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An easy synthetic way to exfoliate and stabilize MWCNTs in a thermoplastic pyrrole-containing matrix assisted by hydrogen bonds

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This work focuses on the design of an engineered thermoplastic polymer containing pyrrole units in the main chain and hydroxyl pendant groups (A-PPy-OH), which help in achieving nanocomposites containing well-distributed, exfoliated and undamaged MWCNTs. The thermal annealing at 100 °C of the pristine nanocomposite promotes the redistribution of the nanotubes in terms of a percolative network, thus converting the insulating material in a conducting soft matrix (60 μΩ m). This network remains unaltered after cooling to r.t. and successive heating cycles up to 100 °C thanks to the effective stabilization of MWCNTs provided by the functional polymer matrix. Notably, the resistivity–temperature profile is very reproducible and with a negative temperature coefficient of -0.002 K^{-1} , which suggests the potential application of the composite as a temperature sensor. Overall, the industrial scale by which A-PPy-OH can be produced offers a straightforward alternative for the scale-up production of suitable polymers to generate multifunctional nanocomposites.

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Introduction

Carbon nanotubes (CNTs) are one of the most popular fillers currently used in polymer nanocomposites (PNCs).¹ CNTs are cylindrical forms of graphitic sheets displaying a single wall (SWCNT) or multi walls (MWCNT) with open or closed ends.² CNTs are inherently multifunctional so they can work as structural supports,³ conductive^{4,5} and sensing platforms in PNCs.⁶ Conductive nanocomposites are currently used in several commercial products.⁷ Among them, PNCs with temperature sensing properties have become very attractive products in the open market due to their possibility of nano-scale tailoring and very low cost of production. Typically, these kinds of materials have shown a resistivity strictly dependent on temperature,⁸ thus opening successful applications in the field of miniaturized and potentially low cost plastic sensors.^{9–11} In some cases, the resistance variation proceeds by the dynamic interconnection/disconnection of the CNT network in the matrix.¹² However, temperature sensing properties of CNTs/polymer composites are demonstrated to be more reproducible when the resistive response is governed by the semi-

conducting features of exfoliated and stabilized CNT networks under thermal solicitations.¹¹

Despite all advantages of electronic temperature-sensing PNCs over conventional thermometers, these kinds of materials are not exempt of problems. A common drawback in the design of resistive sensors based on CNTs/polymer composites is the strong tendency of CNTs to aggregate in bundles during composite processing due to the strong van der Waals interactions between their graphitic surfaces, which make their large-scale utilization problematic.^{7,13,14}

Several strategies to improve CNT dispersion in polymeric matrices have been reported in the open literature (*e.g.* CNT/*in situ* polymerization composites, high-shear melts processing, injection molding *etc.*¹⁵). Among them, particularly attractive is the functionalization of the CNTs surfaces by the covalent attachment of functional groups,^{3,15–17} but disruption of their sp² conducting network even occurs.^{18,19} A non-disruptive strategy used to disperse CNTs by means of conductive polypyrroles has been reported for applications in electronics.^{20,21} In these particular systems, the pyrrole groups get in contact with the sp² network of the CNTs surface *via* supramolecular π–π interactions, which promote the polymer wrapping around the filler. As a result, multi-layers of the polymer form bridges that separate CNTs from each other yielding effective percolation pathways. However, despite the good conductive properties, polypyrroles/MWCNTs composites are highly brittle so that they must be doped with counter ions or coated on flexible polymer

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substrates to improve their mechanical performance (e.g. fracture toughness).^{22,23}

In a previous work, we reported on a simple approach to cross-link MWCNTs by using polyamines synthesized from alternating aliphatic polyketone by the Paal–Knorr reaction.²⁴ The covalent attachment of the polyamines on the MWCNTs occurred *via* covalent (amidation-type) grafting. The melt blending of the polyamine cross-linked MWCNTs with LDPE was attempted to reinforce the mechanical properties of the LDPE matrix. However, the enhancement of modulus and tensile strength of the LDPE matrix, as compared to un-cross-linked MWCNTs, was not achieved due to the poor dispersion of the filler.

Herein, we report on the design of a thermoplastic OH-functionalized alternating aliphatic polypyrrole (A-PPy-OH) matrix, which is capable to exfoliate and stabilize MWCNTs without the need of surface modification of the filler. The thermoplastic polymer is designed by the chemical modification of alternating aliphatic polyketone PK *via* Paal–Knorr reaction with amine compounds. The production of the polymer²⁵ and its chemical modification²⁶ occur in high yield, low cost and fast kinetic using relatively mild conditions in bulk and with water as the only by-product. Specifically, the reaction between the polymer and an OH-amine compound turns the alternating carbonyl backbone of PK into pyrrole units bearing hydroxyl moieties (Fig. 1). The combination of the polymer with MWCNTs produces a malleable and conductive rubber-like nanocomposite displaying electronic temperature-sensing properties. On one hand, the pyrrolic backbone exfoliates and stabilizes bundles of MWCNTs *via* supramolecular π – π interactions with the graphitic surface of the filler during thermal annealing. On the other hand, the OH-functional groups pending from the pyrrole units assist the polymer during the thermal stress by hydrogen bonding interactions in order to keep its dimensional stability and mechanical features. In order to figure out the role of the OH-motifs,

