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Influence of hard segment content and nature on polyurethane/multiwalled carbon nanotube composites

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Abstract

Knowledge of how polyurethanes, PU, complexity affects their derived multiwalled carbon nanotube, MWCNT, composites could shed important clues for preparing future tailored PU/MWCNT elastic, strong and electrically conductive composites. In this regard, hard segment content and nature, along with MWCNT functionalisation, are believed to have great influence on both nanoscale PU/MWCNT self assembling mechanisms and on final composites properties. In this work the effect of PU hard segment content into composites was analysed. According to the results, a preferential interaction of nanotubes with polyurethanes hard segments can be assumed although nanotubes introduction hindered both soft and hard segments crystallisation. In all cases carbon nanotubes percolative network formation seemed to be crucial for obtaining significant reinforcement, being observed at this stage, a reduction of ductility, phenomena which is related to an increase on hard domains interconnections by MWCNT. The hard to soft segment ratio into PU plays a crucial role on determining the stress transfer to MWCNT. In addition, PU hard domains nature has important effect on nanotubes reinforcing character, this fact being related to the different PU intrinsic morphologies as well as different PU-MWCNT interactions.

Keywords:

A. Carbon nanotubes

A. Polymer-matrix composites (PMCs)

B. Mechanical properties

B. Fracture

B. Interfacial strength
1. Introduction

The concept of creating both structural and functional multi-nanophase composites with improved performance is currently under development in a wide variety of metallic, ceramic and polymeric matrices, although the emphasis to date has been on polymeric systems [1-3]. By introducing carbon nanotubes (CNT) into a polymer matrix, many properties can be improved at once, including mechanical [4,5], thermo-mechanical [6], electrical [7,8], thermal [9,10], chemical [11] and optical properties [12].

Polyurethane (PU) is among the most versatile materials today [13-15] and can be tailored according to the desired application requirements choosing carefully the reactants, their ratio and their synthesis procedure [16]. Different combinations of strength, ductility, biodegradability or hardness can be achieved with the proper formulation and preparation. Nevertheless there are still some properties such as thermal stability, electrical and thermal conductivity, stiffness or stress recovery, which could be improved with the addition of CNT. Combination of polyurethane elastomers with CNT might lead to new functional materials with combined properties.

Some researches have focused more on the applicability of PU/CNT composites to different fields such as biomedicine [17-19], adding benefit of the unique properties of CNT, while some others have focused more on the materials science point of view [20]. While in some works the research was focused on understanding the effects of the addition of CNT to a polyurethane matrix through melt blending [21] or solvent mixing [22-24], in some others it has been tried to benefit from the polyurethanes chemistry to prepare grafted nanotubes hybrids which were further polymerised in-situ with the aim of creating CNT cross-linked networks.
[25-30], or to functionalise the nanotubes and then solvent mix them with the polyurethanes to compare the effect of different nanotubes treatments [31,32]. Nevertheless, to authors knowledge, any thorough study of PU nature influence on MWCNT reinforcing character has yet been addressed.

PU/MWCNT composites can be considered as ternary systems formed by the nanofiller, hard segments (HS) and soft segments (SS), each of which can vary in ratio, chemical composition and physical properties. The study of the polyurethanes phase morphology variation (which without filler is governed by HS and SS thermodynamic incompatibility) with the nanotubes inclusion and its effects onto mechanical performance, along with the understanding of thermodynamics of preferential association between nanotubes and PU phases should be considered of great importance to develop tailored PU/MWCNT composites.

In this work, a set of polyurethane/multiwalled carbon nanotubes (PU/MWCNT) with different HS content and nature was prepared by the solvent casting approach and buckypaper infiltration method for medium and high MWCNT concentrations, tested both mechanically and electrically and analysed by scanning electron microscopy (SEM). The morphology was also analysed by means of differential scanning calorimetry (DSC).

