Dielectric study of the glass transition of PET/PEN blends

J Sellarès¹, J A Diego¹, J C Cañadas¹, M Mudarra¹, J Belana¹, P Colomer², F Román² and Y Calventus²

¹ Departament de Física i Enginyeria Nuclear, Universitat Politècnica de Catalunya, Campus de Terrassa, c. Colom 1, E-08222 Terrassa, Spain
² Departament de Màquines i Motors Tèrmics, Universitat Politècnica de Catalunya, Campus de Terrassa, c. Colom 11, E-08222 Terrassa, Spain
E-mail: jordi.sellarès@upc.edu

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Abstract
An analysis of the glass transition of four materials with similar chemical structures is performed: PET, PEN and two PET/PEN blends (90/10 and 70/30 w/w). During the melt processing of the blends transesterification reactions yield block and random PET/PEN copolymers that act as compatibilizers. The blends obtained in this way have been characterized by ¹H-NMR and differential scan calorimetry (DSC). A degree of randomness of 0.38 and 0.26 has been found for the 90/10 and 70/30 copolymers. It is shown by DSC that this copolimerization is enough to compatibilize the blends. The α relaxation, the dielectric manifestation of the glass transition, has been studied by thermally stimulated depolarization currents. The relaxation has been analysed into its elementary modes by means of a relaxation map analysis. The activation energies of the modes of the glass transition do not change significantly between the four materials: in all cases the modes with a larger contribution have around 3 eV and modes with less than 1 eV are not detected. The change in the pre-exponential factor accounts entirely for the relaxation time change from material to material, that is larger as the PEN content increases. The compensation law is fulfilled and compensation plots converge for high-frequency modes. The polarizability decreases as the PEN content increases due to the increased stiffness of the polymer backbone. An analysis of the cooperativity shows that the central modes of the distribution are the most cooperative while high-frequency modes tend to behave more as Arrhenius. The low-frequency modes are difficult to study due to the asymmetry of the distribution of relaxation times. PEN turns out to be the less cooperative material. It is demonstrated how the parameters obtained from the dielectric study are able to reproduce calorimetric data from DSC scans and are, therefore, a valid description of the glass transition.

(Some figures may appear in colour only in the online journal)

1. Introduction

Poly(ethylene terephthalate) (PET) and poly(ethylene naphthalate) (PEN) are two commercially important polyesters that have a similar structure, as can be seen in figure 1. Indeed, the only difference between them is that the single aromatic ring in PET is replaced by a double one in PEN. For this reason, PEN has improved thermal and mechanical properties over PET even though both materials share some desirable properties such as low permeability to gases, good impact strength and resistance to chemical attack. In spite of this fact, commercial applications of PEN are limited because it is more expensive than PET.

PET/PEN copolymers are interesting because they have properties that are intermediate to both materials at a lower cost than pure PEN. PET and PEN are immiscible [1] but when transesterification takes place the copolymers that are formed act as compatibilizers and, therefore, miscible blends can be obtained [2]. In a transesterification reaction a PET repeating unit and a PEN repeating unit interchange their positions

\[
\text{T–E–T + N–E–N} \rightleftharpoons \text{T–E–N + N–E–T},
\]

where T is terephthalate and N is naphthalate.
The glass transition of PET/PEN copolymers has been completely studied through dielectric properties, using mainly dielectric loss spectroscopy. It has been found that the dielectric strength and the width of the relaxation time distribution are sensible to the processing parameters but molecular coupling does not change significantly with the blend composition or the degree of transesterification [4]. The $\alpha$ relaxation, the dielectric manifestation of the glass transition [5], depends strongly on the thermal history [6] because crystallization and physical ageing affects the amorphous phase that is where this phenomenon takes place [7]. The relative content of PET and PEN also affects the $\alpha$ relaxation. Particularly, there is a noticeable reduction in the dielectric strength due to a decrease in the mobility of local segments as the content of PEN increases [8]. With regards to the other relaxations, a subglass relaxation $\beta^*$ can be observed in PEN and copolymers but not in PET [6, 9].

In a previous work [10], we demonstrated that the glass transition could be studied by dielectric means using the thermally stimulated depolarization currents (TSDCs) technique to perform a relaxation map analysis (RMA) [11]. The poling schemes used in the RMA must excite only elementary modes of the relaxation. To this end, non-isothermal window poling (NIW) or window poling (WP) are appropriate choices. Within a RMA, a depolarization current is obtained for each elementary mode and then the parameters of each mode are obtained fitting the data to a relaxation model such as Arrhenius or Tool–Narayanaswamy–Moynihan (TNM). The latter is able to model the structural state of the system and therefore it is suitable to model memory effects or physical ageing. This approach has the advantage that a distributed process such as the glass transition can be analysed as elementary modes.

