Organically-Expanded Graphite/Octadecylamine: Structural, Thermal and Relaxation Evaluation

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ABSTRACT

Natural graphite was chemically modified by an acid mixture (H2SO4/HNO3, 4:1) using sonication. The resulting material was then expanded with octadecylamine by the dispersion method. The sample was characterized by scanning electron microscopy (SEM), Fourier transform infrared spectroscopy (FT-IR), thermogravimetry (TG), wide-angle X-ray diffraction (WAXD) and relaxometry—low-field nuclear magnetic resonance (LFNMR). The SEM images revealed the expansion of the graphite layers. The presence of CH absorption was detected in the infrared spectrum of the expanded sample. The WAXD showed an additional diffraction peak at lower 2θ angle, indicating that intercalation of octadecylamine was successful. The thermogravimetry curve revealed three degradation steps. Two of them could be attributed to different structures (delaminated and exfoliated). The relaxometry showed that the relaxation time was dependent on the frequency and the curve of the graphite/octadecylamine presented two peaks—approximately before 10^6 Hz and after 10^7 Hz. The results permitted inferring that organically modified graphite was achieved.

Keywords: Graphite; Acid Oxidation; Octadecylamine; Sonication

1. Introduction

Due to the intrinsic properties, many synthetic and natural fillers, both modified and unmodified, are employed alone or in micro and nanocomposites. Silica is one of the most used. Hybrid materials containing different amounts of Aerosil® were studied by Tavares et al. [1] to investigate relationship among silica content, molecular structure and intermolecular interactions. Sol-gel silica was synthesized through hydrolysis and condensation of tetraethyl orthosilicate (TEOS) for use in dental composites [2]. Different biopolymer/filler systems were investigated and changes in the relaxation times were detected by proton-spin lattice relaxation [3]. Relaxometry study showed the influence of Viscogel® B8 and silica on relaxation times in poly (lactic acid) nanocomposites [4]. Lamellar α-zirconium phosphate was expanded with octadecylamine and incorporated in linear low density polyethylene (LLDPE) to produce nanocomposites [5]. Alpha-zirconium phosphate was synthesized by refluxing and added to poly (vinyl alcohol) to obtain nanocomposites [6]. Novel fire retardant material using α-zirconium phosphate was prepared [7]. An organic-inorganic hybrid layered zirconium phenylphosphate was prepared and used in a nanocomposite based on poly (ethylene terephthalate) [8].

Graphite is an abundant natural material. Its good electrical conductivity, thermal, chemical and mechanical properties attract research interest in it [9]. Hwang and collaborators [10] prepared supports for PtRu catalysts in direct methanol fuel cells using modified graphite carbon nanofibers. Electrocatalysts were produced by carbon nanotubes and graphite powder [11]. Electrode material for a vanadium redox flow battery was prepared by graphite felt with nanodispersed bismuth [12]. Selective determination of dopamine was performed through functionalized-graphene modified graphite [13]. A graphite-epoxy composite was prepared for electrochemical immunosensing [14]. Expanded graphite was applied in the preparation of a nanocomposite based on styrene-butyadiene rubber [15]. A graphite electrode modified by 8-hydroxiquinolines was used for determination of copper at trace levels [16]. An electrodeposited Pb-modified graphite strip electrode was developed as a trichloro-
2. Experimental

2.1. Materials

Graphite powder was supplied by National de Grafite Ltda. The oxidizing agent—sulfuric acid (98%) and nitric acid (65%)—were supplied by Vetec. Octadecylamine was supplied by Spectrum.

2.2. Graphite Treatment

Natural graphite (NG) was modified in two steps. The first step was based on literature reports [18-20]. To remove the moisture before treatment, the graphite powder was dried at 80°C for 12 hours. A sulphonitric solution (sulfuric and nitric acid mixture, 4:1 v/v) was added to the graphite powder. The suspension was sonicated and stirred at room temperature for 5 hours. In this process, the graphite acid mixture (AM-G) was obtained. After that, the suspended powder was filtered and washed with deionized water until pH 5.5 and dried at 80°C for 12 hours.

The second step consisted of octadecylamine (ODA) functionalization. AM-G was added in a 70/30 v/v alcohol solution and the suspension was maintained without stirring for 24 hours. After that, the suspension was stirring for 24 hours and then filtered and washed with ethanol for removal of non-reacted octadecylamine and dried at 80°C for 12 hours to obtain ODA-AM-G.

2.3. Thermal Analysis

The thermogravimetric analysis was performed with a TGA 92 SETARAM under air atmosphere operating in the range of 30°C - 700°C at a heating rate of 10°C/min.

2.4. Morphological and Structural Evaluation

Scanning electron microscopy (SEM) was performed with a JEOL JSM 5610 LV microscope with 15 kV of voltage, with magnification up to 3000 times.

Wide-angle x-ray diffraction (WAXD) was carried out in a Brüker D8 Advance diffractometer, using CuKα radiation with 40 kV of voltage and current of 40 mA, with 2θ between 2° - 35°.

FTIR spectroscopy was performed with a Thermo Nicolet Nexus Instrument. The samples were analyzed in a medium infrared region (4000 - 600 cm⁻¹) using attenuated total reflectance (ATR) and using KBr disks, with resolution of 2 cm⁻¹ and 640 scans.

2.5. Low-Field Nuclear Magnetic Resonance Relaxometry

The measurements of proton spin lattice relaxation time (T₁H) in NG is not possible due this material have only carbon atoms in its composition. Thus, the acid mixture (AM) was analyzed as the starting point for comparison with the treated materials. So, the measurements of proton spin lattice relaxation time (T₁H) in AM, AM-G and ODA-AM-G were carried in the frequency range between 50 kHz and 300 MHz using different techniques. The measurements of T₁ above 9 MHz (23 MHz, 90 MHz and 300 MHz) were performed in two different devices with inversion-recovery pulse sequence (180°- τ-90°). The variation range of the waiting interval (τ) between the pulses ranged of 0.1 to 10,000 ms and the interval between recycle applied to the end of the pulse sequence was 5 seconds. The devices were a Maran Ultra 23 and a 0.21 - 7.1 T variable magnetic field NMR spectrometer.

