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Preparation of elastomeric nanocomposites using nanocellulose and recycled alum sludge for flexible dielectric materials

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Flexible dielectric materials with environmental-friendly, low-cost and high-energy density characteristics are in increasing demand as the world steps into the new Industrial 4.0 era. In this work, an elastomeric nanocomposite was developed by incorporating two components: cellulose nanofibrils (CNFs) and recycled alum sludge, as the reinforcement phase and to improve the dielectric properties, in a bio-elastomer matrix. CNF and alum sludge were produced by processing waste materials that would otherwise be disposed to landfills. A biodegradable elastomer polydimethylsiloxane was used as the matrix and the nanocomposites were processed by casting the materials in Petri dishes. Nanocellulose extraction and heat treatment of alum sludge were conducted and characterized using various techniques including scanning electron microscopy (SEM), thermogravimetric analysis/ derivative thermogravimetric (TGA/DTG) and X-ray diffraction (XRD) analysis. When preparing the nanocomposite samples, various amount of alum sludge was added to examine their impact on the mechanical, thermal and electrical properties. Results have shown that it could be a sustainable practice of reusing such wastes in preparing flexible, lightweight and miniature dielectric materials that can be used for energy storage applications.

Keywords: Nanocellulose; biodegradable; sustainable; alum sludge; nanocomposite; flexible; dielectric.

1. Introduction

Elastomers are an important class of polymeric materials in which the molecules are lightly cross-linked to form an amorphous structure. They typically undergo large elastic deformation under a small force and therefore have a low Young's modulus and very high elongation at break when compared with other polymers.¹ Some recent research has been reported on the generation of elastomeric nanocomposites with the incorporation of nanofillers which has led to the enhancement of their properties significantly. These include the change of optical properties and color; the improvement of structural stability and surface characteristics; modification of thermal, magnetic and electrical properties and the improvement of mechanical properties.² Various nanosized fillers in the form of fiber, whisker and particle have been investigated, for example carbon fiber,³ chitin whisker⁴ and layered clay.⁵ Over the past few decades, with increasing focus on environmental issues such as climate change, severe environmental pollution and the depletion of fossil fuels and other resources, the use of wood-based cellulosic materials as fillers or reinforcements to replace the conventional fillers in elastomeric composites has attracted great attention.⁶

Cellulose, a naturally occurring polymer, is a polysaccharide consisting of linear chains of D-glucose units.⁷ It is a key structural component in many higher plants, for example, tree, hemp, wheat straw and bamboo. In cellulose structure, elementary units align to form larger bundles of microfibrils and ultimately a cellulosic fiber matrix, complexed with further polymers hemicellulose and lignin bound to the cellulose surface and entrapped within the microfibrils.⁸ As nanoscience and nanotechnology develop, the extraction of cellulose nanomaterials using various methods has been widely reported.^{9,10} Nanocellulose is used as a key

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D. Sun et al.

component for preparing nanocomposites because of its high modulus, strength and lightweight properties, as well as its inherent biodegradable and biocompatible characteristics.¹¹ Therefore, the addition of such a nanosized reinforcement to elastomers would achieve a more sustainable elastomeric nanocomposite with tailored properties of high specific modulus and specific strength yet flexibility and processibility for various applications.

The work presented in this paper uses a common floating plant, water hyacinth (WH) (*Eichhornia crassipes*) as the feedstock from which to derive cellulose nanofibrils (CNFs) as the key reinforcement for the nanocomposites. WH is considered an invasive species: it colonizes water bodies at a rapid pace, blocking sunlight and reducing oxygen levels in the local ecosystem.¹² Huge sums of money are being spent worldwide to selectively remove the weeds by manual harvesting and use them as animal feed or for energy production by direct combustion.¹³ However, it is worth investigating the feasibility of using WH as feedstock for nanocellulose processing due to their high cellulose content.