a reference polymer displaying the same pyrrolic backbone of A-PPy-OH was prepared by the chemical modification of PK with *n*-butylamine, where the hydroxyl group is replaced by a CH₃ group (A-PPy-CH₃). The CNTs exfoliation and their effective dispersion within polymer matrices were investigated by *in situ* resistance measurements and charge-contrast SEM imaging, whereas composites resilience under thermal stress was investigated by DSC analysis. The electrical resistance of composites was eventually evaluated under thermal cycles between r.t. and 100 °C, to explore their potential for the development of sensitive, stable, and reproducible temperature sensors.

Experimental section

Materials

The alternating aliphatic polyketone (PK) was synthesized according to Mul *et al.* 2002.²⁷ The resulting co- and ter-polymer of carbon monoxide presents a total olefin content of 30% of ethylene and 70% of propylene (PK30, MW 2687 Da). 3-Amino-1-propanol (OH) (Acros), butylamine (CH₃) (Sigma Aldrich, 99%), 2,5-hexanedione (Sigma Aldrich 98%), multi-walled carbon nanotubes MWCNTs (O.D. 6–9 nm, *L* 5 μ m, Sigma-Aldrich 95% carbon), DMSO-d₆ (Laboratory-Scan, 99.5%), 1-methyl-2-pyrrolidinone (Sigma-Aldrich, 99.5%) were purchased and used as received.

Model reaction

A model reaction between stoichiometric amounts of 2,5-hexanedione (8.7 mmol) and 3-amino-1-propanol was carried out in order to identify the presence of any side product after the Paal–Knorr reaction.²⁶ The reaction was performed in bulk, in a 100 mL round-bottom flask equipped with a reflux condenser and a magnetic stirrer. The reaction mixture was heated up to 100 °C during 4 h.

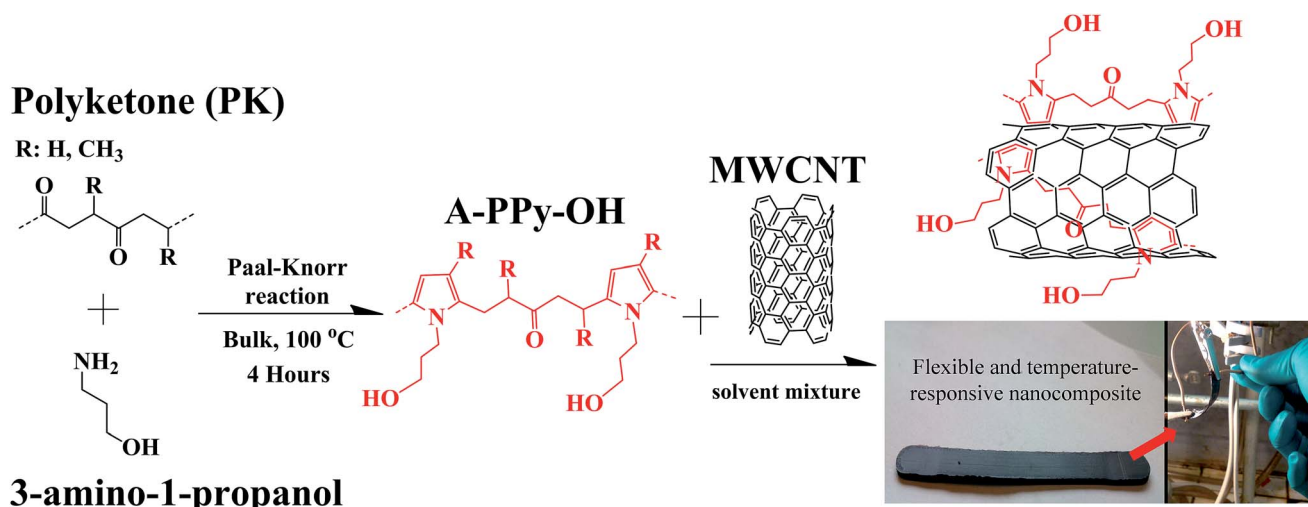


Fig. 1 Schematic representation of PK chemically modified with 3-amino-1-propanol *via* the Paal–Knorr reaction (A-PPy-OH), and the resulting composite after mixing with MWCNTs.



