost and simple processing routes, thus, they are the most widely used composite materials [5]. Low service temperatures, high moisture absorption, enhanced thermal expansion coefficients, and low directional elastic characteristics are some of the reasons for their
limits in specific applications. During the design and development process, composite materials can be
tweaked or tailored to improve their qualities and expand their uses [6–9]. Selection of appropriate
components, component modification, and composite interface engineering are all strategies for tailoring
composite materials. The reasons for tailoring in composite materials is to obtain a trade-off in properties’
requirement for specific applications [10,11].

The emergence of modern rapid analytical techniques and equipment has made it possible to study the
nature, structure and composition of natural and synthetic materials. Materials characterization can be
accomplished using a variety of analytical techniques and equipment [12]. Most of these accessible
materials characterization techniques and tools use probe photon, electron, or ion beams on the material
being studied. The interaction of these probe beams with the material or changes made to the intensity,
energy level, and angular spread of the beam are observed and measured to acquire the analytical
information. The ions, electrons, or photons emitted from the sample under probe beam bombardment may
be examined in various procedures. Only one element of the produced alterations could be identified using
a particular analytical technique, such as the amount of incident light absorbed or the kinetic energy level
and distribution of expelled electrons. In many instances, numerous interrelated processes are at work at
the same time. Chemical composition data (oxide or elemental analysis), chemical state and properties,
phase identification, and structural determination are among the types of information offered by these
procedures (such as length of bonds, atomic lattice sites, and angles).

Common analytical techniques and equipment for quantitative, qualitative and functional chemical
characterizations of composites materials include: scanning electron microscope with an additional energy
dispersive X-ray spectroscopy detector (SEM-EDS); energy dispersive X-ray fluorescent spectroscopy
(EDXRF); and Fourier transform infrared spectroscopy (FTIR) [12–14]. Infrared spectrometry equipment
is an effective tool in analytical laboratories for determining the presence of characteristics molecular
functional groups. The relative IR absorbance or transmittance of a substance is plotted against the
appropriate vibrational frequency in an infrared (IR) spectrum. IR radiation's energy is converted into
atomic vibrations when it is absorbed or transmitted by a substance. A change in the bond's length, known
as stretching, or a shift in the bond angle, known as bending, can be caused by the energy absorbed or
transmitted by an atomic bond. Depending on the stiffness of the connection and mass reduction of the
linked atoms or functional groups, these stretching and bending events resonate at a particular frequency
[15]. Verification and evaluation of chemical components and impure elements are also indispensable in
materials analysis. Hence, elemental quantification and oxide analysis using energy dispersive X-ray
fluorescent spectroscopy (EDXRF) with high reliability rank first for this kind of analysis. SEM is
commonly employed for high field resolution morphological examination of materials. SEM could be
equipped with elemental characterization capability by the attachment of an EDS detector. The chemical
states and composition of unsaturated polyester-based composites with microcrystalline wood charcoal
particle reinforcement were characterized in the current work using these analytical instrumental
techniques.

2. MATERIALS AND METHOD

2.1. Preparation of Wood Charcoal Reinforcement Materials

Dry lumps of wood charcoal obtained locally were pulverized using a grain mill (model No. 116). Standard
BSS 200 sieve size was used for sieving the milled wood charcoal fillers to obtain microcrystalline particles
of 75 μm size.