The goal of this work is to understand in which way the glass transition of PET/PEN blends/copolymers differ. Since both polymers are very similar we expect that some parameters of the glass transition will remain unchanged while some others will change gradually between the PET and the PEN values in terms of the PET/PEN ratio. These parameters will be obtained following the aforementioned TSDC-RMA approach. Throughout this work we will take advantage of the distinctive capability of this approach to study individual modes on its own. Since we will be studying the glass transition using dielectric techniques, we will also check that the parameters obtained reproduce data from calorimetric experiments and, therefore, describe the glass transition satisfactorily.

2. Experimental

2.1. Preparation

A commercial PET, with a number-average molecular weight ($M_n$) of $2.0 \times 10^4$ [12] and a commercial PEN, Kaladex, supplied by Goodfellow, with a density of $1.36 \text{ g cm}^{-3}$ and nominal granule size of 5 mm were used in this work.

Previous to the preparation of the blends, the polymers were conditioned for 12 h at 150 $^\circ$C in a vacuum oven, in order to eliminate the absorbed water and minimize possible hydrolysis during the melting process.
PET/PEN blends, from 90/10 and 70/30 (w/w) mixtures, were prepared by melt extrusion using a twin screw extruder Haake Rheocorder with a screw speed of 50 rpm. Samples of 60 g, approximately, were melted at 290 °C for 5 min. A nitrogen purge was used during processing to prevent oxidative degradation. The extrudate passed through a cooling ice-water bath and was vacuum dried at 70 °C in order to eliminate absorbed water.

PET and PEN pure polymers were processed in the extruder under the same conditions as the blends so finally four materials have been studied: PET, 90/10, 70/30 and PEN.

The samples obtained from the extruder were melted at 290 °C. After 5 min at this temperature they were moulded and quenched to room temperature by two parallel plates refrigerated by water. A thickness of 0.5 mm was obtained for all these sheets.

The sheets obtained in this way had low crystallinity, as indicated by the transparency of the sheets and the lack of milky patches. The crystallinity degree was estimated from differential scan calorimetry (DSC) measurements and found to be less than 10% in all cases. Also, it should be noted that all the measurements can be performed with the same resolution with our equipment. The parameters employed in our experiments are enclosed in table 1.

### Table 1. Parameters used in the NIW RMA.

| V     | R  | \( t_w \) | \( t_p \) | \( \Delta T_p \) | \( T_d \) | \( t_d \) |
|-------|----|----------|----------|----------------|---------|---------|
| 1500 V| 2.5 °C min^{-1} | 0        | 0        | 2 °C          | 20 °C   | 10 min  |
| PET   | 90 °C | 78 °C to 52 °C in 2 °C steps |
| 90/10 | 96 °C | 82 °C to 58 °C in 2 °C steps |
| 70/30 | 102 °C | 90 °C to 64 °C in 2 °C steps |
| PEN   | 132 °C | 122 °C to 94 °C in 2 °C steps |

The composition determination and the sequence distribution analysis of the copolymesters were conducted by \(^1\)H-NMR spectra using a Bruker AMX-300 spectrometer at 25 °C operating at 300 MHz. The samples (20 mg) were dissolved in 1 mL of a mixture of deuterated chloroform (CDCl₃)/trifluoroacetic acid (TFA) (8/1 v/v) and spectra were internally referenced to tetramethylsilane (TMS) [13]. Sixty-four scans were required with 32 K data points and a relaxation delay of 20 s. Integration of the overlapping signals was made by lorentzian deconvolution of the spectra using the Bruker 1D WIN-NMR computer software.

DSC scans were obtained with a Mettler DSC-20 calorimeter controlled by a TC11 processor. Experiments were performed on 20 mg samples sealed on aluminium pans. DSC scans with a heating rate of 2.5 °C min⁻¹ were performed on these samples.

### 2.3. Measurements

The experimental setup for TSDC consists of a custom build forced air oven controlled by a Eurotherm-2416 PID temperature programmer and a Knürr-Heinzinger (6 kV) potential source. The samples are positively biased connecting one electrode to the source while the other one is grounded. The temperature is measured to an accuracy of 0.1 K by a thermocouple sensor located close to the samples. A Keithley-6514 electrometer was employed for the current intensity measurements.

The sheets were cut into appropriate samples for measurement by TSDC. Circular aluminium electrodes with 1 cm diameter were vacuum deposited on each side of the samples.

An outline of a conventional TSDC experiment can be seen in figure 2. At the beginning of the experiment the sample is at an initial temperature \( T_i \). It is brought to a poling temperature \( T_p \) and after a short time \( t_w \) it is poled for a poling time \( t_p \). Then, the sample is cooled down to a deposit temperature \( T_d \). During this cooling ramp, the poling field is switched off at a temperature \( \Delta T_p \) below \( T_p \). After a deposit time \( t_d \) it is heated up to a final temperature \( T_f \) while the depolarization current is recorded. The depolarization current represented in front of \( T \) is also called a spectrum or a thermogram.

