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Evidence of nanostructure development from the molecular dynamics of poly(pentamethylene 2,5 furanoate)

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Abstract

We report on the molecular origin of poly(pentamethylene 2,5-furanoate) particular physical properties. From the structural point of view we found this polymer can develop crystallinity when stored at room conditions for months. On the other hand, we used broadband dielectric spectroscopy (BDS) measurements, to analyze in very detail the local and segmental molecular dynamics of the material subjected to several thermal treatments. In this way we evidenced that the molecular dynamics are sensitive to thermal history over a broad temperature range. This behavior has been attributed to possible inter-chain interactions detected via infrared spectroscopy and rheology measurements in the non-crystallized polymer.

1. Introduction

Thanks to their lightness, versatility and affordable cost, plus the possibility to be appropriately modified for particular applications, plastic materials are widely produced and used.¹ In particular, packaging is the industrial sector with the most significant use of plastics, and also the one in which the final products have the shortest shelf-life.¹ That leads to the accumulation of large quantities of plastic waste in the environment that threatens both the terrestrial and marine habitat. 2 In addition to heavy pollution, traditional plastics also lead to excessive use of non-renewable sources, especially oil.³

The strong academic and industrial commitment to the realization and implementation of new materials suitable to replace traditional plastics, and the current European Union directives, push towards the reuse of plastic products and their recycling.^{1, 4-7} However, to meet the particular needs of a specific type of application, the polymeric matrix is often coupled with other compounds such as additives, paper, metals, as well as with other polymers. $8-10$ This multi-layered nature of a package generally hampers its recyclability.¹⁰⁻¹¹ A possible solution to the problems of the so-wide use of plastics is constituted by biopolymers, *i.e.*, materials that are produced from renewable sources and/or are biodegradable.^{3, 12-13} Therefore, replacing traditional plastics with *mono-material* bioplastics is a desirable solution for a circular economy.

In this context, the scientific community's huge interest in the polyesters based on furandicarboxylic acid is not surprising, as evidenced by a large number of recent literature reports.¹⁴⁻²⁵ The furandicarboxylic acid (FDCA) is a bifunctional aromatic monomer produced from biomass through an eco-sustainable process.²⁶⁻³² It is mainly employed in the synthesis of polyesters^{14-25, 33} and copolyesters with improved functional properties compared to their terephthalic acid-based counterparts, such as excellent thermal stability and processability, excellent barrier properties, and tunable mechanical

response.^{17, 34-52} For example, when compared with its terephthalic analog poly(ethylene terephthalate) (PET), poly(ethylene furanoate) (PEF) is easier to process due to its lower melting temperature and would perform better in applications because of its higher glass transition temperature. Moreover, PEF shows six times better barrier properties for $O₂$ and three times for $CO₂$ than PET, which makes it a very interesting material for the production of soft drink bottles.⁵³⁻⁵⁴ Some authors have highlighted how the properties of this family of polyesters can be suitably modulated by acting on the length of the glycol sub-chain.^{23, 25} Specifically, the different number of methylene groups has a direct effect on the chain mobility and crystallizing capability.^{23, 55-56} These characteristics allow to obtain from amorphous to semicrystalline polymers, which at room temperature can be in the glassy or in the rubbery state. According to these features, a different microstructure can develop, having a direct effect on the macroscopic properties. 25

In the family of FDCA-based polyesters, poly(pentamethylene furanoate), PPeF, is attracting interest.²⁴ PPeF is characterized by a glass transition temperature below room temperature and shows crystallization kinetics so slow that, under normal conditions, it can be considered completely amorphous.^{24, 55-56} Despite these characteristics, PPeF can be processed as films and, very surprisingly, it exhibits an elastic mechanical response at room temperature, as well as exceptional barrier properties.^{24, 55} This behavior has been explained on the basis of particular inter-chain interactions, 24 also proposed for the other polymers of the family. ²⁵ These interactions give rise to a microstructure responsible for the exceptional performances that make PPeF an excellent candidate for the realization of flexible *mono-material* devices, with outstanding gas barrier capabilities.

In this work, we have investigated the molecular origins of the peculiar and interesting properties of PPeF, using a synergistic combination of different experimental techniques. Our results provide new insights on the study of this polymer and differentiate from previous publications, where the material was considered as fully amorphous²³ and with physical properties independent from thermal history.⁵⁶

2. Experimental Section

Samples. PPeF homopolymer has been synthesized according to the procedure described elsewhere and free-standing films have been prepared by compression molding as previously reported. 25

Differential Scanning Calorimetry (DSC). DSC measurements were carried in a TA Instruments Q2000 equipment, with a liquid nitrogen cooling system. Heating/cooling ramps were conducted at a mean rate of 3 K/min. PPeF samples for DSC measurements were prepared by encapsulating *c.a.* 5 mg of material in aluminum pans.

Atomic Force Microscopy (AFM). AFM experiments were carried out using a Bruker Multimode AFM, equipped with a Nanoscope V controller. Height images were captured using the PeakForce Tapping protocol and Tap150Al-G probes (BudgetSensors). The 512x512 pixel/lines images were taken at a 0.05V PeakForce setpoint, and at a 0.3 Hz scan rate. For AFM imaging, we microtomed the PPeF films. In this way, we were able to study the material structure within its volume. The films were cut with a microtome that was equipped with a diamond knife at 0° C. A sample holder designed for crosssection analysis was used. This fixed the sample in the same position throughout the entire process (cutting $+$ imaging).

