Eco-friendly polyvinyl alcohol (PVA)/bamboo charcoal (BC) nanocomposites with superior mechanical and thermal properties

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Abstract

Carbon-based nanofillers, such as carbon nanotubes (CNTs) and graphene sheets are considered as effective nanoreinforcements due to their unique structures and material performance. However, the utilisation of such nanofillers can be hindered owing to a high level of nanotoxicity via human inhalation and high material cost for CNTs, as well as the tendency to form agglomerates of graphene sheets in polymer matrices. Bamboo charcoals (BCs) are eco-friendly and sustainable carbon-based particles, which possess good affinity with polyvinyl alcohol (PVA), one of popular water soluble biopolymers, to achieve excellent properties of PVA/BC nanocomposites. In particular, porous structures of BC particles enable polymeric molecules to easily penetrate with the strong internal bonding. In this study, fully eco-friendly PVA/BC nanocomposite films were successfully fabricated using a simple solution casting method to achieve the high dispersibility of BCs. With the inclusion of only 3 wt% BCs, tensile modulus and tensile strength of PVA/BC nanocomposite films were enhanced by 70.2 and 71.6%, respectively when compared with those of PVA films. Better thermal stability is manifested for resulting nanocomposite films as opposed to that of pristine PVA, which is evidenced by the maximum increase of 17.8% in the decomposition temperature at the weight loss of 80%. It is anticipated that BCs can

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compete against conventional carbon-based nanofillers with a great potential to be developed into eco-friendly nanocomposites used for thin-film packaging application.

**Keywords**: Polyvinyl alcohol (PVA); bamboo charcoals (BCs); nanocomposites; mechanical properties; thermal stability

1 Introduction

The annual global production of plastics is expected to exceed 300 million tons by 2015 [1]. However, the majority of these plastics belong to synthetic polymers such as polypropylene (PP), polyethylene (PE), and polystyrene (PS). Synthetic polymers include a huge amount of non-degradable wastes and cause a serious depletion of landfill capacities. [2] Furthermore, the disposal of synthetic polymers becomes more formidable considering that many of these polymers resist physical and chemical degradation [3]. As such, synthetic polymers can be alleviated by developing biodegradable polymers such as water-soluble PVA, which has received considerable attention due to its favourable mechanical properties, thermal resistance, excellent flexibility [4] and recyclability, and bio-tribological properties [5]. It is also worth noting that these aforementioned properties of PVA make it a suitable material candidate for a wide range of biotechnological applications from tissue engineering, drug delivery, articular cartilage to biosensors[4-6].

In recent years, many researchers have focused on using nanomaterials or nanofillers to reinforce PVA in order to significantly enhance its mechanical and thermal properties, as well as biodegradability. Those nanofillers comprise montmorillonite (MMT) clays[7-11], halloysite nanotubes (HNTs) [12-15], CNTs [16,17], graphene sheets,[18-21] cellulose nanocrystals [22-24], laponite [25] and nanodiamond [26].

Notwithstanding the effective reinforcement role of CNTs, major drawback of CNTs lies in their nanotoxic characteristic to accumulate in cytoplasm and finally destroy human cells in a
suitable inhalational condition in addition to high production cost. [2] On the other hand, graphene sheets can also undergo the limited use owing to their high tendency to form agglomerates[27].

More recently, the emphasis has been laid on material development to utilise bamboo charcoals (BCs) as carbon-based eco-friendly and functional materials. Such BC particles are generally made from rapidly growing bamboo carbonised at the high temperature about 1000°C under the anerobic condition [28]. BCs consist of short stacks of graphene sheets in connection to form 3D networks with innumerable lengthwise and crosswise pores [29-30]. Moreover, their volumetric porosity, mineral constituents and absorption efficiency are about 5, 8 and 10 times more than those of wood charcoals.[30] In particular, the inner surface area of BCs is about 250-390 m²/g, as opposed to 10 m²/g in the case of wood charcoals [31-32]. These remarkable features of BCs lead to various applications including suppliers for negative ions [33] humidity regulators [34], water purification [35], oxidation prevention, anti-bacterial [36] and anti-fungal features and breathability[32]. The humidity control of BCs enables them to inhibit the growth of bacteria and fungi to maintain a fresh food level [34, 37]. This approach is very important for food packaging industries so as to increase the shelf life of products. Additionally, BCs are also capable of absorbing far more infrared energy from environment, which is emitted to support the cell activation for efficient human blood circulation [38]. The elemental inclusion such as calcium, potassium, sodium and iron within BCs is essential for the purposes of food cooking, baking and storage [39, 40].

