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# Preparation of conductive cellulose paper through electrochemical exfoliation of graphite: The role of anionic surfactant ionic liquids as exfoliating and stabilizing agents



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#### ABSTRACT

A facile electrochemical exfoliation method was established to efficiently prepare conductive paper containing reduced graphene oxide (RGO) with the help of single chain anionic surfactant ionic liquids (SAILs). The surfactant ionic liquids are synthesized from conventional organic surfactant anions and a 1-butyl-3-methyl-imidazolium cation. For the first time the combination of SAILs and cellulose was used to directly exfoliate graphite. The ionic liquid 1-butyl-3-methyl-imidazolium dodecylbenzenesulfonate (BMIM-DBS) was shown to have notable affinity for graphene, demonstrating improved electrical properties of the conductive cellulose paper. The presence of BMIM-DBS in the system promotes five orders of magnitude enhancement of the paper electrical conductivity  $(2.71 \times 10^{-5} \text{ S cm}^{-1})$  compared to the native cellulose  $(1.97 \times 10^{-10} \text{ S cm}^{-1})$ . A thorough investigation using electron microscopy and Raman spectroscopy highlights the presence of uniform graphene incorporated inside the matrices. Studies into aqueous aggregation behavior using small-angle neutron scattering (SANS) point to the ability of this compound to act as a bridge between graphene and cellulose, and is responsible for the enhanced exfoliation level and stabilization of the resulting dispersion. The simple and feasible process for producing conductive paper described here is attractive for the possibility of scaling-up this technique for mass production of conductive composites containing graphene or other layered materials.

#### 1. Introduction

Growing global concerns over the environment and sustainability are directing the development of next generation renewable materials. Biomaterials or biocomposites are considered as promising materials over synthetic polymers and can be used in industrial or smaller scale process. With the annual production estimated over  $7.5 \times 10^{10}$  ton per year (Pinkert, Marsh, Pang, & Staiger, 2009), cellulose is hailed as the most abundant renewable material in the world. Cellulose is often extracted from plant cells i.e. wood pulp and can find immediate use in the paper industry or other specific applications. It has several attractive properties including biodegradability, biocompatibility, and superior chemical stability, consequently represents as the most promising resource for producing biocomposites (Abdul Khalil, Ireana Yusra, Bhat, & Jawaid, 2010; Abdul Khalil, Bhat, & Ireana Yusra, 2012).

Introduction of electric fillers into a cellulose paper can switch the electrical properties of the resulting paper and make it possible for applications in flexible energy storage devices, electrodes, and sensors

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Nomenclature		VOCs	Volatile organic compounds
		I(Q)	Scattering intensity
CVD	Chemical vapor deposition	$I_D/I_G$	The ratio between the intensity of D- and G-band
GCP	Graphene/cellulose paper	P(Q)	Scattering form factor
GO	Graphene oxide	R <sub>a</sub>	Polar axis ratio of ellipsoidal micelle
ILs	Ionic liquids R <sub>cvlinder</sub>		Cylindrical micelle radius
NRL	Natural rubber latex	R <sub>disk</sub>	Stacked disk radius
RGO	Reduced graphene oxide	<b>R</b> <sub>sphere</sub>	Spherical micelle radius
SAILs	Surfactant ionic liquids	S(Q)	Structure factor (interparticle interaction)
SANS	Small-angle neutron scattering	Х	Aspect ratio of ellipsoidal micelle
SDBS	Sodium dodecylbenzenesulfonate	ζ-potent	ial Zeta potential
SDS	Sodium dodecylsulfate	σ	Electrical conductivity
SLD	Scattering length density		

(Kang, Li, Hou, Wen, & Su, 2012; Kiziltas et al., 2016; Weng et al., 2011; Ye et al., 2016; Yoon, Jin, Kook, & Pyun, 2006). There is an ever-increasing interest in the utilization of graphene (and its derivatives) as a reinforcing filler in nanocomposites, and its attractive properties make it particularly suitable for the development of novel electrical cellulose paper. However, both cellulose and graphene present problems regarding dispersibility/solubility in aqueous solution (Hernandez, Lotya, Rickard, Bergin, & Coleman, 2009; Lindman, Karlström, & Stigsson, 2010; Medronho, Romano, Miguel, Stigsson, & Lindman, 2012). Without modifications to the chemical structure or the use of dispersing agents, it is rather challenging to obtain a good dispersion of cellulose and graphene in water (Roy, Semsarilar, Guthrie, & Perrier, 2009).

Current pre-treatments of cellulose dissolution typically employ acids and bases, and sometimes involve cuprammonium and xanthate which use relatively harsh processing solvents (traditional dissolution) (Zhang et al., 2009; Zhou & Zhang, 2000; Zhu et al., 2006). Other efforts have used ionic liquids (ILs) to achieve dissolution/dispersion. Ionic liquids (ILs) are considered as environmentally benign solvents replacement for volatile organic compounds (VOCs) due to their low vapor pressures, inflammability and recyclability (Chiappe & Pieraccini, 2005; Earle & Seddon, 2000). Reports have shown that some hydrophilic ILs based on imidazolium is able to dissolve large amounts of cellulose. Readers may refer to the seminal paper of Pinkert and co-workers for reference (Pinkert et al., 2009). Along with this, a number of other works have pointed out that ILs are suitable solvents to produce stable graphene dispersions with high concentrations (Bari et al., 2014; Liu et al., 2008; Nuvoli et al., 2011).