X-ray diffracton (XRD). X-ray diffraction patterns were acquired with a X'Pert PRO diffractometer in reflection mode with copper K_{α} radiation and a fast solid state X'Celerator detector. The 2θ range 5 - 60° was scanned in 0.1° steps, counting 100 s /step. The crystallinity index (X_c) was calculated from the X-ray pattern as the ratio between the crystalline diffraction area (A_c) and the whole area under the diffraction

profile (A_t), as $X_c = A_c/A_t$ · 100. A_c was obtained from the total area of the diffraction profile by subtracting the amorphous halo, which was modelled as a broad bell-shaped peak. The incoherent scattering was subtracted in advanced.

Broadband Dielectric Spectroscopy (BDS). BDS allows the study of molecular dynamics via the analysis of the complex dielectric permittivity, $\varepsilon^*(\omega) = \varepsilon'(\omega)$ – $i\varepsilon''(\omega)$, where $\varepsilon'(\omega)$ is the dielectric constant and $\varepsilon''(\omega)$ the dielectric losses. The complex dielectric permittivity is a function of the applied electric field frequency (ω = $2\pi f$, being *f* the frequency), and the temperature *T*. We performed BDS experiments in a frequency range $10^{-1} \le f$ (Hz) $\le 10^{7}$, using a Novocontrol dielectric spectrometer, equipped with an Alpha dielectric interface. The temperature was controlled by a nitrogen jet (Quatro from Novocontrol), with a temperature stability during every single frequency sweep of ± 0.2 K. PPeF samples for dielectric studies were prepared by melt-pressing, in order to obtain homogeneous films. These samples were pressed between two gold-plated disc electrodes of 40 mm diameter (lower electrode) and 20 mm diameter (upper electrode). We used narrow Teflon spacers (150 μm thick) to avoid possible shortcircuits.

The BDS data analysis was carried out using the Havriliak-Negami (HN) formalism, where the complex dielectric function can be described as 57 .

$$
\varepsilon^*(\omega) = \varepsilon_{\infty} + \sum_{x} \Delta \varepsilon_x \left[1 + \left(i\omega \tau_{HN_x} \right)^{b_x} \right]^{-c_x} + \left(\frac{\sigma_{DC}}{i\varepsilon_0 \omega} \right) \tag{1}
$$

where $\Delta \varepsilon$ is the dielectric strength of the relaxation and τ_{HN} a characteristic relaxation time; *b* and *c* are shape parameters related to the symmetric and asymmetric broadening, respectively. The summation in equation (1) extends over all the processes present in the experimental window at a specific temperature. Also, the last term in this equation accounts for the contribution of charge carriers to the dielectric signal, where σ_{DC} is the direct current (DC) conductivity and ε_0 the vacuum permittivity.

From τ_{HN} , the peak relaxation time (τ_{max}) was calculated using the equation ⁵⁷:

$$
\tau_{\text{MAX}} \equiv \frac{1}{2\pi f_{\text{MAX}}} = \tau_{\text{HN}} \left[\sin \frac{b\pi}{2 + 2c} \right]^{-1/b} \left[\sin \frac{b c \pi}{2 + 2c} \right]^{1/b} \tag{2}
$$

where f_{max} is the frequency of maximum loss, and the rest are the HN-function parameters.

Rheology. Dynamic torsional shear experiments were conducted over the angular frequency range 10^{-2} - 10^{2} rad \cdot s⁻¹, using an ARES torsional rheometer from TA Instruments, under N_2 atmosphere. Parallel plate geometry with 8 mm diameter tools was used. Sample thicknesses were about 0.5 mm and low strain amplitude was used to ensure linear regime.

Fourier Transform Infrared Spectroscopy (FTIR). FTIR spectra were taken using a Jasco 6300 FTIR spectrometer, working in transmission mode. The samples were deposited on a Zinc Selenide window from a solution of the polymer in chloroform. Prior measurement the samples were subjected to a high vacuum (pressure $< 10^{-7}$ mbar) to allow residual solvent evaporation. The thermal treatments were performed inside the spectrometer, using a LinkamTHMS600 stage connected to a T95-LinkPad System controller.

3. Results

3.1. Thermal transitions and nanostructure development on PPeF. Figure 1 and Table 1 present the calorimetric traces and the corresponding DSC results, respectively, of heating ramps from 220 to 420 K of PPeF films subjected to three different thermal