In a nanocomposite system, BC morphology displays a wide range of pore distribution from <1 nm to 1 µm. The walls of basic unit inside BCs, known as parenchyma, are very rough along with their smooth outer surfaces to make the whole charcoals quite hard. A great many pores associated with rough walls inside BCs may be able to promote the effective interfacial bonding between fillers and polymer matrices, thus resulting in improved mechanical
properties of composite materials. This is believed to arise from the penetration of polymeric chains into internal pores of BCs leading to strong mechanical bonding in addition to the hydrogen bonding of pores. Main research work has focused on BCs as microparticles to reinforce polymers in composite systems such as polylactic acid (PLA)/BC composites[41], ultra-high molecular weight polyethylene (UHMWPE)/BC composites [42] and polyaniline (PANI)/BC composites[43]. Up to now, very limited published work is available regarding the effect of BC nanoparticles on mechanical and thermal properties of PVA/BC composite films. In this study, a simple synthetic approach of PVA/BC nanocomposite films was employed based on the solution casting method. The effects of BC content on material characterisation, mechanical and thermal properties of final nanocomposite films were investigated in detail to evaluate the potential of BCs as new reinforcing nanofillers used in the eco-friendly packaging field.

2. Experimental details

2.1 Materials

PVA biopolymer MFCD00081922 (average molecular weight $M_w = 89000-98000$ g/mol and the degree of hydrolysis at over 99%) was obtained from Sigma Aldrich Pty. Ltd, NSW, Australia. BC nanoparticles (particle size: 700-800 nm) used in this study were purchased from Jiangshan Luyi Bamboo Charcoals Co., Ltd, China.

2.2 Fabrication of PVA/BC nanocomposite films

PVA/BC nanocomposites were prepared by a simple solution casting method. Initially 5% PVA aqueous solution for stock was prepared by dissolving 10 g PVA into 190 ml deionised water via vigorous magnetic stirring at 400 rpm and 90°C for 3 h until PVA was completely dissolved. The aqueous suspension of BC nanoparticles was obtained using mechanical
mixing in deionised water at 405 rpm and 40°C for 2 h. This was followed by the ultrasonication with the aid of an ultrasonic cleaning unit ELMA Ti–H–5 at 25 kHz and 40°C with the power intensity of 70% for 1 h. The BC contents in range from 0, 3, 5 to 10 wt% were obtained by adding different amounts of PVA. BC aqueous suspension was then gradually dripped in a dropwise manner into PVA solutions, simultaneously subjected to mechanical mixing at 405 rpm and 40°C for 2h. Then their mixture was stirred at 400 rpm and 90°C for 1 h prior to the subsequent similar sonication for 30 min to achieve uniform dispersion of BC nanoparticles. Finally, 20 ml solution was cast on a glass petri dish and further allowed to dry using an air-circulating oven at 40°C for 48 h. Then PVA/BC nanocomposite films were peeled off from petri dishes and stored in a desiccator with underneath silicon gels prior to further material testing and analysis.

2.3 Material Characterisation

Tensile properties of PVA/BC nanocomposite films were measured on a Lloyd EZ50 universal testing machine with a crosshead speed of 10 mm/min (initial gauge length: 50 mm). For each material batch, six samples were tested and average data with associated standard deviations were calculated accordingly.

The surface morphology of fractured tensile testing samples was observed by a field emission scanning electron microscope (FESEM, Zeiss NEON 40 EsB Cross Beam) at an accelerating voltage of 5 kV. These samples were platinum coated prior to surface morphological analysis.

Thermal stability of nanocomposite films was examined with a TGA/DSC 1 STArE system (METTLER TOLEDO) from 35 to 700°C at a scan rate of 10°C/min under argon atmosphere (flow rate: 25 ml/min).
Carbon, hydrogen, nitrogen and oxygen contents of BCs were measured by a CHNS/O element analyser at 900°C (Perkin Elmer 2400 Series II).

Fourier transform infrared (FTIR) spectrometry (100FT-IR Spectrometer-Perkin Elmer) was employed to characterise PVA, BC nanoparticles and PVA/BC nanocomposites at wave numbers ranging from 650-4000 cm\(^{-1}\) with a resolution of 4 cm\(^{-1}\) using an attenuated total reflectance (ATR) method.

