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Cite this: *RSC Adv.*, 2017, 7, 9998

The impact of graphene nano-plates on the behavior of novel conducting polyazomethine nanocomposites

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The aim of the current work was to synthesize a new series of polyazomethine/graphene nano-plate nanocomposites in the form of PAMs/GNP_{a-e}. Different compositions were fabricated according to the percentage of GNP loading. The composites were formed *in situ* during the polymerization reaction for the formation of pure PAMs. The corresponding pre-monomer (1), aminothiazole monomer (2) and consequently, new conducting PAMs polymer were synthesized. The structures of these compounds were confirmed by elemental and spectral analyses, including FT-IR, ¹H-NMR and ¹³C-NMR. The pure PAMs and PAMs/GNP_{a-e} nanocomposites were characterized using available characterization techniques. X-ray diffraction (XRD), thermal analyses, conductivity tests and scanning electron microscopy (SEM) were also performed. Furthermore, a comparative study was carried out to demonstrate the role of GNP inclusion into the PAMs polymer matrix in the form of nanocomposites. Intense absorption peaks related to GNPs and pure PAMs were observed in the FT-IR spectra of our new PAMs/GNP_{a-e} nanocomposites, and these peaks represent a good reference for the fabrication of such nanocomposites. The thermal degradation of pure PAMs and the PAMs/GNP_{a-e} nanocomposites occurred in two steps. The degradation steps were dependent upon the nature of the desired nanocomposites, which was mainly related to the decomposition of the pure PAMs content. Moreover, SEM images offered more obvious evidence for the formation of the nanocomposites. In addition, an electrical conductivity test showed a complete change in the conductivity behavior of PAMs upon GNP loading, from a very poor state to semiconductor behavior in the PAMs/GNP_{a-e} nanocomposites.

Received 27th December 2016
 Accepted 21st January 2017

DOI: 10.1039/c6ra28756e

rsc.li/rsc-advances

1. Introduction

Organic/inorganic nanocomposites and in particular, those involving a polymer matrix, have attracted wide attention in different fields, especially in the field of material science, towards the development new efficient materials with excellent properties such as low cost, low density and photoconductivity for electronics, in addition to many other fantastic properties reported in the literature.¹⁻⁵ Recently, these nanocomposites have been used instead of steel, for example, due to their low specific weight and high strength. Much more attention has been given to polymer nanocomposites, perhaps due to their improved properties, as reported in the reinforcement of graphene with polymers.⁶⁻⁸ Several attempts have been made to reinforce inorganic particles with polymers at the nanoscale.⁹⁻¹¹

More specifically, carbon nanotubes, graphene oxide and graphene nano-plates are considered to be the most important nano-materials, which exhibit a wide variety of properties in different fields of application. It is one of our main aims to study the effect of nano-fillers on polyazomethine chains, and to investigate graphene nano-plates (GNPs), which have electrical properties sufficient to completely change the conducting behaviors of any material, such as our synthesized polyazomethine. Besides that, GNPs and other nano-fillers such as CNTs have other exceptional properties, leading to new materials with excellent properties such as high specific surface areas, unique size distributions and tube structures in the case of CNTs. Such properties permit nano-fillers to be used in different industrial applications such as sensing, inorganic pollutant remediation, catalysis, solar cells, composites, medical applications, photonics and fuel cells.¹²⁻¹⁹ In the present work, a new series of polyazomethine/graphene nanocomposites of different compositions was synthesized. *In situ* nanocomposite formation was achieved during the polymerization for the preparation of PAMs. The new composites were characterized using characterization techniques including FT-IR spectroscopy measurements, thermal analyses (TGA, DTG and DTA), X-ray diffraction (XRD), conductivity tests, and

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scanning electron microscopy (SEM). Then, a comparative study was carried out to demonstrate the role of GNP inclusion in the polyazomethine polymer chains.

2. Experimental

2.1. Chemicals, solvents and reagents

Diaryl ether and chloroacetyl chloride were purchased from Merck. Anhydrous aluminum chloride was purchased from Sigma-Aldrich. Absolute ethanol, carbon disulphide and concentrated hydrochloric acid were all purchased from BDH. Thiourea and sodium acetate were also purchased from BDH. All the above mentioned chemicals and reagents were used as received without any further purification. Furthermore, graphene nano-plates (GNPs) and terephthalaldehyde were also purchased from Sigma-Aldrich and were used as received.

2.2. Preparation of monomer and pre-monomer

4,4'-Di(chloroacetyl)diaryl ether pre-monomer (1) as well as 4,4'-di(2''-aminothiazol-4''-yl)diaryl ether monomer (2) were prepared as previously reported in our previous studies as follows.^{20,21}

2.2.1. 4,4'-Di(chloroacetyl)diaryl ether pre-monomer (1). 3.2 mL of diaryl ether and 3.18 mL of chloroacetyl chloride were mixed together with 80 mL of carbon disulfide in a 250 mL conical flask. The reactant mixing ratio was 20 : 40 mmol. The whole mixture was cooled to 0 °C under constant stirring using an ice-bath. Simultaneously, 10.68 g (80 mmol) of aluminum chloride was added slowly to the mixture. The reaction was completed after 6 hours; then carbon disulfide was removed from the reaction mixture by evaporation. The residue was transferred into cold hydrochloric acid. A yellow precipitate was obtained and collected by filtration, and was then crystallized from ethanol. The product melting point was measured to be 103 °C and its yield was calculated at 82%.

2.2.2. 4,4'-Di(2''-aminothiazol-4''-yl)diaryl ether monomer (2). In a round bottom flask, 2 g of pre-monomer (1) was mixed with 0.94 g of thiourea in the presence of 50 mL absolute ethanol as the reaction solvent. The mixture was heated and refluxed for 4–6 hours to completion. A cold solution of sodium acetate was first prepared and then transferred to the reaction mixture leading to the formation of a final precipitate. This crude product was collected by filtration. Ethyl acetate was used as a crystallization solvent producing yellowish needles with a melting point of 240 °C and a yield of 87%. The chemical structures of both the pre-monomer (1) and monomer (2) were checked by elemental and spectral analysis, giving correct structures as previously reported.^{20,21}

