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Discontinuity of physical properties of carbon nanotube/polymer composites at the percolation threshold

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ABSTRACT

Evolution of DC conductivity with double-walled carbon nanotube concentration in a polyamide 11 matrix displays the electrical percolation phenomenon. The percolation threshold is observed for this composite at 0.85 wt.%. Thermal, mechanical and dielectric manifestations of the glass transition have been investigated as a function of carbon nanotube content. Thermal and mechanical analyses have shown a depression of the glass transition temperature or its mechanical manifestation at the percolation threshold. Regarding the dielectric modulus, three distinct behaviors (below, at, and above percolation threshold) have been observed. A discussion about the interactions between the polymer matrix and the conductive fillers will be proposed to further the understanding of the discontinuity of physical properties at the percolation threshold.

Keywords: Carbon nanotubes; Polymer-matrix composites; Electrical percolation threshold; Glass temperature depression

1. Introduction

In the past decades, introducing fillers in a polymer matrix has been a promising challenge to obtain multifunctional materials. Several routes have been tried for obtaining a conductive behavior: the introduction of conductive polymers such as polypyrrole [1,2] or conductive particles such as carbon black [3,4], carbon nanofibers [5], carbon nanotubes [6–10], graphene [11], metallic particles [12], and metallic nanowires [13,14]. Polymer based nanocomposites offer the advantage to keep the mechanical properties of the matrix while acquiring interesting conductive properties. With the highest aspect ratio, carbon nanotubes are particularly attractive. Discovered by Iijima [15], CNTs enable the dissipation of electrostatic charges for antistatic applications [16] at a very low filler concentration. Indeed, the insulator-conductor transition, also called percolation threshold [17], can be reached with this type of particle in a semi-crystalline polymer matrix with less than 1% in weight [18]. This percentage depends mainly on polymer matrix, particle dispersion state and particle apparent aspect ratio [19]. A lot of work has been devoted to the evolution of the glass transition temperature (Tg) with filler content. Ash has shown a 20 °C decrease of T_g in poly(methylmethacrylate)/alumina composites [20]. This decrease disappears when the particles are coated, thus making them compatible with the matrix. The influence of interactions between

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particle and matrix has been confirmed by Sun [21] and Rittigstein and Torkelson [22]. Saiter has highlighted a confinement effect of polymer chains in the galleries of clay (intercalated nanocomposite) inducing a strong reduction of the cooperativity and of the glass transition temperature of the polymer matrix [23]. By varying on the interactions between different particles and polymers, they have shown that $T_{\rm g}$ can increase, stay stable or decrease with particle concentration. An increase of T_g values has also been pointed out in other polymer based composites [24–26]. All these authors agreed on the origin of the T_g fluctuations: strong interactions between particles and polymer matrix result in a decrease of polymer chain mobility and an increase of glass transition temperature. In contrast, weak interactions enable an increase of free volume accessible by the polymer chains which lead to a decrease of composite $T_{\rm g}$ values. Only several authors have highlighted a depression (decrease followed by an increase) in their glass transition temperature values. It has only been observed with conductive fillers. Barrau et al. [27] examined on the characterization of glass transition temperature evolution in polyepoxy/polypyrrole and polyepoxy/CNTs composites by using thermal and mechanical analyses. A depression of T_g has been observed in both systems. T_g depression is larger for polypyrrole particles: 15 °C instead of 5 °C for composites with CNTs. A 5 °C depression was also observed in a composite of polycarbonate/CNTs at the percolation threshold [28]. This phenomenon is still misunderstood; however some explanations have been put forward. The influence of fillers on polymer chain mobility can explain a decrease or an increase in Tg but considering the shape

of the $T_{\rm g}$ depression (a decrease follows by an increase) at the electrical percolation threshold, further investigation is required to gain a full understanding of this phenomenon.

A previous work [18] describes the introduction of double-walled carbon nanotubes (DWCNTs) in a semi-crystalline thermoplastic polymer, polyamide 11, by solvent way and by an extrusion process. This polymer matrix has been chosen for its good mechanical properties and its low hydrophilicity (2–3% of water absorption) which implies a good stability of the physical properties whatever environmental conditions. The dispersion of the conductive nanofillers have been optimized following the electrical conductivity and a satisfactory dispersion has been reached since the value of the electrical percolation threshold has been found as one of the lowest for a semi-crystalline thermoplastic matrix.