The matrix used to form the elastomeric nanocomposite is a biodegradable elastomer, acetoxy-polydimethylsiloxane (PDMS). PDMS is used in wide applications including biomedical engineering,¹⁴ energy harvesting¹⁵ and micromanufacturing,¹⁶ exploiting its distinctive characteristics of flexibility, biocompatibility, optical transparency, chemical and thermal stability and ease of manufacturing.¹⁷ A fixed weight ratio of PDMS and CNF was used to prepare the composite.

The key application targeted for the above nanocomposites is to be as flexible dielectric materials for energy storage. Dielectric materials have characteristics of high-energy storage density, low loss and good cycle stability, which are greatly desired in energy storage applications such as electric vehicles, portable electronic devices and renewable energy systems. As automation and data exchange help define manufacturing in the new Industry 4.0 era,¹⁸ portable electronic technology and wearable technology are all in a rapid expansion stage, sharing common features of flexibility, integration, miniaturization and intellectualization.¹⁹ Wearable technology incorporates lightweight, smart electronic devices that are worn close to and/or on the surface of the skin, where they detect, analyze and transmit information from the user to the external device.²⁰ This can facilitate applications including health monitoring²¹ and human-machine interaction.²² Traditional ceramic dielectric materials for energy storage are unable to meet the requirements of lightweight, flexibility and miniaturization in the above applications. As a result, research has focused on processing novel dielectric materials that combine good dielectric properties with lightweight, flexible, low-cost, multifunctional properties, good mechanical performance and high-energy efficiency characteristics.^{23,24} PDMS has been used with nanoparticle reinforcements such as polymer nanowire²⁵ and SrTiO₃ nanoparticles²⁶ to form flexible dielectrics. However, using sustainable materials like

CNF in PDMS for the same application is less well-studied and so forms the basis for this work.

The addition of another component, alum sludge, was also investigated for its impact on the thermal, mechanical and electrical properties of the PDMS/CNF composite. The use of alum sludge in this work follows similar waste management principles as for the WH material. Alum sludge is a by-product of a water treatment process in which aluminum sulfate is prevalently used as the primary coagulant.²⁷ According to the data from a project partner — Scottish Water, there are approximately 10,000 t of alum sludge generated from water treatment work landfilled every year in Scotland. The industry has been facing escalating financial and environmental pressure to develop more sustainable strategies to deal with alum sludge wastes. In the available literature, some work on utilizing alum sludge has been reported, e.g., aluminum recovery or agriculture and land reclamation.²⁸ However, little work can be found in applying it in processing energy materials for enhanced energy density and efficiency. Therefore, the work has focused on the pre-treatment of the alum sludge and its addition to the PDMS/CNF composites at various loading. The impact on the thermal, mechanical and electrical properties of the composites will also be investigated to see if a more sustainable solution for processing flexible dielectric materials of high performance can be achieved.

2. Materials and Methods

2.1. *Materials*

WH stems were harvested from a lake in Taman Tasik Seri Aman, Puchong, Malaysia. The WH stems were carefully separated from the roots and leaves, washed and air dried for 3 days before being ground into smaller size as raw materials for preparing the CNF (Fig. 1(a)). Dewatered alum sludge was obtained directly from a local water treatment plant of Scottish Water (the Scottish Water Rosebery Water Treatment Works, Midlothian, Scotland). The average water content of the dewatered sludge was found to be approximately 76% (Fig. 1(b)).

Sodium hypochlorite solution (NaClO, 6–14% active chlorine), heptane ($CH_3(CH_2)_5CH_3$, 99%) and sodium hydroxide (NaOH) were received from Sigma-Aldrich, UK. Glacial acetic acid (CH_3COOH) was received from Fisher Scientific, UK. The biodegradable elastomer of acetoxy-PDMS (PDMS, Elastosil E43) was received from Wacker Chemicals Ltd., UK. All chemicals were used without further purification.

2.2. Processing methods

The major experimental procedures involved in this work can be categorized into three parts: heat treatment and refinement of alum sludge, the processing of CNF and the forming of cellulose/alum sludge composites in PDMS matrix.



Fig. 1. (a) Dried and ground WH material; (b) dewatered alum sludge sample collected from a treatment plant of Scottish Water.