Functionalization of polyketone with alcohol pendant groups

The solvent-free Paal–Knorr reaction²⁶ between PK and 3-amino-1-propanol was carried out using different molar ratios between the 1,4-di-carbonyl groups of polyketone and 3-amino-1-propanol (Table 1). A reference polymer that displays the same backbone structure, but bearing a CH₃ instead of an OH group was also prepared using butylamine instead of 3-amino-1-propanol in the Paal–Knorr reaction, with the aim of evaluating the effect of hydrogen bonds on the thermal resilience of the composite. These chemical modifications of the polyketone were carried out in a 250 mL round-bottom glass reactor equipped with a reflux condenser, a U-type anchor impeller and an oil bath for heating. First, 60 g of PK (0.455 moles of di-carbonyl unit) were preheated to a liquid state at 100 °C. Then, 3-amino-1-propanol or butylamine was added dropwise to the reactor during 20 min. Next, the stirring speed was set to 600 rpm and the reaction was carried out for 4 h. Initially, the reaction mixture was colourless, but gradually turned to brown due to pyrrole formation on the polymer backbone.^{26,28,29} The resulting mixture was diluted with chloroform and washed 3 times with a 0.2 M NaCl Milli-Q water solution, to remove the unreacted 3-amino-1-propanol, if any. Thereafter, the organic phase was evaporated under vacuum at 50 °C for 24 h. Light-brown powders were obtained as final products. In order to avoid hydration, the samples were sealed in brown glass vials and stored at 6 °C for further characterization. The corresponding samples are coded as A-PPy-OH a_i or A-PPy-CH₃ a_i with a_i being the mol percentage of amine with respect to the carbonyl groups in the feed.

The percentage of conversion of carbonyls (C_{co}) into pyrrole groups can be calculated as follows:

$$C_{co} = \frac{y}{y+x} 100\%$$

where x and y represent the moles of di-ketone and pyrrolic units after conversion, respectively (Fig. 1). y can be calculated as follows:

$$y = \frac{\text{wt}(\text{N})}{A_m(\text{N})}$$

Table 1 Chemical modification of PK with different amounts of 3-amino-1-propanol (OH) and butylamine (CH₃). Samples are coded by stating the target percentage of conversion of carbonyls into pyrrole units considering a maximal 80% (a_i) of modification²⁹

Sample	Mol _{amine} (moles)	C_{co}^a (%)	η^b (%)
A-PPy-OH20	0.091	15	75
A-PPy-OH40	0.182	36	90
A-PPy-OH60	0.274	57	95
A-PPy-OH80	0.365	79	99
A-PPy-CH ₃ 80	0.365	78	98

^a C_{co} is the % of carbonyl conversion (determined by elemental analysis). ^b η is the conversion efficiency of carbonyl groups. In all samples 0.455 moles of di-carbonyl unit (60 g of PK) were used.

where wt(N) represents the grams of nitrogen in the final product according to elemental analysis, and $A_m(\text{N})$ is the atomic mass of nitrogen. x can be calculated as follows:

$$x = \frac{g_{\text{prod}} - y \times M_w^y}{M_w^{\text{pk}}}$$

where g_{prod} represents the grams of product after conversion, M_w^y the molecular weight of the pyrrolic unit and M_w^{pk} the molecular weight of a 1,4 di-ketone unit (131,6 g mol⁻¹). The conversion efficiency η is defined as the ratio between the carbonyl conversion C_{co} and the targeted one according to the amount of polymer and amine compounds provided in the feed (C_{co}^{feed}):

$$\eta = \frac{C_{co}}{C_{co}^{\text{feed}}} 100\%$$

the C_{co}^{feed} is calculated as follows:

$$C_{co}^{\text{feed}} = \frac{\text{Mol}_{\text{amine}}}{\text{Mol}_{\text{d-co}}} 100\%$$

with Mol_{amine} representing the moles of amine compounds and Mol_{d-co} the moles of di-carbonyl units in the feed.

A-PPy-OH40/MWCNTs composite

A-PPy-OH40 and MWCNTs were mixed in *N*-methylpyrrolidone, a solvent reported as an effective dispersant for MWCNTs,¹⁶ using fixed amounts of polymer and different amounts of MWCNT expressed as wt%. For this step, only one of the OH-functionalized polymers was selected aiming at the design of a flexible temperature-responsive nanocomposite³⁰ (Table 2).

In detail, 4 g of polymer were completely dissolved in *N*-methylpyrrolidone (10 vol%). The required wt% of MWCNTs was mixed with the same solvent and sonicated in a bath for 30 min and then poured to the polymer solution in a round bottomed flask at 50 °C for 24 h under stirring. Then, the mixture was rotary evaporated and finally transferred into a vacuum oven (80 °C for 72 h) to ensure the complete removal of the solvent. Rectangular solid samples with different dimensions were prepared by compression-molding at 150 °C for 30 minutes at 40 bar to ensure full homogeneity.