In this work, three sets of polyamide 11/DWCNT nanocomposites are studied and the evolution of their glass transition temperature with DWCNT content is followed through thermal measurements. These experiments are completed by dynamic mechanical analysis led on one set of nanocomposite; the evolution of the dielectric modulus with DWCNT concentration is also examined. The influence of the conductive fillers on the polymer physical properties will be explored.

2. Experimental

2.1. Materials and processing

The Polyamide 11 (PA 11) polymer matrix was provided either in granules or in a powder (30 µm) by Arkema (France). The carbon nanotubes were supplied by Marion Technologies (MT) (France) and by the CIRIMAT (Université Paul Sabatier — France). They were synthesized by catalytic chemical vapor deposition and are both mainly double-walled (around 80%) [29]. Double-walled carbon nanotubes (DWCNTs) are of key interest because they offer the possibility of functionalizing the outer wall which may ensure the connection with the external environment while retaining the remarkable mechanical and electronic properties of the inner nanotube with the lowest density as possible. For this study, both DWCNTs are used as received without any functionalisation.

Two methods of composite elaboration have been used: the solvent and the extrusion way. For the first one, PA 11 granules are first added in dimethylacetamide (DMAc), previously heated at 160 °C. DWCNTs (from Marion Technologies), dispersed by ultrasound in DMAc, were added in the PA 11-DMAc blend. After precipitation, the mixture is filtrated, water washed and vacuum dried at 160 °C during 12 h to eliminate all solvent residues.

For the second one, a twin screw extruder (Minilab II from Haake) was chosen for dispersing DWCNTs (either from MT or CIRIMAT) in the semi-crystalline thermoplastic matrix. Extrusion parameters were previously optimized to ensure the best dispersion [18] and chosen as follow: 30 rpm as screw rotation speed, 210 °C for the mixing temperature (melting temperature of PA 11 + 20 °C [30]), a mixing time of 15 min with screws in a co-rotating configuration. In both cases, nanocomposites were hot pressed at 210 °C in order to realize desired shape for different characterization techniques.

Three sets of composites were then manufactured: PA 11/DWCNT-MT by solvent way (loaded at 0.1, 0.3, 0.7, 1, 1.3, 1.6, 1.8, 2, 3 and 4 wt.%), PA 11/DWCNT-MT by extrusion (loaded at 0.1, 0.3, 0.7, 0.85, 1, 1.3, 1.6, 2, and 4) and PA 11/DWCNT-CIRIMAT (loaded at 0.3, 0.7, 1, 1.3, 1.6, 2 and 4 wt.%).

For each characterization techniques, samples have been manufactured in an appropriate shape and conserved 24 h at ambient temperature. This step allows a physical aging since this last is promoted between $T_{\rm g}$ (which is comprised between 40 and 50 °C for PA 11) and $T_{\rm g}$ –50 °C. It corresponds, in polyamide, to the establishment of a dense hydrogen bonded network. This step is necessary to promote the heat capacity step. This step will be discussed later.

2.2. Standard differential scanning calorimetry (DSC)

Standard DSC measurements were performed using a differential scanning calorimeter (Perkin-Elmer DSC7). The heating rate was 20 °C·min $^{-1}$ from 20 °C to 210 °C. Sample weight was in the range of 5 to 10 mg. The glass transition temperature of samples with identical thermal history was defined by the midpoint of the heat capacity step.

2.3. Dynamic mechanical analysis (DMA)

DMA experiments were carried out by a Rheometrics ARES Scientific strain-controlled rheometer in the torsion rectangular mode. The measurements were performed in the linear viscoelastic range of composites at a fixed angular frequency (1 $\text{rad}\cdot\text{s}^{-1}$) from -150 to 170 °C (at 3 °C·min $^{-1}$). Sample thickness was 1 mm. The dynamic dissipative modulus G'' was plotted as a function of temperature. The anelastic manifestation of the glass transition temperature T_{α} was measured at the maximum of the G'' peaks.