treatments: slow and fast cooling from high temperature and storage at room temperature for 18 months. In Figure 1, the continuous line is representative of the calorimetric response after cooling the polymer from 420 to 220 K at 0.5 K/min (slow cooling). After this thermal treatment, PPeF showed no signs of crystallization/melting. As a matter of fact, we observed only the calorimetric step corresponding to the glass transition phenomenon (T_g) at 292 K. In Figure 1, the DSC trace of a PPeF sample ballistically cooled ($>$ 70 K/min) inside the DSC from 420 to 220 K (fast cooling) is represented by the dotted line. We observed that the subsequent heating run looked nearly identical to that of the slowly cooled sample, presenting the same values for T_g and Δc_p (Table 1). In particular, the Δc_p values in these two cases are close to that previously found by us for an amorphous poly(butylene 2,5-furanoate) (PBF) sample, ⁵⁸ and comparable to the ones reported by Guidotti et al. and Papamokos et al. for amorphous PPeF.²³⁻²⁴ The dash-dotted line in Figure 1 corresponds to the DSC trace of the sample stored in a desiccator, at room temperature, for 18 months (Aged-PPeF). This sample was ballistically cooled down from 300 K to 220 K, in the DSC instrument, to avoid erasing its thermal history. During the subsequent heating scan, we observed a broader T_g step with inflection point at 289 K, a value slightly lower compared to the slow and fast cooled samples, as well as a reduced Δc_p . Also, we detected a multiple endothermic phenomenon, with maxima at 324 K and 337 K, respectively. It must be noted that in previous works, bulk PPeF was considered as a fully amorphous polymer; however, the observed endothermic process has been ascribed to a melting phenomenon as confirmed below by means of other techniques too. In fact, in principle, both the Δc_p reduction as well as the presence of the endothermic peaks indicated the development of crystals during storage. The associated heat value is of the same order of magnitude as those previously found for other furan-based semicrystalline polymers; 23 on the contrary, the melting temperature is fairly lower.

Sample	$T_{\rm g}$ (K)	$\Delta c_{\rm p}$ (J/g·K)	$T_{\rm m}$ (K)	ΔH_m (J/g)	
slow cooled-PPeF	292 ± 1	0.37 ± 0.01			
fast cooled-PPeF	292 ± 1	0.37 ± 0.01			
Aged-PPeF	289 ± 1	0.27 ± 0.02	324 ± 1 K	$23 + 1$	
			337 ± 1 K		

Table 1. DSC results of PPeF samples under different thermal treatments.

Figure 1. DSC curves (exo up) for PPeF samples undergoing different thermal histories: fast (dotted line) and slow cooling from the melt (solid line), storing for 18 months at room temperature (dash-dotted line).

To complement the DSC results, Figure 2 shows AFM height images of slow cooled and Aged-PPeF samples. In both cases, the height images were captured on the films cross-sections, as detailed in the experimental section. The AFM images of slow cooled PPeF, reported in Figures 2a-c, showed a flat surface with a mean roughness of about 1.3 nm (calculated using Figure 2a). The elongated topography features correspond to imperfections probably left by the microtoming process. Focusing the analysis onto a defect-free zone, as the image shown in Figure 2c, the mean roughness decreased down

to 0.3 nm. This value, similar to those reported for amorphous polymer films 59 , is in line with the DSC results (Figure 1, continuous line). Comparable results were found for a fast cooled PPeF sample (data not shown). Figures 2d-f show the AFM images for Aged-PPeF. In this case, we observed a different topography. Now, there were randomly distributed nanostructures on the film's cross-section that did not cover the entire scanned area. The $10x10$ um image (Figure 2d) had a mean roughness of 3.4 nm. This value only showed a small decrease, to 3.1 nm, when calculated for the 2x2 μm image where most of the probed area is covered by the nanostructures (Figure 2f). This represents a 10x increase in the mean roughness if comparing to the slow cooled sample, and it is comparable to those found for semicrystalline terephthalate-based polymer films 60 . Using Figure 2f, we studied the geometrical characteristics of the nanostructures. We found that the features had lengths in the $20 - 100$ nm range and widths in the $10 - 20$ nm range.

Figure 2. AFM height images of PPeF cross-sections: slow cooled PPeF (a-c) and aged-PPeF (d-f). Magnification increases from left to right.

Lastly, Figure 3 shows the XRD patterns for slow cooled PPeF and Aged-PPeF. The slow cooled PPeF sample (dashed lines) showed the diffractometric pattern typical of an amorphous polymer sample, with amorphous halos, positioned at $q = 1.48 \text{ Å}^{-1}$ (2 θ) $= 20.9^{\circ}$) and $q = 3.11 \text{ Å}^{-1}$ (44.6°). The same results were found for the fast cooled PPeF sample (profile not shown). On the contrary, the profile of Aged-PPeF sample (solid line) was characterized by quite distinct features typical of a semi-crystalline material: several peaks overlapped to a bell-shaped background. The presence of a crystalline phase is coherent with the nanostructures detected through AFM analysis, and with the endothermic phenomena observed by DSC. The XRD peak positions are summarized in Table S1 (Supporting Information).

Figure 3. XRD results for slow cooled (dashed lines) and Aged-PPeF (solid lines) samples. Black arrow indicates the position of the maximum related to a mesophase, as reported by Guidotti et al.,²⁴ as presented in the discussion section of this work. The patterns are displaced in the vertical direction for clarity.

3.2. Impact of nanostructure formation on the PPeF molecular dynamics. We have studied, via BDS, the molecular dynamics of PPeF under different thermal protocols, as

described in the following lines. Figure 4 shows the BDS results, as the imaginary part of the complex dielectric permittivity (ε'') as a function of the electric field frequency. Empty triangles refer to the fast cooled PPeF sample (FC-PPeF). In this case, the film was heated inside the cryostat from room temperature up to 453 K, held for 5 min and then the dielectric cell was immersed into a liquid N_2 bath. Once the dielectric cell temperature was stable, it was transferred back to the cryostat, now at a temperature of 153 K. Measurements started at 203 K, heating in isothermal steps. Solid circles are relative to the results obtained for a sample measured in 5 K isothermal steps, cooling from 453 K to 203 K (SL-PPeF). Finally, crossed squares refer to the Aged-PPeF sample. In this case, the film was loaded into the dielectric cell at room temperature, and immediately cooled down to 203 K to start measuring in isothermal steps, heating up to 453 K.

