Mechanical Behavior of Nano-Scaled Graphene Oxide Reinforced High-Density Polymer Ethylene for Orthopedic Implants

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Abstract: The application of bio-composite materials is in trends recently, for both commercials as well as the scientific community. In order to achieve inherent strength and efficient performance, biocompatible materials are often modified. This is done with an aim to replace organic cells/tissues if required. In recent times, the polymer composites are used as an alternative for body implants than traditional metal-based components. Cost inefficiency and corrosion resistivity are the major problems associated with metallic implants. Polymer composite offers similar advantages as that of metallic based components. Moreover, it can also be molded in the requisite shape, are cost-efficient, have a high corrosion rate, and less weight. This paper aims to fabricate a biopolymer nanocomposite with efficient fatigue, mechanical, and wear properties, which is suitable for human implant applications. The studies evident effective mechanical and efficient tribological properties, demonstrated by implants for knee and hip replacement. The graphene oxide employed in the experiment study retains antibiotic properties and is also tends to be bio-compatible. Nano graphene oxide, along with high-density polyethylene at varying wt.% of 0.5-2.5%, was reinforced, which resulted in better tribological outcomes. This composite material is environment friendly and has the potential to be used for numerous applications.

Keywords: nano-graphene oxide; high-density polymer ethylene; mechanical behavior; fatigue testing; wear testing.

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1. Introduction

The HDPE (high-density polyethylene) has a wide array of applications in various fields, including biomedical, daily household items, and defense as these fields require high-performance materials, and the HDPE tends to be cost-effective, corrosion resistance, durable and tough [1-4]. Apart from the benefits, certain disadvantages like its abrasion resistance and creep decrease the efficiency of polymer material [5-7]. Amidst this rundown, Ramanathan et al., in their paper, proposed the use of nanofillers in order to improve the mechanical properties for polymer-based materials [8]. Several researchers have concluded that the addition of nanoparticles as reinforcements can enhance the various properties of materials, which they initially lack [9-13]. There has also been a tremendous amount of increment in the use of nanoparticles for numerous biomedical applications in recent years [14-21].

Despite this, the ideal material for using as fillers in polymers has always been a matter of debate in the research world. Owing to its tensile strength, CNT (carbon nanotube) as
reinforcement has been considered an ideal candidate by researchers; however, due to numerous aspects, CNT’s are still in the development phase [22-24]. The complex structure of CNT’s obstructs the matrix to reinforcement load transfer, as it fails to hold on to the surface of the polymer matrix [25]. Other than this, the dispersion of CNT in a viscous solution is also a significant challenge, limiting the scope of polymer applications. The fabrication of CNT’s resolves this issue. However, at the same time, it also disturbs their confinement structure, which leads to a declining trend in the tensile strength of the polymer [26-28]. Amidst this scenario, the functionalized graphene as an alternative replacement material is adequate [29]. The graphene retains similar properties as of CNT’s [30]. By its flat structure, at matrix, the strength transformation by diverse direction is viable and it also successfully reduces the matrix slippage trouble. This is an advantage, as the tensile strength of CNT is robust along the length only [31-32]. Several research studies reported an increase in mechanical properties of graphene-based polymers, integrated with nanocomposites [33-37].

Yoon et al., in their article, discuss the improvement of mechanical & thermo-mechanical properties of a PLGA nanofiber mesh, reinforced with graphene oxide. Only 1-2 wt.% of graphene oxide was integrated into the polymer [38]. A different study demonstrated enhanced flexural strength of the HDPE based nanocomposites, upon the integration of paraffin wax along with graphene; in comparison to the uncoated polymer surface [39, 40]. Interaction between graphene and polymer matrix, along with adequate dispersion and orientation of the graphene with polymer matrix, is still an obstacle, as it hinders the load transformation from the matrix to reinforcement. Due to multilayer graphene slippage and inadequate interphase between graphene and polymer matrix, understanding of strength for graphene-based applications is still not adequately understood [41-43].

This paper focuses upon the determination of optimum composition and methodology for the fabrication of a biopolymer nanocomposite along with efficient fatigue, mechanical & wear properties. The fracture trend of the composite is also discussed. The focal aim of the entire study was to fabricate a clinically relevant polymer material, which can be used for the fabrication of orthopedic implants.

2. Materials and Methods

2.1. HDPE composites processing.

Specimens were fabricated at a temperature range from 120°C - 160°C, the polyethylene high-density composite samples were obtained. Unlike nano polymers; the combination of resin reinforcement for HDPE is not obtained directly. After the complete homogenous blend of the solvent, the nanopowder from the solvent matrix tends to be redistributed in the polymer matrix, followed by the evaporation; this is achieved by distributing the nanoparticles of optimum dimensions in the solvent through appropriate technique. The HDPE pellets used for this study were in the form of pellets, and graphene oxide nanoparticles obtained for the study were in the form of long flakes. Both these materials were obtained from Sigma Aldrich, Hyderabad, India. The properties of both these materials HDPE and nano-graphene oxide are shown in Table 1 below.
### Table 1. Properties of high-density polyethylene and nano-graphene oxide (as said by the supplier)

| Material                      | Yield Strength (MPa) | Ultimate Tensile Strength (MPa) | Elongation (%) | Young’s Modulus |
|-------------------------------|----------------------|---------------------------------|----------------|-----------------|
| High-density polyethylene     | 7.005                | 34.74                           | 48             | 547.48          |
| Nano graphene oxide           | 34.378               | 130                             | 0.6            | 2400            |