2.2.1. Heat treatment of alum sludge

The dewatered sludge was firstly dried in an oven over a period of 24 h at 105°C before being further processed. Once dried, the resulting crumb particle size was reduced using a planetary ball mill. Sieve separation was used and particles smaller than 250 μ m were retained. Particles larger than this were returned to the ball mill for further processing until all material passed through a 250 μ m mesh. The sieved material was then heat treated in a box furnace for 7 h at 300°C, 800°C, 1300°C and 1500°C using a heating rate of 50°C/ min and left to cool overnight to degrade thermally unstable materials. This experiment was to determine a suitable heat treatment temperature for the sludge powder to be prepared and used in the subsequent composite forming work.

2.2.2. Cellulose nanofibril processing

A simple and less-energy intensive method was used to process CNF as adapted from the previous work²⁹: dried WH was soaked in water to form slurry before being subjected to a delignification process using NaClO solution. Additionally, the material was processed in 1% NaOH solution to continue the extraction process. Both reactions were completed at room temperature and the material was washed using ultrapure water after each chemical reaction. Upon the completion of the above, a 0.5% cellulose slurry was prepared. To obtain nanofibrils, the cellulose slurry was further treated in a high shear homogenizer (PSI-20, Adaptive Instruments, UK) by passing it through a 200 μ m Z-shaped interaction chamber for 1 pass, at 700 bar. The produced nanocellulose was eventually solvent exchanged from water into heptane to aid the composite forming process. The solid content of the final slurry is approximately 3.5%.

2.2.3. Composite forming

The composites were prepared by first mixing the CNF slurry with the PDMS using 70:30 weight ratio. This value

was determined by trials of other ratios (e.g., 80:20 and 60:40 weight ratio), in which they can either be of insufficient PDMS as the matrix or excessive PDMS to obstruct the flowability of the mixture. The mixing was completed in heptane solvent in order to achieve a free flow condition of the mixture. Subsequently, the alum sludge powder that was heat treated at 800°C was added to the mixture. The wt.% of the alum sludge powder with respect to the mixture was 10%, 15% and 20%, respectively. The mixture of approximately 40 g in weight was vigorously mixed using an acoustic mixer (LabRAM, Resodyn, USA) and a rotor-stator mixer (T 25 digital ULTRA-TURRAX, IKA, UK), in order to achieve a good dispersion. The mixture was cast in a Petri dish and left to cure at ambient condition. Finally the dried film was removed from the mold and denoted as PDMSCNF, PDMSCNF10AS, PDMSCNF15AS and PDMSCNF20AS, for the composite samples with 0%, 10%, 15% and 20% alum sludge, respectively.

2.3. Characterization

2.3.1. Characterization of CNF and alum sludge

The morphological features of the CNF and alum sludge samples were examined using scanning electron microscopy (SEM) (FESEM, Hitachi, UK). Thermogravimetric analysis and its derivative thermogravimetric (TGA/DTG), was conducted on the raw alum sludge crumb (before being heat treated) using Netzsch simultaneous thermal analyser, STA 449 (Netzsch, Germany). 20–25 mg of each sample was heated in an aluminum oxide pan from 30°C to 900°C at a heating rate of 10°C·min⁻¹ under a constant nitrogen of 50 mL·min⁻¹. DTG curves were obtained by performing a first derivative on the percentage weight loss data from TGA using Origin 2020b version.

The X-ray diffraction (XRD) patterns were recorded with a Bruker D8 Advance X-ray diffractometer (Germany) by using CuK α radiation ($\lambda = 0.1542$ nm), a parallel beam with Gobel mirror and a Dynamic Scintillation detector. The accelerating voltage was of 40 kV, the current of 30 mA and scanning range between 5° and 40° (2θ).

2.3.2. Characterization of composite films

The morphology of the composite films was examined using the SEM. Mechanical testing was completed to investigate the strength and ductility properties of the composite. A universal testing machine was used for this test in which the material was pulled to break at a rate of 50 mm/min. 12 specimens were prepared for each sample, using which the mean values and standard deviation of tensile strength and percentage elongation at break were calculated, respectively.