2.3. Synthesis of polyazomethine conducting polymer

The new desired conducting polyazomethine polymer was synthesized using ideal polycondensation procedures as follows: a mixture of monomer (2) 0.732 g and terephthalaldehyde 0.268 g was suspended in 40 mL of absolute ethanol inside a three-necked flask under an inert nitrogen atmosphere. Drops of piperidine were added to the mixture at room temperature, and

the mixture was stirred during each addition. After the additions were complete, the reaction mixture was heated to ~80 °C with constant stirring for 10–12 hours. In the early stages of the reaction, the desired polymer product began to precipitate and the amount increased with increasing reaction time. The reaction was completed after 10–12 hours. A yellow solid was separated out, collected and washed with simple hot organic solvents during filtration. The new product then was dehydrated under reduced pressure (1 mmHg) at 70 °C for two days. Anal. calcd for C₂₆H₁₆N₄S₂O: C, 67.24; H, 3.45; N, 12.07; S, 13.79. Found: C, 65.80; H, 3.83; N, 13.05; S, 14.14. IR (ATR) ν cm⁻¹: at 3080 (w, CH aromatic stretching), at 1610 (s, C=N azomethine group), at 1620–1590 cm⁻¹ (m, phenylene rings), at 1255–1260 (s, ether bond), and at 750 (w, C-S carbon sulfur single bond). ¹H-NMR (DMSO-d₆, ppm): δ = 8.2–8.3 (s, s, 2H, 2CH=N azomethine), 7.73–7.55 (s, s, 2H, aromatic H thiazole ring) 8.1–7.87 and 7.18–6.92 (m, 12H, H of aromatic diaryl ether and H of *p*. phenylene). ¹³C-NMR (DMSO-d₆, ppm): δ = 175.2, 163, 159.4, 151.6, 136.9, 134.8, 133, 128.77, 127, 125.2, 118 and 110, in agreement with the submitted structure.

2.4. Synthesis of PAMs/GNP_{a-e} nanocomposites

In a typical procedure, a new series of PAMs/GNP_{a-e} was synthesized using an *in situ* polymerization procedure depending mainly on a solution polycondensation process. The polymerization process was carried out first at room temperature for 30 min. Then it was heated up to 80 °C under a nitrogen atmosphere for 12 hours and the remaining procedure was carried out in the same way as that for the synthesis of pure PAMs described above. The produced nanocomposites possessed a fixed weight and ratio of monomer (2) and terephthalaldehyde. Different loadings of GNPs were introduced (wt by wt%), with respect to the total weight of the amino-thiazole monomer (2). GNP loadings of 2%, 5%, 10%, 20% and 30% were employed for each composite. The PAMs/GNP_{a-e} nanocomposites were synthesized as follows: the calculated amounts of GNP nanoparticles were suspended into a mixture of monomer (2) 0.732 g and terephthalaldehyde 0.268 g in 40 mL of absolute ethanol. The whole mixture was ultrasonicated for 15 min. After that, drops of piperidine were added to the mixture and it was stirred during the addition. Simultaneously, a nitrogen atmosphere was allowed to flow over the reaction. Then, the reaction mixture was heated to ~80 °C with constant

Table 1 The suggested symbols and chemical compositions for pure PAMs and PAMs/GNP_{a-e} nanocomposites

| Products | Monomer (1) (weight, g) | Terephthalaldehyde (weight, g) | GNP% (weight, g) |
|-----------------------|----------------------------|-----------------------------------|---------------------|
| Pure PAMs | 0.732 g | 0.268 g | — |
| PAMs/GNP _a | 0.732 g | 0.268 g | 2% (0.02) |
| PAMs/GNP _b | 0.732 g | 0.268 g | 5% (0.05) |
| PAMs/GNP _c | 0.732 g | 0.268 g | 10% (0.10) |
| PAMs/GNP _d | 0.732 g | 0.268 g | 20% (0.20) |
| PAMs/GNP _e | 0.732 g | 0.268 g | 30% (0.30) |



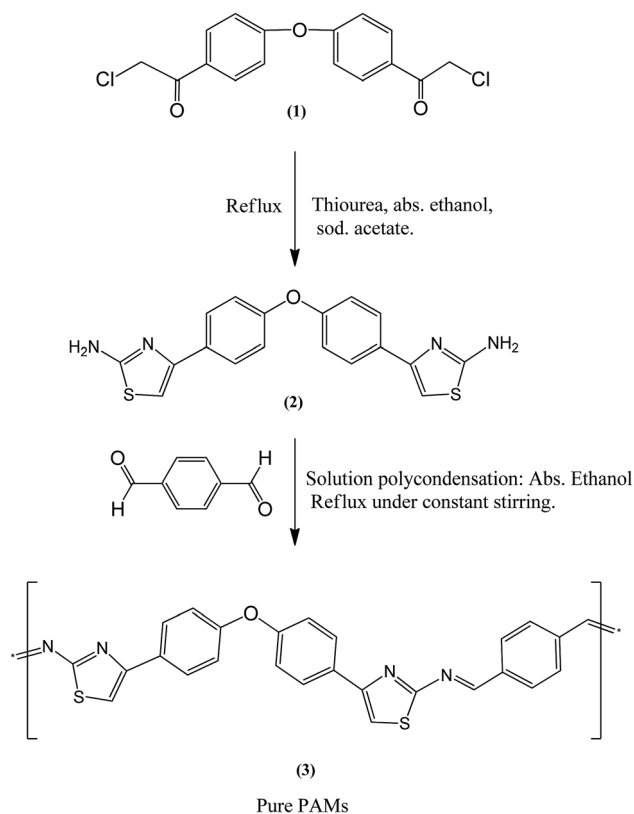


Fig. 1 Synthesis procedure for the pre-monomer (1), monomer (2) and pure polyazomethine PAMs.

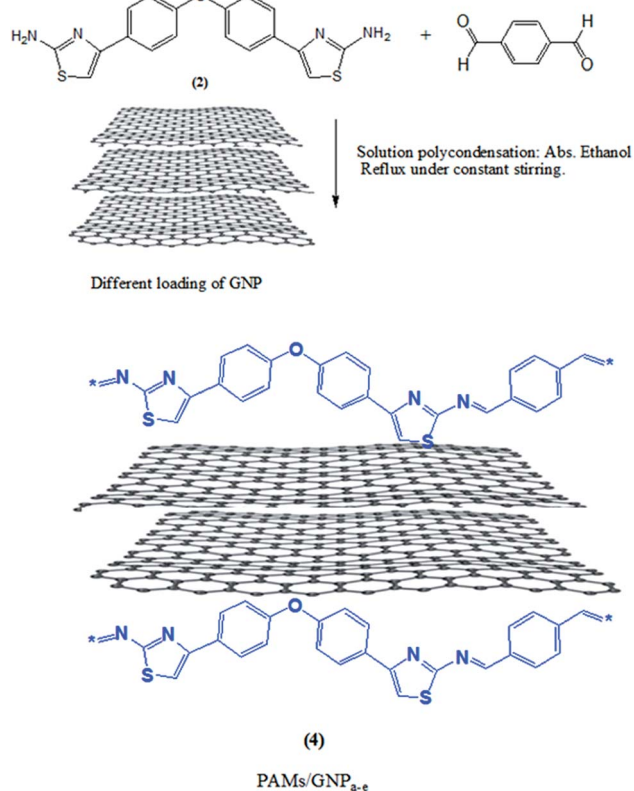


Fig. 2 Synthesis procedure of the polyazomethine nanocomposites PAMs/GNP_{a-e}.