XRD analysis was conducted via Bruker D8 Advance Diffractometer. The X-ray source was Ni-filtered Cu-K\(\alpha\) radiation at the accelerating voltage and current of 40 kV and 40 mA with the X-ray spectra recorded in range of 10\(^{\circ}\) -50\(^{\circ}\) at the scan rate of 0.015\(^{\circ}\)/sec.

3 Result and discussion

3.1 BC characterisation

Chemical composition of BCs generally consist of carbon, oxygen, hydrogen, nitrogen and a small amount of ash. [44]. In this study, the carbon content of BCs is 80.04%, which is in good accordance with previous results obtained by Li et al, [44]. It is suggested that BCs could be regarded as good carbon-based fillers for effective reinforcements in nanocomposite systems. Oxygen, hydrogen and nitrogen contents for BCs are less pronounced (O\(\%\): 6.93\%, H\(\%\): 2.54\% and N\(\%\): 0.52\%). Moreover, surface area and pore volume have also been detected for BCs in order to evaluate BC absorption capacity within polymer matrices. Experimental results reveal that all three major surface areas including Brunauer–Emmett–Teller (BET) surface area \((S_{BET})\), micropore area \((S_{micro})\) and external surface area \((S_{ext})\) are 527.41, 390.84 and 136.57 m\(^{2}\) g\(^{-1}\), respectively. As such, BCs demonstrate much better absorption capability within polymer matrices, thus resulting in more efficient load transfer from matrices to fillers as mentioned elsewhere [45].
The FTIR of BC particles was carried out to investigate the surface functionality of as-received particles, Figure 1(a). The peak at about 2439.6 is assigned to the C≡H stretching [46]. Whereas the peaks at about 1566.6 cm⁻¹ represent C=C vibration in an aromatic system [45]. The peak at 1695.9 cm⁻¹ is corresponding to the C=O reign primarily for ionisable carboxyl groups as an indicator of surface hydrophilicity [47]. The band existing at 1111.1 cm⁻¹ is ascribed to the axial deformation of C-O band. Moreover, the band at 872.1 cm⁻¹ is referred to as the C-H bending (in plane). The out of plane –OH bending is also designated by the band appearance at 743.6 cm⁻¹ for BCs. Finally, missing –OH peak at 3350 cm⁻¹ in BC spectra infers that BCs possess much lower moisture and alcohol contents [48].

Moreover, XRD pattern of BCs, as depicted in Figure 1(b), reveal the existence of two broad peaks. The board peaks at 2θ = 22.9° is corresponding to the sharp peaks of graphite, which is assigned to the (002) diffraction plane [49]. Moreover, second broad peak at 2θ = 43.6° characterises 2D in-plane symmetry along with graphene layers, corresponding to (100) diffraction plane.

### 3.2 PVA/BC Composite film properties and characterization

FTIR analysis was carried out to evaluate possible interactions between PVA and BCs, as illustrated in Figure 2 (a), indicating that the assigned –OH stretching and –C–OH stretching bands can be used to form hydrogen bonding. The wave number band ranging from 3100-3500 cm⁻¹ is assigned to the strong hydroxyl band for free and hydrogen bonded alcohols, which shifts to a lower number after the incorporation of BCs into PVA matrices. For instance, the wave number at 3271.5 cm⁻¹ for pure PVA film shifts to 3262.0 cm⁻¹ in the case of PVA/3 wt% BC composites, which is in good agreement with previous result for PVA/graphene nanocomposites [41]. This phenomenon is attributed to the dissociation of hydrogen bonding between hydroxyl groups in PVA and subsequent formation of hydrogen bonding between –OH groups within BCs and PVA matrices [50, 51]. Moreover, the wide
peak range between 2940 and 2845 cm\(^{-1}\) is known to be associated with (–CH\(_2\)–) asymmetric
and symmetric stretching, respectively. The increasing tendency of peaks implies that BC
nanoparticles are strongly bonded to PVA matrices owing to the close connection formed by
hydrogen bonding, which is similar to PVA/graphene oxide (GO) nanocomposites [52].