The thermal properties of the four composite films were examined using the above TGA/DTG instrument and the same test conditions. For dielectric and energy storage properties, the samples were coated with silver paste (Johnson Matthey, No. E110) on both sides of the film. Frequency-dependent permittivity and dielectric loss in the frequency range of 30 Hz–1 MHz were measured using a precision LCR meter model, Agilent 4184A. The polarization–electric field (P–E), and current–electric field (I–E) loops of coated samples were measured by the ferroelectric hysteresis measurement tester (NPL, UK) at 10 Hz.

3. Results and Discussion

CNF were successfully obtained using both chemical and mechanical steps. In the first step, lignin was removed in the acidified NaClO solution, followed by the removal of hemicellulose in the NaOH solution. This can be observed by color change in the material (from dark brown to white) in the process. The resultant material was then subjected to high shear processing to fibrillate the material into nanofibrils. Figure 2 shows the microstructure of high aspect ratio CNF obtained from the above process. The long nature of the fibrils results in the formation of a web-like network structure, similar to that observed in other published works.³⁰ The enlarged image in Fig. 2(b) clearly shows the morphology of individual fibrils, which can be estimated as approximately 20 nm in diameter. Due to the entangled nature of the fibrils, it is not possible to measure their length accurately; however, it can be estimated from Fig. 2(a) that they can be in the order of several micrometers long.

It is also evident from the circularly highlighted area in Fig. 2(a) that a small proportion of larger fiber bundles were present, having a diameter of approximately several hundred nanometers. This could be the result of the incomplete fibrillation of material during the homogenization step. Previous work has shown that the degree of fibrillation is determined by various aspects, including the nature of the feedstock, processing time and the amount of shear force applied on the materials; the latter two aspects are directly correlated to the amount of energy consumed in the process. In this work, the material was processed using a 200 μ m interaction chamber for 1 pass (1 cycle), which shows a significant improvement in energy consumption efficiency than in previous work where the material is passed through a 100 μ m chamber for up to 5 cycles. Subsequent experiments in this work have identified that the processed material is suitable for the composite forming process.

The TGA/DTG thermogram elucidating the thermal stability of the raw alum sludge crumb is shown in Fig. 3(a). A noticeable amount of weight (approximately 46%) has been lost when the sample was heated up to 900°C. Three steps can be seen in the weight loss profile: (i) up to 100°C, approximately 4% of weight is lost, which could mainly be associated to the physically absorbed moisture (remaining free/unbound water in the porous structure of unheated sludge sample that can be seen in Fig. 3(b)). (ii) There is a major weight loss between 105°C and 500°C. This loss could possibly be because of the removal of the remaining bonded water and the decomposition of some organic compounds (e.g., polyaluminum chloride hydroxide sulfate, one of the key components in the sludge, usually decomposes at about 200°C



Fig. 2. (a) FESEM images of CNFs showing the web-like network structure; the arrow highlights a large fiber bundle with incomplete fibrillation; and (b) the enlarged image of the rectangular area in (a), showing the morphology of individual nanofibrils.



Fig. 3. (a) TGA/DTG thermograms of the raw alum sludge; (b) the alum sludge without heat treatment and (c) the sludge which was heat treated at 800° C.

(Ref. 31)). It is reported that the decomposition of polyacrylamide, an anionic organic polymer (e.g., Magnafloc®) that is commonly used to improve the dewatering of alum sludge in the water treatment process, can occur in between 350°C and 550°C (Ref. 32); this could be another aspect for the above weight loss in our work. (iii) Above 500°C, the curve started plateauing, showing insignificant weight loss in this range. All these observations are correlated with the thermal event shown in the DTG curve in the same temperature range. In summary, the alum sludge used in this work contains various compounds that are thermally unstable; thus, it is necessary for them to undergo some heat treatment to remove the volatile substances before being used in forming the composites.

