Polymer based graphene/zinc oxide nano crystal (GZnNC): an outstanding thermoelectrical energy conversion material

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Abstract

This work presents the synthesis of a new material, graphene/zinc oxide nano composite (GZnNC) by employing ultrasonication techniques where nano-ZnO and graphene nano-sheet have been dispersed in ethanol followed by microwave irradiation. The GZnNC was well characterized by XRD, HRTEM, FTIR, and Raman spectroscopy. Also, polymer based GZnNC has been subjected to the measurement of energy harvesting/thermoelectric properties. Present study includes PVAc, PVAc/PEDOT: PSS, and PEDOT: PSS based compositions with concentration variation of GZnNC/graphene and measurement of thermoelectric properties like electrical conductivity, Seebeck coefficient, power factor (PF), thermal conductivity and figure of merit (ZT). PEDOT: PSS/GZnNC composite showed the twelve-fold increase in electrical conductivity and two times increase in Seebeck coefficient as compared to the PVAc-graphene composite. Interestingly, the calculated power factor for PEDOT: PSS/GZnNC composite increases up to 50 times as compared to PVAc/graphene composite. Thermal conductivity gets reduced to 3.01 W/mK. Hence, figure of merit is reached up to 0.0051. This value is comparatively very high compare to the existing nanocomposites.

Introduction

Thermoelectricity is an impending area which is related to smart materials transforming electricity into heat and vice versa. In recent times, thermoelectric materials have expanded incredible research wares as green and clean energy birthplace that assistances in harvesting of energy from surplus heat. The competence of thermoelectric materials is articulated as $ZT = \frac{S^2 \sigma T}{\kappa}$, where $'S'$ represented by Seebeck coefficient i.e. thermopower, $\sigma$ is expressed by electrical conductivity, $T$ symbolize thermal conductivity and $'T'$ stands for absolute temperature [1]. Currently, few inorganic materials like silver-doped Cu$_2$Se and Cu$_2$Te, SrTiO$_3$ by Pr doping, copper telluride magnesium silicide, Bi$_2$Te$_3$ nanowires, n-type SiGe alloys, Cu$_3$S and β-Zn$_5$Sb$_3$ [2-10] exhibit enriched thermoelectric enactment. Out of them, Bi$_2$Te$_3$ is originated as the superlative ambient thermoelectric materials [11-14]. Organic compounds like polymers having conjugation are flattering more prospective [15-18] material owing to their enhance thermoelectric power factor and figure of merit. Besides their low-slung thermal conductivity (0.1–0.5 W/mK), [19-20] tranquil processibility, flexibility, nontoxicity, strength and inexpensiveness as compared to chalcogenides sort them more advantageous. Numerous other recompenses like thermoelectric paint ended armor vehicle for furtiveness, mechanical robustness for better permanence and cloths based on flexible fabricated thermo electric materials for energy mowing from human body warmth. Such submissions are very challenging to realize in the event of chalcogenide based thermoelectric. In polymer based composite, greater power factor, i.e., S'0σ, can be realized by couple of mechanisms: 1. Blending the polymer [21-23] with diverse conducting nano fillers [24-26] similar to CNT [27-28] and graphenes [29-32]. Since the award of Noble prize on the topic of graphene, numerous endeavors have been made by diverse investigators in the direction of jeer out its expedient properties [33]. Graphene has a panorama to attract devotion due to its exclusive possessions like ambipolar electric field effect, ambient condition quantum effect, and extraordinary carrier mobility [2]. Extreme reduction of such distinguishing properties of graphene is possible with escalation in numeral of graphene layers till it reaches to three-dimensional arrangement (graphite, a bulk form of graphene, with multilayer structure) [35,36]. Ten or less than ten of such graphitic layers reveal distinctive possessions as compared to the bulk arrangement [37]. Such extraordinary properties sort graphene as one of the best engaging material for its submissions in innumerable fields such as composite, electronic and advance machine-driven resonator [38,39]. Graphene has an elite capability of snowballing the thermoelectric effect by employing the appropriate metal oxide/ metal nanoparticles on its surface. Consequently, the number of operative site on surface of graphene upsurges. In the field of nanotechnology, it has been used as a substrate. Investigations on the thermoelectric possessions of such composites are instituted them to be viable on chalcogenides. However, their adeptness is quiet subordinate [21,24,25]. Out of many polymers, (3,4-ethylenedioxythiophene): poly (styrene sulfonate) (PEDOT: PSS), [16,40] poly (3-hexylthiophene) (P3HT), [41] and polyaniline (PANI) [31,42-44] are frequently used owing to their inherent great electrical possessions like ambipolar electric field effect, ambient condition quantum effect, and extraordinary carrier mobility [2].