2. Experimental

2.1 Preparation of PU matrices

Polyurethane matrices were synthesised in our laboratory by the two shot polymerisation approach with 10, 14, 30, 40, 50 wt% hard segment formed by 1,6-hexamethylene diisocyanate (HDI, Bayer, Desmou H) and 1,4-butanediol (BD) (Aldrich, 99 % purity) and with 30 wt% of hard segment content formed by 4,4'-
diphenylmethane diisocyanate (MDI, Bayer Desmou 44) and BD. Soft segment
consisted on a polydiol, formed by polycaprolactone and polycarbonate blocks (PCL-b-
PHMC-b-PCL, Ravecarb 111, Polimeri Europa). The preparation and properties of the
polyurethanes is described elsewere [33]. The prepared polyurethanes are named
indicating the employed diisocyanate in their hard segments and weight percent of hard
segment in the polyurethane, as HDI-10 or MDI-30 for PU with 10 wt% hard segment
based on HDI and 30 wt% hard segment based on MDI, respectively. The molar
composition of all the polyurethanes is listed in Table 1.

2.2 Nanotube acid pre-treatment.

To favour nanotubes dispersion into solvents and polymer matrices, multiwalled
carbon nanotubes (Nanocyl 3100, Belgium) were always purified with a 2 h strong acid
treatment, consisting on a mixture of nitric and sulphuric acids in a ratio HNO₃:H₂SO₄
of 1:3, in a sonic bath, obtaining acid treated multiwalled carbon nanotubes, MWCNT-
COOH with average lengths of 740 nm, as described elsewere [34].

2.3 Preparation of PU/MWCNT composites

Composites with acid treated carbon nanotubes contents up to 20 wt% were
prepared by solvent casting (SC) method [34]. Higher MWCNT loading composites
were prepared by buckypaper (BP) infiltration of different concentrated polyurethanes
tetrahydrofuran (THF) solutions and quantities, always with a controlled pressure of
200 ± 20 mbar. For polyurethanes above 30 wt% of hard segment the solvent employed
was a mixture of THF:N,N-dimethylformamide (DMF) (1:1). Infiltration times were
shorter for diluted solutions and smaller solution quantities. Typically 50 mL of 1 mg
mL⁻¹ solution were infiltrated in about 12 h, while 50 mL of 2 mg mL⁻¹ needed around
24 h, and 100 mL of a 1 mg mL⁻¹ solution required around 20 h to infiltrate. When
around 12 h of infiltration time was achieved, infiltrating pressure was reduced down to
50 ± 10 mbar in order to speed up the process.

2.4 Characterisation methods

Final nanotube volume fraction into the composites, φ, was related to the nanotubes weight fraction, \( X_{\text{MWCNT}} \), polymer density, \( \rho_{\text{Polymer}} \), and nanotubes density, \( \rho_{\text{MWCNT}} \), using the following equation [35]:

\[
\phi = \frac{X_{\text{MWCNT}}}{X_{\text{MWCNT}} + (\rho_{\text{MWCNT}}/\rho_{\text{Polymer}} - (\rho_{\text{MWCNT}}/\rho_{\text{Polymer}})X_{\text{MWCNT}}}.
\]

The films density was calculated by measuring precisely materials dimensions with a calliper (Mitutoyo, DIGIMATIC CD-15CP) and a low-torque digital micrometer (Mitutoyo) of pieces of about 7 mm x 0.15 mm x 2.5 mm and by weighting them on a thermo-gravimetical analysis balance (Mettler Toledo, TGA/SDTA 851) with precision down to ± 0.001 mg. Typical weight values ranged between 0.5 and 4 mg.

Electrical DC measurements were carried out using a two probe current intensity voltage scan from 0 to 5 V employing a semiconductor analyser (Keithley 4200-SCS). Resistance was calculated from the slope of current intensity versus voltage (I-V) curves and averaged from three measurements. Distances between silver paint printed electrodes and specimen widths were measured with a digital calliper while thicknesses were measured using a low-torque digital micrometer in order to normalise resistance values into conductivity (S cm\(^{-1}\)) values.

Tensile tests were performed with a constant crosshead speed of 100 mm min\(^{-1}\) for solvent cast composites and low MWCNT volume fraction infiltrated buckypapers, and with a crosshead speed of 1 mm min\(^{-1}\) for all buckypapers, with an initial crosshead distance of 8.5 ± 0.5 mm. Sample sizes were in the range of 80-140 \(\mu\)m in thickness and of 2.5 mm in width. The equipment used (MTS insight 10) was provided with pneumatic grips (Advantage Pneumatic Grips) and with a loading cell of 250 N. The
results were averaged from a minimum of 3 specimens.

SEM analysis was performed using a JEOL JSM-7000F, with an accelerating voltage of 20 kV and a distance of 9.0 ± 0.5 mm. Solvent cast composites with a MWCNT content lower than 20 wt\%, were coated with a thin gold layer of about 5-10 nm thickness with a Bal-Tec SCD-004 sputtering equipment.