Local relaxations. Figure 4a shows the results at 243 K ($T < T_g$). At this temperature, all the samples presented a broad relaxation, characterized by a single maximum around 10^4 Hz. The intensity and the asymmetry of the curves were both dependent on thermal history. Specifically, FC-PPeF was the sample with the highest intensity and higher asymmetry. Then, SL- and Aged-PPeF samples showed a progressive decrease in intensity and a more symmetric peak shape. To model these results, we used Cole-Cole (CC) functions, derived from eqn (1) by imposing $c_x = 1$, and $\sigma_{DC} = 0$ S/cm, plus a power-law at low frequencies to account for upcoming relaxations (for further details, refer to Figure S2). For samples showing a nearly symmetrical local relaxation, SL- and Aged-PPeF, we used a single CC function. In the case of the FC-PPeF sample, the experimental data were fitted using a sum of two Cole-Cole functions to account for its asymmetry towards low frequencies. The faster component was called β_1 , and the slower one β_2 , in line with our previous report on poly(alkylene 2,5-furanoate)s'

molecular dynamics.^{56, 58} Lines in Figure 4a are representative fitting curves. In Figure 5, we show the temperature dependence of the resulting fitting parameters, for all three samples.

Figure 4. BDS results for FC-PPeF (\triangle) , SL-PPeF (\bullet) and Aged-PPeF (\boxtimes) at different temperatures: (a) 243 K, (b) 303 K, (c) 333 K, (d) 348 K. Continuous lines correspond to the total fits, as detailed in the text.

Figure 5a presents the Arrhenius relaxation map, where the local relaxations are located in the range $3.5 \le 1000/T$ (K⁻¹) ≤ 5.0 . The temperature dependence of the local relaxations were well described using the Arrhenius equation:⁵⁷

$$
\tau_{\text{MAX}}(T) = \tau_0 \exp\left[\frac{E_A}{RT}\right] \tag{3}
$$

where τ_0 is a pre-exponential factor, E_A the activation energy of the process, and R the ideal gas constant. The Arrhenius fits are shown in Figure 5a by the different continuous lines, while the fitting parameters are summarized in Table 2. The β_1 process showed in all cases a similar pre-exponential factor around 10^{-15} s, and activation energies slightly depending on the thermal treatment. We observed that FC-PPeF presented the highest E_A value, which decreased for the SL-PPeF sample. Finally, the semicrystalline Aged-PPeF polymer showed the smallest E_A value (46 kJ/mol). The β_2 relaxation, only observed in FC-PPeF, presented a much higher E_A value compared to the faster component. In this case, the pre-exponential factor was too low as compared to the reciprocal of a vibrational frequency $(\sim 10^{-13} \text{ s})$.

Figure 5b shows the relaxation strength as a function of the temperature. We observed that the β_2 component, only detected for FC-PPeF, decreases rapidly with increasing temperature, until it is no longer well-resolved above 270 K. On the contrary, the strength of the β_1 component is essentially the same for FC-PPeF and SL-PPeF showing in both cases a slight increase with temperature. Finally, the Aged-PPeF sample presented markedly smaller relaxation strength but with a similar temperature dependence.

Figure 5. Temperature dependence of the fitting parameters used for modeling the BDS results: FC (\triangle) , SL (\bullet) , Aged (\boxtimes) . (a) Relaxation plot, (b) relaxation intensity, (c) standard deviation of the Gaussian distribution of decimal logarithm of relaxation times for local relaxations, (d) shape parameters of the HN equation used to model the segmental relaxation (*b*, open symbols; *bc*, filled symbols).

			α		
Sample	τ_{0} (S)	E_A (kJ/mol)	$\tau_{0-\text{VFT}}$ (s)	D	$T_{\rm VFT}$ (K)
FC-PPeF (β_1)	$(2.2 \pm 0.4) \cdot 10^{-16}$	52 ± 1	10^{-12}	5.1	251 ± 1
FC-PPeF (β_2)	$(1.2 \pm 0.9) \cdot 10^{-24}$	104 ± 2			
SL-PPeF	$(1.3 \pm 0.3) \cdot 10^{-15}$	49 ± 1			251 ± 1
Aged-PPeF	$(2.9 \pm 0.3) \cdot 10^{-15}$	46 ± 1			250.0 ± 0.5

Table 2. Arrhenius and VFT parameters for all PPeF samples.

To analyze the shape of these local relaxations we calculated the standard deviation, $\sigma_{\rm G}$, of the distribution of Debye relaxations according to a Gaussian function of $log_{10}(\tau)$, which corresponds to the symmetrical broadening of the CC functions fitting the data $(b \text{ in eqn } (1))$:

$$
\sigma_{\mathcal{G}} = \frac{5 \cdot 10^{-4}}{b^4} - 4 \cdot \log_{10}(b) \tag{4}
$$

A detailed description of eqn (4) is provided in the Supporting Information (Figure S4). The results obtained from this calculation are presented in Figure 5c, as a function of inverse temperature. We observed that all the local relaxations presented the same trend, *i.e*., the relaxation peaks were narrower as temperature increased. The broadness of the local relaxations decreases linearly as a function of $1/T$ from about 3.2 to 1.8 decades. Such a linear dependence is expected by assuming that the secondary relaxation is originated by superposition of single Debye processes with different time scales resulting from a Gaussian distribution of energy barriers separating equivalent local equilibrium states.