An RMA was performed on a sample of each material using two different poling schemes. In the first scheme, NIW, there is no isothermal poling stage \( (t_w = t_p = 0) \) so the poling field is applied entirely during a cooling ramp. It is switched on at a polarization temperature \( T_p \) and removed when the sample reaches a temperature \( T_p - \Delta T_p \). If \( \Delta T_p \) is small enough, the depolarization current behaves almost as coming from an elementary mode. NIW poling has the advantage that all the measurements can be performed with the same thermal history even if they have different \( T_p \). The value of \( \Delta T_p \) employed (2 °C) is the smallest one that still yields a depolarization current that can be recorded with an acceptable resolution with our equipment. The parameters employed in our experiments are enclosed in table 1.

As saturation is usually not attained using NIW poling, another RMA was performed for each material using the window poling (WP) scheme [14], to obtain data about polarizability. Within this scheme we pole the sample isothermally at \( T_p \) and switch off the electric field at the end of the isothermal poling stage, just when the sample cooling begins \( (\Delta T_p = 0) \) [15,16]. As most poling temperatures...
are just below $T_p$, the sample will undergo physical ageing during the poling stage. A moderate poling time (5 min) was employed to limit physical ageing during the poling stage and, therefore, ensure that the fictive temperature and, consequently, the relaxation time do not change too much during the isothermal stage. This minimizes the effect of the different thermal history of the samples in each measurement. The parameters employed in our experiments are enclosed in table 2.

In a previous work [10] it was found that NIW with $\Delta T_p = 2^\circ$C and WP with $t_p = 5$ min gave similar values of the relaxation time of each mode and of their relative weight, in the case of PET. The agreement between these results is an indication that, on the one hand, $\Delta T_p$ is small enough to obtain spectra from mainly one elementary mode and, on the other hand, $t_p$ is small enough so that there is no significant change on the relaxation time due to physical ageing during the isothermal stage.

To check the validity of the modelling of the glass transition, DSC scans were performed with different cooling rates. These scans were also made with the Mettler DSC-20 calorimeter with samples prepared in the same way as for the characterization of the blends. In these experiments each sample is heated at the same initial temperature as in the TSDC experiments. Then it is cooled at a cooling rate $R$, of 1.25, 2.5 or $5^\circ$C min$^{-1}$ until it arrives to $20^\circ$C. The sample is kept for 10 min at this temperature and then it is heated at a rate of $2.5^\circ$C min$^{-1}$.

### 3. Results and discussion

#### 3.1. Characterization of the blends results

In the $^1$H-NMR spectra, the peak of the ethylene links N–E–T (between naphthalate and terephthalate) appears at approximately 4.84 ppm, while those for N–E–N links (between two naphthalate units) and T–E–T links (between two terephthalate units) appear at approximately 4.89 ppm and 4.79 ppm, respectively. The evaluation of the extent of transesterification and the resulting sequence distribution were made from the areas under these peaks [17, 18]. Table 3 shows the obtained results of the composition and of the sequence distribution of the blends.

Molar composition of the blends, $x_{\text{TET}}$ and $x_{\text{EN}}$, was determined from the aromatic proton resonances observed in $^1$H-NMR spectra. The mole fraction ($P$) of dyads of T–E–N, T–E–T and N–E–N units, $P_{\text{EN}}$, $P_{\text{TET}}$ and $P_{\text{NET}}$, are obtained from the intensities or areas of these three peaks ($A_{\text{TET}}$, $A_{\text{NET}}$ and $A_{\text{EN}}$, respectively)

\[
P_{\text{TET}} = \frac{A_{\text{TET}}}{A},
\]

\[
P_{\text{EN} = \frac{A_{\text{EN}}}{A},
\]

\[
P_{\text{NET}} = \frac{A_{\text{NET}}}{A},
\]

where $A = A_{\text{TET}} + A_{\text{NET}} + A_{\text{EN}}$.

The degree of randomness ($\chi$) is determined by the equation:

\[
\chi = \frac{P_{\text{NT}} + P_{\text{TN}}}{P_{\text{NT}}^2} = \frac{P_{\text{NET}}}{P_{\text{NT}}^2},
\]

where $P_n = (P_{\text{NET}}/2) + P_{\text{EN}}$ is the mole fraction of naphthalate units, $P_t = (P_{\text{NET}}/2) + P_{\text{TET}}$ is the mole fraction of terephthalate units, $P_{\text{NT}}$ is the probability of finding an N unit next to a T unit and $P_{\text{TN}}$ is the probability of finding a T unit next to an N unit. From probability theory $P_t = 1 - P_n$. Therefore,

\[
\chi = \frac{P_{\text{NET}}}{P_{\text{NT}}(1 - P_n)}. \tag{5}
\]

According Shi and Jarabin [2, 17], if $\chi = 1$, the N and T units take a totally random distribution, that is, the copolymer formed is a random copolymer. If $\chi < 1$, the N and T units tend to cluster in blocks of each one of the units, that is, the sequence length is long and the copolymer formed is a block copolymer. If $\chi > 1$, the length of the sequences becomes shorter and the copolymer tends to form an alternating copolymer. If $\chi = 2$, an alternating copolymer is formed. If $\chi = 0$, the system is a mixture of homopolymers. The randomness degree of the blends used in this work are 0.38 and 0.26, respectively. This fact indicates the presence of a block copolymer in both samples as consequence of the transesterification reaction. The randomness degree decreases with the increasing of PEN content.