DSC analysis of solvent cast composites was performed with scanning rates of 10 °C min\(^{-1}\) from -60 °C to 200 °C using nitrogen as purge gas and an electric intracooler as cooling source. Heat flow was normalised to polyurethane weight fraction into composites.

The non-isothermal crystallisation experiments were carried out taking the samples to 220 °C at 20 °C min\(^{-1}\), leaving to melt for 4 min at that temperature and cooling down to 20 °C at 2.5, 5, 10 and 20 °C min\(^{-1}\) of cooling rate.

3. Results and discussion

3.1 MWCNT percolation network formation

Sequentially increasing nanofiller addition to a hosting matrix leads to a situation in which at a particular concentration the filler forms a continuous percolated network. Upon this particular concentration a dramatic variation in some properties is expected due to the formation of two co-continuous phases which affect in different ways to the final composite properties. Detection of the critical MWCNT percolation volume fraction, \(\phi_p\), in a HDI-10 polyurethane, was performed graphically from the sharp increase of the electrical conductivity at \(\phi_p\), as seen in Fig. 1. The bending point, upon which the increase of properties was sharper, occurred at \(\phi = 0.025\) (4 wt\%). The nanotubes disposition in relation to the PU matrix changes from below to above percolation threshold concentration, upon which nanotube-nanotube contacts become
more and more important.

Electrical behaviour was modelled using the classic power scaling law for percolative systems [36]. The green curve has been drawn using this model with \( \{ \sigma_0, \phi_p, t \} = \{ 20 \text{ S cm}^{-1}, 0.025, 2.73 \} \) \( \sigma_0 \) being the electrical DC conductivity of the 100 wt% MWCNT buckypapers and \( t \) the value obtained from the slope of the experimental \( \log \sigma_{\text{DC}} \) vs. \( \log (\phi - \phi_p) \) curve. This curve has been extrapolated towards \( \phi = 0 \), introducing the conductivity of the neat polyurethane as initial value of \( \sigma_{\text{DC}} \).

Composites prepared by the infiltration method presented slightly higher electrical conductivities than those based on the solvent cast approach, probably due to a more direct contact between nanotubes in the network junction points.

3.2 Influence of hard segment content.

The effect of MWCNT-COOH addition on the elongation to break, \( \varepsilon_{\text{max}} \), normalised tensile strength, \( \sigma/\sigma_0 \), and elastic modulus, \( E/E_0 \), of composites based on polyurethanes based on HDI diisocyanate with different hard segment content is gathered in Fig. 2. The common feature for all types of composites is that mechanical properties evolution was virtually invariable or very small up to the nanotube percolation threshold concentration.

As seen in Fig. 2a-c this spike on the property versus \( \phi \) behaviour, occurred at higher nanotube concentration for composites whose matrices contained higher amount of hard segment. The ductility decrease results further suggests a preferential interaction of nanotubes with polyurethane hard segments. Being the soft domains responsible of polyurethanes extensibility, it is understandable that polyurethanes with bigger amount of soft segment might have nanotubes hosted within soft domains at lower nanotube concentrations than polyurethanes with higher amounts of hard segment. This fact is
also observed in Fig. 3a, where the decrease in ductility is represented for several polyurethane hard segment contents against nanotube weigh fraction. In polyurethanes with higher hard segment content the decrease in extensibility occurred at higher nanotube loadings because the nanotubes are primarily interacting with hard domains, which do not contribute primarily to material extensibility.

Modulus enhancements as calculated for each HDI based polyurethane composite series are gathered in Fig. 3b. It is seen that the tendency of this value (slope of $E_c$ vs. $\phi$) trends to increase with polyurethane hard segment content, with a dramatic increase for HDI-30 system. These values, when normalised over the matrices elastic modulus, $E_m$, provide the relative modulus enhancement (slope of $E_c$ vs. $\phi$ divided over matrix elastic modulus), which values are gathered in Fig. 3c. As seen, the enhancements tend to decrease with hard segment content.