2.4. Dynamic dielectric spectroscopy (DDS)

DDS measurements were carried out using a Novocontrol broadband spectrometer. The composites were manufactured with a thickness large enough (between 1 and 1.5 mm) for avoiding the alignment of CNTs in the pellet plan. By this way, a homogeneous dispersion of nanofillers in the three space directions was promoted. The measurements were done in the frequency range from 10^{-2} to 10^6 Hz on aged samples. The complex dielectric permittivity was measured in the temperature range $-150\,^{\circ}\text{C}/+150\,^{\circ}\text{C}$, by steps of 2 °C. The complex dielectric permittivity was deduced from impedance measurements. In order to analyze high temperature relaxations, the complex dielectric modulus formalism (M* = ϵ^{*-1}) is required [31]. For conductivity measurements, the dynamic complex conductivity σ^{*} is deduced from the complex permittivity ϵ^{*} :

$$\sigma^* = i\omega \epsilon_o \epsilon^* \tag{1}$$

where ω is the angular frequency, and ε_0 is the permittivity of vacuum.

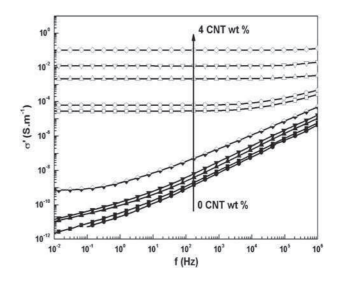


Fig. 1. Frequency dependence of the real part of the complex conductivity, at 20 °C, for PA 11/CNT nanocomposites with different CNT contents: 0 wt.% (●), 0.1 wt.% (■), 0.3 wt.% (△), 0.7 wt.% (▼), 0.85 wt.% (\diamondsuit), 1 wt.% (\square), 1.3 wt.% (\bigcirc), 1.6 wt.% (\triangle), 2 wt.% (∇) and 4 wt.% ().

3. Results and discussion

3.1. Isothermal electrical conductivity

The isothermal electrical conductivity of PA 11/DWCNT-MT nanocomposites elaborated by extrusion is investigated. Fig. 1 shows the frequency dependence of the real part, $\sigma'(\omega)$, of the complex electrical conductivity σ^* , for different DWCNT contents, measured at 25 °C. Regarding the neat polymer matrix, its conductivity increases with frequency. At 10^{-2} Hz, the electrical conductivity is around 10^{-11} S·m⁻¹ while it reaches $10^{-6} \,\mathrm{S} \cdot \mathrm{m}^{-1}$ at $10^6 \,\mathrm{Hz}$. This dependence is characteristic of an insulating material. Composites with 0.1, 0.3 and 0.7 wt.% of CNTs follow the same behavior. From composites loaded at 0.85 wt.%, $\sigma'(\omega)$ becomes independent of frequency at the lower frequencies of measurement. Above a characteristic onset angular frequency ω_c , the dynamic conductivity follows a power law. This type of behavior was observed in a wide series of disordered solids; it was investigated by Jonscher [32] and later on by Dyre [33]. Consequently, it was designated as the "Universal Dielectric Response" (UDR). It is commonly described by the following equation:

$$\sigma'(\omega) = \sigma(0) + \sigma_{AC}(\omega) = \sigma_{DC} + A\omega^{s}$$
 (2)

where σ_{DC} is the static conductivity (when ω approaches 0, σ' is identified as the dc conductivity), A and s with 0 < s < 1 are only temperature dependent. Eq. (2) can also be written:

$$\sigma'(\omega) = \sigma_{DC} \left[1 + \left(\frac{\omega}{\omega_c} \right)^s \right] \tag{3}$$

with ω_c , the critical frequency.

When CNT content increases from 0.85 to 4 wt.%, the dc conductivity σ_{DC} increases as well as the frequency independent region becomes broader (i.e. ω_c increases). Then, σ_{DC} reaches $10^{-1}~\rm S\cdot m^{-1}$ i.e. 10^{11} times higher than the value for neat PA 11. This step is indicative of the establishment of percolation paths. The electronic structure of CNTs [34–36] brings charge carriers that move between localized states of different energy levels accessible by hopping [37] or tunneling [38].

3.2. Glass transition temperature

DSC measurements have been carried out on PA 11/DWCNT-MT nanocomposites elaborated by extrusion. The corresponding DSC thermograms obtained at the first run on aged samples are depicted

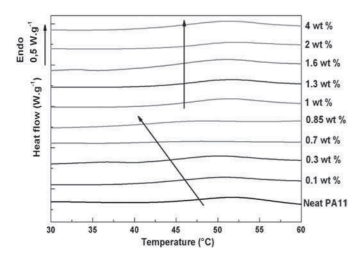


Fig. 2. DSC thermograms of neat PA 11 and PA 11/DWCNT-MT nanocomposites prepared by extrusion loaded at 0.1, 0.3, 0.7, 0.85, 1, 1.3, 1.6, 2 and 4 wt.%. Arrows show the evolution of the glass transition temperature.