Ionic liquids offer tuneability and there are a huge number possibilities to alter their properties by modifying chemical structure (Chiappe & Pieraccini, 2005). A notable advancement in the field of ILs is the discovery of anionic surfactant ionic liquids (SAILs) (Brown et al., 2011, 2012). These compounds may act as solvents with the ability to self-assemble, offering unique opportunities to solving problems between two incompatible materials. Without doubt, SAILs will be useful

Table 1



in tailoring the incompatibility between graphene and cellulose to fabricate electric cellulose paper. To the Authors' knowledge, such a system has not been tested either for graphene or nanocomposites processing.

Ever since graphene hit the headlines, there has been a large body of work devoted to its synthesis and properties. Sophisticated techniques such as chemical vapor deposition (CVD) or the tedious mechanical exfoliation do tend to produce high quality graphene films (Whitener & Sheehan, 2014), however, these methods are not suitable for bulk material applications. A low-cost process to make graphene by exfoliating graphite layers may result readily-dispersed graphene that can find applications in polymer composites, conductive inks, and supercapacitors. The process can be performed efficiently through electrochemical treatment in surfactant solutions that provides colloidal stability (Alanyalıoğlu, Segura, Oró-Solè, & Casañ-Pastor, 2012; Kakaei & Hasanpour, 2014; Suriani et al., 2016). Interest in this technique stems from the simple experimental setup: graphite rods as working electrodes, immersed in a surfactant solution as the electrolyte, and a DC power supply. Although many studies on electrochemical exfoliation of graphite in surfactant solution are successful, they all focus on exfoliation effectiveness, and graphene or composite characteristics. Unfortunately, the surfactant aspect is relegated to just a stabilizing agent.

This present work reports the facile fabrication of nanofibrillated cellulose paper from kenaf (*Hibiscus cannabinus* L.) by directly exfoliating graphene in a cellulose/SAILs dispersions, which is then followed by concentration of the suspension to obtain graphene/cellulose paper (GCP). Kenaf is well known as a cellulosic source with both economic and ecological advantages. After a decrease in wood resources for papermaking, one of the most viable replacements is kenaf. Aside from the ability to be processed into pulps, it can be used for building materials or biocomposites. It has many advantages over wood, such as being inexpensive and easy to grow in a wide range of climatic conditions. Hence, it is important to explore the most promising applications of kenaf-based materials for improved value (Abdul Khalil et al., 2010, 2012).

This paper aims to investigate the effect of modifying the counterion of conventional surfactants, thus turning them into SAILs, towards their ability in stabilizing aqueous systems consisting of graphene and cellulose fiber. The SAILs are designed by exchanging a conventional sodium counterion of commercial surfactants sodium dodecylsulfate (SDS) and sodium dodecylbenzenesulfonate (SDBS) with a large organic 1-butyl-3-methyl imidazolium (BMIM) cation. The structures of the surfactants and SAILs are given in Table 1. Here, the surfactant and SAILs performances are studied in terms of the GCP electrical conductivity. The results showed that the applied strategy gives a subtle effect on the ability of the SAILs to stabilize the aforementioned system over their conventional surfactants. Significantly, this study provides new alternatives for generating environmentally friendly and economic graphene-compatible compounds considering that the starting materials are cheap and the synthesis process is simple.

# 2. Materials and methods

#### 2.1. Materials

Sodium dodecylsulfate (SDS) and sodium dodecylbenzenesulfonate (SDBS) was purchased from Sigma Aldrich without further purification. 1-Butyl-3-methyl-imidazolium chloride (99%) was obtained from Merck and used as received. Nanofibrillated kenaf cellulose was supported by Forest Research Institute Malaysia.

#### 2.2. SAILs synthesis

The SAILs were synthesized and purified following on previous work by Brown et al. (Brown et al., 2012) The synthesized SAILs were characterized by  ${}^{1}$ H NMR spectroscopy; all are consistent with the

expected values (see Supplementary Material).

# 2.3. Graphene/Cellulose paper (GCP) preparation

Nanofibrillated kenaf cellulose (2.5 g) was first dispersed in surfactant/SAILs solution (50 mL) for 2 h under vigorous stirring until a stable dispersion had been formed. High purity graphite rods with a diameter of 10 mm and 150 mm in length were used as the electrodes. The as-prepared cellulose/SAILs (or surfactant) mixture was then used as the electrolyte for the exfoliation process at a constant voltage of 7 V (GW INSTEK GPS 3030DD). The electrochemical exfoliation was carried out at room temperature for 24 h. The obtained dispersions were then subjected to mechanical stirring and sonication for 1 h to form homogeneous mixtures of graphene oxide (GO)/cellulose/stabilizer. To the resulting mixture, a suitable amount of hydrazine hydrate (0.1 mL hydrazine / 10 mL GO dispersion) was added and the reaction carried out under reflux at 90-100 °C for 24 h. After the reaction was complete, the solids wee then filtered and left to dry on the filter paper and left to dry overnight in an oven at 50 °C. A dark grey composite paper containing reduced graphene oxide (RGO) henceforth labeled as GCP was obtained by peeling the paper material from the filter paper substrate. The route to prepare GCP is summarized in Fig. 1. Details on the amount of materials used for the preparation of GCP can be found in Supplementary material.