The interfacial shear strengths, $\tau$, as calculated from the slope of $\sigma$ vs. $\phi$ data for each polyurethane series, using a common Bowyer-Bader model for aligned fibres in the manner described elsewere [34], are gathered in Fig. 3d. As can be seen, the maximum value was achieved utilising HDI-30 as a matrix. The bell type pattern of the data can be related to the match between polyurethanes surface tension, $\gamma$ as calculated by water contact angle measurements [33], that occurs around a polyurethane composition of 30 wt% hard segment (42.6 mJ m$^{-2}$), with that of the carbon nanotubes of 40 mJ m$^{-2}$ reported by Bergin et al [37].

The ductility reduction fact can be explained taking into account the nanofiller and hard domain relative sizes. These particular polyurethanes hard domains have an average diameter of ~ 15 nm with interdomain distances in the range of 14-32 nm [16] (see Fig. 1). On the other hand the nanotubes have been measured to have diameters of
~ 12 nm and lengths of ~ 740 nm on average [34]. Therefore nanotube length may be more than enough to interconnect different hard domains and therefore hinder soft segments ductility, as represented in Fig. 4. This fact strongly supports the idea that nanoconfinement into polyurethane hard domains is an important factor to retain the ductility [38], as has been suggested to be the case for natural silk [38,39].

The fracture surface of composites with 20 wt% MWCNT-COOH and HDI-10, HDI-30 and HDI-50 matrices were studied by SEM. Results are gathered in Fig. 5. The increase of matrix hard segment content led to more brittle surface pattern, characterised for a more homogeneous fringe sizes and distributions. In the first row (Fig. 5a,b) the HDI-10 system is shown at different magnifications, revealing high deformations and tubular structures protruding from the bulk, which could be due to bundled nanotubes aligned along the stress direction, wrapped by HDI-10 matrix. No isolated nanotubes were observed for the HDI-10 system. In HDI-30 system (Fig. 5c,d) no isolated nanotubes were observed, but some brighter dots protruding from the bulk were observed, which could be related to nanotubes tips appearing along the surface. In HDI-50 system the nanotubes were clearly seen (Fig. 5f) protruding perpendicularly to the fracture surface. This result suggests that despite the closer values between Hildebrand global solubility parameters [40], δ of MWCNT-COOH to polyurethane hard segments (Table 2), the overall matrix surface tension may be the determinant factor to account for the stress transfer between the polyurethane matrix and the nanotubes.

As mentioned in the introduction section, PU composites can be regarded as ternary systems in which their complexity arises not only by the presence of a nanofiller but also for the changes in morphology of PU hard phase and soft phases with a third component addition. Therefore to study morphology evolution with MWCNT addition calorimetric measurements were performed. Heating scans of composites prepared with
the polyurethanes based on HDI with 10, 30 and 50 wt% hard segment contents are shown in Fig. 6a-c, respectively. In Fig. 6a it can be seen that the endotherm appearing with low hard segment content PU around 50 °C, related to hard segments interacting with soft segments, which is activated at hard segment glass transition, \( T_{\text{Hard::Soft}} \) is decreased as nanotube content increases, while another peak appears at higher temperatures, \( T_{\text{Hard::MWCNT}} \), corresponding to interaction of hard segments with nanotubes. This area is coloured with dark brown. This is related to the lower mobility of hard segments and the hindrance to develop hard segments-soft segments interactions. The increase in the transition endset suggests some hard segment-nanotube interactions stabilise the dynamic thermal properties of the composites, as has been reported in our previous paper [34]. The crystallisation exotherm of soft segments, coulored with light blue, with a peak at -28 °C, decreased as nanotube content increased, suggesting also a restricted mobility of soft segments. In PU with higher hard segment content such as HDI-30 and HDI-50, the hard segments are able to develop more phase separated structures which have higher temperatures endotherms associated with hard segment crystals. In Fig. 6b,c for HDI-30 and HDI-50, respectively, this endotherm is coloured with light orange and in both cases is reduced with nanotubes incorporation while it is seen a concomitant increment of the heat capacity change related to amorphous hard segments at the glass transition at \( T_{\text{g Hard}} \). This fact suggests nanotubes interacted with hard segments reducing their crystallising capability and leading to more amorphous hard phase. The transition corresponding to amorphous hard segments increased, once again confirming the presence of more independent segments in a non-crystalline state as nanotube content increased (further confirmation by DSC in the supporting information, Fig.S1).