stirring for 12–14 hours. The desired nanocomposites were separated out by filtration, washed with deionized water and hot ethanol in sequence, and then dried under vacuum at 70.0 °C for two days. Table 1 represents the suggested symbols and chemical compositions for pure PAMs and PAMs/GNP_{a-e}.

2.5. Identification and characterization techniques

The melting points of the pre-monomer (1) and monomer (2) were determined on a Gallen-kamp melting point apparatus with a digital thermometer. Elemental analysis (C, H, N) was performed on a CHN rapid analyzer. Fourier transform infrared (FT-IR) spectra were obtained using an ATR smart part technique in the wave number range 4000–400 cm⁻¹ on a Nicolet-6700 Thermo Fisher Scientific FT-IR spectrophotometer. ¹H-NMR and ¹³C-NMR spectra were recorded on a Bruker Advance 800 MHz using DMSO-d₆ as a solvent. TMS was used as the interior reference. The X-ray diffractograms of the pure PAMs and PAMs/GNP_{a-e} were collected using a Philips X-ray PW1710 diffractometer, and Ni-filtered CuK α radiation in the 2 θ range from 10 to 80°. Thermal analyses (TGA, DTG and DTA) measurements were obtained on a TA 2000 thermal analyzer. All measurements were carried out in an air flow under the same conditions with

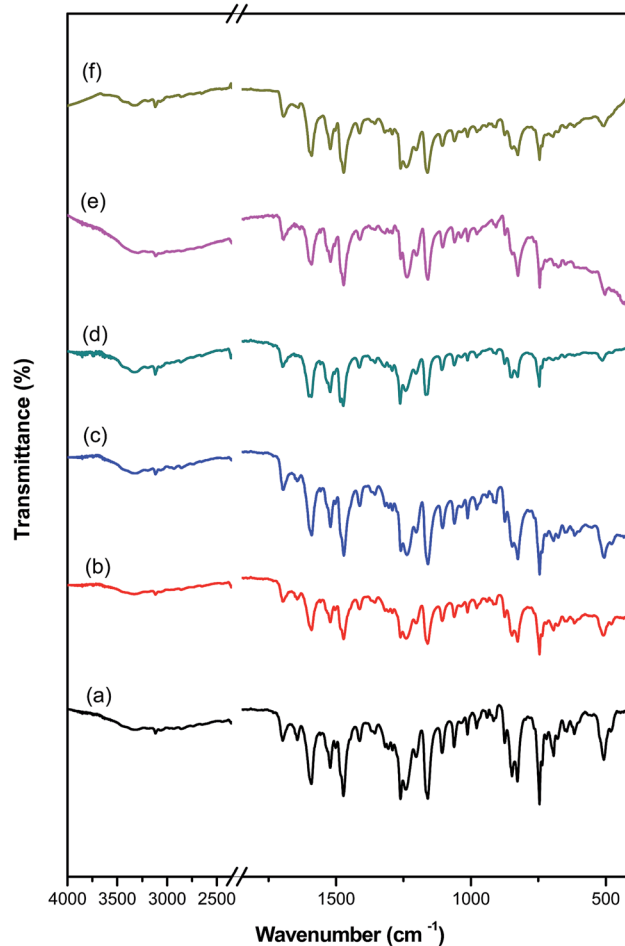


Fig. 3 FT-IR spectra of pure polyazomethine and its nanocomposites: (a) pure PAMs, (b) PAMs/GNP_a, (c) PAMs/GNP_b, (d) PAMs/GNP_c, (e) PAMs/GNP_d and (f) PAMs/GNP_f.



a heating rate $10^{\circ}\text{C min}^{-1}$. The morphological features of the pure PAMs and PAMs/GNP composites were imaged using SEM on a Jeol JSM-5400 LV instrument. Such SEM images were recorded using a Penta Z Z-50P Camera with Ilford film. The microscope was operated at an accelerating voltage of 15 kV and a 10 mm working distance from the carbon film. The electrical conductivity was measured using a conductivity cell. The samples were converted into compressed disks (under a pressure of up to 2 tons per cm^2) of 1 cm in diameter and about 1–2 mm in thickness. Silver paste was used to coat the compressed disks on both sides. To test for ohmic contact, a Hioki LCR high tester 3531 was used, employing a two-probe method, as a function of temperature up to 900 K and frequency (10 kHz to 5 MHz).

3. Results and discussion

3.1. Conducting PAMs synthesis and fabrication of PAMs/GNP_{a-e} nanocomposites

Experimentation led us to conclude that ethyl alcohol is the preferable solvent for the polymerization of PAMs. PAMs

polymer is strong enough to be precipitated as a product when the polymerization process has started. A new conducting polyazomethine polymer based on diaryl ether was synthesized by the reaction of a thiazole based monomer (2) and terephthalaldehyde using a polycondensation method.^{21–23} Prior to the polymerization reaction, the 4,4'-di(chloroacetyl) diaryl ether pre-monomer (1) and the 4,4'-di(2''-aminothiazol-4''-yl)diaryl ether monomer (2) were prepared and their chemical structures were analyzed by obtaining both elemental and spectral data as reported in our previous studies.^{20,21} The synthetic route for the synthesis is given in Fig. 1. The chemical structure of this new conducting polymer was obtained using elemental analysis, FT-IR and NMR spectroscopy. The structure was confirmed by its corresponding FT-IR and NMR spectral data. FT-IR results obtained using an ATR technique show several major distinctive absorption peaks, including azomethine peaks, CH stretching of aromatics and C–O–C ether peaks. Moreover, the $^1\text{H-NMR}$, and