2.2. Development of Wood Charcoal/Polyester Composites

The wood charcoal (WC) particles reinforced polyester composites were produced gradually using the open
mould method. Each sample had a total composite volume of 98.21 cm³ after the volume of the polyester
resin and the microcrystalline wood charcoal fillers were adjusted. A calibrated glass cylinder and an
electronic weighing scale were used, respectively, to determine the necessary amounts of polyester matrix
and wood charcoal filler. To avoid filler agglomeration and promote faster and more homogeneous blending
of the reinforcement particles in the matrix, the measured matrix and reinforcement materials were mixed in a 200ml beaker and stirred for 10 minutes with a long glass rod. The catalyst, methyl-ethyl-ketone peroxide (MEKP), was introduced using a disposable syringe at a volume ratio of 0.2ml catalyst to 10ml polyester resin and swirled continuously for two minutes. Cobalt naphthanate, an accelerator, was injected using a different disposable syringe at a volume fraction of 10ml polyester resin to 0.1ml accelerator. The mixture was then stirred for an additional two minutes. The composite mixture was then carefully poured into the individual test specimens’ prepared wooden molds, where they were left to cure for 24 hours before being taken out of the molds. Before pouring, a petroleum jelly release agent was put to the wooden mold to make it easier to remove the composites from the molds. Six different weight fractions (5, 10, 15, 20, 25 & 30wt. %) of the WC reinforcements were used to develop a total of six composite compositions as well as the unreinforced polyester matrix.

2.3. Functional Characterization

Agilent Technologies' Cary 630 FTIR scientific instrument was used to analyze the FTIR of produced composites in accordance with ASTM E168 specification. The transmittance method was used to record the spectra during the Fourier transform infrared spectroscopy (FTIR) investigation, and an 8 cm$^{-1}$ resolution FTIR spectrum was acquired in the range of 650 to 4000 cm$^{-1}$ wave number. It was able to ascertain the details of the chemical bonds (functional groups).

2.4. Chemical Composition Analysis

The elemental and oxide content of the wood charcoal was quantitatively examined and analyzed using the XRF technique. A “Minipal 4” EDXRF spectrophotometer model, capable of detecting elements lying between the atomic numbers of sodium (Na, Z = 11) and Uranium (U, Z = 92) with great resolution and quick analysis was used in this case.

2.5. SEM-EDS Examination

Etching of the prepared composite samples was carried out at room temperature for about 10s in HNO$_3$ and HF solution, mixed in a volume ratio of 1:12, and then analyzed with a Phenom Pro X Model SEM (Phenom world, Eindhoven, Netherlands) fitted with an EDS detector, which gives the SEM analytical capabilities. To boost the electrical conductivity of the SEM samples, they were placed on a conductive carbon impression created by the adhesive tape and coated for 5 minutes. An accelerating voltage of 15 kV was used to scan the samples. The SEM image is electronically sent to an EDS detector with appropriate elemental mapping software for automatic identification and quantification of the primary, secondary, and trace elements in the polyester composite samples.

3. THE RESEARCH FINDINGS AND DISCUSSION

3.1. Characterization of the Wood Charcoal Fillers by XRF Method

The EDXRF spectrometer was employed to ascertain the elemental and oxide chemical composition of the microcrystalline wood charcoal filler. In wood charcoal powder, high peaks of calcium, iron, zinc, and copper can be seen in Figure 1.
The presence of high quantities of these elements was also confirmed by the oxide analysis in Table 1. It is opined that the presence of high peaks of these hard metal oxide should improve the mechanical characteristics of microcrystalline wood charcoal reinforced polyester composites. As a result, it is increasingly used as a particle reinforcement for composites based on metal and polymer matrix.

**Table 1. XRF oxide analysis of the wood charcoal reinforcement**

<table>
<thead>
<tr>
<th>Compound (Oxide)</th>
<th>Peaks (cps/mA)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CaO</td>
<td>4309</td>
</tr>
<tr>
<td>Fe$_2$O$_3$</td>
<td>3456</td>
</tr>
<tr>
<td>CuO</td>
<td>2017</td>
</tr>
<tr>
<td>ZnO</td>
<td>1636</td>
</tr>
<tr>
<td>SiO$_2$</td>
<td>531</td>
</tr>
<tr>
<td>Al$_2$O$_3$</td>
<td>496</td>
</tr>
<tr>
<td>MgO</td>
<td>123</td>
</tr>
<tr>
<td>P$_2$O$_5$</td>
<td>308</td>
</tr>
<tr>
<td>TiO$_2$</td>
<td>173</td>
</tr>
<tr>
<td>MnO</td>
<td>439</td>
</tr>
<tr>
<td>K$_2$O</td>
<td>724</td>
</tr>
<tr>
<td>WO$_3$</td>
<td>52</td>
</tr>
<tr>
<td>SrO</td>
<td>33</td>
</tr>
<tr>
<td>ZrO$_2$</td>
<td>22</td>
</tr>
<tr>
<td>BaO</td>
<td>21</td>
</tr>
<tr>
<td>V$_2$O$_5$</td>
<td>15</td>
</tr>
<tr>
<td>Cr$_2$O$_3$</td>
<td>12</td>
</tr>
<tr>
<td>Na$_2$O</td>
<td>5</td>
</tr>
<tr>
<td>Rb$_2$O</td>
<td>5</td>
</tr>
<tr>
<td>Ga$_2$O$_3$</td>
<td>4</td>
</tr>
<tr>
<td>PbO</td>
<td>2</td>
</tr>
</tbody>
</table>
3.2. Functional Analysis by Infrared Radiation (FTIR)