Further confirmation of nanotube effects on domains crystallinity was
demonstrated by dynamic DSC crystallisation measurements. In Fig. 7a the nucleating effect of nanotubes over polyurethane hard segments is seen as an increase in the crystallisation exotherm peak of about 11 °C in these cooling scans of composites prepared with HDI-50. This fact suggests that hard segments interacted with nanotubes and at the same time they reduced their crystallinity degree. Performing similar experiments under sub-zero temperature the same nucleating effect was found for soft segments in HDI-10.

In Fig. 7b typical modified Avrami plots [41] for non-isothermal crystallisation experiments, carried out at 2.5 °C min⁻¹ cooling rate, are presented, indicating the differences in slope, \( n_1 \), of the curves, which varied from 38 for the pure HDI-50 to 14 for the composite containing 3 wt% of nanotubes. This indicates a dramatic change in hard segment crystallisation mechanism [42]. The y-axis intercept, related with crystallisation rate, increased from -57 to -19, from the HDI-50 to the nanocomposite, indicating a big increase in hard segment crystallisation rate with the addition of MWCNT. This fact contradicts results obtained for PA6/MWCNT [43] and Nylon6/Foliated graphite [44] composites in which a decrease in both crystallisation rate and crystalline degree were found. Differences could arise from the difference in the matrix employed. Fig. 7c represents schematically the proposed nucleating effect of nanotubes over hard segments at the same time they also reduce the hard segment crystalline degree by avoiding hard segment mobility and self assembling (Melting enthalpies are shown in the support information, Fig. S1).

All the results from both the solvent casting method and buckypaper infiltration approach were now gathered to try to create a ternary composition diagram, as exposed in the introduction. Fig. 8 presents a ternary diagram in which the compositions of SC composites made up with HDI-10, HDI-30 and HDI-50 are presented along with some
of the derived high loading infiltrated BP nanocomposite compositions. Obtained data for some representative systems is also included. It is seen that a wide range of properties can be obtained and varied by the proper selection of the polyurethanes hard segment to soft segment ratio, but also by the addition of MWCNT. This tuneability of mechanical properties permits these ternary systems (e.g. HDI-BD hard segments, PCL-b-PHMC-b-PCL and MWCNT) to fulfil the demand of a different range of applications.

The complexity of using ternary diagrams to relate composition with properties, is highlighted when some other aspects such as processing techniques [34], thermal treatments [45], solvent choice [46] or nanotubes functionalisation and aspect ratio are considered. But as will be discussed next, variation of polyurethane hard segment nature also has important influence on some nanocomposite final mechanical performance features such as ductility or mechanical reinforcement.

3.3 Influence of hard segment nature

The neat HDI based polyurethanes had high modulus/hard segment content ratio. With just 10 wt% hard segment into the polyurethane the tensile modulus was 8 ± 0.6 MPa. There are other examples in literature [15] in which such a modulus value is just obtained using about 30 wt% of aromatic MDI and BD as hard segment constituents, and using a polyol of about the same molecular weigh as in this work. This is related to the higher crystallinity of HDI/BD hard segment based polyurethanes in comparison to those based on MDI/BD [33]. Therefore an advantage of using this high crystalline polyurethane with low hard segment contents is that while providing fairly good modulus it retains the benefits of a highly ductile material.

For comparison of MWCNT-COOH reinforcement into a HDI based polyurethane with those based on MDI, a hard segment ratio of 30 wt% was chosen.
The effects of nanotube loading on mechanical properties for both HDI-30 and MDI-30 polyurethanes are gathered in Fig. 9. Table 3 gathers the reinforcement parameters along with surface tension values of composites prepared with HDI-30 and MDI-30 as matrices and different nanotubes contents. It is seen that the relative modulus enhancement is higher for the MDI based polyurethane than for the based on HDI, this fact being related to the differences between matrices crystallinity.

Surprisingly MDI-30 polyurethane retained its ductility for higher nanotube volume fractions than HDI-30, while modulus enhancement began to occur at lower nanotube volume fractions than for HDI-based polyurethane. It is noteworthy that for the series of HDI-based polyurethanes with varied hard segment content the percolation threshold effects affecting both ductility and modulus enhancement scaled up with polyurethanes crystallinity (increase in hard segment content). MDI-30 has a lower degree of crystallinity than HDI-30 but the effects on ductility were perceived at higher nanotube concentrations than for HDI-30.