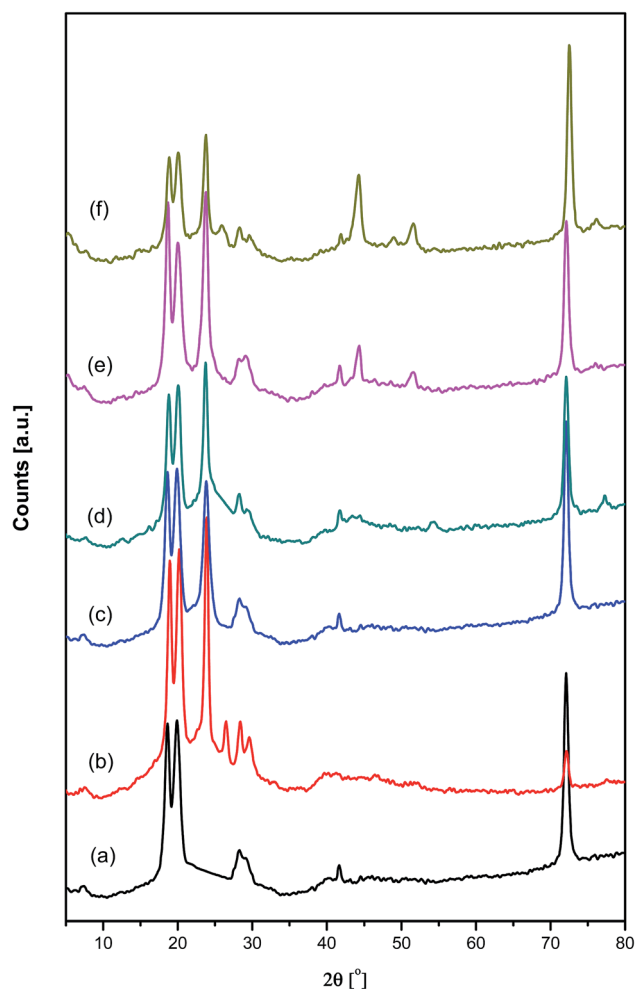


Fig. 4 XRD diffraction patterns of pure polyazomethine and its nanocomposites: (a) pure PAMs, (b) PAMs/GNP_a, (c) PAMs/GNP_b, (d) PAMs/GNP_c, (e) PAMs/GNP_d and (f) PAMs/GNP_f.

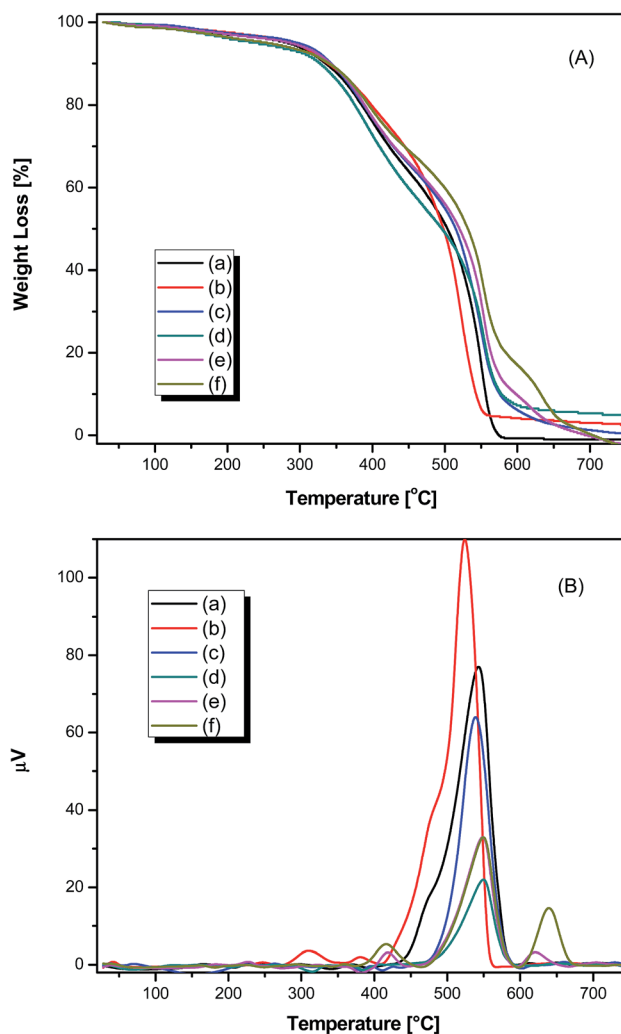


Fig. 5 TGA (A) and DTG (B) thermograms of pure polyazomethine and its nanocomposites (a) pure PAMs, (b) PAMs/GNP_a, (c) PAMs/GNP_b, (d) PAMs/GNP_c, (e) PAMs/GNP_d and (f) PAMs/GNP_f, in air at a heating rate of $10^{\circ}\text{C min}^{-1}$.



^{13}C -NMR results show distinctive symmetrical peaks which are in agreement with the submitted structure as explained in the experimental section.

During the synthesis of the designed products, the polymerization process was adjusted in order to optimize the fabrication process. In a typical procedure, a new series of PAMs/GNP_{a-e} nanocomposites was fabricated by an *in situ* polymerization procedure.²⁴ The fabrication procedures are similar to the earlier aforesaid procedures for PAMs. This fabrication method is mainly based on different loadings of GNPs in the presence of a fixed weight aminothiazole monomer (2) and terephthalaldehyde. The fabrication procedure begins at room temperature, followed by heating under a nitrogen

atmosphere. Fig. 2 shows a schematic representation of this fabrication procedure.

3.2. Identification and characterization of PAMs/GNP_{a-e} composites

As diverse identification and characterization techniques, FT-IR spectroscopy, wide angle XRD diffraction, TGA thermal analyses and SEM scanning electron microscopy were employed in order to identify and to characterize the structures of the fabricated PAMs/GNP_{a-e} nanocomposites.

The FT-IR results give clear evidence for the bonding interaction between PAMs and the PAMs/GNP_{a-e} nanocomposites. FT-IR spectroscopy confirms that pure PAMs physically interact with the PAMs/GNP_{a-e} nanocomposites in the composite materials. This type of interaction is considered an important method of bonding between any polymer matrix and its imbedded nano-filler. The FT-IR spectroscopic analysis for pure PAMs and PAMs/GNP_{a-e} nanocomposites formulations was performed between 4000 and 400 cm^{-1} . Fig. 3 displays the FT-IR spectra for pure PAMs and the PAMs/GNP_{a-e} nanocomposites. The spectrum in Fig. 3a shows intensive characteristic absorption peaks for pure PAMs. The peak at 1610 cm^{-1} is attributed to C=N in the azomethine groups. Another peak at around 1255–1260 cm^{-1} is assigned to C–O ether bonds. Furthermore, the peaks at 3080 cm^{-1} and 1620–1590 cm^{-1} are due to the CH stretching of aromatic, phenylene rings, while that at 750 cm^{-1} is due to C–S carbon sulfur single bonds.