XRD patterns for pure PVA and PVA/BC composite films are shown in Figure 2(b). Pure
PVA films reveal a strong maximum diffraction peak at \(2\theta = 19.7^\circ\), corresponding to the
total crystalline plane (101) of PVA. [53]. Moreover, XRD peaks for PVA/BC composites
present a slight shift to larger diffraction angles, which is mostly likely due to the interaction
between PVA and BCs.

Mechanical properties of PVA/BC nanocomposite films in term of tensile strength, tensile
modulus and elongation at break are shown in Figure 2(c) and Table 1. It can be seen that the
tensile modulus of nanocomposite films increases with increasing the BC content. PVA/BC
nanocomposites reinforced with 3wt% BCs have achieved a tensile modulus of 3.54 GPa
with an modulus increase of 70.2% compared to that of neat PVA. This result can be
interpreted by the restriction of BCs to the chain motion of PVA matrices, thus decreasing
PVA deformation capacity in the elastic region to make nanocomposites much stiffer [50-52].

In general, tensile strength is an indicator in mechanical properties to evaluate the interfacial
bonding between fillers and polymer matrices within composite materials. As illustrated in
Figure 2(c), tensile strengths of PVA/BC nanocomposites increased at BC contents from 0 to
3 wt%, and then they decreased at the higher content levels up to 10 wt%. With the inclusion
of 3 wt% BCs, tensile strength of nanocomposites was increased by 71.6% when compared
with that of pure PVA, which is ascribed to the good interfacial bonding between well-
dispersed BC nanoparticles at low contents and PVA matrices. This is because uniformly
dispersed BCs generate large surface areas to promote effective load transfer from nanofillers
to PVA matrices. In addition, porous structures of BCs enable the mobility of polymeric
chains into internal pores and gaps of nanoparticles, leading to strong mechanical adhesion between nanofillers and PVA matrices. When the BC content is beyond 3 wt%, tensile strengths of nanocomposites declined remarkably though they were still higher than that of neat PVA. This strength reduction trend is most likely due to the occurrence of more BC aggregates, acting as high stress concentration sites prone to mechanical failure with much weaker interfacial bonding. The elongation at break appeared to significantly decrease with increasing the BC content, Table 1. The reduced extensibility of PVA matrices lies in the resistance of rigid BCs, which tends to be more severe when BC aggregates are formed at high content levels. Due to the BC agglomeration, partial particle-particle and PVA-particle separations may occur, thus obstructing the stress propagation when tensile stress is applied [41].

The BC dispersion status within continuous PVA matrices has been examined by scanning electron microscopy (SEM). SEM images on tensile fracture surfaces of neat PVA and PVA/BC nanocomposite films are depicted in Figure 3. Neat PVA possesses fairly smooth fracture surfaces without any obviously preferential orientation, Figure 3(a). Whereas, fracture surfaces of PVA/BC nanocomposites tend to be much rougher with a clear sign of embedded BC nanoparticles. In particular, layered structures and well dispersed BCs are presented on fracture surfaces of PVA nanocomposites reinforced with 3 wt% BC nanoparticles, Figure 3(b). Good interfacial bonding between BCs and PVA matrices is clearly demonstrated for an effective load transfer, which is in good accordance with previously obtained maximum tensile strength of corresponding nanocomposites in Figure 2(c). Another plausible reason is associated with BC porous structures, assisting in the mobility of polymeric chains into internal pores and BC gaps. It leads to strong adhesion between polymer matrices and BC fillers. As shown in Figures 3(c) and (d), with increasing the BC content from 5 to 10 wt%, the interparticle distance can be significantly reduced.
owing to large amounts of particle agglomerates. As a result, the formation of local networks of BC particles that work as stress concentration sites prone to mechanical failure takes place, thus lowering tensile strengths of nanocomposites depicted in Figure 2(c).

It is interesting to note that mechanical performance of PVA/BC nanocomposites is not the only aspect for the property enhancement. Thermal stability of such nanocomposites has also been improved significantly with increasing the BC content. TGA and DTG results for pure PVA and its nanocomposites with the BC contents of 3, 5 and 10 wt% are depicted in Figure 4 and Table 2. Both pure PVA and PVA/BC nanocomposites can decompose in a three-step process according to previous literatures [20, 41]. First of all, the evaporation of trapped water occurs at the temperature range from 50-176°C. The second stage at 200-400°C involves the elimination reaction of water as well as the formation of polyacetylene structures. Finally, further degradation of polyene residues to yield carbons and hydrocarbons takes place at 400-550°C. TGA curves of nanocomposites have gradually shifted toward a higher temperature range when the BC content increases from 0 to 10 wt%, as evidenced by unanimously higher $T_{5\%}$, $T_{80\%}$ and $T_d$ values. Such a phenomenon indicates that the presence of BC nanoparticles can delay PVA thermal degradation, which may act as material barriers to hinder volatile degradation products [26, 43].