Contrarily, for MDI-30 systems the strength reached a maximum around the percolation threshold concentration and trend to decrease above it. This fact could be related to the lower surface tension that MDI-based polyurethanes present [33] leading to favoured but weak nanotube-nanotube interactions. Despite having MDI/BD hard segment higher solubility parameter than the HDI-BD hard segments (Table 2) and its composites higher interfacial shear strength (Table 3), the un-match between the materials overall surface tension with that of the nanotubes could be the determinant factor influencing the loss in reinforcement at high nanotube loadings.

The higher observed absolute value of reinforcement, dσ/dϕ, listed in Table 3 for MDI-30 composites could be related to a stronger interaction between aromatic MDI/BD hard segment with nanotubes surface via π-π stacking in addition to common
polar and hydrogen bond interactions, as comparison with the aliphatic HDI/BD hard segments.

4. Conclusions

Control of the complexity of PU chemistry and physical properties of their derived composites may lead to new high performance hierarchically assembled composites. Addition of nanosized fillers with outstanding properties, such as MWCNT could be regarded as an interesting approach for the purpose of creating new elastomeric materials with enhanced mechanical and electrical properties.

The reduction of ductility with MWCNT addition could be related to the relative MWCNT big sizes in comparison to PU hard domains, being, as demonstrated also by other groups, an important factor to target polyurethanes hard domains in order to obtain improvement on mechanical properties without a consequent reduction in ductility and toughness, therefore resembling the performance of materials such as natural silk.

5. Acknowledges

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References


FIG. CAPTIONS

Fig. 1. MWCNT/PU Percolative network detection and DC electrical conductivity. (■) Data obtained with solvent cast technique produced composites. (▲) Data obtained for buckypapers. The green line represents the general percolative model.

Fig. 2. Mechanical properties of composites: a) Strain at break, b) normalised strength and c) normalised elastic modulus of composites prepared with HDI polyurethanes as a function of hard segment and nanotube concentration.

Fig. 3. a) Extensibility as a function of hard segment content and nanotube weight fraction, b) Modulus enhancement as a function of polyurethane hard segment. c) Normalised modulus enhancement as a function of polyurethane hard segment content. d) Interfacial shear strength as calculated using a Bowyer-Bader model for aligned short fibres, as a function of polyurethane hard segment content.

Fig. 4. a) Model of a polyurethane network formed by hard domains (yellow) interconnected by the soft flexible chains (brown ropes), non stretched and b) stretched along the blue row direction. c) Long nanotubes inclusion interacting with hard segments at room temperatures may reduce the ductility of soft segments, as depicted in (d).

Fig. 5. Scanning electron microscopy analysis of fracture surfaces for composites with 20 wt% MWCNT and different HS contents: a, b) HDI-10; c, d) HDI-30 and e, f) HDI-50. Scale bars in first and second column are of 10 μm and 1 μm, respectively.

Fig. 6. DSC heating scans of composites with the indicated weight fraction (wt%) of MWCNT. a) HDI-10, b) HDI-30 and c) HDI-50 as composites matrices.

Fig. 7. a) Non-isothermal crystallisation scans of HDI-50 composites with the indicated carbon nanotube content. b) Modified Avrami plots for a non-isothermal crystallisation of HDI-50 and a HDI-50 nanocomposite with 3 wt% of MWCNT, at 2.5 °C min⁻¹ cooling rate. c) Sketch of the nucleating effect of nanotubes and crystallinity reduction.

Fig. 8. Ternary composition diagram of solvent cast and infiltrated buckypapers composites. Mechanical properties of composites are summarised as function of carbon nanotubes, hard segment, and soft segment weight fractions. Values of composites elastic modulus, tensile strength and elongation to break along values of density and DC conductivity for some materials are included.

Fig. 9. Mechanical properties versus nanotube loading on polyurethanes based on HDI (▲) and MDI (○). The dotted blue and red lines corresponds to the Bowyder-Bader model prediction for the HDI-30 and MDI-30 polyurethanes based composites respectively.
TABLES

Table 1. Summary of prepared matrices chemical composition and reactants molar ratio.