For ease of comprehension and presentation, the vibrational frequency of an IR photon is generally translated to wavelength and then inverted. A wavenumber (λ) is a unit of measurement that is symbolized by cm$^{-1}$. Three different regions make up the infrared spectrum: far infrared (0-400 cm$^{-1}$), mid infrared (400 – 4,000 cm$^{-1}$), and near infrared (4000-14,285 cm$^{-1}$) [15]. The mid-infrared range contains a substantial percentage of molecule structure information and functional groups for most practical applications. Specialized information, such as lattice vibrations, is provided by the near- and far-infrared regions. There are four spectrum sections in the mid-infrared region: the fingerprint region (400-1500 cm$^{-1}$), the double bond region (1500 – 2000 cm$^{-1}$), the triple bond region (2000 – 2500 cm$^{-1}$), and the X–H stretching region (2500 – 4000 cm$^{-1}$). The FTIR operator uses the signals that emerge in these to identify most molecular structures, functional groups, and organic bonds by comparing them to the infrared correlation chart [15,16].

![FTIR spectra of microcrystalline WC (15wt%) particles reinforced polyester matrix composite](image)

**Figure 2.** FTIR spectra of microcrystalline WC (15wt%) particles reinforced polyester matrix composite

The FTIR spectra of the polyester matrix reinforced with 15 weight percent particulate wood charcoal is shown in Figure 2. Strong stretching in O-H groups resulted in a high transmittance of 94.904 in the FTIR spectra for the band at 3477.6 cm$^{-1}$. This H-bonded O-H stretch is generally found in the frequency region of 3500 – 3200 cm$^{-1}$ on the infrared correlation chart for alcohols and phenols [15]. This band for unsaturated polyester has been observed by other researchers at 3448 cm$^{-1}$ [17]. The C-H bond could be responsible for the medium intensity stretching measured at 2881.2 cm$^{-1}$ with a transmittance of 59.880 (this is usually found in both alkanes and aldehydes) [18]. The high transmittance of 93.684 at 1744.4 cm$^{-1}$ could be due to strong C=O stretching from unsaturated ketones and aldehydes. Researchers elsewhere found the same intense stretching vibration in unreinforced polyester at 1728 cm$^{-1}$ [17]. The band at 1654.2 cm$^{-1}$ could be linked to a medium bend of N-H functional groups present in amines. A moderate intensity CH$_3$ asymmetrical bending of C-H groups [18] and moderate stretching of the C-C bond observed in aromatics may be matched to the band at 1464.8 cm$^{-1}$ with a transmittance of 76.723. The significant stretching vibration of C-O is connected to the FTIR spectrum range 1282.2 cm$^{-1}$ (normally observed for carboxylic acids, alcohols, ethers and esters functional groups). The C-N stretching vibration is matched to the medium
transmittance IR band at 1241 cm\(^{-1}\). C-O stretching seen at 1282.2 cm\(^{-1}\) is considered to be the cause of weak molecular interactions between CH\(_2\) groups and reinforcements [17]. The FTIR band range 1144.3-1103.3 cm\(^{-1}\) may be related to the moderate stretching vibrations of C-N seen in functional groups of aliphatic amines. FTIR band 1341.8 cm\(^{-1}\) revealed moderate symmetrical stretching of N=O observed in nitro compounds at a transmittance of 63.676. The moderate intensity stretching vibration of the C-Cl bond observed in alkyl halides is usually fixed at 850 – 550 cm\(^{-1}\) range of frequency on the infrared correlation chart [15], but this moderate intensity stretching vibration of the C-Cl bond was seen at 842.4 cm\(^{-1}\) with a transmittance of 51.347 in the current investigation. The medium IR band at 946.7 cm\(^{-1}\) has been linked to O-H bending vibrations, which are common in carboxylic acids. The significant IR transmittance band at 719.4 cm\(^{-1}\) could be due to C=O stretching vibrations present in saturated and unsaturated esters.