The number average sequence length of N and T units in the polymer are given by:

\[
n_{\text{EN}} = \frac{2P_n}{P_{\text{NET}}}, \quad n_{\text{TET}} = \frac{2P_t}{P_{\text{NT}}}. \tag{6}
\]

In table 3 it can be seen that a very low value of 2.9 is obtained for the 90/10 blend. As the PEN content is larger in the blend, the number average sequence length of EN increases (2.9 for the 90/10 sample and 5.3 for the 70/30 sample) and the value of ET decreases (30.5 for the 90/10 sample and 14.7 for the 70/30 sample). In this table we also present for comparison, in brackets, the theoretical values calculated on the basis of a random comonomer distribution using the copolyester composition data given in this table.

In figure 3 the DSC scans of the materials before being processed into sheets is presented. Approximate $T_g$ values can be deduced from this figure. These values are presented in table 4. A single glass transition is observed on all the plots. This indicates that there has been enough polymerization to compatibilize the blends as expected from $^1$H-NMR results.

### Table 2. Parameters used in the WP RMA.

| V  | R   | $t_w$ | $t_p$ | $\Delta T_p$ | $T_1$ | $T_2$ | $T_3$ |
|----|-----|-------|-------|--------------|-------|-------|-------|
| 1500 V | 2.5 $^\circ$C min$^{-1}$ | 1 min | 5 min | 0 | 20 $^\circ$C | 10 min |
| PET/PEN | $T_1$, $T_t$ | $[T_p]$ |
| PET | 90 $^\circ$C | 78 $^\circ$C to 48 $^\circ$C in 2 $^\circ$C steps |
| 90/10 | 96 $^\circ$C | 82 $^\circ$C to 54 $^\circ$C in 2 $^\circ$C steps |
| 70/30 | 102 $^\circ$C | 90 $^\circ$C to 60 $^\circ$C in 2 $^\circ$C steps |
| PEN | 132 $^\circ$C | 122 $^\circ$C to 90 $^\circ$C in 2 $^\circ$C steps |
Table 3. Composition and sequence distribution of PET/PEN blends.

| Blend w/w | Molar composition       | Dyads (mol%)       | Number average (Sequence length) | Randomness χ |
|-----------|-------------------------|--------------------|----------------------------------|--------------|
| PET/PEN   | $x_{ET}$, $x_{EN}$      | $P_{ET}$, $P_{EN}$ | $n_{ET}$, $n_{EN}$              |              |
| 90/10     | 92.2, 7.8               | 88.4, 6.0          | 5.6, 0.6                         | 30.5, 2.9    | 0.38          |
| 70/30     | 74.5, 25.5              | 68.4, 10.0         | 21.5, 6.5                        | 14.7, 5.3    | 0.26          |

Figure 3. DSC scan of as-received material. Heating rate 2.5°C min⁻¹: (a) PET, (b) 90/10 PET/PEN, (c) 70/30 PET/PEN, (d) PEN.

Table 4. Approximate values of dynamic $T_g$ at 2°C min⁻¹ obtained from DSC.

| PET  | 90/10 | 70/30 | PEN |
|------|-------|-------|-----|
| 76   | 80    | 87    | 118 |

3.2. TSDC-RMA results

Figure 4 shows the depolarization currents obtained by TSDC/NIW measurements that make up a RMA. In all cases, we obtain almost elementary spectra, this is, spectra that records mainly the response of a single mode. These spectra represent the modes of the relaxation. We will call optimal poling temperature ($T_{po}$) to the poling temperature employed to obtain the spectrum that yields the maximum current value of all the spectra, and temperature of the maximum ($T_{max}$) to the temperature at which that maximum current takes place [5].

The first observation is that for each material $T_{max}$ follows the same pattern as $T_g$ in figure 3. In fact, it is known that $T_{max}$ corresponds to the value of $T_g$ at the rate of the heating ramp [5]. Figure 4 only shows one glass transition temperature in each sample, as it was observed by DSC. This fact supports the suitability of the copolymerization-blending process carried out. In figures 4(a)–(c) (samples with some PET content), the presence of a small $\rho$ peak can be seen at about 8°C above $T_{max}$.