The XRD patterns of untreated alum sludge and samples with various heat treatments are shown in Fig. 4. Due to the large background signal, the crystallinity of the raw alum sludge is quite poor, and it is difficult to identify a distinct peak. The spectrum for the 300°C shows a peak for hexagonal quartz SiO₂. When the sample was heat treated at 800° C, more peaks can be seen which could be associated to cubic γ -Al₂O₃. Similar results have been reported elsewhere^{33,34} and these structures could possibly be formed in the oxidation process at the elevated temperatures. A broader peak in the spectrum at 800°C indicates that the sample is of low crystallinity at this stage in comparison with heat treatment at 1300°C and 1500°C, at which the peaks became sharper and more numerous. These peaks may be associated with orthorhombic corundum Al₂O₃ and mullite Al₆SiO₁₃. Since no significant difference can be seen between the two spectra of 1300° and 1500°, it is believed that from 1300°C, the compound has achieved its crystalline phase. Based on the results in Figs. 3 and 4, it is reasoned that alum sludge heat treated at 800°C developed an oxide structure with reasonable thermal stability and is thus suitable for use in the formation



Fig. 4. XRD spectra for the raw and heat treated alum sludge samples at 300°C, 800°C, 1300°C and 1500°C, respectively.

of composites. Although the heat treatment at 1300° and 1500° would result in a material of higher crystallinity, particle coalescence would make the sample nonhomogeneous and difficult to disperse in the matrix when preparing the composites.

A representative SEM micrograph showing the morphology of the PDMSCNF10AS film can be seen in Fig. 5(a). Alum sludge particles appear embedded in the matrix with a broadly even distribution, although clusters of smaller particles are evident, shown in more morphological detail in the enlarged image in Fig. 5(b). An image with greater magnification (Fig. 5(c)) shows the CNF is also embedded in the PDMS matrix homogeneously with random orientations. The CNF structure can be recognized as identical when compared with the one shown in Fig. 2. The microstructure of other film samples was similar to PDMSCNF10AS (except the different content of alum sludge particles) and these are not shown.



Fig. 5. (a) SEM micrograph of cross-sectional structure of the PDMSCNF10AS composite film; (b) enlarged image showing the alum sludge particles; (c) enlarged image showing the CNF in the matrix.

The strength and percentage elongation at break obtained from the mechanical tests on composite samples are summarized in Fig. 6. It can be seen that the addition of alum sludge has adversely affected the tensile strength of the composites: the PDMSCNF composite with no alum sludge has a tensile strength of approximately 0.65 MPa, but this reduces to roughly 1/3 with 20% alum sludge. This may be attributed to particle heterogeneity and poor interfacial bonding between the particles; the PDMS matrix thus providing a weak point of the material when under external stress. During the preparation of the composite mixture, the compound was always of high viscosity, especially when the alum sludge content was increased. This made it difficult to homogenize the components and may have resulted in uneven local dispersion in the composite film: a possible contributory factor in lowered tensile strength and greater standard deviation in sample sets. An increase in volume fraction of particles to enhance stiffness in a polymer matrix composite may be expected to decrease toughness/flexibility and this is confirmed by elongation at break data, shown in Fig. 6(c). The PDMSCNF composite under stretch shows the flexibility of the polymer matrix with cellulose fiber reinforcement. The reduction of percentage elongation at break broadly correlates with volume fraction of the alum sludge in the composite, a trend apparently bucked by the data for PDMSCNF10AS which shows higher elongation at break than the PDMSCNF sample without alum sludge. Standard deviation is however comparatively high in these two sample sets and the data may be significantly improved by increasing the sample size and optimizing experimental conditions for improved particle



Fig. 6. Engineering tensile stress versus engineering tensile strain curve for the composite film; the two embedded images show the tensile samples and the composite film before testing.