## 2.4. GCP characterization

The electrical conductivity of GCP with dimensions of  $15 \times 15$  mm and thickness ~ 0.10 mm was measured by a four-point probe method (Keithley 2636 A). The measurement was repeated 5 times to ensure accuracy. The morphologies and microstructure of GCP were observed under field emission electron microscope (FESEM, Hitachi SU8020). Raman spectroscopy was carried out using a Renishaw InVia micro Raman system spectrofotometer with a 514 nm argon-ion laser source. Five regions were measured for each paper. To visualize the embedded microstructure of cellulose paper using HRTEM (JEOL 2100F), the



Fig. 1. Schematic of the GCP preparation.

samples were ultramic rotomed with a diamond knife to give sections with nominal thickness  $\sim\!80\,\text{nm}.$ 

## 2.5. Zeta potential measurements

Zeta potential measurements were performed by ELSZ-1000 Zetapotential and Particle Size Analyzer (Photal OTSUKA ELECTRONICS) with Smoluchowski equation as zeta potential conversion equation and 1 peak Lorentz fitting. Measurements were carried out with a flow cell at sampling time 400  $\mu$ s, cumulative number 7, measuring angle 15°, temperature 25 °C, pin hole size 50  $\mu$ m, cell constant 70.000 cm<sup>-1</sup>. Properties of aqueous mixtures (refractive index 1.3328, viscosity 0.8878 cP, and permittivity 78.3 Fm<sup>-1</sup>) were used for calculation of zeta potential. Zeta potential values were finally obtained as average values of 10 runs for each sample.

# 2.6. Small-angle neutron scattering (SANS)

Small-angle neutron scattering (SANS) studies were carried out on the time-of-flight LOQ instrument at ISIS, UK. The accessible *Q* range was 0.007–0.23 Å<sup>-1</sup>, arising from incident neutron wavelengths of  $\lambda = 2.2$ –10 Å. Absolute intensities for I(Q) (cm<sup>-1</sup>) were determined to within 5% by measuring the scattering from a partially deuterated polymer standard. Neutrons are scattered by short-range interactions with sample nuclei, the 'scattering power' of different components being defined by a scattering-length density (SLD),  $\rho$  (cm<sup>-2</sup>). The samples were prepared in 2 mm path-length quartz cells and held on a thermostatted automatic sample changer at 25 °C. Data have been fitted using the SASView interactive fitting program, fixing scattering length density differences as calculated and fitting for micellar volume fraction and appropriate structural parameters as required by the different scattering laws. Scattering length density of surfactants, SAILs and graphene are given in Supplementary material.

#### 3. Results and discussion

## 3.1. Electrical conductivities of graphene/cellulose paper (GCP)

To obtain an ideal conductive network, it is required that the graphene nanofillers be well separated despite the ever present van der Waals forces, and that a high dispersity of the nanofiller be maintained in the resulting cellulose nanocomposites. Poor dispersion of reinforcing nanostructures at the nanoscale, and weak interfacial interactions, may result in composites with limited enhancement in electrical conductivity. In an attempt to meet these requirements, many researchers have tried to develop conductive composites with various fabrication methods. Being electrically non-conductive in nature, here nanofibrillated kenaf cellulose (1.97  $\times 10^{-10}$  S cm<sup>-1</sup>) was converted to an electrically conductive nanocomposite by incorporation of reduced graphene oxide (RGO) stabilizer surfactant and SAILs.

Uniform dark grey composite papers were obtained as thin discs with a diameter of 7.00 cm and 0.15–0.19 mm thickness. To evaluate the efficiency of surfactants in exfoliating graphite and stabilizing the graphene/cellulose dispersion system, a series of surfactant concentrations were used ranging from 0.050 M to 0.100 M. The GCP electrical conductivities as a function of surfactant concentration are summarized in Table 2 and Fig. 2. It is important to note that the electrical measurements reported herein are for one-sided samples, however, both surface produce identical results due to good dispersion.

Clearly, replacement of the sodium counterions (Na<sup>+</sup>) with imidazolium cations subtly changes the ability of the SAILs in stabilizing system comprising of RGO and cellulose. GCPs prepared with SAILs have electrical conductivities ( $\sigma$ ) a little higher ( $\sim$ 1 order of magnitude) than if prepared with its sodium analogue (see Table 2). The optimum electrical conductivity enhancement, attaining five orders of magnitude enhancement, was obtained by 0.1 M BMIM-DBS  $(2.71 \times 10^{-5} \text{ S cm}^{-1})$ . The effectiveness of these ionic liquids and cellulose combinations for direct exfoliation of graphite is very interesting. A similar concept, but using tip sonication, was recently reported (Ye et al., 2016). Although the electrical properties were studied in terms of charge-discharge capacities (and cannot be used as a direct comparison for this study), the resulting composites exhibited attractive potential for application as anode materials. A previous study utilizing regenerated cellulose with DMAC/LiCl reported an electrical conductivity of  $3.7 \times 10^{-6} \text{ S cm}^{-1}$  while another using NaOH as a processing aid yielded a cellulose composite with  $1.1 \times 10^{-6} \text{ S cm}^{-1}$  (Feng, Zhang, Shen, Yoshino, & Feng, 2012; Zhang, Liu, Zheng, & Zhu, 2012). It is therefore important to mention that the fabrication approach here is a facile, fast, inexpensive and versatile process to constructing materials from various complementary materials.

This increased electrical properties observed here may be influenced by the good combination of cellulose/ionic liquid as graphite exfoliant and dispersion stability. Due to its insolubility in water (pH 7) and recalcitrance to dissolve in organic solvents, cellulose has often been viewed as a hydrophobic material (Lindman et al., 2010; Medronho & Lindman, 2014, 2015). Recently, however, it is suggested that cellulose may display amphiphilic properties that can affect its dissolution/dispersion (Lindman et al., 2010; Medronho & Lindman, 2015). The hydrogen bonding of the heterocyclic ring BMIM counterions which is assumed to compensate for the hydrophilic character probably plays an important role for the interaction with the hydrophilic parts of cellulose (Brown et al., 2012).