Fig. 2. In the temperature range 30–60 °C, a heat capacity step is pointed out. It is associated with the glass transition of the amorphous phase of polyamide 11. A weak endothermic peak is superimposed to the heat capacity step and helps to its analysis. It is associated with a breaking of physical bond characteristics of physical aging. The heat capacity step decreases in temperature when DWCNT content increases up to 0.85 wt.%. For the nanocomposite loaded at 1 wt.%, this temperature increases and is equivalent to the neat PA 11 one. It stays stable for higher DWCNT concentration. The DSC thermograms obtained at the second run are not displayed here since the endothermic peak is not observable anymore. Between the first and the second run, the kinetic is too fast to allow the complete rebuilding of previous physical bonds. Consequently, the heat capacity step becomes unexploitable.

The evolution of the glass transition temperature (T_g) of PA 11/DWCNT-MT composites, as a function of DWCNT weight percent, are plotted in Fig. 3. The σ_{DC} conductivity corresponding to the σ' values taken at 10^{-2} Hz (Fig. 1) are also indicated as reference. For PA 11/DWCNT-MT elaborated by extrusion, the percolation threshold value was found at 0.93 wt.% of DWCNT.

The $T_{\rm g}$ value for neat PA 11 is around 46.5 °C and decreases to 42 °C for composite with 0.7 wt.% of DWCNTs. Tg stays low for composite till DWCNT contents of 0.85 wt.%; T_g value increases (to 45-46 °C) for composites with CNT contents higher than 1 wt.%. A sharp depression is observed around 0.85 wt.%, corresponding to the percentage at which the percolating path is created. At the electrical percolation threshold, glass transition temperature of the PA 11/DWCNT composites drops by 4.5 °C. This change is significant since the precision of measurement on T_g is $\pm 1^\circ$. The same temperature depression has been observed in polyepoxy [27] and in polycarbonate [28]. Neither the nature of the matrix (thermoset or thermoplastic), nor its chemistry (polyamide or polycarbonate) looks to have an influence on the depression value. Dispersed DWCNTs modify the glassy state physical structure until the creation of the percolation path. It appears as a decrease of the glass transition temperature. Such a decrease vanishes upon the establishment of the conductive path.

For evacuating an experimental artifact and confirming the previous result i.e. a depression of the glass transition temperature at the electrical percolation threshold, two other sets of composites have been studied: PA 11/DWCNT-MT elaborated by solvent way and PA 11/DWCNT-CIRIMAT elaborated by extrusion. For both, the DC conductivity was extracted from the real part of the complex electrical conductivity $\sigma'\left(\omega\right)$ measured at 25 °C and taken at 10^{-2} Hz.

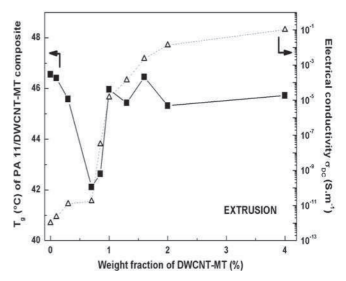


Fig. 3. Variation of T_g obtained by DSC for PA 11/DWCNT-MT nanocomposites prepared by extrusion as a function of CNT weight content (filled square and solid line). The variation of σ_{DC} extracted from σ' at 10^{-2} Hz is indicated as reference (open triangles and dashed line).

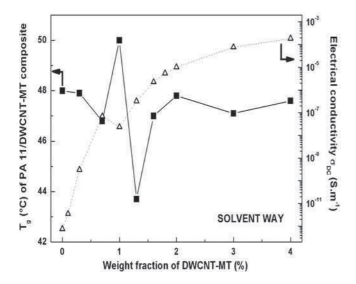


Fig. 4. Variation of T_g obtained by DSC for PA 11/DWCNT-MT nanocomposites prepared by solvent way as a function of CNT weight content (filled square and solid line). The variation of σ_{DC} extracted from σ' at 10^{-2} Hz is indicated as reference (open triangles and dashed line).