Characterization

Elemental analysis was performed with an Euro EA elemental analyzer. It was used to establish the percentage of nitrogen in the modified polymers. ¹H NMR spectra were recorded on

Table 2 Experimental conditions of A-PPy-OH40 mixed with different wt% of MWCNTs

Polymer	Polymer (g)	MWCNT (wt%)
A-PPy-OH40	4	2
A-PPy-OH40	4	4
A-PPy-OH40	4	6
A-PPy-OH40	4	8
A-PPy-OH40	4	10



a Varian Mercury Plus 500 MHz apparatus using DMSO- d_6 as solvent. ATR-FT-IR spectra were recorded using a Thermo Nicolet NEXUS 670 FT-IR. Differential scanning calorimetry DSC thermograms were recorded on a TA-Instrument DSC 2920 under N_2 atmosphere. The samples were weighed (10–17 mg) in an aluminium pan, which was then sealed. The samples were first heated from -20 to 180 °C and then cooled down to -20 °C. Four cycles were performed from -20 to 180 °C, with heating and cooling rates set to 10 °C min^{-1} . GPC measurements were performed with a HP1100 Hewlett-Packard instrument. The equipment consists of three 300×7.5 mm PLgel 3 μm MIXED-E columns in series and a GBC LC 1240 IR detector. The samples were dissolved in THF to obtain a final concentration of 1 mg mL^{-1} . THF was used as eluent at a flow rate of 1 mL min^{-1} at 40 °C. The calibration was done using polystyrene as standard and the data were determined using PSS WinGPC software. Scanning electron microscope images were acquired on a Philips XL30 Environmental SEM FEG instrument as previously reported.^{31,32}

In situ resistance measurements were performed during thermal annealing (at 100 °C) on samples (7.28 mm long, 5.85 mm wide, 1.25 mm thick) constrained between copper plates, connected to a multimeter and placed inside a chamber provided with a heater and a temperature controller (± 0.1 °C). Dynamic mechanical thermal analysis DMTA was conducted on a Rheometrics scientific solid analyzer (RSA II) under an air environment using the dual cantilever mode at an oscillation frequency of 1 Hz and a heating rate of 3 °C min^{-1} between 0 and 25 °C. The samples for DMTA analysis were prepared by compression-molding of 500 mg of the composite into rectangular bars (6 mm wide, 1 mm thick, 54 mm long) at 150 °C for 30 min under a pressure of 40 bar.

Results and discussion

Model component preparation

Model compounds are useful to characterize the structure of complex polymer systems *via* effective and rapid investigation techniques such as 1H NMR spectroscopy. Here, a model compound was prepared by reaction of 2,5-hexanedione (*i.e.* representative for the di-carbonyl moieties along the PK backbone) with 3-amino-1-propanol. This Paal–Knorr reaction resulted in the formation of a pyrrole unit, bearing a hydroxy terminated alkyl chain (Fig. 2).

Paal–Knorr chemical modification of PK

The reactions between PKs and 3-amino-1-propanol (Fig. 1) were carried out according to the different molar ratios as reported in Table 1. Notably, these functionalized PKs display a total carbonyl conversion (C_{co} , measured by elemental analysis) close to the target conversion (relative efficiency, $\eta \geq 90\%$, see Table 1), with a slightly lower value for the A-PPy-OH20 sample only. This result validates the robustness and versatility of the Paal–Knorr reaction of PK. The formation of the desired modified PKs containing pyrrole units with $-OH$ motifs

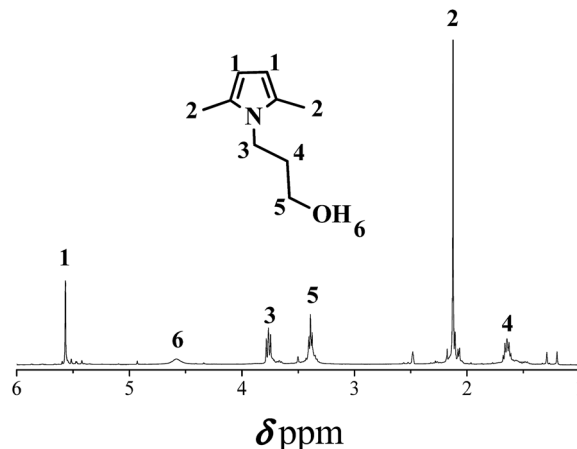


Fig. 2 1H NMR spectrum of model compound prepared by reaction of 2,5-hexanedione with 3-amino-1-propanol. The signal of protons associated to the pyrrole ring is at 5.6 ppm, CH_2 units give signals at 1.6, 3.4 and 3.8 ppm and the $-OH$ group gives a broad signal at 4.6 ppm.

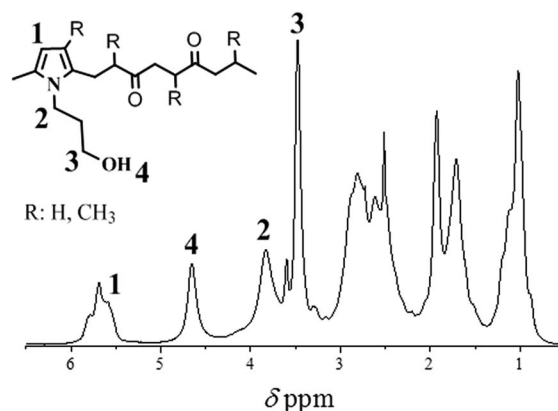


Fig. 3 1H NMR spectrum of A-PPy-OH40 in DMSO- d_6 .

was confirmed by 1H NMR spectroscopy (A-PPy-OH40 sample, Fig. 3).