Segmental relaxation. At temperatures above 290 K, the segmental (α) relaxation peak appeared in the BDS experimental frequency window. This is the main relaxation observed for all the samples in Figures 4b-d. The FC- and SL-PPeF samples presented intense peaks, with small but noticeable differences. Systematically, for all temperatures, the SL-PPeF showed a lower peak frequency. The Aged-PPeF sample presented the most different α -relaxation. In this case, at 303 K and 333 K, the relaxation peak showed a much lower intensity and a wider shape, while its maximum stayed at similar frequency, compared to the other two samples. At 348 K, when $T > T_{\text{m}}$, the relaxation peak intensity increased reaching the same value (also in shape and position) as the FC-PPeF sample. The segmental relaxation peaks were modeled using a HN

function with a DC conductivity contribution, eqn (1), (see lines in Figure 4b-c). In the fitting procedure a CC function was used to account for the secondary relaxation contributions at high frequencies. The data, accompanied by the fitting components, are presented in Figure S3.

The temperature dependencies of the fitting parameters are shown in Figure 5. As already commented, there are small variations in the peak maxima between FC- and SL-PPeF presented in Figure 5a. We described the relaxation times for all the samples using the Vogel-Fulcher-Tamman (VFT) law: 57

$$
\tau_{\text{MAX}}(T) = \tau_{0-\text{VFT}} \exp\left[\frac{DT_{\text{VFT}}}{T - T_{\text{VFT}}}\right] \tag{5}
$$

where τ_{0-VFT} is a pre-exponential factor, *D* a parameter related to the so-called fragility (steepness around T_g), and T_{VFT} the so-called Vogel temperature (usually 30-60 K below T_g). We found first the τ_{0-VFT} and *D* values by fitting the data in a broad temperature and frequency range, as detailed in Ref $⁶¹$. The obtained *D* value is comparable to that</sup> previously found for PPeF by Papamokos and collaborators 23 . The T_{VFT} values that resulted by maintaining τ_{0-VFT} and *D* fixed in the VFT equation are summarized in Table 2. T_{VFT} values were around 250 K varying in line with the calorimetric glass transition temperatures, and similar to that previously reported $(247 \text{ K}, \text{Ref}^{23})$.

The α relaxation strength, shown in Figure 5b, was almost constant for both FCand SL-PPeF, being the latter slightly lower for all temperatures. The intensity of the Aged-PPeF segmental relaxation peak was much smaller but increased with temperature, as the sample enters in the melting range observed in DSC. Above the melting range, it remained constant and equal to that of FC-PPeF film. Finally, the shape parameters

(Figure 5d) were similar for both the FC- and SL-PPeF samples, being weakly temperature dependent. The Aged-PPeF sample showed a non-monotonous marked temperature dependence, related to the transformation from the semicrystalline solid state at low temperatures to the melt.

4. Discussion

PPeF showed no signs of crystallization even after slow cooling from the melt, by both DSC and XRD, unlike the results presented for other furan-based polymers with a shorter glycolic subunit, as PEF , $62-63$ PTF 64 and PBF, 58 as well as for some of them with longer glycolic chains.²³ Nonetheless, we observed that bulk PPeF undergoes cold crystallization at room temperature with extremely slow kinetics, slower than those detectable in usual laboratory times. This result is in line with recent AFM observations of PPeF crystallization in thin film geometry, where crystalline structures started appearing after 3 months of storage at room temperature. ⁵⁵ To provide further information about the crystallization behavior of bulk PPeF, in our current work we performed DSC and XRD studies of a freshly prepared sample as a function of the storage time, as presented in Figure S1. There, it is observed that the PPeF film requires about 4 months to develop some detectable crystalline structures. This could be the reason why in previous reports this polymer has been considered as fully amorphous in the bulk.^{23-24, 56} Crystallinity values (X_c), calculated from XRD scans, were 3 ± 1 %, 7 ± 2 % and 17 ± 2 % for samples with storage times of 4, 10 and 12 months, respectively. Concerning XRD results, we highlight that in the article by Guidotti and collaborators, the authors detected a sharp and intense reflection at $q = 1.26 \text{ Å}^{-1}$ for an amorphous PPeF sample.²⁴ This peak was observed to be independent of thermal history of the sample, and its nature was attributed to the formation of a mesophase. 24 In the Aged-PPeF sample, we do not observe a reflection at this specific angle; however, there is a really close one located at $q = 1.23$

 \AA ⁻¹ (black arrow in Figure 3b) that could mask it. Contrary to the previous work, this reflection disappears once the polymer sample is heated above the calorimetric T_m suggesting it is related to the crystal phase.

After cold crystallization, the amorphous phase of the PPeF is affected in different ways by the presence of surrounding crystals. DSC results showed a broader glass transition of Aged-PPeF in the same temperature range. The Δc_{p} also showed a decrease in comparison to that corresponding to the other thermal treatments. From this result, one can calculate the mobile amorphous fraction in the Aged-PPeF sample as $\chi_{a,m}$ = $\Delta c_p^{\text{Aged-PPer}}/\Delta c_p^{\text{FC-PPer}} = 0.74$. This result would indicate that about 26% of the sample should be comprised by immobile phases, as crystals and rigid amorphous fractions. The crystallinity index obtained from XRD ($X_c = 17$ %) takes into account only the amount of material in crystal form and can be taken as a support for this result.