3.3. Modelling of TSDC-RMA

Modelling of TSDC data have been performed assuming that, if the thermal window is narrow enough, the response will be due almost entirely to just one elementary mode. As such, it will be well described by a single relaxation time ($\tau$). The intensity current that corresponds to an elementary mode $i$ of the $\alpha$ relaxation is given by [19]

$$I_i(t) \propto \exp \left[ - \int_{t_0}^{t} \frac{dt'}{\tau_i(t')} - \ln[\tau_i(t)] \right].$$

In this equation, the dielectric relaxation time is considered to be the same as the structural relaxation one. The parameters that correspond to each mode are obtained fitting a calculated depolarization current $I(T)$ obtained from equation (8) to experimental data. There are several models for $\tau$ that can be used to modelize the current.
a distribution of relaxation times (DRTs). This DRT can be
modeled like TNM. Later the aspects that go beyond this model with a more capable
modelization of the Vogel–Tammann–Fulcher [21] or TNM are more commonly
used to modelize the relaxation. Nevertheless, both include
the Arrhenius model as limit case and have three adjustable
parameters, one more than Arrhenius, so we can use Arrhenius
as a convenient approximation to get the big picture and study
the sample by the polarization temperature [11].

The existence of multiple modes is modelized through
a low-frequency mode, the optimal
contribution of that mode [10]. Within an RMA we can identify
the weight of a mode that quantifies the
relaxation in mind. Nevertheless, both include
the Arrhenius model as limit case and have three adjustable
parameters, one more than Arrhenius, so we can use Arrhenius
as a convenient approximation to get the big picture and study
the sample by the polarization temperature [11].

In spite of the fact that the Arrhenius model describes
the α relaxation in a simple way (and with a considerable
ease of interpretation), well-known non-linear phenomena like
thermal history effects and physical ageing cannot be explained by
this model. The simplest extension to the Arrhenius model that incorporates this parameter is the Tool–Narayanaswamy–
Moynihan (TNM) model [23]:

$$\tau(T, T_i) = \tau_0 \exp \left( \frac{E_a}{RT} \right) \exp \left[ \frac{(1-x)E_a}{RT_i} \right]. \quad (10)$$

In this model the separation between $T$ and $T_i$ is introduced
through a non-linearity parameter $x$ ($0 \leq x \leq 1$). The other two parameters are the pre-exponential factor $\tau_0$ and
the activation energy $E_a$. The evolution of $T_i$ with time and
temperature is introduced assuming an ideal-viscous return to
equilibrium that depends on the relaxation time $\tau(t)$ by the equation

$$\frac{dT_i}{dt} = \frac{1}{\tau(t)}(T - T_i). \quad (11)$$

Equations (10) and (11) are coupled. As a consequence, numerical methods are required to integrate them [10]. Some values of $E_a$, $\tau_0$ and $x$ obtained during the fitting processes are reproduced in table 6. The selected values follow the same criteria as those of table 5. The fit of the TNM model for the optimal polarization mode of PET and PEN is also shown in
figure 5. There is a slight but noticeable improvement over
the fit with the Arrhenius model. It can be seen that the
TNM model deals better with the asymmetry of the current
peak, not surprisingly since it was conceived with the structural
relaxation in mind.

3.4. Discussion of the TSDC-RMA modelling outcome

We will begin discussing the results of the Arrhenius model.
In figure 6 it can be seen the weight of each mode $N$ against

Table 5. Some representative parameters obtained from the
mathematical fit of TSDC/WP curves to the Arrhenius model.

| $\tau_0$ (s) | $E_a$ (eV) | $E_p$ (kJ mol$^{-1}$) | $P$ (C m$^{-2}$) | $T_p$ (K) |
|-------------|------------|----------------------|----------------|-------------|
| PET         |            |                      |                |             |
| 0.189 × 10$^{-59}$ | 4.28     | 413                 | 2.33 × 10$^{-4}$ | 351.15     |
| 0.863 × 10$^{-42}$ | 3.05     | 294                 | 3.20 × 10$^{-4}$ | 347.15     |
| 0.362 × 10$^{-13}$ | 1.07     | 103                 | 6.10 × 10$^{-5}$ | 333.15     |
| 0.361 × 10$^{-61}$ | 4.44     | 428                 | 1.17 × 10$^{-4}$ | 355.15     |
| 0.400 × 10$^{-42}$ | 3.10     | 299                 | 1.78 × 10$^{-4}$ | 351.15     |
| 0.564 × 10$^{-14}$ | 1.13     | 109                 | 3.98 × 10$^{-5}$ | 337.15     |
| 0.122 × 10$^{-58}$ | 4.37     | 422                 | 1.41 × 10$^{-4}$ | 363.15     |
| 0.370 × 10$^{-44}$ | 3.32     | 320                 | 2.23 × 10$^{-4}$ | 359.15     |
| 0.185 × 10$^{-15}$ | 1.26     | 122                 | 5.99 × 10$^{-5}$ | 345.15     |
| 0.264 × 10$^{-49}$ | 4.03     | 389                 | 6.60 × 10$^{-5}$ | 393.15     |
| 0.115 × 10$^{-33}$ | 2.80     | 270                 | 1.05 × 10$^{-4}$ | 387.15     |
| 0.218 × 10$^{-16}$ | 1.43     | 138                 | 2.93 × 10$^{-5}$ | 373.15     |

Figure 4. Plot of the TSDC spectra obtained using NIW: (a) PET, $T_p$ from 78 to 64 °C in 2 °C steps, (b) 90/10 PET/PEN, $T_p$ from 82 to 68 °C in 2 °C steps, (c) 70/30 PET/PEN, $T_p$ from 90 to 76 °C in 2 °C steps, (d) PEN, $T_p$ from 122 to 108 °C in 2 °C steps.