The 1H NMR spectrum shows analogous signals to those found in the model compound (Fig. 2), *i.e.* the pyrrole units at 5.8 ppm, the CH_2 in the pendant group attached to the pyrrole unit at 3.5 and 3.8 ppm, and the hydroxyl group at 4.65 ppm.

An increase in the ratio between the amine compound and the polyketone in the Paal–Knorr reaction leads to the expected higher degree of conversion of the carbonyl groups of polyketone backbone into pyrrole units, as shown by the increasing C_{co} values in the A-PPy-OH a_i series from A-PPy-OH20 to A-PPy-OH80 (Table 1). The increase in the concentration of pyrrole units along the polymer backbone promotes polymer rigidity, as demonstrated by the enhancement in the T_g values of the functionalized polymers (Fig. 4).

The linear increase of the T_g with C_{co} is not only ascribed to the presence of pyrrole units in the backbone, but mainly to effective interactions between the $-OH$ groups of the polymer chains. This hypothesis is suggested by the much higher T_g value of A-PPy-OH80 (58 °C) compared to that of A-PPy- CH_3 80



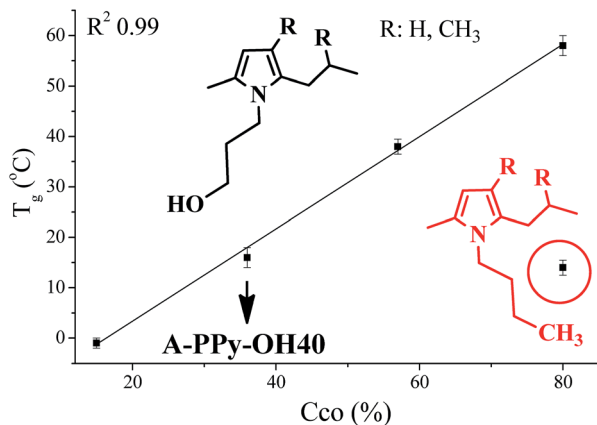


Fig. 4 T_g of the polymers of the A-PPy-OHa_i series (in black) and of A-PPy-CH₃80 (in red), plotted as a function of the degree of carbonyl conversion (C_{co}). The T_g values were measured by DSC in three temperature cycles from -20 to 180 °C.

(14 °C), which is characterized by the same conversion degree but contains only a paraffinic moiety as pendant group. The role of $-OH$ groups in promoting secondary interactions among macromolecules was further investigated by FT-IR spectroscopy (Fig. 5).

The characteristic band attributed to the $-OH$ groups stretching modes (3600 – 3100 cm^{-1}) becomes progressively wider and shifts to lower energies as the degree of functionalization of the A-PPy-OHa_i polymers increases. This is likely due to the higher number of hydrogen donors (OH) in the polymer chains. According to the literature, hydrogen donors undergo highly directional interactions with their hydrogen acceptors.^{33–35} In our case and according to Fig. 5, the intensity of the peak varies with the hydrogen bonding density, which depends on the degree of polymer modification. However, if a shift of the

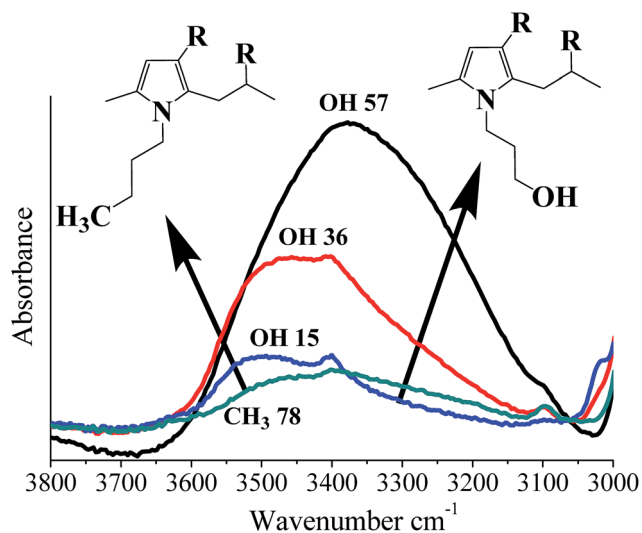


Fig. 5 FT-IR spectra of the A-PPy-OHa_i series and A-PPy-CH₃80. Numbers indicate final % of carbonyl conversion after Paal-Knorr reaction.