From the molecular dynamics point of view, PPeF showed clear changes in the local relaxations depending on thermal history. The FC-PPeF sample presented an asymmetric response more extended towards lower frequencies, which was modeled by the sum of two CC components. This result is in line with our previous BDS measurements on several melt-quenched poly(alkylene 2,5-furanoate)s.^{56,58} In these latter works, the faster component was related to the $O - C$ bond of the ester oxygen to the aliphatic carbon, while the slower one to the $C - CA$ link between the ester group carbon and the furan ring. The activation energies found for these two relaxations, in FC-PPeF, indicate that the two local motions used to describe the sub-glass relaxation should correspond to those previously described. Concerning the relaxation strength of these two components, we noticed that $\Delta \varepsilon_{\beta_1} \gg \Delta \varepsilon_{\beta_2}$, indicating that the whole β relaxation was dominated by the faster component, a result also in line with previous reports.^{56,58}

After slow cooling from the melt, a single CC component was sufficient to describe the β relaxation of PPeF (SL-PPeF sample). This CC function related quite well to the β_1 relaxation component of the fast cooled sample, considering the obtained parameters (Figure 5). From the relaxation map, we observed that E_A only showed a small decrease, coinciding with the one reported by Papamokos et al., where a singlecomponent local relaxation was observed for PPeF.²³ Also, Soccio et al. found a single local relaxation for several poly(alkylene 2,5-furanoate) samples, including PPeF, cooled at 3 K/min from the melt.⁵⁶ To evaluate the dependence of the local relaxation asymmetry with cooling rate, we subjected the PPeF to different thermal protocols. The corresponding results are presented in Figure 6.

First, Figure 6a shows the BDS results at 218 K ($T \ll T_{\rm g}$) for the FC- and SLsamples, as well as for two additional thermal protocols represented by solid squares and solid stars. These addition protocols correspond to PPeF samples cooled down from 453 to 153 K, at 50 K/min and 3 K/min, respectively, values that lie in between the rate of fast and slow cooling tests. In both these new cases, the BDS measurements were carried out on heating from 203 K, in 5 K isothermal steps. We observed that the dielectric loss asymmetry, indicative of the intensity of the β_2 process, was reduced as cooling rate was reduced. Thus, as the system cools down from the melt, the blocking of the molecular motions related to the β_2 relaxation increased with the time that the system was allowed to equilibrate. From the molecular point of view, our present work, in combination with literature reports,^{56, 58} show that in a well-equilibrated sample there is a blocking of the motion involving the $C - CA$ link.

Second, trying to have a better understanding of the origin of this unusual phenomenon, we performed the following experiment. The PPeF sample was heated to 453 K and melt-quenched into liquid N_2 , as for the FC-PPeF sample discussed so far.

Then, the sample was measured in 5 K isothermal steps up to 218 K. The data collected at this temperature are depicted in Figure 6b by the pink circles with the highest intensity. Then, the BDS experiments proceed as follows. The sample was heated to 223 K, a new BDS measurement was performed isothermally, as usual, and then the sample was cooled down to 218 K to be re-measured. This protocol was repeated by increasing the so-called "temperature of treatment" ($T_{treatment}$) 5 K higher each time, and always cooling down to 218 K to check the response evolution. A schematic representation of the protocol is presented in Figure S5. We observed a monotonic decrease of the low frequency losses, at 218 K, as *T*treatment increased. In the inset of Figure 6b, we show the maximum of the ε'' signal (ε''_{MAX}), at 218 K, after each *T*_{treatment}. We observed that for *T*_{treatment} \geq 258 K, ε''_{MAX} reached a constant value, somehow indicating that the system did not further evolve. These results seem to be directly connected to the continuous decreasing of the relaxation strength of the β_2 component as temperature increases and the fact that it is no longer detectable at $T_{\text{treatment}} \geq 270 \text{ K.}$

Figure 6. Dielectric losses at 218 K, (a) after cooling from 423 K at different rates: FC-PPeF (\triangle) , 50 K/min (\blacksquare) , 3 K/min (\star) , SL-PPeF (\bullet), and (b) after progressive annealing. Inset in panel (b) shows the evolution of ε''_{MAX} as a function of treatment temperature.

Considering that the low frequency loss component of the local relaxation arises from a rather specific molecular motion, it becomes really interesting that its evolution takes place below T_g where the polymer is in a low-mobility state. It is worth noticing that this "local equilibration", occurring well below the glass-transition range, does not result in significant differences in the DSC signals of FC- and SL- samples, both showing approximately the same glass transition temperature and Δc_p values. Concerning the segmental relaxation, we only detected tiny differences, as illustrated in Figure 4. Interestingly, the SL-PPeF sample presents a small excess of losses in the low frequency flank of the α -relaxation loss peak, with respect to that of the FC sample. As a result, the peak position appears shifted towards lower frequencies and the full-width-at-halfmaximum of the loss peak slightly increases. These effects persist until a rather high temperature, 343 K, which is well above T_g . This result would be indicative of the occurrence of some structural development in the SL-PPeF, which is prevented (at least in part) by fast cooling.