Fig. 3b–e show the FT-IR results for PAMs/GNP_{a-e}. The immersion of GNPs into the pure PAMs polymer matrix

Table 2 Thermal behavior of pure PAMs and PAMs/GNP_{a-e} nanocomposites

| Products | CDT _{max} ^a (°C) | Temperature (°C) for various percentage decompositions ^b | | | | |
|-----------------------|--------------------------------------|---|-----------------|-----------------|-----------------|-----------------|
| | | T ₁₀ | T ₂₀ | T ₃₀ | T ₄₀ | T ₅₀ |
| Pure PAMs | 545 | 332 | 380 | 418 | 465 | 503 |
| PAMs/GNP _a | 538 | 340 | 400 | 450 | 475 | 505 |
| PAMs/GNP _b | 542 | 341 | 388 | 423 | 468 | 510 |
| PAMs/GNP _c | 554 | 340 | 388 | 425 | 470 | 512 |
| PAMs/GNP _d | 555 | 348 | 396 | 438 | 490 | 528 |
| PAMs/GNP _e | 555 | 350 | 400 | 445 | 500 | 534 |

^a Determined from DTA curves. ^b The values were determined by TGA at a heating rate of 10 °C min⁻¹.

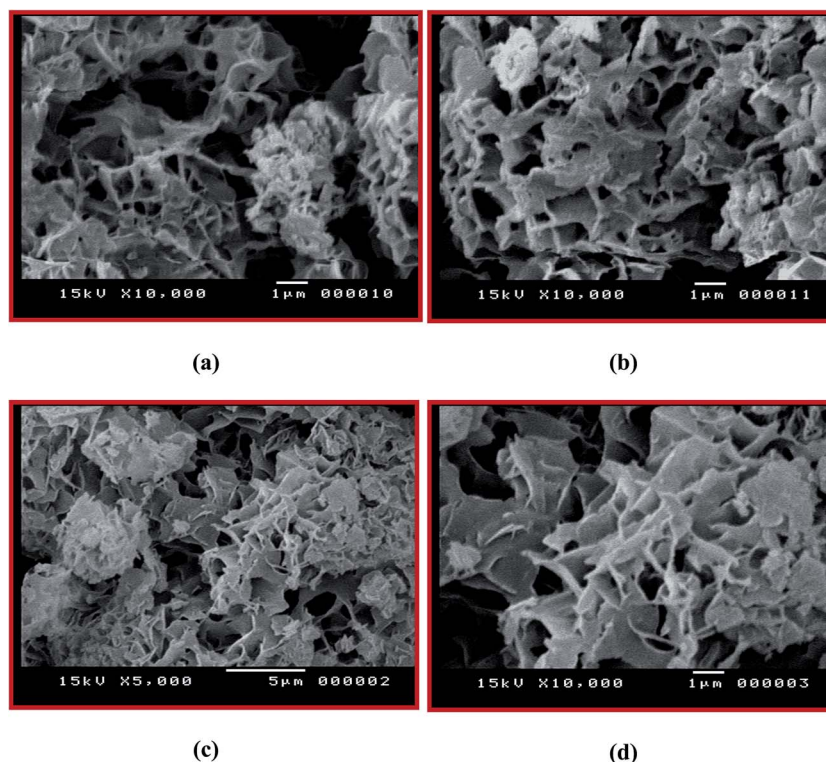


Fig. 6 SEM images of the PAMs/GNP_b nanocomposite (a, $\times = 5000$ and b, $\times = 10\,000$) and PAMs/GNP_c nanocomposite (c and d, $\times = 10\,000$).



somewhat changes its IR spectrum. The nanocomposite formation was confirmed by FT-IR spectra (Fig. 3b–e). The spectra of our new nanocomposites show intensive absorption peaks related to GNPs.^{25–28} Furthermore, all of the adsorption peaks related to pure PAMs can be clearly observed in the spectra of the PAMs/GNP_{a–e} nanocomposites as shown in Fig. 3b–e.

Fig. 4 shows the XRD diffraction patterns for pure PAMs and the fabricated PAMs/GNP_{a–e} nanocomposites. The XRD patterns were obtained over the range of $2\theta = 4–80^\circ$. The 2θ scan for pure PAMs shows three crystalline diffraction peaks. These three crystalline peaks appear at nearly $2\theta = 18.8^\circ$, 20.0° and 72.5° , and are due to the higher degree of crystallinity and highly ordered crystal structure of this new conducting PAMs material. Moreover, the corresponding diffraction patterns show irregular halo peaks located within the range of $27.2–30.3^\circ$ and at $2\theta = 41.6^\circ$, indicating a minimal level of crystallinity. These diffraction peaks are considered to be intermediate between

organized and unorganized interferences. These peaks are only revealed in the PAMs diffractogram. Furthermore, XRD diffraction patterns for the PAMs/GNP_{a–e} nanocomposites with different GNP loading clearly indicate the formation of our targeted composite products. The previously mentioned patterns exhibit a slight overlap of the diffraction peaks, which corresponds to the PAMs and GNP loading ratios. The GNPs present a strong diffraction peak at around $2\theta = 25.56^\circ$ (002) characteristic of carbon-based materials, which corresponds to graphite planes. Two more tiny peaks are present at about $2\theta = 44.27^\circ$ and $2\theta = 52.67^\circ$.^{29,30} All of the previously mentioned peaks can be clearly seen in the XRD pattern for the PAMs/GNP_{a–e} nanocomposites, as shown in Fig. 4. Furthermore, no more organized and/or unorganized bands that may be related to another crystal structure, impurities and/or phases have been found in the diffractograms of our composites.