In turn, the measurements of T₁ in the range of 50 KHz until 9 MHz were done at room temperature using a fast field cycling (FFC) device developed at the physics department of Instituto Superior Técnico, Portugal. In this type of relaxometry, the analyses were proceeded through the sequence of field cycles. In each cycle, the sample remained subjected to the B₀ field during a time τ. After the B₀ → B₀ transition, a pulse of radiofrequency was applied to the sample in resonance with the Lamor detection frequency. The free induction decay was detected and the sample was left to relax to equilibrium for a stabilization time at a T₁ value five times longer. The cycle time was always greater than the stabilization time. The initial amplitude of the free induction decay signal was proportional to the magnetization M₀ (τ). The M₀ (τ) decay can in general be multi-exponential, with different relaxation times. All measurements were done according to the work developed by Sousa and collaborators [21] and Preto and collaborators [22].

3. Results

The results provided by SEM, FT-IR, WAXD, TG and LFNMR are discussed hereafter.

3.1. SEM Results

Figure 1 shows the SEM images (magnification—3000x) of the NG, AM-G and ODA-AM-G. In comparison with NG, morphological changes can be seen in the AM-G image, evidencing partial lamellar expansion. The image the sample modified with octadecylamine (ODA-AM-G) indicated that partial lamellar exfoliation was successful. This permits inferring that the insertion of the hydroxyl (OH) and carboxyl groups (COOH) on the surfaces of the
graphite layers and their distension was promoted by acid oxidation and sonication. The presence of OH and COOH groups enabled the modified graphite to react with octadecylamine and to expand and exfoliate its layers.

3.2. FTIR Results
The FTIR spectra of the samples are shown in Figures 2(a) and (b). As expected, the NG spectrum did not present any absorption in the medium infrared range. In the AM-G (Figure 2(b)) spectrum, OH (3500 cm$^{-1}$) and COOH (1700 cm$^{-1}$) could be detected. The bands at 3333, 3252 and 3161 cm$^{-1}$ (amine N-H bonds) and in the range of 2918 - 1366 cm$^{-1}$—related to the C-H and N-H bonds—are characteristic of octadecylamine (Figure 2(a)). The ODA-AM-G spectrum showed absorptions related to ODA, indicating the insertion of amine in the graphite galleries, obtaining modified graphite.

3.3. WAXD Results
Figure 3 shows the WAXD results. The NG and AM-G
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diffractograms presented a single 2θ peak at 26.5° but the diffractogram of the treated graphite showed a broadened and less intense diffraction peak. After acid treatment and sonication processes, the insertion of the OH and COOH reactive groups promoted some defects in the crystal distortion and orientation of the graphite lamellar stacking. In the diffractogram of the graphite modified with octadecylamine (ODA-AM-G), although the 2θ peak at 26.5° still remained, it was less intense than for the AM-G and a new diffraction peak around 21° was observed. This suggests that ODA insertion occurred and at least a partial delaminated structure was achieved. If any exfoliated structure was produced, it could not be detected by WAXD.

3.4. TGA Results

Figure 4 shows the thermogravimetric curves. The degradation curve of NG showed a first degradation around 100°C, attributed water absorption and degradation of the carbon chain around 700°C. The AM-G curve also revealed carbon chain degradation near 700°C. Three different degradation steps were observed in the ODA-AM-G curve. The first was located between 200°C - 350°C, the second was around 350°C - 600°C and the final one was above 700°C. The first two could be attributed to the degradation of the octadecylamine inserted in different types of structures of the modified graphite (delaminated and exfoliated). The final step represented the degradation of the graphite carbon chain. The results confirmed that organically modified graphite was produced.

3.5. Relaxometry Results

The relaxometry results are presented in the Figure 5. The acid mixture (AM sample) showed a straight line parallel to the frequency axis, being T1H quasi constant for all frequencies. For octadecylamine, the variation of relaxation time versus frequency presented a sigmoidal curve (S shape). For the graphite modified with acid mixture (AM-G sample), the curve revealed a rise from 10^5 Hz to 10^7 Hz, after which it remained constant. The profile was due to the presence of carboxyl and hydroxyl groups inside the graphite galleries, which can be chemically linked or adsorbed on the surface. The relaxometry curve of graphite modified with octadecylamine (ODA-AM-G) also showed that the relaxation time was dependent on the frequency. The profile of the graphite/octadecylamine curve indicated two rises. The first started at 10^5 Hz and finished at approximately 10^6 Hz. There was a plateau between 10^6 - 10^7 Hz but the point near 10^7 Hz suggested that a second rise occurred. It could be assumed that at least two kinds of structures were present in the graphite/octadecylamine sample. The first rise could represent the exfoliated structure where the octadecylamine molecules are less constricted while the second one could be attributed to the delaminated structure—octadecylamine molecules are more squeezed between the graphite layers. The result supports the observations from the WAXD and TG analyses.

4. Conclusion

Natural graphite was modified by the action of an acid mixture. The presence of hydroxyl and carboxyl groups on the graphite lamellar surface allowed its chemical modification through reaction with octadecylamine. Organically modified graphite was obtained. The techniques of characterization allowed inferring that a mix of structures—delaminated and exfoliated—was achieved.

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Figure 5. LFNMR for AM, AM-G, ODA-AM-G and ODA samples.

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REFERENCES