From a general viewpoint, gradually increasing the concentration of surfactant led to modest increases in the electrical conductivities. Again reiterating the importance of surfactant concentration in exfoliation and keeping the exfoliated layer stably dispersed (Lotya et al., 2009; Mohamed et al., 2018; Suriani, Nurhafizah, Mohamed, Zainol, & Masrom, 2015; Wang, Yi, & Shen, 2016). It was hypothesized that the chosen surfactant concentrations (0.05–0.1 M) are sufficient to form a network that provides electrical pathways, considering the significant  $\sigma$ from neat cellulose, and that the presence of RGO enveloping the cellulose network did assist the development of the electrical conductivity of the paper. This observation agrees with previous research on graphene/natural rubber latex (NRL) composites, showing surfactant at a concentration of 0.1 M produces composites with the highest electrical conductivity (Suriani et al., 2015, 2016). It is instructive to compare these  $\sigma$  values with those systems without surfactants or SAILs. An attempt was made to exfoliate graphite using cellulose dispersion without surfactant or SAILs, however it was unsuccessful, and the resulting nanocomposite electrical conductivity could not be determined using four point probe measurement. As shown in Table 2 the presence of surfactant alone does not significantly affect the cellulose paper

Electrical conductivities of nanofibrillated kenaf cellulose paper with and without RGO and the zeta ( $\zeta$ )-potential values of RGO dispersion stabilized by surfactants and SAILs.

Surfactant	Zeta ( $\zeta$ )-potential of	Surfactant concentration (M)				
	RGO dispersion (mV)		Electrical conductivity (S cm <sup>-1</sup> )			
		0.050	0.075	0.100		
With RGO						
SDS	$-43 \pm 4$	$2.82  imes 10^{-8}$	$2.08  imes 10^{-8}$	$3.09  imes 10^{-7}$		
SDBS	$-40 \pm 8$	$1.80  imes 10^{-7}$	$4.67 \times 10^{-7}$	$1.48  imes 10^{-6}$		
BMIM-DS	$-36 \pm 1$	$3.30  imes 10^{-8}$	$4.74 \times 10^{-7}$	$5.86 \times 10^{-6}$		
BMIM-DBS	$-59 \pm 1$	$2.17  imes 10^{-7}$	$1.23  imes 10^{-6}$	$2.71  imes 10^{-5}$		
Without RGO						
SDS	-	$6.03 \times 10^{-9}$	$5.76 \times 10^{-9}$	$9.32 \times 10^{-9}$		
SDBS	-	$5.73 imes10^{-9}$	$9.32  imes 10^{-9}$	$4.59  imes 10^{-9}$		
BMIM-DS	-	$1.37 imes10^{-9}$	$1.64  imes 10^{-9}$	$4.04  imes 10^{-9}$		
BMIM-DBS	-	$3.34\times10^{-9}$	$4.38\times10^{-10}$	$2.55  imes 10^{-9}$		

Table 2



**Fig. 2.** Electrical conductivities of the nanofibrillated kenaf cellulose paper, nanofibrillated kenaf cellulose with surfactant/SAILs paper and GCP containing surfactants and SAILs. The error bars are given for five experimental measurements.

electrical conductivity. Clearly, the enhancements of electrical properties of cellulose papers are due to the RGO dispersed in the system.

#### 3.2. Raman spectroscopy

Carbon allotropes show their fingerprints under Raman spectroscopy mostly by D, G, and 2D bands that lie around 1350, 1580, and 2700 cm<sup>-1</sup>, respectively (Dresselhaus, Jorio, Hofmann, Dresselhaus, & Saito, 2010; Ferrari et al., 2006). Identification of these features allows characterization of graphene layers in terms of the number of layers present or presence of defects as also observed in either GO and RGO (Ferrari et al., 2006). The band near 1580 cm<sup>-1</sup> arises due to the in plane vibration of the sp<sup>2</sup> hybridization of carbon atoms. Meanwhile, the D band appears due to the presence of disorder in atomic arrangement or edge effect of graphene, ripples, and charge puddles (Dresselhaus et al., 2010). The ratio between the intensity of D- and Gband (I<sub>D</sub>/I<sub>G</sub>) has been widely used an indicator of the amount of disorder in RGO sheets. Comparison of Raman spectra between the GCPs and graphite is shown in Fig. 3.

The Raman spectrum of the pristine graphite is expected display a prominent G peak at 1581 cm<sup>-1</sup> and a very low intensity of D peak at 1354 cm<sup>-1</sup> hence the I<sub>D</sub>/I<sub>G</sub> value of 0.03. The Raman spectrum of the GCPs also contains a G band that is now broadened and shifted to around 1585–1594 cm<sup>-1</sup>. In addition, the D band at 1347-1350 cm<sup>-1</sup> becomes more prominent therefore giving an increased D/G intensity ratio compared to that in graphite. These alterations suggest changes in sp<sup>2</sup> domains upon reduction of the resulting GO through electrochemical exfoliation (Dresselhaus et al., 2010; Ferrari et al., 2006). It is seen here that the  $I_D/I_G$  decreases monotonically when the electrochemical exfoliation was carried out with SAILs, suggesting less structural damage (defects) when creating graphene compared to those using SDS or SDBS surfactant. Looking to the paper electrical properties, cellulose paper containing graphene with lowest  $I_D/I_G$  (those stabilized with BMIM-DBS) give the highest electrical conductivity. As the properties of the resulting composite crucially depend to the quality of RGO and the dispersion stability itself, it is evident that the efficiency of the surfactant performance should be correlated with surfactant chemical structure modification.