Table 3 GPC measurements of the polymers of the A-PPy-OHa_i series and of A-PPy-CH₃80

Experiments	$M_n (\times 10^3)$	$M_w (\times 10^3)$	PDI
A-PPy-OH20	3.2	7.6	2.4
A-PPy-OH40	3.0	7.0	2.3
A-PPy-OH60	2.9	6.7	2.3
A-PPy-OH80	2.7	6.2	2.3
A-PPy-CH ₃ 80	3.4	6.1	1.8

peak is observed, it might be attributed to the balance between different competing intermolecular hydrogen bonds.³³ So that, the red-shift of about 100 cm^{-1} observed in Fig. 5 suggests that one single directional interaction of the $-OH$ group can be excluded. This fact can be explained by the mobility of hydrogen bonding so that its interaction with different groups on the surface of the polymer could also take place as the conversion increases and the most probable hydrogen acceptor is decreased (*i.e.* carbonyl groups). Moreover, we cannot exclude at this stage that the broad band at about 3400 cm^{-1} could be also attributed to a slight but effective hydration of the sample. In any case, the contribution of the OH groups in getting higher T_g values compared to the paraffinic counterparts results crucial since GPC measurements of all polymer systems showed no significant differences between their molecular weights (Table 3).

A-PPy-OH40/MWCNTs composites

Polymer nanocomposites were designed with the aim to target a flexible, light and temperature-responsive system. In this attempt, A-PPy-OH40 was chosen among the A-PPy-OHa_i polymers because its T_g is close to room temperature, and this is expected to provide the desirable flexibility of the composite at easily accessible temperatures. Moreover, this feature supports the use of the A-PPy-OH40/MWCNTs system as a resistive sensor for temperature variations close to the physiological regime. The A-PPy-OH40/MWCNT composites were prepared by mixing

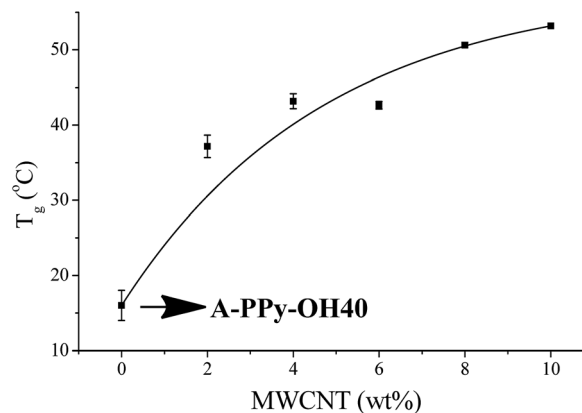


Fig. 6 T_g of the composites obtained by mixing A-PPy-OH40 with different wt% of MWCNTs. The T_g values were measured by DSC in three temperature cycles from -20 to 180 °C.



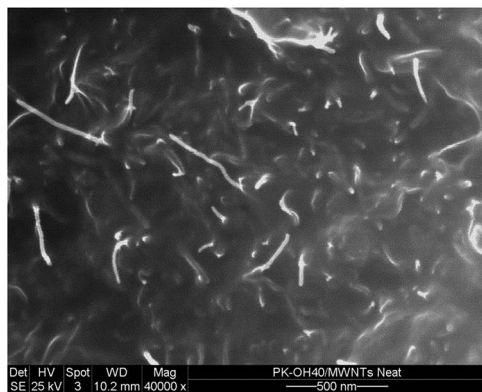


Fig. 7 SEM morphological study of 4 wt% MWCNT/A-PPy-OH40 nanocomposite.

A-PPy-OH40 with different amounts of MWCNTs (Table 2) in *N*-methylpyrrolidone. The presence of MWCNTs causes an increase in the T_g of the composite materials compared to the parent polymer (Fig. 6).

According to the literature, the increase of T_g values with MWCNTs loading in polymeric systems is found to be related to the increment in their viscosity^{36,37} due to the interfacial interaction between matrix and filler at molecular level. Functional aromatic groups included in the backbone or as pendant groups in polymer chains, get in contact with the graphitic surface of the filler *via* supramolecular π - π interactions. This behavior hinders the mobility of the polymer chains and hence increases the T_g .³⁸⁻⁴⁰ Conductive polypyrroles have been reported as good dispersant agents for CNTs due to the interfacial connection of the filler with the pyrrole groups *via* π - π interactions. In our case, the backbone of the polymer contains 36% of pyrrole units that possibly promote effective interactions with the graphitic structure of CNT. In order to determine MWCNTs exfoliation, the 4 wt% MWCNT/A-PPy-OH40 nanocomposite was analyzed by SEM (Fig. 7).

The SEM image clearly shows the single (unbundled) nanotubes as the dominant species, thus confirming the good interactions between MWCNTs and the A-PPy-OH40 matrix.

With the aim to further improve the dispersion of the MWCNTs in the polymer matrix in terms of a fibrous percolative structure, the rubber-like composite containing 4 wt% of MWCNTs was subjected to thermal annealing at a constant temperature of 100 °C for 3 h and then cooled down to r.t. During the annealing, *in situ* resistivity on the compressed mold specimen was measured in order to evaluate the effect of the thermal treatment on the formation of the conductive MWCNTs network within the polymer matrix. Before annealing, the resistivity of the polymer was infinite but it decreased sharply to $\approx 60 \mu\Omega$ m during the first minutes of thermal annealing and well persisted after cooling (Fig. 8).