Fig. 4 shows the evolution of the DC conductivity and the glass transition temperature as a function of DWCNT content for PA 11/DWCNT-MT obtained by solvent way. Regarding the electrical conductivity, the eight decade gap between insulator $(10^{-12}~\rm S\cdot m^{-1})$ –conductor $(10^{-4}~\rm S\cdot m^{-1})$ behaviors highlights the electrical percolation phenomenon. It appears at 1 wt.% of DWCNTs. However, the conductor path is established over a wide DWCNT percentage range i.e. the increase of the conductivity is not abrupt as it is in composites elaborated by extrusion. This is typical of a heterogeneous dispersion where the percolation path is created between CNT clusters and not between CNT themselves. The glass transition temperature of neat PA 11 is equal to 48 °C. It is 1.5 °C above the one represented in Fig. 3. It comes from that granules of PA 11 have been used instead of PA 11 powder, employed for both other sets of composite.

The T_g shows a depression for the composite loaded at 1.3 wt.% of DWCNT. This decrease is around 4.5 °C compared to T_g of neat PA 11. This result is of the same order of magnitude than the depression observed for PA 11/DWCNT-MT elaborated by extrusion confirming the existence of a T_g depression at the electrical percolation threshold. It has to be noticed that the composite loaded at 1 wt.% is characterized by a T_g 2 °C higher than neat PA 11. At this concentration, the DC conductivity is lower than the one of the composite loaded at a lower DWCNT content (i.e. 0.7 wt.%) which reflects a heterogeneous dispersion of carbon nanotubes as mentioned above. This observation testifies that the T_g value of PA 11/DWCNT is in close relationship with the carbon nanotube space arrangement in the vicinity of the percolation threshold.

PA 11/DWCNT-CIRIMAT nanocomposites elaborated by extrusion have been analyzed by DDS and DSC. The evolution of both DC conductivity and glass transition temperature as a function of DWCNT content is pointed out Fig. 5. The electrical conductivity shows an abrupt increase from 10⁻¹² S·m⁻¹ to 10⁻¹ S·m⁻¹ which is typical of the percolation phenomenon. The percolation threshold is evaluated at 0.6 wt.% of DWCNT. A decrease of the percolation threshold value from 0.93 wt.% (PA 11/DWCNT-MT) to 0.6 wt.% (PA 11/DWCNT-CIRIMAT) is observed depending on the origin of the double-walled carbon nanotubes. Since the same manufacturing process was used for both sets of nanocomposites, the filler dispersion state is considered as the same. A general relationship between the percolation threshold of various objects and their associated excluded volume has been established by Balberg et al. [19]. In the

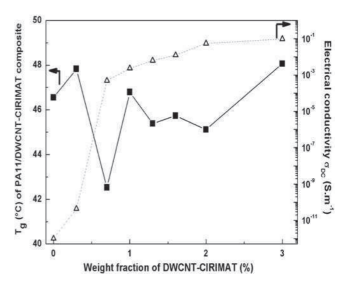


Fig. 5. Variation of T_g obtained by DSC for aged PA 11/DWCNT-CIRIMAT nanocomposites prepared by extrusion as a function of CNT weight content (filled square and solid line). The variation of σ_{DC} extracted from σ' at 10^{-2} Hz is indicated as reference (open triangles and dashed line).

isotropic case of randomly distributed long sticks (length L and radius r) the proposed relation is:

$$(L/r) f_c \approx 3$$
 (4)

where L/r is the aspect ratio and f_c, the critical volume fraction of the sticks at the percolation threshold. According to Balberg's work, the decrease of the electrical percolation threshold value in case of PA 11/DWCNT composites is attributed to an increase of the DWCNT aspect ratio. For the same CCVD process, CIRIMAT double-walled carbon nanotubes possess a higher aspect ratio than DWCNT from Marion Technologies. DWCNTs from CIRIMAT are manufactured at a laboratory scale while DWCNTs from MT are industrially produced.

The PA 11/DWCNT-CIRIMAT composite loaded at 0.7 wt.% shows a $T_{\rm g}$ value of 42.5 °C which is 4 °C lower than neat PA 11 one's. This composite is situated at the electrical percolation threshold. This result confirms experimentally both the previous ones.