#### 3.3. FESEM observation

The micromorphologies of the nanofibrillated kenaf cellulose and GCPs are shown in Fig. 4. Scanning electron microscopy (SEM) images of nanofibrillated kenaf cellulose showed networks of randomly oriented thin fibrils, as shown in Fig. 4a'. The fibers have tap-like

morphology with ~25–50 nm wide. A clearly dispersed RGO sheets was displayed as flakes or tactoids over the surface of cellulose (Fig. 4b–e). Two different domains can be distinguished: a network of cellulose microfibrils and incorporated RGO distributed among them. The bright regions in the images were attributed to the RGOs on account of their high conductivity that can form conducting paths through and/or across cellulose fibers to render the obtained GCP electrically conductive (Yu et al., 2007).

In respect to GCPs with SDS and SDBS, higher magnification images (Fig. 4b' and c') revealed bulk pieces of aggregates throughout cellulose matrix. Interestingly, GCPs with BMIM-DS and BMIM-DBS demonstrated the presence of a nanodispersion of individual RGO flakes suggesting increased exfoliation and dispersion quality over the commercial surfactant. Note that an enlarged view of GCPs with BMIM-DBS offers clearer illustration of RGOs inside the cellulose matrix, in which edges of the upright RGO flakes can be seen (Fig. 4e'), unlike those observed with SDS and SDBS.

The uniform dispersion of RGOs in a polymer matrix is one of the most important requirements for achieving a uniform conductivity throughout the composites (Mohamed et al., 2016; Stankovich et al., 2006; Yu et al., 2007). Here, it can be seen that the higher electrical conductivity enhancement of GCPs with SAILs arises from the well dispersed graphene in the cellulose matrix. It is reasonable to assume that the SAILs are able to mediate a better exfoliation and dispersion of graphene sheets and lessen the tendency of aggregation between the RGO flakes which is beneficial to form 3D-continuous conductive network (Mohamed et al., 2016).

# 3.4. HRTEM

Further morphological characterization was carried out by imaging ultrathin sections of the GCP stabilized with 0.1 M BMIM-DBS under high resolution transmission electron microscopy (HRTEM). In general, TEM is frequently used to image nanosize materials to atomic scale resolution where a transmitted electron beam passes through the ultrathin sample (Singh et al., 2011). TEM observations at low magnification (Fig. 5a) provide the overall dispersion imaging of RGO in cellulose. It can be seen from Fig. 5b that the spaces between adjacent cellulose fibers are filled with the graphene sheets (Cataldi et al., 2015). It is therefore evident that the RGOs are finely dispersed throughout the cellulose matrix as has been previously revealed by FESEM observations. The enlarged view (Fig. 5c) shows dark lines on the edge of the RGO which indicate the existence of multilayer of graphene in the



Fig. 3. Raman spectra of the graphite (a) and GCPs stabilized surfactant and SAILs (concentration: 0.1 M): (b) SDS, (c) SDBS, (d) BMIM-DS, (e) BMIM-DBS.



Fig. 4. FESEM images of nanofibrillated kenaf cellulose (a and a'), GCP: with SDS (b and b'), with SDBS (c and c'), with BMIM-DS (d and d') and with BMIM-DBS (e and e'). The area for higher magnification imaging are marked in red square. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article).



**Fig. 5.** TEM images of the GCP stabilized BMIM-DBS: (a) typical morphologies at low magnification (b) higher magnification. Grey areas are cellulose fibers. Note that many RGO sheets are embedded throughout cellulose matrix (c) edge view of RGO sheets dispersed in cellulose matrix.

cellulose matrix (Kang et al., 2012). The reason for good dispersion is probably attributed to the enhanced interfacial interaction between RGO and cellulose due to the presence of SAILs (Bari et al., 2014; Liu et al., 2008). The possible mechanism behind this unique interaction will be discussed in the following section.

# 3.5. Studying graphene dispersion stability: zeta potential measurement

As shown above the quality of graphene dispersions can be assessed by microscopy. The important question is which surfactant properties control dispersion quality? Adsorption of ionic surfactants onto graphene imparts an effective charge onto the coated graphene. It is known that the mechanism for the stabilization of graphene dispersions by ionic surfactants is electrostatic repulsion between graphene surfaces owing to adsorbed surfactant molecules. Electrostatic repulsion is generally quantified by the electric potential in the vicinity of the surface of the coated graphene sheets, which is known as the zeta ( $\zeta$ )potential (Hunter, 1981). A surfactant-coated graphene is typically surrounded by a tightly bound layer of adsorbed surfactant ions, which, in turn, is surrounded by a more diffuse zone of mobile counterions (Lotya et al., 2009; Sun et al., 2008). The zeta potential is the potential just beyond the layer of bound surfactant ions, which is at the hydrodynamic slip plane. Graphene coated with charged molecules will display  $\zeta$ -potentials with a sign reflecting the charge of the adsorbed molecules. Therefore, graphene coated with anionic surfactant should give a negative  $\zeta$ -potential value.

In general, the dividing line between stable and unstable suspensions is taken as +30 or -30 mV with particles having zeta potentials outside these limits are normally considered stable (Hunter, 1981; Rosen, 2004). Thus, to achieve stable graphene-coated surfactant dispersions, the  $\zeta$ -potential should be maximized so that the sheets will tend to repel each other, and there will be no tendency for the particles to come together. It is therefore worthwhile to consider how to achieve graphene-coated surfactant with very high  $\zeta$ -potential. The most obvious way is by increasing the effective charge of the coated graphene. This can be done by surfactant structure modification, increasing surfactant concentration, pH and so on (Hunter, 1981; Rosen, 2004).