Key words: graphene, zinc oxide, energy harvester, thermoelectric, power factor, figure of merit

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Among various notorious stable iron oxide stoichiometries, Fe₂O₃ is oxide and doped zirconium oxide exhibited remarkable TE properties. viz. used as a substrate. Graphene in conjunction with few metal oxides packing and competence of thermoelectric material. lesser thermal conductivity which assistances to upsurge the filler packing and competence of thermoelectric material.

To upsurge the electrical conductivity radically, graphene has been used as a substrate. Graphene in conjunction with few metal oxides viz. iron oxide, [48] nickel oxide, titanium dioxide, [49-52] cadmium oxide and doped zirconium oxide exhibited remarkable TE properties. Among various notorious stable iron oxide stoichiometries, Fe₂O₃ is an oxide that has been investigated for the situation of TE property. Few reports advocate that Fe₂O₃ can be an auspicious transition metal oxide for TE application at higher temperatures as it exhibits high TPFs at elevated temperatures and ambient. Iron oxide thin films have been revealed to exhibit highest Seebeck coefficient of 1650 µV/K in the temperature expance of 270-290 K. Electrical conductivity of 3.5 × 10⁵ S/m has been described in the identical thermal conditions, occasioning to a great TPF [52] of 1.5 × 10⁴ µW/m.K². It has been mentioned that nano ZnO [50,51] can perform as efficient of thermoelectric material. In GZnNC, nano ZnO was adorned over the graphene sheet throughout its exfoliation. The staking environment of graphene sheet abridged extremely after nano ZnO adornment which reduces the tendency of graphene to graphite transformation process. Present work demonstrates the modest preparation process of GZnNC with adequate characterizations and its submission as thermoelectric material for waste energy harvesting. The composition, morphology, and structure of the as-prepared GZnNC have been characterized by Raman spectroscopy, Fourier Transform Infrared Spectroscopy (FTIR), Field Emission Scanning Electron Microscope (FESEM), High-Resolution Transmission Electron Microscopy (HRTEM) and X-ray diffraction (XRD). Further, thorough study of thermoelectric properties (electrical properties, Seebeck coefficient, power factor, thermal conductivity and figure of merit) of PVAc, PVAc/PEDOT: PSS and PEDOT: PSS with different fillers like GZnNC, graphene have been evaluated and presented extensively.

**Experimental details**

**General**

We have purchased graphitic flakes from Reinste nano ventures, ethanol (absolute) from AR, Merck and graphite powder, H₂SO₄, NaNO₃, NaOH, H₂O₂ (30%), 0.5M zinc nitrate [Zn(NO₃)₂], 6H₂O, 1M NaOH, polyvinylpyrrolidone from Sigma-Aldrich. The chemicals received were of analytical grades and used as such without further purifications and characterization. Milli-Q water (18 MQ cm⁻¹) has been used to make aqueous solutions. Sonication and microwave irradiation were conducted by using ultrasonication bath (35 kHz, Kudos) and microwave reactor (make: Raga), respectively. FESEM and HRTEM were carried out with Quanta 200, FEI and TACHNIAI F-30, FEI with 300 kV field emission gun (FEG), respectively were dispersed in methanol with ultrasonication. The dispersions of nano ZnO, graphene and GZnNC were placed on TEM grid, and the solvent was evaporated to dry. FTIR and Raman’s traces were collected by Nicolet 5700, Thermoscientific and Invia reflex micro-Raman, Renishaw, respectively. X-Ray diffraction analysis was conducted by X-Ray Diffractometer D8 Advance, Bruker in conjunction with Cu Kα source at an angle (20) expansion of 2-90° through a scan rate of 2°/min.

**Synthesis of ZnO nano-particles**

ZnO nanoparticles were synthesized by the previously reported method but with slight modification [53]. The synthesis was performed by preparing the alkali solution of zinc nitrate by dissolving 0.5M zinc nitrate [Zn(NO₃)₂, 6H₂O] and 1M NaOH, in distilled water (100 ml). The solution was continuously stirred at 75°C maintaining the pH 12. The reaction mixture was stirred for 5-6 hrs at 75°C after addition of polyvinylpyrrolidone (PVP, 0.2 gm) to obtain the white suspension of ZnO nanoparticles. The suspension was separated by centrifugation followed by washing with distilled water and ethanol. The white precipitate obtained was annealed at 400°C for two hours in a muffle furnace for obtaining highly nano-crystalline ZnO nanoparticles.