Fig. 7. TGA (a) and DTG (b) thermograms of the PDMSCNF composites with various weight ratio of alum sludge; a — PDMSCNF; b — PDMSCNF10AS; c — PDMSCNF15AS; d — PDMSCNF20AS. The peak degradation temperatures on Fig. 7(b) correspond to the degradation of CNF (\sim 300°C) and PDMS (\sim 550°C).

homogeneity, dispersion and interfacial interactions with the matrix.

Thermal stability studies were carried out and the results overlayed in the TGA (TGA, Fig. 7(a)) and DTG (DTG, Fig. 7(b)) curves for the four PDMSCNF composites with various alum sludge content. The TGA graph shows a relatively flat line below 300°C, indicating minimum weight loss due to solvent evaporation. Two degradation troughs can be observed at around 300°C and 550°C in the associated TGA and DTG curves. These are assigned to the degradation of



Fig. 8. Frequency dependence of the (a) dielectric constant and (b) dielectric loss for the PDMSCNF composite films with alum sludge; (c) dielectric constant and (d) dielectric loss measured at 1 kHz as a function of the alum sludge content.



Fig. 9. *P–E* loop and *I–E* loop for the PDMSCNF composites with various content of alum sludge.

CNF and PDMS, respectively. As the content of alum sludge increases, the degradation temperatures for CNF and PDMS are affected differently (Fig. 7(b)): the peak degradation temperatures of CNF were shifted to higher temperatures, while the peak degradation temperatures of PDMS shifted to slightly lower values. This is more pronounced in the PDMSCNF15AS and PDMSCNF20AS samples, indicating that a greater amount of alum sludge affected the degradation temperatures more significantly.

It is worth noting that the addition of alum sludge at different weight loadings to the composites increased the amount of char remaining at 600–900°C. Char formation is a desired property required for fire-retardant composite materials. The high residual char of these composites make them desirable in electrical devices which are prone to short-circuiting.³⁵ The increased amount of alum sludge in the composite could improve the fire-retardant property of the composite when they are used in energy storage applications.

Figure 8 shows the frequency dependence of the dielectric constant and dielectric loss for the PDMSCNF composite films with various alum sludge content. In the frequency range of 30 Hz–1 MHz, the dielectric constant drops gradually as frequency increases. This drop is due to the diminishing influence of dipolar polarization. As the alum sludge content increases, the dielectric constant of the PDMSCNF composite films rises, which is because of the greater dielectric constant of Al_2O_3 and mullite-based sludge (6.7–10) compared to pure cellulose (2–4) at this frequency range.^{36–38}

Figure 9 displays the P-E and I-E loops of the composite samples to explore the energy storage properties. In the same electric field, as the amount of the alum sludge increases, polarizability increases and electrical resistivity decreases. The rise in polarizability is consistent with the measured dielectric constant as shown in Fig. 8(a). This is owing to the presence of comparatively high dielectric constant of the alum sludge. Moreover, the reduction in electrical resistivity may not be due to intrinsic factors, but rather to poor adhesion between the electrode and the composite films with the alum sludge in the high electric fields. This coincides with the observation in Fig. 8(b) that the dielectric loss of PDMSCNF at low frequencies is higher than the PDMSCNFAS films. These film samples will need enhanced silver paste electrodes in the future, which will result in a larger applied electric field and thus a greater energy storage property.

4. Conclusions

This work has demonstrated the use of waste materials to reinforce nanocomposites for flexible dielectric materials. Two waste material sources were used; dried WH and alum sludge and these were mixed with a biodegradable elastomer PDMS matrix. A simple and less-energy intensive method was used to extract CNF from dried WH and the as-received alum sludge was heat treated and refined to be suitable for the composite forming process. The results showed that incorporating these reinforcements into a polymer matrix to produce a functional composite is possible and that the addition of alum sludge particles in the PDMS/CNF composite has significant effects on the material properties of the composite. Mechanical, thermal and electrical properties are differentially affected by alum particle reinforcement, allowing a tailored approach to maximize desirable features while compromising less desirable ones for any specific application. This work has provided a strong basis for showing the potential of reusing waste materials in preparing flexible, lightweight and miniature dielectric materials that can be used for sustainable energy storage applications.

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