2.2. Ultrasonic vibrations.

The inspiration for using ultrasonic vibrations was obtained through studying various experimentations performed by other researchers to increase the particle size, surface area, and proper mixture of the nanoparticles in the composites [44-48]. The steps refereed are in respect to the specimen containing 0.5% nano composition, along with 50ml xylene as a solvent and 0.5 g of nano-graphene oxide particles as solute. The specimen was precisely weighed via a weighing balance, followed by the placement of the specimen in an ultrasonic vibrator, partially submerged into water. Relatively, 20000 Hz of ultrasonic vibration frequency was produced for a time of 30 minutes, in order to obtain nano dimensions of the graphene oxide via vigorous agitation. Ideally, nanoparticles were supposed to obtain the dimensions of around 60-70(nm) x 60-70(nm) x 4-5(nm), given they are precisely distributed. The electron microscopy scanning technique was used for determining the definite dimensions of the nanoparticles. The obtained nano-solvent mixture would be used as the reinforcement component.

2.3. Melt blending.

Many researchers in recent years have used the melt blending process for an efficient blending of different materials and enhancement in properties of the materials [49-54]. For the melt blending process, 100g of HDPE was placed in a stainless steel container followed by a heat treatment via a hot plate eater. Promptly upon the indication of heat input being turned on, xylene was added. The maintained matrix-polymer volume ratio was 5:1. Thus 500 ml of xylene w.r.t pellets were added in the stainless-steel container. The solvent, xylene, tends to dissolve the HDPE pellets rapidly due to the heat treatment provided earlier. At 140°C, the HDPE pellets started melting. Minutes before this point, the nano powder-xylene mixture was integrated. The vaporizing temperature for xylene and the melting point of HDPE pellets are similar. As the melting process of pellets started, the compatibility of nano-graphene oxide for polymer increased, and the vaporization phase of xylene provided ideal transformation rate and distribution of nanoparticles upon the polymer matrix. During this process, a temperature range of 150-180°C was maintained, as the vaporization phase for xylene occurs at this range only. Within a few minutes HDPE, along with ideally distributed nano-graphene oxide, was obtained, as the xylene used was effectively vaporized in the atmosphere.

2.4. Vacuum molding.

The composite is obtained in a semi-fluid state; initially, the composite was manually moved into a compression mold plate, followed by the squeezing process in order to obtain plate specimens. A field of blowholes around the center was detected for the obtained specimens. This obstruction was partially reduced by the preheating treatment. In order to completely eradicate this colony of blowholes, the quenching method was employed, for which
the mold plates were tightly confiscated and immersed in water for several minutes and were then cooled for the next 24 hours. These treatments had a considerable effect on the blowhole area. However, it was observed that in the presence of atmospheric air, the blowhole tends to form. In order to eliminate this phenomenon, a vacuum treatment was contemplated. For this, an apparatus consisting of a vacuum pump, along with a vacuum box, was built. The composite at a vacuum pressure of 0.8 bar was transported from the stainless-steel container to the molding plates, which were tightly confiscated, before the vacuum sustentation for 5 minutes, followed by the cure treatment of the same time frame. After this procedure, the specimens were further quenched, which resulted in a defect less specimen along with a smooth surface finish.

The technique mentioned was employed for the manufacturing of other samples having 0.5%, 1%, 1.5%, 2%, & 2.5% wt.% of nano-graphene oxide.

3. Results and Discussion

To verify the reliability of the proposed composite polymer, certain characteristics such as bonding stability, fatigue, strength, and wear resistance were evaluated. This inspection allowed the assessment of HDPE for clinically relevant applications. In order to detect the distribution of nanoparticles, micro-structural bonding, reinforcement dimensions, and its stability inside the matrix, scanning electron microscopy was employed. In order to determine the composite failure, several loads were periodically engaged, through which the results for fatigue tests were obtained. For a composite polymer to be clinically relevant, its wear properties are crucial. Fixed travel distances, load, and speed parameters were employed for testing the effect of incrementing nanoparticles composition on wear rate.