In order to detect the presence of any structural feature in PPeF, we have performed linear rheological experiments by means of dynamic torsional shear tests in the temperature range 313 - 393 K, i.e. above its T_g . The results, depicted in Figure 7, show that the time-temperature superposition $(TTS)^{65-67}$ can be reasonably applied to obtain a master curve, with shift factors following the expected Williams-Landel-Ferry (WLF) law⁶⁷ (see line in the inset). From this master curve, it is inferred that the PPeF melt does not behave as a conventional linear polymer, since it is found that Newtonian flow cannot be attained even at temperatures higher than T_g+100 K. Instead, both G' and G'' present a parallel behavior at low frequencies approaching to a power law with exponent 2/3. This finding reminds that of randomly branched polymers,⁶⁸ for which at the gel point G' ~ G" ~ $\omega^{2/3}$. This result could be related to previous ones showing that

amorphous PPeF presents non-conventional mechanical and gas transport properties at room temperature. 24

Figure 7. Master curve obtained from the dynamic experiments on PPeF using (linear) torsional shear: G' empty symbols, G'' solid symbols. The inset presents the horizontal shift factors used and the line shows the WLF equation fitting the data.

Given the linear nature of the synthetized polymer, the obtained results suggest the presence of some kind of supramolecular structures in the PPeF molten state. Literature results on PEF have pointed out the presence of specific intermolecular interactions between neighboring furan moieties, as hydrogen-bonding and/or $\pi-\pi$ stacking.^{23-25, 69} If this type of interactions exists in the PPeF melt, as recently proposed,²⁴ they could give rise to a network structure, which would explain the observed anomalous behaviors and particularly the PPeF rheology.

In order to detect any additional evidence of such specific intermolecular interactions in the amorphous PPeF melt, we have performed FTIR experiments, at 230 K, on PPeF films to compare the absorption spectrum of FC- and SL-PPeF samples. We found that there are minor but significant and reproducible differences. As presented in Figure 8, FTIR shows that a weak additional absorption at 670 cm^{-1} appears in the SL-PPeF only, together with some excess of absorption in the ranges 700-750 cm⁻¹, 1500-1550 cm⁻¹ and 3000-3450 cm⁻¹. Note that the absorption at these higher frequencies is mainly originated by the C-H stretching of the furan rings that are assumed to be involved in the specific interactions. All these results support the idea that there exist some specific interlinkages that would give rise to local supramolecular structures in PPeF molten state and that the development of such specific interactions depends on the way the polymer is cooled from high temperatures.

Figure 8. Effect of the cooling protocol on the room temperature infrared absorption of a PPeF film. The three selected ranges where significant differences are observed are presented. The scheme presented as inset illustrates one of the possible interactions responsible for such FTIR differences, as suggested in Ref 24 .

The presence of local supramolecular structures by slow cooling PPeF, as suggested by BDS, Rheology and FTIR measurements, would impact to some extent the segmental dynamics. According to our findings, the slower component of the α relaxation times is enhanced, which could be explained by some induced correlation in dipole orientation between neighboring entities around the sites where the specific interactions are stronger. The same picture could explain the behavior observed in the secondary relaxation (see Figure 6). By fast cooling PPeF, the development of specific interactions would be partially prevented. As evidenced by the results of the experiment summarized in Figure S5, by heating the FC-PPeF sample to 243 K, the local molecular mobility could be sufficient to allow some conformational changes favoring specific interactions, which would result in a reduction of the slower secondary relaxation component.

The above arguments could also explain the fact that the reduction in the β relaxation intensity observed in the semicrystalline polymer (Aged-PPeF) is far larger than that expected. Particularly, as thoughtfully discussed in previous publications, $58, 70$ - 72 polymer crystallinity affects the local relaxation, mostly in their relaxation strength without other significant changes, usually. In this way, one could expect the following relation:

$$
\chi_{\rm c} \approx 1 - \frac{\varepsilon_{\beta-\text{Semicrys}}^{\prime\prime}}{\varepsilon_{\beta-\text{Amorph}}^{\prime\prime}}\tag{4}
$$

where χ_c would be a measure of the crystallinity, and $\varepsilon''_{\beta-\text{Semicrys}}$ and $\varepsilon''_{\beta-\text{Amorph}}$ the local relaxations for the semicrystalline and amorphous samples, respectively. However, the results from Aged-PPeF showed a quite smaller local relaxation as compared to SL-PPeF. For instance, at 243 K, we found: $1 - \frac{\varepsilon_{B}^{\prime\prime} -$ Semicrys $\frac{\epsilon_B^{\mu}$ -Semicrys ≈ 0.58 a value much higher than that estimated by XRD (17 %).

This much larger reduction of the secondary relaxation strength could be reflecting the structural features discussed above that would modify local packing prior crystal formation. However, in the case of Aged-PPeF this effect can also be due to partial chain orientation occurring during the film preparation.²⁴ In order to test the genuine effect of crystallinity on the local relaxation, we performed new BDS measurements on an already measured PPeF sample that was subsequently maintained between the two electrodes for 2 years. The sample capacitor was left in a desiccator, at room temperature, and previously the new BDS experiments it was subjected to high vacuum for 48 hours to remove any possible trace of moisture. The new BDS measurements were performed starting from 183 K up to 373 K, in 5 K isothermal steps. Then, the sample was cooled in 10 K isothermal steps, down to 183 K, and re-measured on heating again up to 373 K. We also performed corresponding DSC measurements by collecting *c.a.* 5 mg of the polymer that was surrounding the smaller electrode, prior the new BDS experiments. The obtained results are presented in Figure 9, together with the calorimetric traces of the PPeF fast cooled-down from the melt.