This behaviour indicates that thermal treatment at 100 °C effectively reorganizes MWCNTs structures within the polymer bulk in a well-defined percolative network. These remarkable changes easily occurred upon a mild thermal treatment, which is however effective in promoting polymer matrix mobility.

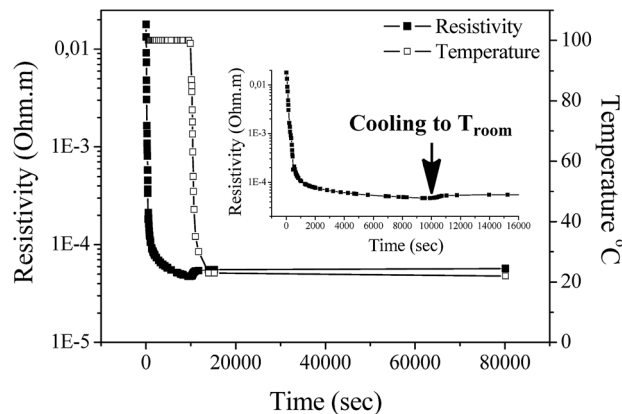


Fig. 8 *In situ* electrical measurement of 4 wt% MWCNT/A-PPy-OH40 composite during thermal annealing at 100 °C. In the inset, a magnification of the first annealing instants.

A morphological study of the composite analyzed by SEM in contrast mode³² corroborates the electrical measurements of the nanocomposite before and after annealing (Fig. 9). The micrographs taken from the surface of freshly teared samples revealed that bundles of MWCNTs are still present in the A-PPy-OH40 before annealing as evidenced at high magnification (Fig. 9B). It might be expected that after the sample is subjected to compression molding at 150 °C for 30 min (see experimental procedure), debundling and good dispersion of MWCNTs should be achieved. However, it is not observed in the micrograph (Fig. 9B). A possible explanation is that during compression moulding the system did not have enough time to create a well percolative network composed by debundled MWCNT. Conversely, the annealing at 100 °C, very close to the softening point, would allow the same polymer matrix mobility and provide more time to the aromatic moieties of the polymer

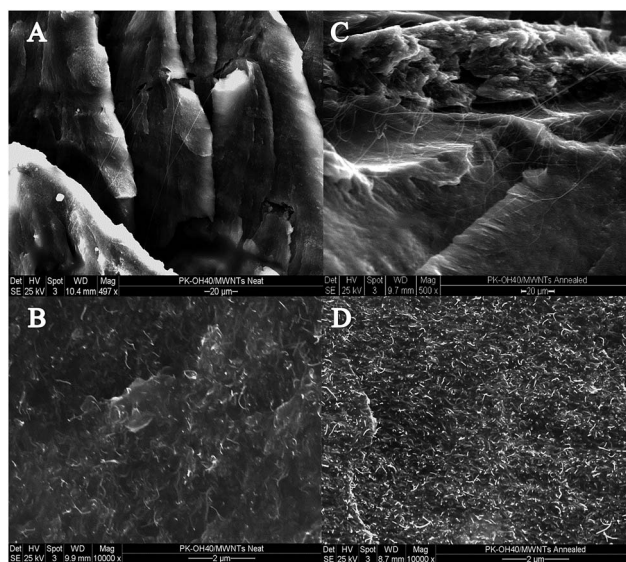


Fig. 9 SEM morphological study of 4 wt% MWCNT/A-PPy-OH40 composite after compression-molding at 150 °C for 30 min (A and B) and after thermal annealing (C and D) at 100 °C.



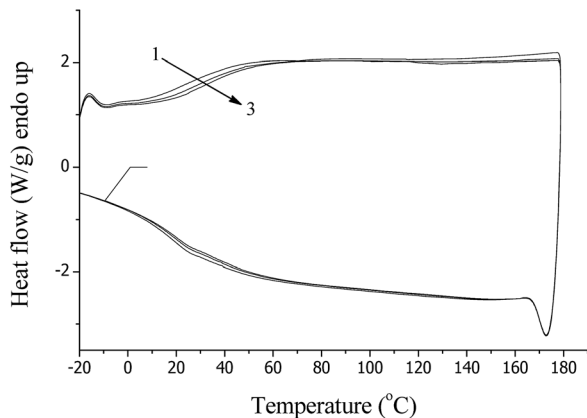


Fig. 10 Three consecutive DSC thermal cycles (indicated by the arrow) of 4 wt% MWCNTs/A-PPy-OH40 composite after annealing.

chains to well interact with the graphitic surface thus inducing a perfect and effective exfoliation (Fig. 9C). This indicates that the thermal treatment at 100 °C favors MWCNTs debundling and promotes their homogenous dispersion in the polymer matrix (Fig. 9D) in a percolative network.

The thermal history of the 4 wt% MWCNT/A-PPy-OH40 nanocomposite after thermal annealing was evaluated by DSC analysis in order to establish the resilience of the material under the investigated temperature regime (Fig. 10).