**Conversion of graphene from graphite flake**

The synthesis of graphene involves three stages [48,54]. In an initial step, graphene oxide was prepared from the flakes of graphite by Hummers method. Thermally expanded graphene oxide (TEGO) was produced in the second stage by thermal enlargement at 1050°C (Ar, the 30s). In the third step, graphene nanosheet was achieved by hydrogen reduction of TEGO for two hours at 400°C.

**Preparation of Graphene/ZnO nano-composite (GZnNC)**

GZnNC was obtained by scattering 50 mg graphene through ultrasonication in ethanol after addition of pre-dispersed ZnO nanoparticles to the dispersion of graphene. The resultant dispersed combination was subjected to ultrasonication for two hours tailed by drying at room temperature to remove the solvent. GZnNC was placed over a petri dish and exposed to microwave irradiation for two minutes to have better exfoliation after drying (Scheme 1).

**Thermoelectric properties**

Seebeck coefficient or thermopower, electrical conductivity, thermal conductivity are the three main thermoelectric properties and useful to calculate the power factor (PF) and figure of merit (ZT). Sample preparation and its characterization were carried out by the following a methodology. Nanocomposites accompanied by polymer were made through ultrasonic dispersion of fillers (graphene and GZnNC) and PVAc water emulsion for 30 min at 35 kHz. After ultrasonication, mechanical stirring sustained for four hours at 235 rpm to get proper dispersion. Complete drying was performed by placing the sample in a vacuum oven at 105°C for 4 hrs after casting in an aluminum tray. The dried samples were exposed to hot pressing at 120°C in conjunction with 5 min preheating time and 3 min compression (10 T of pressure approx.). The synthesized polymer nanocomposite sheets were endangered to the determination of thermoelectric possessions. Polymer nanocomposite based on PVAc/PEDOT: PSS and PEDOT: PSS were also processed and subjected to the test (Scheme 2).

**Results and discussion**

**Characterization of GZnNC**

**XRD and FTIR analysis:** The presence of ZnO nanoparticles and graphene in the GZnNC was confirmed by X-Ray diffraction scan of the samples (Figure 1). The XRD spectrum of ZnO nanoparticles obtained by sol-gel process displayed somewhat wide peaks owing to its nano-sized nature. The 2 peaks present at 25.25° (101), 37.8° (004),
47.9° (200), 53.59° (105) and 62.36° (204) confirmed the wurtzite crystal structure of ZnO. The XRD spectra of GZnNC sample showed a broad peak at 2θ =25.8° indicated a haphazard packing of graphene sheets and related to (002) plane of graphite. The FT-IR data of the GZnNC (Figure 2a) did not disclose any evidence concerning the existence of organics in the product and the corresponding peaks for graphene oxide were preoccupied. However, peaks were present at 3400 cm⁻¹ corresponding to –OH group due to moisture and a peak at 1675 cm⁻¹ due to skeletal vibration in sheets of graphene. The sharp peak at 2355 cm⁻¹ could be due to the presence of adsorbed air-borne CO₂ molecules. The GZnNC sample did show a peak at around 1380 cm⁻¹ which could be due to the carbon moiety carried by the ZnO nanoparticles. This carbon moiety can be formed after the sintering process of ZnO nanoparticles which resulted in decomposition of PVP capping, producing carbon residue in the sample.

Raman analysis: Figure 2 (b), (c) and (d) represents raman touches of GZnNC. The Raman spectrum of pristine graphene showed the characteristic in-plane vibration peak (G band at 1580 cm⁻¹) which originates from sp² carbon atoms along with primary and second order overtone of different in-plane vibration peaks D band (1350 cm⁻¹) & 2D (2690 cm⁻¹). It was observed that the pristine graphene was composed of multi stacked layers as the intensity of 2D band peak was less than the G band peak. The calculated ratio for \( \frac{I_{2D}}{I_G} \) (~0.9) for GZnNC sample was more as compared to the pristine graphene sample (~0.4). Thus, the stacking of graphene sheets decreased in the nanocomposite sample which could be a result of the presence of ZnO nanoparticles between the graphene sheets. Also, the ratio of \( \frac{I_G}{I_D} \) was 2.6 for pristine graphene and more than 5 for GZnNC sample. The defect intensity increased in the GZnNC sample as compared to pristine...
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Figure 2. a) FT-IR spectrum of GZnNC (b) Raman traces of GZnNC sample (c) and (d) represent comparative bar diagram of $I_{2D}/I_{G}$ and $I_{G}/I_{D}$ with GZnNC and graphene, respectively.