In the Arrhenius model the relaxation time is assumed to
obey the equation:

$$\tau(T) = \tau_0 \exp \left( \frac{E_a}{RT} \right), \quad (9)$$

where the activation energy $E_a$ and the pre-exponential factor $\tau_0$ have a simple physical interpretation [20]. Some values of $E_a$ and $\tau_0$ obtained from the fits are reproduced in table 5. These values correspond to a low-frequency mode, the optimal
polarization mode and a high-frequency mode.

The fit of the Arrhenius model for the optimal polarization
mode of PET and PEN is shown in figure 5. The fit is pretty
satisfactory in spite of the fact that the Arrhenius model is not
the best model for a cooperative relaxation. Models such as
Vogel–Tammann–Fulcher [21] or TNM are more commonly
used to modelize the $\alpha$ relaxation. Nevertheless, both include
the Arrhenius model as limit case and have three adjustable
parameters, one more than Arrhenius, so we can use Arrhenius
as a convenient approximation to get the big picture and study
later the aspects that go beyond this model with a more capable
model like TNM.
its activation energy $E_a$. The weights are normalized so the estimated maximum weight is equal to 1. A fifth-order polynomial regression is used to estimate the shape of the distribution. We can hardly see any change in these curves between the samples.

The activation energy that corresponds to the mode with maximum weight is around 3.5 eV in all the cases except for the PEN sample that shows a slightly lower value (3.1 eV). The high temperature side of the curve also differs in the PEN sample from the other ones, although this fact is probably due to the presence of the $\rho$ peak in the samples with PET, that may influence the calculations. The shape of the distribution is noticeably asymmetric, with a steeper decrease on the low-frequency side. The only remarkable differences between materials appear in the lowest $E_a$ value obtained in each curve that show a progressive increase as PEN content does, from 1.0 to 1.4 eV.

This can be seen better in figure 7, where the poling temperature at which each mode obtained is plotted against its activation energy. The low-frequency part of the relaxation (higher $E_a$) is difficult to study because there are too few points due to the asymmetry of the relaxation. Instead, the high-frequency part of the relaxation (lower $E_a$) has plenty of points that allow us to see that there is a cutoff activation energy. This fact means that there are no chain segments activated below certain value of $E_a$. This lower value of the activation energy increases with PEN content, from 1.0 to 1.4 eV as stated above.

So what is the reason why the relaxation time becomes larger and the glass transition shifts to higher temperatures as the PEN content increases? A plot of $E_a$ in front of $\tau_0$ is called a compensation plot and usually gives a straight line when plotted on a log-linear scale. The cause of this relationship is unclear. Some authors think that it is characteristic of cooperative relaxations [24] while others point out that it will appear whenever there is a sharp increase in the activation energy in the DRT [25]. Nevertheless, we are going to use the compensation plot just to compare the behaviour of the relaxation time of the materials. Compensation behaviour in polyesters has already been reported [26].

In the compensation plot presented in figure 8 we can see that each polymer has its own compensation slope. The change in this slope is what enlarges the relaxation time because it gives a larger pre-exponential factor if activation energies are

Table 6. Some representative parameters obtained from the mathematical fit of TSDC/WP curves to the TNM model.

| $\tau_0$ (s) | $E_a$ (eV) | $E_x$ (kJ mol$^{-1}$) | $x$ | $P$ (C m$^{-2}$) | $T_p$ (K) |
|-------|--------|----------------|-----|-------------|--------|
| PET   | 0.259 $\times 10^{-81}$ | 5.78 | 558 | 0.70 $\times 2.15 \times 10^{-4}$ | 351.15 |
|      | 0.702 $\times 10^{-57}$ | 4.07 | 393 | 0.73 $\times 2.94 \times 10^{-4}$ | 347.15 |
|      | 0.362 $\times 10^{-13}$ | 1.07 | 103 | 1.00 $\times 6.10 \times 10^{-5}$ | 333.15 |
|      | 0.278 $\times 10^{-8}$ | 6.05 | 584 | 0.69 $\times 1.07 \times 10^{-4}$ | 355.15 |
|      | 0.700 $\times 10^{-55}$ | 3.98 | 384 | 0.76 $\times 1.67 \times 10^{-4}$ | 351.15 |
|      | 0.564 $\times 10^{-14}$ | 1.13 | 109 | 1.00 $\times 3.98 \times 10^{-5}$ | 337.15 |
| 90/10 |      | 0.271 $\times 10^{-77}$ | 5.70 | 550 | 0.74 $\times 1.33 \times 10^{-4}$ | 363.15 |
|      | 0.137 $\times 10^{-59}$ | 4.41 | 425 | 0.72 $\times 2.07 \times 10^{-4}$ | 359.15 |
|      | 0.186 $\times 10^{-15}$ | 1.26 | 122 | 1.00 $\times 5.99 \times 10^{-5}$ | 345.15 |
| 70/30 |      | 0.575 $\times 10^{-71}$ | 5.70 | 550 | 0.63 $\times 5.95 \times 10^{-5}$ | 393.15 |
| PEN   | 0.220 $\times 10^{-44}$ | 3.61 | 348 | 0.75 $\times 9.88 \times 10^{-5}$ | 387.15 |
|      | 0.219 $\times 10^{-16}$ | 1.43 | 138 | 1.00 $\times 2.93 \times 10^{-5}$ | 373.15 |