The similarity in each consecutive thermal cycle demonstrates the resilient character of the material without any sign of thermal degradation till 180 °C. The thermal traces also indicate that the same thermodynamic response of the composite remains upon heating, thus indicating no phase separation between the components. Considering that the T_g of the composite is around 40 °C, it is worth noting that the sample does not display any endothermic transition during the annealing at 100 °C. This is another prove of the role played by the -OH groups in keeping the dimension stability of the composite.

Fig. 11 shows a picture with two different materials after being subjected to dynamic mechanical thermal analysis DMTA (from r.t. to 35 °C). A-PPy-OH40/MWCNT is capable to keep its dimensions. Contrarily, A-PPy-CH₃80/MWCNT gets completely deformed. This suggests that the presence of hydrogen bonding plays a relevant role in giving dimensional stability to the material.

The resistivity response of the polymer/MWCNT nanocomposite towards consecutive temperature cycles was monitored between r.t. and 100 °C (Fig. 12).



Fig. 11 Pictures of two different nanocomposites displaying their differences in dimension stability (damping capability of the material) after being tested by dynamic mechanical thermal analysis DMTA (data about modulus not showed for brevity).

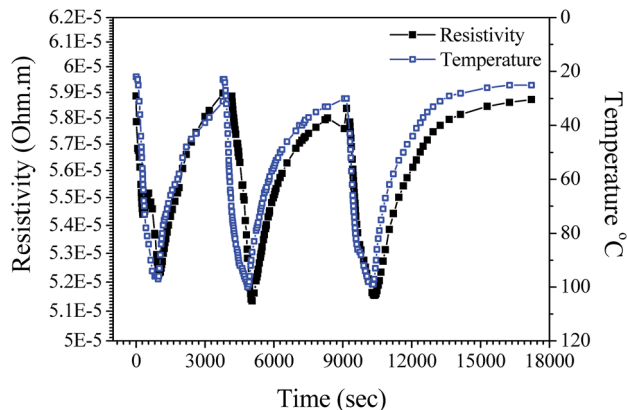


Fig. 12 *In situ* electrical measurement of 4 wt% MWCNT/A-PPy-OH40 composite during three consecutive thermal cycles between r.t. and 100 °C.

Reproducible resistance variations were observed, with maximum amplitudes of 51 to 59 $\mu\text{ohm m}$ within the temperature interval of 80 °C. Moreover, the resistivity-temperature profile was very reproducible and with a negative temperature, thus proving that the percolation network does not experience any significant changes during the heating-cooling cycles.

The electrical response with temperature shows a negative temperature coefficient of resistance^{41,42} of -0.002 K^{-1} (average of three temperature cycles), an absolute value that is comparable to the highest values found in metals ($0.0037\text{--}0.006 \text{ K}^{-1}$, 0.00385 K^{-1} for a Pt100 sensor) and similar to other CNT/polymer nanocomposites reported in the literature.^{6,11,12}

In summary, by this approach we have demonstrated an easy method (industrially scalable) to modulate the thermo-mechanical and electrical conductive performance of the A-PPy-OH40/MWCNT nanocomposite by adjusting the pyrrole group content and hydrogen bonding density of the polymer. Remarkable, the ability of the polymer in dispersing and stabilizing unmodified MWCNTs, *via* physical interactions, represents a technological advance in composite science considering the nanoscale and low cost of production at which the temperature sensor material can be tailored to be used in electronic applications.

Conclusions

We have demonstrated that a thermoplastic pyrrole-containing matrix is an effective dispersant for MWCNTs exfoliation. The polymer was prepared *via* the Paal-Knorr modification of an alternating aliphatic polyketone (PK) with OH-amine compound. The chemical reaction turns the waxy PK into a flexible rubber-like OH-functionalized pyrrole-containing polymer with modulable T_g depending on the amounts of -OH groups. The polymer is able to generate non-covalent functionalization of the MWCNT graphitic materials through effective π - π interactions, and that the exfoliation process does not do significant damage to the one-dimensional CNT



structure. SEM micrographs and DSC traces demonstrate that the A-PPy-OH/MWCNTs nanocomposite is capable to undergo continuous thermal cycles from -20 to 180 °C without any sign of interphase modification and matrix degradation. Notably, the conductive CNT network is maintained after several temperature cycles (from r.t. to 100 °C) proving the remarkable stability of the MWCNT homogeneous dispersion within the polymer matrix. Measurements repeated over three successive heating cycles revealed highly reproducible resistivity variations with negative temperature coefficient of about -0.002 K $^{-1}$, an absolute value comparable to the values found in metals. Overall, this data consistently support the use of A-PPy-OH/MWCNT nanocomposite as a soft and highly reproducible resistive sensor for temperature variations. It is certainly expected that the novelty of this approach, together with the industrial scale by which A-PPy-OH can be produced, will pave the path toward the scale-up of the proposed material at industrial level.

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