| FESEM and HRTEM analysis: Field emission scanning electron microscopy (FESEM) images of samples at different magnification (Figure 3a and 3b) highlighted the presence of exfoliated graphene sheets. The FESEM images of GZnNC samples (Figure 3c) indicated the uniform presence of ZnO nanoparticles over and around layers of graphene. It has been clearly observed that ZnO nanoparticles are existent in combined form in the GZnNC. The exfoliated nature of graphene layers in conjunction with nanosized distorted shaped ZnO will ensure enormous surface availability of the GZnNC. The great surface area of GZnNC can be vivacious for its desired thermoelectric nature. HRTEM of the GZnNC studies were conducted to identify the shape and size of ZnO nanoparticles in GZnNC (Figure 3d and 3e). The distorted shaped ZnO nanoparticles were confirmed by HRTEM images as various pebble shaped ZnO nanoparticles with a particle size in the expanse of 20-100 nm were formed after sintering. The exfoliation of graphene was also evident for the printing graphene samples (Figure 3d) which helped in providing a platform for a large number of ZnO nanoparticles to assemble on its surface (Figure 3e). It was interesting to observe that distorted rod-like, hammer-like ZnO nanoparticles were also present on the graphene sheets in GZnNC sample (Figure 3d and 3e). The presence of such shaped ZnO nanoparticles would facilitate a significant improvement in surface area in GZnNC sample which as mentioned above can play a vital role in thermoelectric applications. The clear visibility of lattice fringes with spacings $d=0.281$ nm (corresponding to 100 crystal plane) not only confirmed the presence of hexagonal ZnO but revealed its nanocrystalline nature.

Characterization of thermoelectric properties of polymer-GZnNC composite

Seebeck Coefficient/Thermoelectric Power Measurements: To measure the thermopower using temperature, samples [54] have been prepared with a dimension of 30 mm(l) × 6 mm(w) × 1 mm(t) from polymer nanocomposite film and positioned on a thermally insulated fiberglass. At one termination of the sample along with a thermally conductive epoxy (electrical insulating 2763 Stycast), whereas the other end, a copper piece (sink of heat) made a connection with the Peltier cooling site. The temperature gradient and voltage fall through the film was dignified using thermocouples organized in series using a couple of copper wires. To make sure that the thermal gradient and the voltage fall were being dignified at the same location, two minor copper films were devoted to the polymer-GZnNC film with electrically/thermally conducting silver epoxy. The voltage wires and the thermocouple were devoted to these copper films. The thermoelectric energies were scrutinized on temperature alteration by Keithley 2182A. Peltier cooling module has been employed to change the base temperature. The thermoelectric power was dogged by two self-determining means:
1. After accomplishment of a steady state through a smeared current to
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Figure 3. FE-SEM images of graphene sheets a) with magnification of 20 KX; b) with magnification of 50 KX; c) FESEM images of GZnNC nanocomposite with magnification of 60 KX. HR-TEM images of d) graphene-zinc oxide nano-composite (GZnNC) with a scale bar of 50 nm e) rod-like distorted ZnO nano particles present in the GZnNC nanocomposite (scale bar is 20 nm).

Scheme 2. Synthesis and characterization procedure of polymer based GZnNC nanocomposite.
the heater. 2. By fitting the linear correlation for V vs. ΔT response to a heating pulse. The deviance among both methods and between diverse experimentation was always smaller than 5%.

**Electrical Resistivity Measurements:** Owing to the high electrical conductive nature of the composite, delta mode four probe method was used to measure electrical resistivity. The lowest conceivable current was obtained (100mA) by Keithley 6220 and the corresponding voltage was scrutinized with nanovoltmeter of Keithley 2182A. The minutest possible current was used to evade heating of the sample at small temperatures. Polymer nanocomposite sample using a dimension 8 mm × 3 mm × 1 mm has been prepared and subjected to measure electrical conductivity.

**Thermoelectric properties measurement set up:** Figure 4. represent thermoelectric measurement set up. Figure 4a, 4b and 4c depicted the XYZ movement set up. Figure 4d correspond to actual homemade set up i.e., Keithley 2182A and Keithley 6220 for electrical conductivity and Seebeck coefficient measurement. Figure 4e and 4f represent Seebeck coefficient and electrical conductivity measurement set up respectively.