The DWCNT percentage corresponding to the T_g depression shifts to lower values at the same time than the value of the electrical percolation threshold meaning that the observed depression is directly linked to the establishment of the conductive path. Furthermore, the percolation phenomenon is associated with a specific arrangement of the conductive fillers in the matrix. In parallel, a decrease of the glass transition is attributed to a plasticization of the polymer macromolecules. Consequently, the specific arrangement of the double-walled carbon nanotubes at the percolation threshold induces an increase of the molecular mobility by a rise of the free volume accessible by the polymer chain segments. It results in a decrease of the glass transition temperature at the electrical percolation threshold. Once the percolating path is created, the homogeneous dispersion of the conductive fillers does not disturb anymore the polymer chain segment mobility and the neat polymer T_g recovers its original value.

DSC experiments have been completed by DMA and DDS analysis led on PA 11/DWCNT-MT composites to follow the evolution of the mechanical and the dielectric manifestation of the glass transition with the DWCNT content.

3.3. Dynamic mechanical relaxation

In order to confirm the results obtained by DSC on PA 11/DWCNT-MT composites, the mechanical manifestation of the glass transition

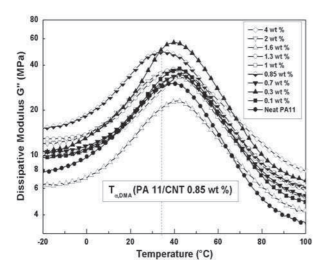


Fig. 6. Dissipative part of the shear modulus *G*" as a function of temperature of neat PA 11 and PA 11/DWCNT-MT nanocomposites prepared by extrusion loaded at 0.1, 0.3, 0.7, 0.85, 1.1.3, 1.6, 2 and 4 wt.%

was evaluated using DMA. The dissipative part of the shear modulus G'' of neat PA 11 and PA 11/DWCNT-MT composites was recorded during the first temperature scan (aged samples). It is depicted in Fig. 6. The dissipative modulus of neat PA 11 as well as composites highlights a peak which is associated with the anelastic manifestation of the glass transition. In this temperature range, the polymer shifts from a glassy to a rubbery behavior. The temperature characteristic of the peak's maximum is called $T_{\alpha, DMA}$ and remains constant for all DWCNT concentration excepted for the composite loaded at 0.85 wt.%.

Fig. 7 reports the evolution of the anelastic manifestation of the glass transition temperature ($T_{\alpha,DMA}$) obtained on aged sample as a function of weight fraction of DWCNTs introduced in the polymer matrix.

A sharp depression of 5 °C occurs for the composite situated at the electrical percolation threshold i.e. 0.85 wt.% of CNTs, confirming results obtained by DSC. $T_{\alpha,DMA}$ is quite stable for other composites (around 40 °C). Compared to DSC, no depression is observed for the composite loaded at 0.7 wt.%. Regarding DMA and DSC experiments, the temperature range in which the α relaxation on G modulus occurs is

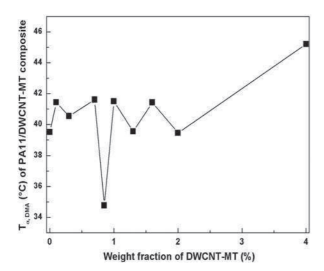


Fig. 7. Variation of the anelastic manifestation of the glass transition T_{α} obtained by DMA for PA 11/DWCNT-MT nanocomposites prepared by extrusion as a function of CNT weight content.

larger than the heat capacity step one. Consequently, it is more difficult to evaluate a sharp change in temperature on $T_{\alpha M}$ due to the DWCNT presence than on T_{α} .

The depression of $T_{\alpha,DMA}$ is only observed during the first scan, i.e. on aged sample. The DMA thermograms obtained at the second run are not shown here since $T_{\alpha,DMA}$ stays constant whatever DWCNT content.

In polyamides, physical aging is due to the establishment of a dense hydrogen bonded network which occurs between $T_{\rm g}$ and $T_{\rm g}$ -50 °C i.e. at room temperature. The first DMA run has been obtained on aged samples. Between the first and the second run, the kinetic is too fast to allow the whole rebuilding of previous physical bonds. This leads to conclude that the specific arrangement of double-walled carbon nanotubes at the percolation threshold acts on the hydrogen bonded network formation constituted upon physical aging.

It is interesting to note that the same temperature depression of T_{α} is observed with DWCNTs dispersed in amorphous matrix like epoxy [25]. Furthermore, for conductive fillers with a different morphology (polypyrrole i.e. different aspect ratio and conductivity) introduced in the same matrix, a three times higher depression has been recorded. The characteristics of the conductive fillers might explain the magnitude of this thermal phenomenon at the electrical percolation threshold.