It can be seen from Table 2 that for all graphene dispersions, the  $\zeta$ potential values are lower than -30 mV, underlining the good stability of graphene-surfactant colloidal dispersions. It can be clearly seen here that BMIM-DBS appears to be the most negatively charged of all dispersants, with the increase of nearly 16 mV compare to the parent, SDBS, giving the most stable system out of the surfactants tested in this study. However, the strategy of changing sodium into immidazolium does not always offer significant colloidal stability improvement (in terms of electrostatic stabilization). Noting that for BMIM-DS, the ζpotential is subtly lower than SDS surfactant. Although when comparing the electrical properties and FESEM observations, BMIM-DS suggest higher affinity compared to SDS, giving moderately higher electrical conductivity and less agglomerated RGO dispersed in the cellulose. Looking only at the  $\zeta$ -potential measurements, all the results here are together too small to explain the ability of the SAILs in enhancing the nanocomposite properties.

#### 3.6. Small-angle neutron scattering

The nature of surfactant, concentration, and type of interaction are known to play a crucial role in the dispersion behavior of classical colloids. Learning from CNT-aided surfactant dispersion studies, surfactants showed various self-assembly structures which are responsible for the stabilization of the dispersions (Lin et al., 2016; Vaisman, Wagner, & Marom, 2006; Yurekli, Mitchell, & Krishnamoorti, 2004). Here, in an attempt to understand the different aggregation behavior of surfactants in solution and adsorbed on graphene, and to learn the nature of surfactant – graphene interaction for stabilization, small-angle neutron scattering (SANS) has been employed. SANS is the ideal method to provide information regarding the size, shape of surfactant self-assembly structure over the nanometer range that is complementary with microscopy technique. In general, the scattering intensity I(Q) is related to the shape, volume, and contrast of the nanoscale structures present in a sample. The contributions to I(Q) for a specified shape and size of particle is described by its form factor, P(Q). A comprehensive introductions to small-angle scattering have been published elsewhere (Feigin & Svergun, 1987; Hollamby, 2013). To provide a comparable study, SANS data were collected for all the RGO-stabilized surfactants dispersion and surfactants solutions at similar surfactant concentration and temperature.

The scattering profiles for the surfactants and SAILs (without RGO) were recorded in dilute aqueous phases and are given in Fig. 6. SANS profiles for the sodium analogue - SDS and SDBS, were indicative of charged micelles (see Fig. 6a), showing an obvious charge repulsion S (Q) peak following the Hayter-Penfold model (Hayter & Penfold, 1983); others show scattering characteristic of spherical micelles (Feigin & Svergun, 1987). The fitted micellar dimensions, R<sub>sphere</sub> for SDS and SDBS both are  $\sim 22$  Å (Table 3). It is known that both SDS and SDBS forms spherical micelles (Brown et al., 2012; Paul et al., 2005; Yurekli et al., 2004; Wang et al., 2016). Yurekli and his co-workers reported the formation of spherical micelles with  $R_{sphere}$  (18 Å at 0.5 wt%; 17 Å at 1 wt%) for SDS even with different concentrations (Yurekli et al., 2004). Magid, Li, and Butler (2000) also reported spherical micelles with radius of ~ 23 Å for 0.07 M SDS in D<sub>2</sub>O at 40 °C. The R<sub>sphere</sub> differences here can be omitted considering experimental error ( $\pm 2$ Å) and may be related to the difference surfactant concentration itself, as micelle shape and size are may be affected by surfactant concentrations, salt addition, or temperature (Feigin & Svergun, 1987; Hayter & Penfold, 1983). Earlier studies on SDBS using quasielastic light scattering give evidence of the formation sphero-cylinder micelles with R<sub>sphere</sub> of 22 Å (Cheng & Gulari, 1982). Recent SANS study also reported the formation of 22 Å spherical micelles of SDBS surfactant at 25 °C (Mohamed et al., 2018).

Just like normal surfactants, SAILs also showed the ability to selfassemble in aqueous solutions to form micelle (Brown et al., 2011, 2012). The scattering profiles here are consistent with spherical micelle for BMIM-DS. The curves could be fitted with charged spherical micelles with Hayter-Penfold charge repulsion S(Q) giving micellar radius of 25 Å. This behavior is analogous to other quaternary ammonium ILs which showed a maintained shape and size of micelle when substituting sodium for tetraalylammonium cations (Brown et al., 2011). Javadian et al. (Javadian, Nasiri, Heydari, Yousefi, & Shahir, 2014) however reported an increase of  $\sim 10$  Å of the micelle radius (assuming that micelles in spherical form conform the Stokes-Einstein equation) when changing sodium into imidazolium counterion measured using light scattering. Imidazolium-based ionic liquids were previously fitted as spherical micelles (Bowers, Butts, Martin, Vergara-Gutierrez, & Heenan, 2004). Recent research on imidazolium-based SAILs properties reported multilayer stacks model for BMIM-DS and retained the aggregation shape over various concentration (0.5-4.0 wt%) (Brown et al., 2012). Moving to BMIM-DBS, this SDBS analogue surprisingly forms cylindrical micelles with a radii of 19 Å and 140 Å in length. In comparison to BMIM-DS, the micelle shape and size transition is much more pronounced. A sphere-to-cylinder transition was previously reported on copolymer ILs comprised of poly(butadiene) and poly(ethyleneoxide) (He, Li, Simone, & Lodge, 2006). They suggested that the transition is encouraged by the preferred interfacial curvature of spherical micelles, when the IL chemical structure lacks a hydrophilic portion (shorter PEO chain). That idea might be in line with the SANS data reported here, considering the presence of a hydrophobic phenyl ring on BMIM-DBS will make it overall less hydrophilic than BMIM-DS. However, none of the other fitted parameters show any obvious trend down the anion series, making it difficult to draw firm conclusions about relationships between cation structure and aggregation.