**Thermal diffusion measurement:** Thermal diffusivity is another important property, required for such purposes as design applications under transient heat flow conditions, determination of safe operating temperature, process control, and quality assurance. The flash method is used to measure values of thermal diffusivity, α, of a wide range of
solid materials. It is particularly advantageous because of simple Figure 4: (a), (b) and (c) represent the schematic diagram of homemade set up. (d) represent actual figure. (e) Schematic representation of Seebeck coefficient geometry, small specimen size requirements, rapidity of measurement and ease of handling, with a single apparatus, of materials having a wide range of thermal diffusivity values over a large temperature range. Under certain strict conditions, specific heat capacity of a homogeneous isotropic opaque solid sample can be determined when the method is used in a quantitative fashion (see later section). Thermal diffusivity results, together with specific heat capacity ($C_p$) and density ($\rho$) values, can be used in many cases to derive thermal conductivity ($\lambda$), according to the relationship:

$$\lambda = \alpha C_p \rho$$

Power factor (PF) and figure of merit (ZT): Power factor (PF) and figure of merit (ZT) were calculated by following the given equation. PF=S$\sigma$, ZT= $S^2\sigma T/\kappa$, where $S$ represent Seebeck coefficient or thermopower, $\sigma$ = electrical conductivity, $T$= absolute temperature, and $\kappa$= thermal conductivity.

In this contrary, to find out the effect of conducting fillers on polymer matrix, seven compositions have been made. The comparative bar diagram of Seebeck coefficient, electrical conductivity and power factor at different PVAc/GZnNC composition (C1 to C7) have been highlighted in S1. According to the graph, electrical conductivity and power factor increases abruptly, but an insignificant upsurge in Seebeck coefficient was perceived. From composition C1 to C7, GZnNC concentration increases systematically to find out the effect of filler. Composition C7 shows maximum PF value. When graphene was used as filler in place of GZnNC. Due to conducting nature of graphene, electrical conductivity increases but a marginal change was observed in Seebeck coefficient. Hence, calculated PF increases with graphene concentration (S2). But the overall PF value is relatively low. Figure 5 depicts the variation of Seebeck coefficient, electrical conductivity, thermal conductivity, power factor and figure of merit in PEDOT: PSS/GZnNC composites. To identify the optimum composition, several compositions have been made and evaluated. The comparative chart indicates C2 composition is the optimized composition of this system which experiences maximum Seebeck coefficient, electrical conductivity and minimizes thermal conductivity. Hence, power factor and figure of merit reach to the maximum value.

When graphene is used in place of GZnNC, electrical conductivity, increases to a certain level then decreases. Seebeck coefficient decreases with graphene concentration but PF follows slightly increasing
trend than decreasing trend (Figure S4). When compositions have been made in combination of PVAc/PEDOT:PSS, all the three properties were found to be inferior in compare to PEDOT:PSS based composition (Figure S5). In the course of studying the thermoelectric properties, PEDOT:PSS/GZnNC composites displayed 12 fold increase in electrical conductivity and two fold of Seebeck coefficient was perceived as compared to composite of PVAc-graphene. The calculated power factor for PEDOT:PSS/GZnNC composite increased to 50 folds than composite of PVAc-graphene. Thermal conductivity is found to be 3.01 W/mK. In GZnNC, nano zinc oxide was decorated over 2D graphene sheet in GZnNC, attendance of nano zinc oxide particle assistances to abolish thermally conductive network but intact electrical network. When GZnNC is employed as conducting filler, it not only decouples σ and S, but also enhances both the parameter simultaneously which is very significant finding. Though the Seebeck coefficient enhancement is minimally about electrical conductivity for PEDOT:PSS/GZnNC composite.

Conclusion

This work presents a cheap and ecofriendly method for the synthesis of Graphene-Zinc oxide nanocomposite (GZnNC) by dispersing nano zinc oxide (ZnO) and graphene nano-sheets (GNS) via ultrasonication in ethanol took by irradiation with microwaves. GZnNC nanocomposite performed as an effective energy harvesting/ thermoelectric material. Throughout the study of thermoelectric possessions, PEDOT: PSS/GZnNC composite displayed 12 fold increase in electrical conductivity and two fold increase in Seebeck coefficient as compared to PVAc-graphene composite. The resultant power factor for PEDOT: PSS/GZnNC composite increases up to 50 folds than PVAc-graphene composite. Thermal conductivity is found to be 3.01 W/mK with the ZT of 0.005. In GZnNC, nano zinc oxide was decorated over 2D graphene sheet. GZnNC not only decouples the σ and κ, but enhance both η and S. Hence PF and ZT reaches to a maximum value.

Supplementary information

[Tabulated data of thermoelectric properties from different polymer composite, Comparative bar diagrams, Summary of thermoelectric properties of the best composite of inorganic and organic materials have included here]. See DOI: 10.1039/x0xx00000x

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