### 3.3. SEM Morphology and Energy Dispersive x-ray Spectroscopy (EDS) Examination

The elemental contents of the composites were disclosed by the EDS mapping and spectra provided in Table 2 and Figure 3. Ca, K, C, Si, Ag, Cl, P, S, and Al were the major constituents in the composites, with traces of Mg, Na, Fe, and Ti in the 15% wood charcoal composite. The quantities of the various chemical ingredients in the composites was controlled by the filler. Figure 3 also shows a SEM micrograph of a 15 wt% wood charcoal reinforced polyester composite. On the micrographs, there are fewer regions where wood charcoal particles agglomerates can be seen. Agglomerations of reinforcement particles in certain zones of composite materials have been attributed to the generation and concentration of stress in these zones, which could act as fracture initiation sites [19]. There is substantial interfacial adhesion, as observed in the micrograph. The porous structure of WC and presence of volatile elements could explain the improved filler-polymer matrix interfacial bonding seen in the SEM image [9,20–23]. Moreover, the filler phase has a homogeneous distribution inside the matrix phase.

#### Table 2. EDS elemental table of 15 wt% wood charcoal particles reinforced polyester matrix composite

<table>
<thead>
<tr>
<th>Element Symbol</th>
<th>Weight Concentration (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>C</td>
<td>77.49</td>
</tr>
<tr>
<td>Ca</td>
<td>5.82</td>
</tr>
<tr>
<td>Ag</td>
<td>4.39</td>
</tr>
<tr>
<td>K</td>
<td>2.67</td>
</tr>
<tr>
<td>Si</td>
<td>2.65</td>
</tr>
<tr>
<td>Cl</td>
<td>1.82</td>
</tr>
<tr>
<td>S</td>
<td>1.59</td>
</tr>
<tr>
<td>P</td>
<td>1.55</td>
</tr>
<tr>
<td>Al</td>
<td>0.93</td>
</tr>
<tr>
<td>Mg</td>
<td>0.39</td>
</tr>
<tr>
<td>Na</td>
<td>0.50</td>
</tr>
<tr>
<td>Fe</td>
<td>0.00</td>
</tr>
<tr>
<td>Ti</td>
<td>0.00</td>
</tr>
</tbody>
</table>
Figure 3. SEM micrograph and EDS Spectra for 15 wt% microcrystalline wood charcoal reinforced polyester matrix composite

4. CONCLUSION

The key chemical components in the polyester matrix composites reinforced with microcrystalline wood charcoal, as determined by various characterization tools and methodologies, are iron, copper, zinc, and calcium, while the predominant oxides in the wood charcoal are CaO, Fe₂O₃, CuO, and ZnO, respectively. Owing to the homogenous distribution of filler particles in the polyester matrix, the porous nature, and the content of volatile compounds described in wood charcoal, SEM micrographs revealed significant filler-matrix interfacial adhesion produced with more wood charcoal. Slight shifts in the assigned peaks for unsaturated polyester were seen in the infrared radiation spectrum acquired in the 600 – 4000cm⁻¹ wave number range, which was induced by interactions between the wood charcoal particles and the functional chemical species in polyester molecules.

CONFLICTS OF INTEREST

No conflict of interest was declared by the authors.

REFERENCES