Again, SANS was used to investigate the aggregate structure in the



**Fig. 6.** SANS profiles for surfactant solutions of (a) SDS and SDBS, (b) BMIM-DS and BMIM-DBS and RGO dispersions with (c) SDS and SDBS, (d) BMIM-DS and BMIM-DBS. [Surfactant] = 0.03 M and T =  $25 \degree$ C. Lines are model fits for charged spherical, ellipsoidal, cylindrical micelles (with Hayter-Penfold *S* (*Q*)) or a stacked disk model. Characteristic error bars are shown for the lowest intensity samples.

Table	3				
Model	fit	parameters	for	SANS	data.ª

Sample	Model	R <sub>sphere</sub> (Å)	R <sub>a</sub> (Å)	R <sub>disk</sub> (Å)	R <sub>cylinder</sub> (Å)	$X \pm 0.2$	Length (Å)
SDS	Sphere	22	-	-	-	-	_
SDBS	Sphere	22	-	-	-	-	-
BMIM-DS	Sphere	25	-	-	-	-	-
BMIM-DBS	Cylinder	-	-	-	19	-	140
SDS + RGO	Sphere	24	-	-	-	-	-
SDBS + RGO	Ellipsoid	-	31	-	-	2	-
BMIM-DS + RGO	Ellipsoid	-	40	-	-	2	-
BMIM-DBS + RGO	Stacked disk	-	-	143	-	-	-

<sup>a</sup> [surf.] = 0.030 M. Charged micelles were fitted with interparticle structure factor *S*(*Q*) for Hayter-Penfold model.

RGO-stabilized surfactants. SANS profiles for RGO dispersions with SDS, SDBS, BMIM-DS and BMIM-DBS are shown in Fig. 6c and d. The SANS data of an RGO dispersion with SDS were still characteristic of charged spherical aggregates with a Hayter-Penfold S(Q), being reminiscent of the results found for pure SDS. The presence of RGO does not have the effect of increasing the micelle size (considering the experimental error), being  $R_{sphere} = 24$  Å. There is a very small reduction of scattering intensity I(Q) throughout the Q range compared to pure SDS. This kind of observation has been previously reported with similar surfactants, suggesting the presence of weak interaction between surfactant and graphene surfaces (Mohamed et al., 2018; Yurekli et al., 2004). On the other hand, there were evolutions in micelle shape and size RGO dispersions with SDBS and BMIM-DS. Both systems have scattering profiles that can be adequately fitted with charged ellipsoidal micelles and a P(Q) with polar axis ratio ( $R_a$ ) and aspect ratio X, multiplied by Hayter-Penfold electrostatic repulsive model. The micellar radii undergo significant changes, giving R<sub>a</sub> of 31 Å for SDBS and 40 Å for BMIM-DBS. Consistent with RGO dispersions stabilized by SDS, the scattering profiles exhibit lower I(Q), evident of surfactant adsorption.

Moving to BMIM-DBS, the data reveals  $Q^{-2}$  regimes of scattering consistent with a disk-like model (Feigin & Svergun, 1987; Hollamby,

2013), giving a fitted disk radius of 143 Å. In an earlier study of graphene nanoplatelets dispersions with anionic aromatic surfactants, a disk type transition was also found for surfactants with highest dispersion efficiency (highest electrical conductivity enhancement) (Mohamed et al., 2018). Recalling the electrical properties and microscopic observation, BMIM-DBS offers an increased dispersion quality compared to the rest of the stabilizers. That result is consistent with SANS data obtained here. In that work it was assumed that the surfactant adsorbed and wrapped the graphene surfaces giving an appearance as disk-type aggregated structure (Mohamed et al., 2018). Taking all the results together, it is clear that RGO dispersions are present, further ideas for the mechanism on how the surfactant adsorbs on graphene surfaces, and the possible interaction between surfactant and cellulose will be discussed in the following section.

# 3.7. The role of surfactants, SAILs and cellulose for stable graphene dispersions: proposed mechanism

There are many ideas in the literature about the amphiphilic properties of cellulose and that dispersion/dissolution of cellulose in water/solvents are closely related to this duality (Lindman et al., 2010; Medronho & Lindman, 2015). Despite extensive studies on cellulose



Fig. 7. Schematic illustration the role of anionic surfactant ionic liquids during exfoliation and the interactions with graphene and cellulose particles.

dissolution using ionic liquids or common surfactants (Lindman et al., 2010; Medronho & Lindman, 2015; Pinkert et al., 2009), the role of individual ionic species involved in dissolution is yet to be fully understood. Regarding the mechanism of dispersion of cellulose in surfactant and SAILs solutions to be used for exfoliating graphite (see Fig. 1), it is assumed that the surfactant/SAIL headgroups interact with the hydrophilic part of cellulose through hydrogen bonding to render cellulose dispersion in aqueous phase (Lindman et al., 2010; Medronho & Lindman, 2015). Cho and his group, in their study utilizing imidazolium based ILs, suggested that the major driving force for dissolution is the presence of hydrogen bonds between the cationic part of ILs (imidazolium moiety) and the cellulose hydroxyl (Cho, Gross, & Chu, 2011). The cellulose-SAILs and surfactant combination here acted as an electrolyte for electrochemical exfoliation. This is reasonable considering that, when mixing, the cellulose/surfactant or cellulose/SAILs will behave as a typical electrolyte (Lewis & Robinson, 1970).