Figure 5. Fits of the Arrhenius model (continuous line) and the TNM model (dashed line) to experimental data of the optimal polarization mode (circle) for PET and PEN.
Figure 6. Weight of each mode against its activation energy as obtained from the NIW RMA fits to the Arrhenius model: (circle) PET, (square) 90/10 PET/PEN, (diamond) 70/30 PET/PEN, (triangle) PEN. The weights of each material are normalized so the estimated maximum value is 1.

Figure 7. Poling temperature at which each mode is obtained against its activation energy as obtained from the NIW RMA fits to the Arrhenius model: (circle) PET, (square) 90/10 PET/PEN, (diamond) 70/30 PET/PEN, (triangle) PEN.

the same. Therefore, the change in the relaxation time is mainly due to a change in the pre-exponential factors, not in the activation energies.

A possible interpretation of these results can be the following one. We can assume that the activation energy of the relaxation, $E_a$, is mainly associated with the energy necessary to break hydrogen bonds between close chains, and/or to regain flexibility in bonds of the main polymer chain. In the framework of this assumption, all the materials (PET, PEN and their copolymers) would show similar values of $E_a$ due to the high similarity in their chemical structure. Once some segments of the polymer chain regain mobility in this way, the increased complexity of PEN monomers will cause the chain segments to move slower resulting in an increased relaxation time, as our results show.

Also, figure 8 hints the existence of a point where the compensation plots of each material could join or, at least, come very close. This would mean that whatever mechanism
is responsible for the increase in the pre-exponential factor is more effective at low frequencies (high activation energies).

Since saturation cannot be attained using the NIW poling scheme, an additional RMA was performed by WP with 5 min polarization time to obtain data about polarizability (see section 2 for details). A correction factor has been applied to take into account deviations of the samples’ thickness from the mean 0.5 mm value. Spectra obtained in these experiments almost exactly resembles those obtained using the NIW poling scheme, although in this case delivering twice as much depolarization current as in the previous case. Polarization against poling temperature is presented in figure 9 for the four materials. A consistent trend is found compatible with a lower polarizability due to an increased stiffness when PEN content increases.

In spite of the interest of these results, the Arrhenius model does not take into account physical ageing since the relaxation time has no dependence on the history of the sample. To quantify this effects the TNM model introduces a non-linearity parameter, \( x \), that quantifies the cooperativity of the modes.

The analysis of the non-linearity parameter has to cope with some uncertainty in the fits but it reaches similar values (around \( x = 0.7 \)) in all the samples, as can be seen in figure 10. The modes activated around the optimum polarization temperature and above (\( E_a \) of 3 eV and above) are the most cooperative. As we approach the high-frequency part of the relaxation (lower \( E_a \)) the cooperativity shrinks until it becomes arrhenius-like in all the cases. This happens earlier as PEN content increases, maybe due to the bulky naphthalene ring of the PEN chain, that makes it more difficult for long chain segments to move in a cooperative way. The analysis of the other parameters coincides with the Arrhenius results and is not discussed again.

We can gain further insight on this matter if we represent the weight of each mode in terms of the value of its non-linearity parameter, as seen in figure 11. It can be seen that in PEN the weight of the non-cooperative modes is almost as large as the weight of the cooperative ones. In spite of this fact the non-linearity of the most important modes is similar in all the materials with a value around 0.7.

3.5. Simulation of DSC curves

Compared with dielectric loss spectroscopy, TSDC data are well suited to comparison with calorimetric data. The depolarization current is recorded when the sample is recovering structural equilibrium, just as happens in DSC. Instead dielectric loss experiments with an equivalent frequency, around \( 10^{-6} \) Hz [27], take too much time to be useful for studying the dynamics of the structural relaxation. Moreover, the thermal history of a TSDC experiment and a DSC history are very similar if not the same. This means that given a complete enough model the normalized calorific capacity around the glass transition can be reproduced from the obtained parameters. TNM is fine for this purpose since it takes into account the structural state of the system through the fictive temperature [10] and therefore it can reproduce memory effects or physical ageing.

The normalized calorific capacity is obtained from the fictive temperature [28] through

\[
\frac{c_p^n}{c_p} = \frac{dT_f}{dT}.
\]
Figure 9. Polarization of each mode against its poling temperature as obtained from the WP RMA: (circle) PET, (square) 90/10 PET/PEN, (diamond) 70/30 PET/PEN, (triangle) PEN.