The thermal properties of pure PAMs and the PAMs/GNP_{a–e} nanocomposites were probed using TGA, DTG and DTA in air

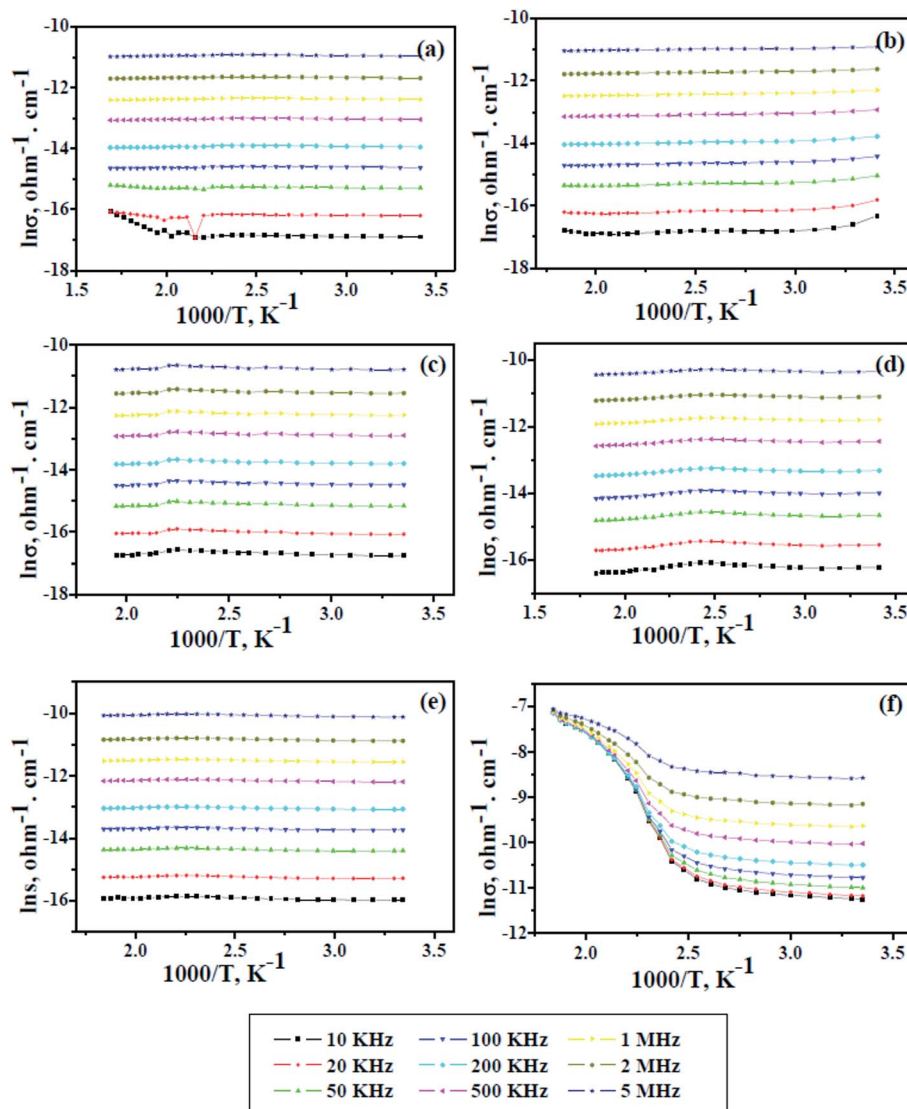


Fig. 7 Conductivity and alternate of temperature for pure PAMs (a) and PAMs/GNP_{a–e} nanocomposites (b–f) as a function of applied frequency.



with a $10\text{ }^{\circ}\text{C min}^{-1}$ heating rate as illustrated in Fig. 5. The thermogravimetric (TG) curves of all samples exhibit the same pattern of decomposition and show a small weight-loss that almost finishes before $170\text{ }^{\circ}\text{C}$. Such behavior is due to the removal of moisture and attached solvents. The TG curves are given in Fig. 5a. The T_{10} - T_{50} values represent the temperatures for variable % weight losses at 10%, 20%, 30%, 40% and 50% as illustrated in Table 2. The initial decomposition of these products occurs a few degrees before T_{10} . For pure PAMs, the degradation mainly occurs in two steps. The first step starts at $272.19\text{ }^{\circ}\text{C}$ and ends at $458.39\text{ }^{\circ}\text{C}$, whereas the second degradation step starts at $458.39\text{ }^{\circ}\text{C}$ and ends at $748.66\text{ }^{\circ}\text{C}$. For the other composite PAMs/GNP_{a-e} products, the degradation also occurs mainly in two steps. The average degradation rate for the first step is comparatively faster than that observed for the second step. The degradation steps are dependent upon the nature of the fabricated composites, and mainly involve the degradation of the polyazomethine polymer chains. Such degradation

involves scission of azomethine bonds, as well as scission of ether linkages, followed by the release of loose concise chains depending on the type of composite material, leading to the formation of burnt carbon particles as final products.³¹⁻³³ Upon closer look at the TG curves, it is easy to detect that the fast degradation step in all samples starts at temperatures higher than $300\text{ }^{\circ}\text{C}$, even for pure PAMs. Thus, it is quite acceptable to claim that these products are thermally stable up until quite high temperatures. The maximum composite degradation temperature (CDT_{max}) corresponds to the temperature at which the detected weight loss reaches its maximum. CDT_{max} can be easily detected from the DTA curves shown in Fig. 5b. CDT_{max} for all the samples are similar and appear in the same range of $538\text{--}555\text{ }^{\circ}\text{C}$, as illustrated in Table 2. All measured values are in agreement with the values obtained from the TGA and DTG curves. PAMs/GNP_a shows the lowest CDT_{max} of all the fabricated samples ($538\text{ }^{\circ}\text{C}$), while PAMs/GNP_{d,e} shows the highest CDT_{max} ($555\text{ }^{\circ}\text{C}$). The order of the CDT_{max} values is as follows:

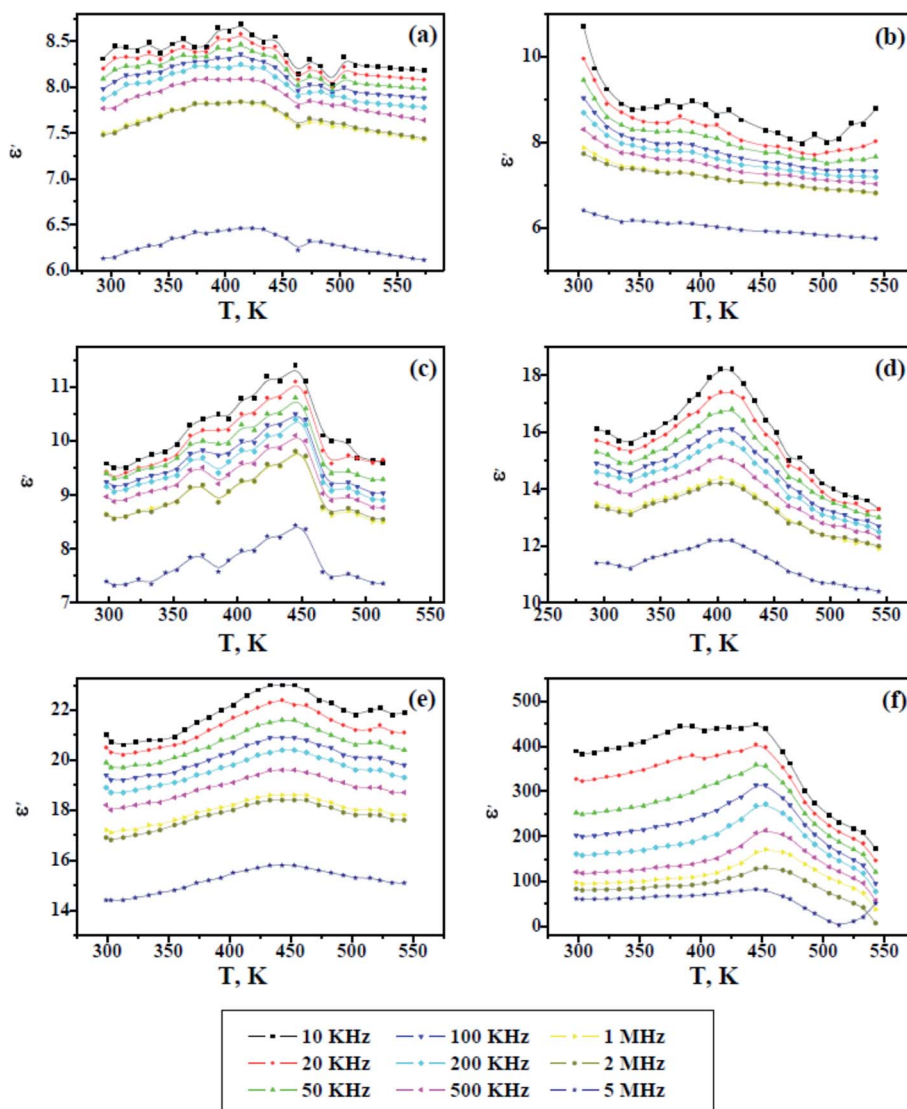


Fig. 8 Dielectric constant versus temperature for pure PAMs (a) and PAMs/GNP_{a-e} nanocomposites (b-f).



PAMs/GNP_{d,e} > PAMs/GNP_c > PAMs > PAMs/GNP_b > PAMs/GNP_a. The thermal stability of the fabricated products mainly depends on the amount of GNP that acts as reinforcements within the PAMs polymer matrix. This observation clearly appears at T_{50} and at all other temperatures for all composites except PAMs/GNP_a.

One more obvious proof for the formation of the composites is their SEM characteristics, which were analyzed using scanning electron microscopy. The morphological aspects of the PAMs/GNP_b and PAMs/GNP_c nanocomposites as selected examples, were examined using a Jeol JSM-5400 LV SEM testing instrument. The SEM test is described in detail in the experimental part. Our main aim for this experiment is to show the morphological changes in the PAMs as a result of GNP insertion into its polymer matrix. The pure PAMs surface shows that it possesses a coral reef-like structure with micro-holes in the form of porous spongy shapes, as clearly observed in b. In the

SEM images for PAMs/GNP_b, the GNPs are inserted inside the polymer matrix and fill in some of its spongy tiny apertures. This can be seen in the low and high magnification images, $\times = 5000$, 10 000 (see Fig. 6a and b respectively). Moreover, the PAMs/GNP_b images display the same features, with more stacking of GNPs over the coral reef structure indicating a higher GNP loading, $\times = 10\ 000$ (Fig. 6c and d).

3.3. Conductivity measurements

The electrical conductivity of pure PAMs and the GNP-based nanocomposites PAMs/GNP_{a-e} was measured as a function of temperature as shown in Fig. 7. In addition, the test was also performed over a wide range of frequencies, starting at 10 kHz and ending at 5 MHz, to probe the frequency dependence. This due attributed to the grain boundaries and mediator polarization at contacts of the tested materials such as polymers as well as polymer composites.³⁴ In such types of materials, the