The same phenomenon of T_g depression (a decrease followed by an increase) upon percolation phenomena has been observed in inorganic composites [39]. The last system is very different since it is constituted by germanium telluride glasses containing silver particles (always with conductive particles). A 15 °C depression was observed for composite with 5% of silver particles, corresponding to their percolation threshold.

An abrupt increase followed by a decrease of the mechanical compliance appears in Alumina/Nickel composites [40]. A sliding effect between the amorphous ceramic matrix and the metallic particles at their percolation threshold seems to be responsible for this increase. An abrupt variation (sharp depression) in effective magnetic permeability has also been revealed in butadiene-nitrile rubber/iron composites [41]. Large conducting clusters, observable just below the percolation threshold, undergo eddy current due to the variation of the magnetic field. Eddy currents cause the appearance of a Joule effect in the composite. Magnetic moments are affected as well as the effective magnetic permeability. These examples show that several phenomena at the electrical percolation threshold are still misunderstood. Establishment of the electrical percolating path is the common factor in these abrupt variations. It shows that a

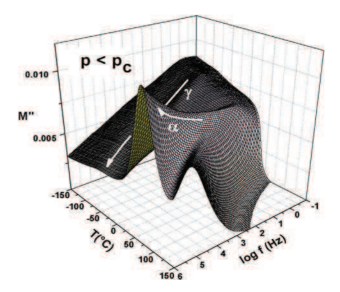


Fig. 8. 3D relaxation map of the dielectric modulus obtained by DDS for PA 11/DWCNT nanocomposite prepared by extrusion with 0.3 wt.% of CNTs.

particular interest has to be devoted to the mobility and interactions of charge carriers with the matrix.

3.4. Dynamic electrical relaxation

3.4.1. 3D electrical relaxation map

Fig. 8 shows the evolution of the imaginary part of the dielectric modulus as a function of temperature and frequency for a DWCNT concentration lower than $p_c\colon 0.3$ wt.%. The relaxation map remains analogous for DWCNT concentration $p < p_c$. The γ relaxation involves a mobility of the methylene sequences between amide groups. The α relaxation mode (named α ,DDS) at $T_{\alpha,DDS}$ is the dielectric manifestation of the glass transition. Above this relaxation, two peaks appear: the relaxation of the constrained amorphous phase is hidden by a Maxwell–Wagners–Sillars mode [14] inherent to the relaxation of free carriers at the interface of heterogeneities in the material and finally, conductivity.

The conductive composite behavior is different (not depicted here). Above the electrical percolation threshold, a percolating path is created. The charge relaxation is the most prominent phenomenon and hides dipolar relaxations. It appears as an increase of modulus at high frequencies on the 3D relaxation map.

3.4.2. Isothermal electrical modulus

Fig. 9 exhibits the evolution of the dielectric modulus at 25 °C for each DWCNT concentration and is extracted from the 3D relaxation map. One peak appears for insulator composites: it corresponds to the dielectric manifestation of the α mode. This peak shifts to higher frequencies when DWCNT concentration increases. The amorphous phase is stiffened by the presence of DWCNTs. For conductive composites, one peak appears as well but at higher frequencies, it is associated with the conductivity phenomenon and highlights a change in the electrical behavior of the composite. This peak also shifts to higher frequencies with DWCNT concentration. The frequency at which the peak appears depends on the mean distance between conductive clusters (which decreases when DWCNT content decreases), the charge carrier mobility and the temperature. DC conductivity is possible at low frequency until the critical frequency ω_c (the same as discussed in Fig. 1) is reached. Above this frequency, the period between two successive changes in the electrical field sign is too short to allow charge carrier transport between conductive fillers. At the percolation threshold, i.e. composite loaded at 0.85 wt.%, the electrical modulus shows a particular behavior resulting from the interaction between dipolar relaxation and conductivity.

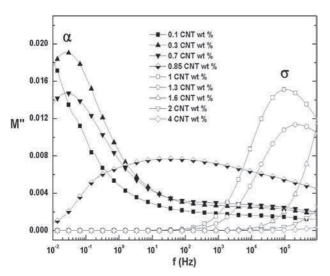


Fig. 9. Variation of the electrical imaginary modulus at 25 $^\circ$ C, as a function of frequency for PA 11/DWCNT nanocomposites prepared by extrusion with different CNTs contents.