Figure 9. (a) DSC results (exo up) of PPeF: aged for 2 years at room temperature inside a BDS capacitor (dotted line), fast cooled from the melt (continuous line) and aged for 3 months at room temperature (dashed line). (b) Dielectric loss spectra, at 243 K, of the PPeF aged for 2 years at room temperature in a BDS

capacitor (empty squares), fast cooled from 370 K after melting (solid squares) and aged for 3 months at room temperature (solid triangles).

The DSC results showed the glass transition temperature at around 289 K for both heating runs, and the Δc_p values were the same (0.37 J/g·K) within error. Considering the endothermic melting peaks above 310 K, we calculated a melting enthalpy of 3 J/g, much lower than the one observed for the Aged-PPeF sample (23 J/g). This new value would correspond to a 2% crystallinity, extrapolating from DSC and XRD results. However, comparing BDS results from the local relaxations we obtained $1 - \frac{\varepsilon_{B}^{\prime\prime} -$ Semicrys $\frac{\varepsilon_{\beta-\text{Semicrys}}^{\prime\prime}}{\varepsilon_{\beta-\text{Amorph}}^{\prime\prime}} \approx 0.33.$ This clearly probes that most of the reduction in the PPeF local-relaxation intensity cannot be accounted by the freezing of local molecular motions in the crystalline phase, pointing to the relevance of the specific interactions affecting local packing.

To further corroborate this result, we performed additional BDS measurements on the same PPeF sample, which was subsequently maintained between the two electrodes for 3 months after melting. The BDS measurements were performed using the same protocol used before and were accompanied by corresponding DSC measurements. The obtained results are also presented in Figure 9. As can be seen, after 3 months the DSC trace showed no signs of crystallization, but the local relaxation presented an evident reduction of intensity, about half of that found after 2 years of aging. Furthermore, this clear change in the local dielectric relaxation was accomplished by only a very minor variation of the main dielectric relaxation intensity in the vicinity of T_g , comparable to the differences observed above between the fast and slow cooled samples (Figure S6). Therefore, these results evidence that there are specific changes in PPeF occurring both below and above *T*g; particularly, the changes occurring at room temperature over a period

of a few months affect significantly the local dielectric relaxation intensity prior to the development of any detectable crystalline phase.

Conclusions

PPeF has a very peculiar behavior compared to the other members of the homologous family, showing no sign of crystallization in regular time scales, but developing a measurable crystallinity when stored at room temperature for months or years depending on sample preparation methods. The crystalline phase so obtained melts at about 333 K, *i.e*. at a temperature much lower than those observed for other furancontaining polyesters, which is indicative of a defective character of the crystals, despite the significant crystallinity. BDS analysis has shown that PPeF molecular dynamics, both at local and segmental range, are sensitive to how fast the material is cooled from high temperatures. The evidenced behavior has been attributed to the development of illdefined structures in the melt, likely related with the presence of specific intermolecular interactions involving the furan rings. Evidences of such local structures have been observed by tiny but reproducible changes in the infrared absorption spectrum. Intermolecular interactions could also be involved in the particular rheological behavior of PPeF, showing a gel-like behavior at temperatures as high as T_g+100 K. Moreover, structural changes occur in PPeF during room temperature storage in months' scale, which reduce significantly the local molecular mobility prior the development of any crystalline structure.

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Supporting Information

Table S1. XRD reflections measured for Aged-PPeF.

Figure S0. XRD peak deconvolution and peak position of Aged-PPeF sample.

Figure S1. DSC (left) and XRD (right) scans of PPeF samples as a function of storage time.

Figure S2. BDS data and fits at 243 K for (a) FC- PPeF, (b) SL-PPeF, (c) Aged-PPeF

Figure S3. BDS data and fits at 303K, 333K, and 348K

Figure S4. A simple physical model of the secondary relaxation is the assumption that the dipolar entities have to overcome an energy barrier separating equivalent positions and that the barriers are distributed following a Gaussian function (of standard deviation σ_E). If the relaxation times follows the Arrhenius equation (Eq. 3 of the manuscript) the secondary relaxation process can be described as the superposition of Debye processes according to a Gaussian distribution of $log_{10}(\tau)$ with a temperature dependent standard deviation $\sigma_{G}(T) \sim \sigma_{E} \frac{\log e}{p_{T}}$ $\frac{\log e}{RT}$.

The symmetrical broadening of the corresponding CC fitting function, b in eqn (1) in the main manuscript can be directly connected with the standard deviation σ_G . To calculate this relationship, we generated $\varepsilon''(\omega)$ functions from the superposition of Debye processes according to a Gaussian distribution of $log_{10}(\tau)$. The obtained curves were fitted to CC functions to obtain the *b* parameter that corresponded to each σ_G (circles in Figure S4). The continuous line in Figure S4 is the best fit to eqn (4) in the main manuscript.

Figure S6. Dielectric loss spectra at (a) 303 K, (b) 333 K, and (c) 348 K for a PPeF capacitor aged for 3 months before melting (solid stars), the same sample after fast cooled (empty triangles) and slow cooled from the melt (solid circles).