4 Conclusion

PVA/BC nanocomposite films were successfully prepared with a simple eco-friendly solution casting method. The SEM images confirm homogenous BC dispersion has been achieved, especially at the low BC content from 3 to 5 wt% in PVA/BC nanocomposites. Significant enhancement of mechanical properties of nanocomposites was detected at low BC content levels in range from 0 to 3 wt%. The addition of BC nanoparticles from 0 to 3 wt% was found to significantly increase tensile strength and tensile modulus of PVA/BC
nanocomposites by 71.6 and 70.2%, respectively. TGA results reveal that the incorporation of BC nanoparticles can greatly improve the thermal stability of nanocomposite films. The increase in the BC content leads to consistently higher $T_{5\%}$, $T_{80\%}$ and $T_d$. It is anticipated that such PVA/BC nanocomposites with superior mechanical and thermal properties prepared in this study can become fully biodegradable and biocompatible composite materials used in future widespread applications.

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**References**


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Fig. 1. Characterisation of individual BC particles (a) FTIR spectrum and (b) XRD pattern.

Fig. 2. (a) FTIR spectra, (b) XRD patterns and (c) mechanical properties of PVA/BC nanocomposite films at different BC contents: tensile modulus and tensile strength.

Fig. 3 SEM micrographs of fracture surfaces: (a) PVA, (b) PVA/ 3 wt% BC nanocomposites, (c) PVA/ 5 wt% BC nanocomposites and (d) PVA/ 10 wt% BC nanocomposites.

Fig. 4 Thermograms of PVA/BC nanocomposite films: (a) TGA curves and (b) DTG curves.
Figure 1

(a)

(b)
Figure 2
Figure 3
Figure 4
Table 1 Mechanical properties of PVA and PVA/BC nanocomposite films

<table>
<thead>
<tr>
<th>Material sample</th>
<th>Tensile modulus (GPa)</th>
<th>Tensile strength (MPa)</th>
<th>Elongation at break (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PVA</td>
<td>2.08 ± 0.53</td>
<td>70.32 ± 2.9</td>
<td>14.60 ± 2.3</td>
</tr>
<tr>
<td>PVA/BC/3wt%</td>
<td>3.54 ± 0.52</td>
<td>120.64 ± 4.3</td>
<td>6.93 ± 2.6</td>
</tr>
<tr>
<td>PVA/BC/5wt%</td>
<td>4.11 ± 0.39</td>
<td>95.06 ± 3.12</td>
<td>5.98 ± 2.19</td>
</tr>
<tr>
<td>PVA/BC/10wt%</td>
<td>4.16 ± 0.41</td>
<td>80.69 ± 4.4</td>
<td>4.88 ± 1.26</td>
</tr>
</tbody>
</table>
Table 2 Thermal properties of PVA and PVA/BC nanocomposite films

<table>
<thead>
<tr>
<th>Material sample</th>
<th>$T_{5%}$ (°C)</th>
<th>$T_{80%}$ (°C)</th>
<th>$T_d$ (°C)</th>
<th>Residue at 700°C (wt%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PVA</td>
<td>200.15</td>
<td>363.5</td>
<td>275.23</td>
<td>3.4</td>
</tr>
<tr>
<td>PVA/BC/3 wt%</td>
<td>256.22</td>
<td>383.3</td>
<td>281.81</td>
<td>7.7</td>
</tr>
<tr>
<td>PVA/BC/5 wt%</td>
<td>258.78</td>
<td>410.12</td>
<td>287.93</td>
<td>9.7</td>
</tr>
<tr>
<td>PVA/BC/10 wt%</td>
<td>265.9</td>
<td>428.03</td>
<td>326.03</td>
<td>13.2</td>
</tr>
</tbody>
</table>

Note $T_{5\%}$ and $T_{80\%}$ refer to decomposition temperatures at 5 and 80% mass loss taking place in TGA curves, respectively. $T_d$ is the maximum degradation temperature in DTG curves.