3.4.3. Dielectric relaxation times

For insulator composites (0.3 and 0.7 DWCNT wt.%) as well as the 0.85 DWCNT wt.% sample, relaxation times have been extracted from the 3D relaxation maps displayed above. For the fitting of each isotherm, the determination of the relaxation time $\tau_{\rm M}$ is performed using the Havriliak–Negami equation:

$$M^* = M_{\infty} - \frac{M_{\infty} - M_0}{[1 + (i\omega\tau_M)^{\alpha}]^{\beta}} \tag{5} \label{eq:5}$$

where M_0 and M_∞ are the electrical modulus at null and infinite frequency respectively and the parameters α and β are such that: $0<\alpha<1$ and $0<\beta<1.$

Fig. 10 displays the Arrhenius diagrams of the α relaxation mode. It has been used to compare qualitatively the influence of DWCNTs on the molecular mobility of polymer chains.

Relaxation times are well described by a Vogel–Fulcher–Tammann equation. This behavior is characteristic of the α relaxation associated with the glass transition. The relaxation kinetic decreases as soon as 0.1 wt.% of CNTs are dispersed in the polymeric matrix. The relaxation temperature associated with the dielectric manifestation of the glass transition decreases with the introduction of carbon nanotubes which is in total agreement with DSC and DMA observations below the percolation threshold i.e. the corresponding shift of the relaxation peak is of the same order of magnitude than the one of T_g and $T_{\alpha,DMA}$. Relaxation temperatures slightly decrease between the various CNT contents from 0.1 to 0.85 wt.%. It is coherent with a plasticization phenomenon for which the macromolecule mobility is modified by the presence of carbon nanotubes. For higher DWCNT concentrations, the nanocomposites become conductor and cannot be further studied by dynamic dielectric spectroscopy.

4. Conclusion

Thermal, mechanical and electrical analyses have been combined to characterize the percolation phenomenon in PA 11/CNTs composites. The originality of this work is to study a phenomenon related to changes in the polymer physical structure linked to the establishment of a percolating path with conductive fillers by different experimental devices. The conductivity of PA 11/CNT composites follows the "Universal Dynamic Response" and these composites are characterized by an electrical percolation threshold at 0.85 wt.% of CNTs. A sharp glass transition temperature depression (5 °C) has been observed by DSC and DMA around this insulator-to-conductor transition. It has been

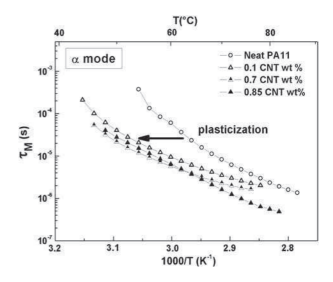


Fig. 10. Arrhenius diagram of the α relaxation mode for PA 11/DWCNT nanocomposites prepared by extrusion with different CNT contents, below the percolation threshold.

confirmed by studying two other sets of PA 11/DWCNT nanocomposites which both undergo a depression of this thermal transition at the percolation threshold. Analogous phenomena are observed in other polymer/CNTs composites. It is important to note here that this depression is only observed after physical aging. Consequently, in the PA 11/DWCNT composite, the specific arrangement of the double-walled carbon nanotubes at the electrical percolation threshold induces an increase of the hydrogen bonded network molecular mobility by a rise of the free volume accessible by the polymer chain segments. It results in a decrease of the glass transition temperature at the electrical percolation threshold. Once the percolating path is created, the homogeneous dispersion of the conductive fillers does not disturb anymore the polymer chain segment mobility and the neat polymer $T_{\rm g}$ recovers its original value.

Dynamic dielectric spectroscopy shows three types of behavior for the loss modulus. Below the percolation threshold, the relaxations are governed by the dielectric relaxations of the polymeric matrix. Above the insulator-to-conductor transition, the percolating path is generating charge relaxations. At the percolation threshold, an insulating/conductor hybrid behavior appears showing both previous relaxations.

The dynamic of dielectric PA 11/CNT composites shows a plasticization of the α relaxation mode associated with the glass transition. This evolution reflects a plasticization due to a decrease of the hydrogen bond density in the polymeric matrix and is coherent with results obtained by DSC and DMA.

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