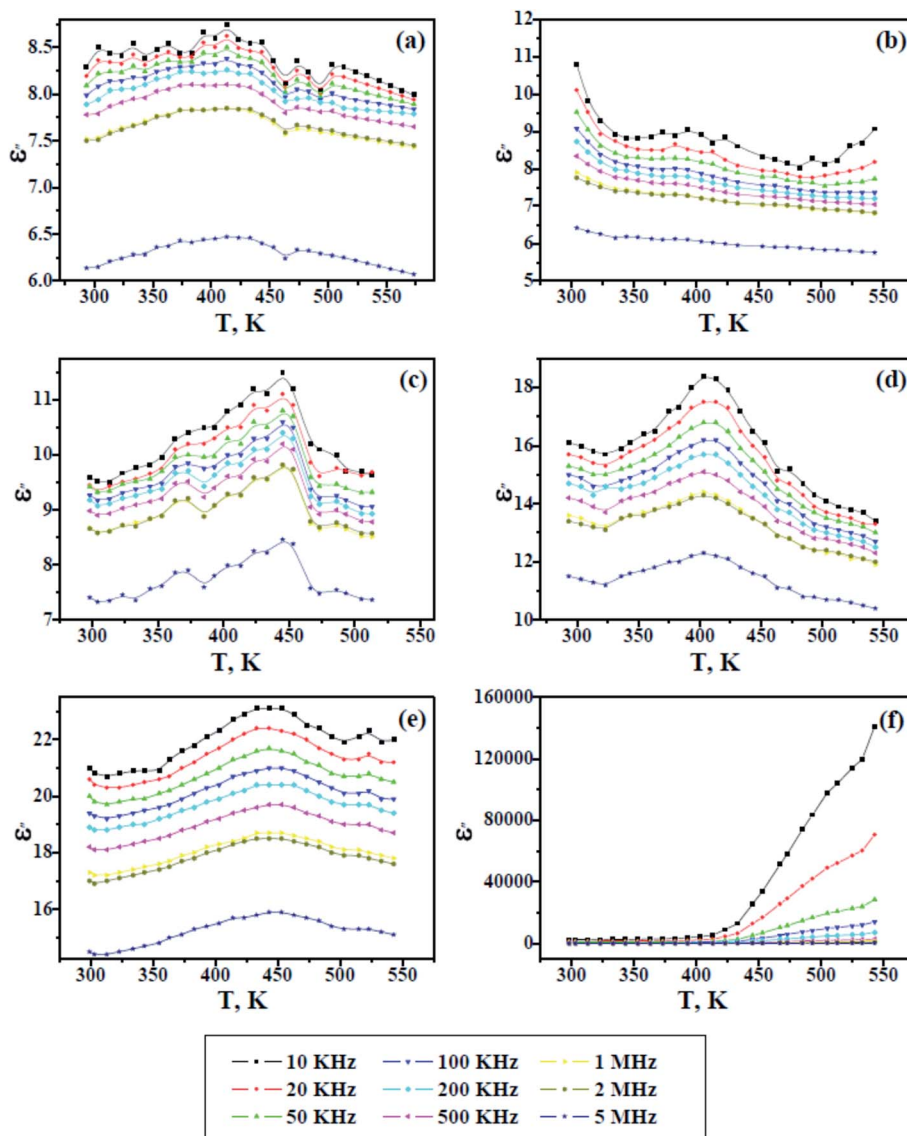


Fig. 9 Dielectric loss versus temperature for pure PAMs (a) and PAMs/GNP_{a-e} nanocomposites (b-f).



conductivity is clearly dependent on the frequency at which the sample is measured. The data in Fig. 7 prove that all the measured samples show a significant increase in their conductivity values with a gradual increase in the applied frequency. The frequency dependence conductivity behavior of pure PAMs gives a poor $\ln \sigma$ value in the range of 10^{-16} to 10^{-11} $\text{ohm}^{-1} \text{cm}^{-1}$ over the applied frequency range. Almost the same $\ln \sigma$ values are obtained for the PAMs/GNP_{a-e} composites, except for PAMs/GNP_e, which gives better value in the same range as semiconductors (10^{-7} $\text{ohm}^{-1} \text{cm}^{-1}$). The results in Fig. 7a show that the $\ln \sigma$ value remains constant while the temperature is increased to a certain limit, and then it somewhat comparatively increases. This is due to the prevalence of electron-phonon clashes because of their increase in free movement. This phenomenon is clearly observed at 10, 20 and 50 kHz. Similar observations are detected for all composite formulations as shown Fig. 7b–e, except for PAMs/GNP_e, which gives an almost constant $\ln \sigma$ value at lower temperatures, and then shows a sharp increase when the temperature increases as shown in Fig. 7f. This observation is in agreement with semi-conducting behavior. Such behavior is also observed at all applied frequency values for the PAMs/GNP_e composite. The thermal stability observed previously in Fig. 5 and the data listed in Table 2 and the enhanced electrical conductivity shown in Fig. 7 indicate a mutual interaction between PAMs and the immersed GNPs. The close proximity of GNPs as an electron acceptor with PAMs as an electron donor in the nanocomposites would lead to charge transfer in the nanocomposites, resulting in enhanced electrical conductivity. Similar results have been reported for other graphene-based nanocomposites.³⁵

3.4. Dielectric constant and dielectric loss

Fig. 8 and 9 show the change in the dielectric constant (ϵ') and dielectric loss (ϵ'') versus the temperature of PAMs and the PAMs/GNP_{a-e} composites at constant frequencies. The values of ϵ' and ϵ'' inform us about the ability of the material to store and dissipate charge, respectively, when exposed to an electric field. Two currents are induced in the material after being affected by an electric field: a conduction current, *i.e.* the presence of free electrons and a displacement current, *i.e.* the presence of bound charges. The bound charges contribute towards ϵ' , whereas the free electrons contribute towards ϵ'' . It can be generally observed in Fig. 8 and 9 that the trend of ϵ' and ϵ'' are similar and their values decrease with increasing frequency at all studied temperature values. This behavior is attributed to the irregularities of the induced dipoles as a consequence of their slow response to field variations at high frequencies.

On the other hand, the values of ϵ' and ϵ'' increase with increasing temperature up to a certain temperature range (400–480 K), which is believed to be related to the T_g values of the composites, after which the values ϵ' and ϵ'' start to decline upon further heating. This result reflects the fact that the molecular mobility of polymeric material at T_g or above, which at its maximum mobility, facilitates dipole orientation and thus increases the dielectric properties. Further heating above the observed T_g range results in lower values for ϵ' and ϵ'' , indicating

that higher temperatures cause excessive oscillation, leading to a disordering of the molecular dipoles in the material. In contrast to the other samples, the dielectric loss observed in Fig. 9f increases with increasing temperature, indicating the effectiveness of the GNP content in this sample. This is because the GNP content increases the conductivity by virtue of the induced conduction current, which gives rise to a higher dielectric loss upon increasing the temperature at lower frequencies.^{36–39}

4. Conclusions

A new group of PAMs/GNP_{a-e} nanocomposites was fabricated using an *in situ* polymerization technique. Monomer (2) was polymerized with terephthalaldehyde in the presence of dispersed GNPs at different loadings under a nitrogen atmosphere. Prior to the polymerization, the corresponding pre-monomer (1) monomer (2) and new PAMs were synthesized in moderate yields. The structures of this new polymer and the new products were identified and characterized using FT-IR spectroscopy, X-ray diffraction, thermal analyses, SEM microscopy and electrical conductivity characterization methods. Furthermore, a comparative study was conducted to show the impact of GNP inclusion inside the PAMs polymer matrix. The XRD and FT-IR measurements of our new PAMs/GNP_{a-e} nanocomposites show intense absorption peaks related to GNP and pure PAMs, offering good evidence for the formation of our desired products. The pure PAMs and PAMs/GNP_{a-e} nanocomposites demonstrate higher thermal stability, whereas their initial decompositions start at over 300 °C. The CDT_{max} values for all the products are similar and were detected in the range of 538–555 °C. The CDT_{max} values for PAMs/GNP_{d,e} (555 °C) and PAMs/GNP_a (538 °C) represent the highest and lowest values respectively. The pure PAMs surface exhibits a coral reef structure with micro-holes that have been blocked by GNPs during their immersion into the PAMs polymer matrix. This can be observed in the SEM images of PAMs/GNP_b and PAMs/GNP_c. The PAMs/GNP_e nanocomposite exhibits the best conductivity value (10^{-7} $\text{ohm}^{-1} \text{cm}^{-1}$), which lies in the same range as those of semiconductors.

Acknowledgements

This project was funded by the Saudi Basic Industries Corporation (SABIC), and the Deanship of Scientific Research (DSR) at King Abdulaziz University, Jeddah, under grant no S/15/325/34. The authors, therefore, acknowledge with thanks SABIC and DSR for technical and financial support.

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