<table>
<thead>
<tr>
<th>Abbreviations</th>
<th>Isocyanate</th>
<th>Wt% Hard segment</th>
<th>Polyl: BD: Diisocyanate</th>
</tr>
</thead>
<tbody>
<tr>
<td>HDI-10</td>
<td>HDI</td>
<td>10</td>
<td>4:1:5</td>
</tr>
<tr>
<td>HDI-17</td>
<td>HDI</td>
<td>17</td>
<td>1:1:2</td>
</tr>
<tr>
<td>HDI-30</td>
<td>HDI</td>
<td>30</td>
<td>1:3:4</td>
</tr>
<tr>
<td>HDI-40</td>
<td>HDI</td>
<td>40</td>
<td>2:9:11</td>
</tr>
<tr>
<td>HDI-50</td>
<td>HDI</td>
<td>50</td>
<td>1:7:8</td>
</tr>
<tr>
<td>MDI-30</td>
<td>MDI</td>
<td>30</td>
<td>1:2:3</td>
</tr>
</tbody>
</table>

Table 2. Hildebrand solubility parameters of MWCNT-COOH along those calculated for the polyurethanes segments.

<table>
<thead>
<tr>
<th>Material</th>
<th>ψ (J^1/2 cm^3/2)</th>
</tr>
</thead>
<tbody>
<tr>
<td>MWCNT-COOH</td>
<td>≥ 25°</td>
</tr>
<tr>
<td>- (HDI-BD)-</td>
<td>23.0°</td>
</tr>
<tr>
<td>- (MDI-BD)-</td>
<td>25.3°</td>
</tr>
<tr>
<td>PCL-b-PHMC-b-PCL</td>
<td>20.9°</td>
</tr>
</tbody>
</table>

°Assumed to be higher than DMF (which is a good pristine nanotubes solvent), for the introduction of polar groups upon acid treatment.

°°Calculated using Feedors molar volumes and Hoftyzer-Van Krevelen method.
Table 3. Reinforcing parameters and surface tension values of MDI-30 and HDI-30 matrices loaded with multiwalled carbon nanotubes

<table>
<thead>
<tr>
<th>Matrix</th>
<th>$\frac{dE}{d\phi}$ (MPa)</th>
<th>$(\frac{dE}{d\phi})E_m^{-1}$</th>
<th>$\frac{d\sigma}{d\phi}$ (MPa)</th>
<th>$(\frac{d\sigma}{d\phi})\sigma_m^{-1}$</th>
<th>$\tau$ (MPa)</th>
<th>$\gamma$ (mJ m$^{-2}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>MDI-30</td>
<td>842 ± 62</td>
<td>80.2</td>
<td>289 ± 61</td>
<td>12.2</td>
<td>5.0</td>
<td>13.8</td>
</tr>
<tr>
<td>HDI-30</td>
<td>1068 ± 74</td>
<td>10.3</td>
<td>221 ± 16</td>
<td>14.3</td>
<td>3.8</td>
<td>42.6</td>
</tr>
</tbody>
</table>
FIGURES

Fig. 1
Fig. 2

(a) Plot of $\varepsilon_{\text{max,c}}$ (\%) vs wt%.
(b) Plot of $\sigma_c/\sigma_m$ vs $\phi$.
(c) Plot of $E_c/E_m$ vs $\phi$. 

Legend:
- HDI-10
- HDI-14
- HDI-30
- HDI-40
- HDI-50
Fig. 3

(a) Graph showing the ratio of maximum strain \(\varepsilon_{\text{max}} / \varepsilon_{\text{max,PU}}\) as a function of hard segment content (wt%).

(b) Graph showing the change in tensile strength \(dE/d\gamma\) with hard segment content (wt%).

(c) Graph showing the change in (dE/d\gamma)\textsuperscript{-1} with hard segment content (wt%).

(d) Graph showing the change in elongation at break \(\varepsilon_\text{b}\) with hard segment content (wt%).

Fig. 4

(a) Image showing the distribution of hard segments at 0.25 wt%.

(b) Image showing the distribution of hard segments at 0.5 wt%.

(c) Image showing the distribution of hard segments at 1.5 wt%.

(d) Image showing the distribution of hard segments at 3 wt%. 

(e) Image showing the distribution of hard segments at 6 wt%.

(f) Image showing the distribution of hard segments at 12 wt%.
Fig. 5
Fig. 7

(a) Heat flow vs. temperature for different wt% of HDI-50. 

(b) Plot of log(-\ln(1-X_c)) vs. log(t) for HDI-50 and HDI-50 + 3 wt% COOH.

(c) Schematic representation of a segmented polymer with soft and hard segments.
Fig. 8

Fig. 9