3.1. Scanning Electron Microscopy.

The SEM (scanning electron microscope) employs a concentrated electron beam to scan and obtain images of the sample provided, with a resolution greater than 1 nm. These electrons interact with specimen atoms, which indicates the surface composition and its topography through information obtained via various signals. In order to obtain an image, the raster scan pattern is adopted via an electron beam. The position of an electron beam, along with a detected signal is used for an image generation. This technique is able to analyze the specimen in harsh environmental conditions, high-low vacuum, and even at cryogenic temperatures. SEM efficiently detects the secondary electrons emitted by the specimen’s atoms because of the electron beam. However, the electron detection is influenced by the specimen’s topography. By virtue of sample scanning, coupled with the detection of secondary electrons, an image of the specimen’s surface topography is created. The scanning electron microscope (JEOL, Model JSM 6510LV, Japan) was used for the above experimentation is shown in Figure 1 below. Figure 2 depicts the HDPE microstructure distribution, along with nanopowder graphene oxide. The uniform dispersion of nanopowder and the HDPE is way significant, as it influences the fatigue strength of the composite material. In order to verify this, Figure. 2 is zoomed up to 3 levels. It is evident from the image that material is evenly distributed, with no sign of agglomerates, indicating the apprehension of desired properties and enhanced fatigue life.
3.2. Fatigue test.

For any given material, its fatigue life is very crucial as it influences the longevity and reliability of various application components. Fatigue failure of the implant is one of the significant obstructions regarding biomaterials, and the integration of nano reinforcements improves material efficiency, as evidenced in many studies. As depicted in Figure 3, a multiaxial fatigue testing machine (Instron, Model 8801, United States) was used for testing the respective composite specimens, with an aim to analyze the impact of nano composition integration on fatigue properties of a material. Various propositions of nano-graphene oxide were processed for 5 different specimens, for load tensile strength of 60%, 70%, and 80%, with uniaxial loading along their length. Figure 4 depicts and compares the S-N curve of each specimen. It is clear from the graph that an increment in the nano reinforcement composition increases the fatigue life of the specimen. However, the curve obtained constantly pushes towards a northeast corner, illustrating a constant increment in the fatigue life which is theoretically not correct because, after a certain period, the accumulation due to nano composition is certain, which would decrease the magnitude of load that the sample could bear before failure. A sample prepared with 0.5% of graphene oxide demonstrated internal blowholes, which increased with the given time. It is also evident from the S-N Curve (Figure 4) that there is a constant improvement in incrementing the nanoparticles from 0.5% to 1% and similarly from 1% to 1.5% & 2%. However, beyond 2% there is not a great improvement as visible on increasing from 1.5% to 2%.
3.3. Wear test.

The longevity and reliability are two crucial factors that determine the efficiency of a mechanical component. As the component ages, it becomes more prone to wear and tear, which makes the wear test of the proposed composite a requisite, as it should be relevant to the clinical standards. DUCOM pin on the disc was utilized for testing wear characteristics (DUCOM, T25, India), as shown in Figure 5. The experiment for analyzing the wear test of the composite was carried out using an abrasive grinding wheel, connected with a counterweight in order to sustain the wear upon the specimen. 100 RPM for the abrasive disc was set at the beginning of the experimental study. Five composite samples with varying composition, i.e., 0.5%, 1%, 1.5%, 2%, and 2.5% of nanoparticles, along with HDPE were selected for analyzing the wear properties, which helps to characterize wear curve because of which the ideal composition
content can be identified. At a full length of the experiment procedure, the parameters were constant, mainly, the load was fixed at 10 N, along with 100 RPM, and wear of 12 minutes was intended. The results obtained from the wear test demonstrated an increment in the wear resistance, along with an increment in the composition of nanoparticles, which can be observed in Figure 6. A sharp decline in the wear trend was observed upon incrementing the weight of reinforcement’s concentration from 1.5-2.5 wt.% shown in Figure 7. A similar result was obtained here as that of fatigue test, where we can clearly see that there is a constant improvement in increasing nanoparticles from 0.5% to 2%. However, beyond that, the result is not much significant or worthy to be considered. It is clear that the wear properties of composites could be enhanced by incrementing the composition of nanoparticles.

Figure 5. Wear Testing Machine DUCOM, T25, India.

Figure 6. Wear vs. % of nano-graphene oxide.

Figure 7. Effect of % of nano-graphene oxide on wear.
4. Conclusions

HDPE composites upholding a thickness range of 10 mm were fabricated via a vacuum compression molding technique coupled with water quenching. This method was both effective and efficient for the fabrication of biomaterial of clinical standards. Also, the ultrasonic dispersion method was employed for the distribution of nanoparticles of graphene oxide, done through a melt blending mechanism. The constant temperature throughout the experiment study cannot be maintained during the melting and mixing procedure, which resulted in a colony of blowholes around the partially concave surfaces at the center region of the specimens. Significant improvement in the fatigue & wear properties of the composite specimen were observed, along with an increment in the tensile strength as well; upon integrating nano-graphene oxide particles. For specimens having 1.5%-2% nanoparticle composition, the distribution of nano-phase reinforcement was ideal, and no accumulation was observed in its microstructure. It can be concluded that 2% wt.% of nanoparticles provides the best results in every testing conditions. The results were not much significant in the difference between the case of 2% and 2.5%. The polymer matrix and the reinforcement both tend to be biocompatible, but the tests were conducted in regard to the wear and fatigue resistance of the composite, and several other factors also contribute to the final assessment for the proposed composite material. If this composite tends to perform as efficiently upon other biological and mechanical stability parameters, it is safe to remark that it could be considered as an ideal alternative bioimplant.

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Conflicts of Interest

The authors declare no conflict of interest.

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