Literature strongly emphasize that graphene can be prepared using electrochemical methods and stably dispersed in water for a certain period of time when coated by adsorbed surfactants (Abdelkader, Cooper, Dryfe, & Kinloch, 2015; Alanyahoğlu et al., 2012). In general, electrochemical exfoliation routes to graphene are based on the intercalation of surfactant ions between the layers of graphite rod electrode due to the flow of electrical current. These ions induce expansion of the interlayer space of graphite and thus facilitate exfoliation (Alanyahoğlu et al., 2012; Parvez et al., 2014). Taking all the results together, it is clear that BMIM-DBS offer increased dispersion quality as compared to others down the series. To understand the molecular interaction underlying this graphene-compatibility, a mechanism is proposed.

Applying bias voltage (7 V) in the electrolyte leads to oxidation at the grain boundaries and edge sites of graphite (Parvez et al., 2014). This results in the opening up of graphite edge sheets. The BMIM-DBS tails are then adsorbed within the edge sheets and initiate the expansion of graphite layers (see Fig. 7). As a result, exfoliated black powder of expanded graphite oxide was dispersed in the solution after applying electrical current for 24 h. A subsequent peeling process involving low shear forces (ultrasonication) are thought to separate the graphitic material into multilayer flakes. The materials produced required reduction for deoxygenation of the oxygen-containing functional groups. Typically reduction of graphite oxide substantially reduces the dispersibility in water. Remarkably however, visual observation showed that the resulting dispersion was stable for over 1 month. In this case, BMIM-DBS played a dual role: as an exfoliant/intercalant and a stabilizer for the reduced graphene oxide in the mixture (Alanyahoğlu et al., 2012; Guardia et al., 2014).

It is believed that the distinct aggregation behavior of BMIM-DBS is a consequence of the increased RGO surfaces being occupied by BMIM-DBS, wrapping up RGO sheets for stabilization. A previous study revealed that an anionic tri-chain aromatic surfactant namely TC3Ph3 (sodium 1,5-dioxo-1,5-bis(3-phenylpropoxy)-3-((3phenylpropoxy)carbonyl) pentane-2-sulfonate) showed a similar disk-type aggregation due to a full-coverage of surfactant-wrapping the graphene surfaces (Lin et al., 2016; Mohamed et al., 2018). By analogy to that model, it was surmised that BMIM-DBS occupy the graphene faces as much as possible with the tails in contact with the graphene surfaces, driven by the hydrophobic effect (Matarredona et al., 2003; Mohamed et al., 2016). The presence of phenyl rings in the chemical structure undoubtedly provides graphene-affinity as has been noted in extensive literatures (Lin & Xing, 2008; Matarredona et al., 2003; Mohamed et al., 2016; Tkalya, Ghislandi, de With, & Koning, 2012). There is also a possibility of hydrophobic interaction between the hydrophobic part of cellulose and graphene surface considering that cellulose has amphiphilic properties (Lindman et al., 2010; Medronho & Lindman, 2015; Medronho et al., 2012). Indeed, the adsorbed BMIM-DBS along with cellulose shields the attractive van der Waals forces between graphene sheets. The most preferential arrangement of head and tails of BMIM-DBS and cellulose is that BMIM-DBS initially adsorbed with the tail lying flat approaching the graphene surfaces to maximize the hydrophobic interactions and begin wrapping graphene until the surface is saturated (see Fig. 7). The SAIL headgroups are hydrogen bonding with cellulose hydroxyls while the hydrophobic part is weakly interacting with graphene leading to a more stabilize dispersion.

# 4. Conclusions

Graphene and graphene based materials still show great promise in many technological applications, but their large-scale production and processing by simple and cost effective means still constitute significant issues in the path of widespread implementation. Here, this study used a straightforward method for the preparation of ready-to-use material that is based on electrochemical exfoliation of graphite in a mixture of anionic surfactant ionic liquids and cellulose as the electrolyte. The advantages of this approach over existing methods to produce similar nanocomposite materials are: lower environmental hazards (the medium is water), economically efficient starting materials of ionic liquids, and simple instrumental setup (Parvez et al., 2014; Suriani et al., 2015, 2016).

Optical microscopy and Raman spectroscopy shows the uniform dispersion of reduced graphene matrix on nanofibrillated kenaf cellulose. Zeta potential measurements reveal increased dispersion stability for BMIM-DBS but firm conclusions on the relationship with counterion type cannot be drawn since BMIM-DS shows a lower zeta potential value. In dilute aqueous phases SANS data indicated transitions in aggregation structure in the presence of graphene for all compounds except SDS. Particularly interesting is how BMIM-DBS induces a cylinderto-disk transition, all consistent with fully covering surfactant-wrapping of graphene surfaces, deviating from the spherical-to-ellipsoid aggregates seen for the rest compounds (Lin et al., 2016; Matarredona et al., 2003; Mohamed et al., 2018). It does appear that aromatic ring coupled with bulkier counterion type is the best combination, and contributes to the increased graphene compatibility.

The unique combination of surfactant ionic liquids with polymer as a greener exfoliating medium may pave the way to development of novel nanocomposite processing routes. Furthermore, these economical and efficient SAILs alone may find applications as both exfoliating agents and dispersants for other layered or two dimensional materials such as metal dichalcogenides and metal-organic frameworks (Mas-Balleste, Gomez-Navarro, Gomez-Herrero, & Zamora, 2011). This graphene conductive cellulose paper produced here may also find applications in electronic devices as flexible supercapacitors, electrodes, or sensors (Hou, Xu, & Li, 2018; Weng et al., 2011).

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#### Appendix A. Supplementary data

Supplementary material related to this article can be found, in the online version, at doi:https://doi.org/10.1016/j.carbpol.2018.08.040.

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