Figure 10. Non-linearity parameter for each mode against its activation energy as obtained from the NIW RMA fits to the TNM model: (circle) PET, (square) 90/10 PET/PEN, (diamond) 70/30 PET/PEN, (triangle) PEN.

Within the TNM model, the fictive temperature is calculated at the same time as the relaxation time since equations (10) and (11) are coupled. In this way, a normalized calorific capacity is obtained for each mode. The normalized calorific capacity of the system can be assumed to be the weighted average of the normalized calorific capacity of each mode, with acceptable results [10]. Again, the weights are the product of the polarization of the mode and the polarization temperature [11].

Figure 12 presents a comparison between DSC data and a calculation of the normalized calorific capacity using the parameters obtained from TNM fits, for the four materials studied. The scale and the origin of the Y-axis for experimental data have been chosen to allow a better comparison with the normalized data provided by the calculations. For the same reason, the direction of the exothermic behaviour has been taken as downward and only the part close to the glass transition has been plotted. As can be seen in the figure, there is agreement on the glass transition region.

The most striking feature of this approach is the ability to reproduce memory effects, as demonstrated in figure 13. In this figure, the results of the calculation of the normalized calorific
Figure 11. Weight of each mode against its non-linearity parameter as obtained from the NIW RMA fits to the TNM model: (circle) PET, (square) 90/10 PET/PEN, (diamond) 70/30 PET/PEN, (triangle) PEN.

Figure 12. DSC scans (continuous line, left scale) compared with predicted calorific capacity (dashed line, right scale): (a) PET, (b) 90/10 PET/PEN, (c) 70/30 PET/PEN, (d) PEN. The two vertical scales have been adjusted in order to ease the comparison.

capacity are compared with DSC data for several cooling rates but the same heating rate. The thermal history is the same in the TSDC experiments with the NIW poling scheme described in table 1 except for the rate of the cooling ramp that has three different values (1.25, 2.5 and 5 °C min⁻¹). The trends given by the varying cooling rate are well-reproduced giving us a demonstration of the capabilities of the TNM model. Let us recall that the Arrhenius model alone is not able to cope with memory effects since the relaxation time depends only on the temperature of the sample at a given time and therefore it would give the same results whenever the heating ramp is the same.

4. Conclusions

From the characterization of the blends follows that enough transesterification took place during the melt processing to compatibilize the PET/PEN blends. As a consequence these
materials present a single glass transition and in TSDC spectra there is a single peak that corresponds to the $\alpha$ relaxation.

$T_{\text{max}}$, the temperature at which the maximum intensity appears when the sample is poled at $T_{\text{po}}$, follows the same pattern as $T_g$ obtained from calorimetric scans. This is not surprising, since the $\alpha$ relaxation is the dielectric counterpart of the glass transition.

The RMA procedure allows us to analyse the $\alpha$ relaxation as elementary modes. Each depolarization current (spectrum) is mainly the response of an elementary mode that can be fitted to obtain the activation energy and the pre-exponential factor of that mode. Within the TNM model also a non-linearity parameter is obtained.

Activation energies almost do not change, although there is a slight and non-significant decrease in the $E_a$ of the optimally poled mode for PEN with regards to the other materials. The change in the relaxation time is due mainly to changes in the pre-exponential factor. The overall shape of the distribution of relaxation times is asymmetric with a much steeper decrease on the low-frequency side.

The compensation law is fulfilled in all cases with a characteristic slope that increases with the content of PEN. An interpretation has been given based on the energy needed to activate the process and the frequency at which it takes place. In any case, for high frequencies the compensation plots of the four materials tend to join.

There is an activation energy cutoff on the high frequency side. The energy of this cutoff increases with the content of PEN. On the other hand, the low-frequency side is difficult to study because it overlaps with the $\rho$ relaxation and also because of the asymmetry of the distribution of relaxation times.

Polarizability decreases with larger PEN content as expected because of the increased stiffness of the polymer backbone.

The non-linearity parameter reaches values around 0.7 for modes obtained poling at temperatures close to the $T_{\text{po}}$. This is another value, together with activation energy, that does not seem to differ between the four materials studied. Instead, the way the cooperativity changes as we consider modes with a higher frequency depends on the PEN content. High-frequency modes are always Arrhenius-like ($x = 1$). As the content of PEN increases the weight of these modes and also of the modes with an intermediate $x$ is larger in comparison with the weight of the central modes of the distribution. This is particularly visible in pure PEN.

It should be noted that this analysis is possible thanks to the capability of TSDC to excite mainly one elementary mode when using a proper poling scheme. Unlike in approaches based on dielectric loss spectroscopy, we do not have to make any assumption on the shape of the distribution of relaxation times or use any kind of deconvolution.

Another plus is that TSDC experiments can be made with the same thermal history as DSC experiments and therefore the return to structural equilibrium that takes place can be compared using an appropriate model. This allows us to reproduce calorimetric data, even memory effects, so we can rely on dielectric results as a proper description of the glass transition of PET/PEN blends.

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