Interplay between structure and relaxation in polyurea networks: the point of view from a novel method of cooperativity analysis of dielectric response†

Nerea Sebastián,ab Christophe Contal,c Antoni Sánchez-Ferrer d,* and Marco Pieruccini*e

The influence of structural constraints on the relaxation dynamics of three polyurea networks with a varying degree of crosslinking, has been studied by means of a thorough analysis of broadband dielectric spectroscopy measurements. Two different relaxation processes are observed, namely, a fast process involving the soft poly(propylene oxide) chains, and a slower and much broader process associated with the immediate surroundings of the hard crosslinkers. Microphase separation in soft and hard domains characterizes the systems in the presence of hydrogen bonding. In this case, different confinement conditions are explored by varying the soft chain length; overall, so called "adsorption" effects dominate. With respect to both cooperativity and the rearrangement energy threshold in fast relaxation, it is found that the enhancement of configurational constraints is similar to cooling, but only on qualitative grounds. An upper bound of the hard domains' interface thickness, in which the slow relaxation is believed to take place, is estimated from the analysis of the fast relaxation in the system characterized by the highest degree of confinement, taking into account the results of the structural analysis. Dropping the hydrogen bonding mechanism, phase separation does not occur anymore and the configurational constraints at the ends of the soft chains are reduced, leaving just those imposed by the rigid crosslinkers. This leads to a significant increase in cooperativity on approaching the glass transition, and to a complex behavior that is thoroughly discussed in comparison with those observed in the micro-segregated systems.

1 Introduction

Molecular dynamics in soft matter is a main subject of debate in condensed matter physics; this does not only have to do with potential industrial interests, but also with knowledge in its most general and fundamental sense. Indeed, notwithstanding the considerable efforts that have been spent over many years in the field, basic aspects of this branch of physics still remain poorly understood. Among others, the mechanisms leading to the formation of a glass from a supercooled liquid yet represent a matter of lively debate. This situation is also a bit confusing, given the use of glasses for centuries.

When a liquid is cooled down and crystallization is avoided, its viscosity progressively increases and, sooner or later, it will form a glass: a solid-like state in which long range molecular order is absent. On approaching this state, the central relaxation time of the configurational motion often shows a super-arrhenian temperature dependence, that is, the apparent activation energy increases on cooling. This behaviour hinted at the possibility that the configurational transitions responsible for the viscosity might involve an ever-increasing number of rearranging units as the glassy state is approached. The quest for a characteristic length of the glass transition thus ran initially through seeking its signature in structural features. Later, it was realized that this length would more appropriately relate to dynamic correlations.

Probing this length, though, is not trivial and several approaches have been (and are being) developed in order to
grasp it from relaxation experiments. Thus, Donth \textsuperscript{1,2} provided a fluctuation theory whereby a characteristic size of cooperatively rearranging regions (CRRs) in a supercooled liquid – i.e., domains where rearrangements may occur without inducing any configurational change in the rest – could be estimated from the temperature fluctuations accompanying the energy exchanges between configurational and non-configurational degrees of freedom.\textsuperscript{3} Some of the most direct evidence for the increase in the CRR size on cooling has been reported in recent investigations on polystyrene and poly(methyl methacrylate), based on modulated calorimetry.\textsuperscript{4}

Another theoretical approach has been developed by Berthier and co-workers, who focused instead on the generalized (4-point) susceptibility \( \chi_{4}(\mathbf{r},T) = \langle \phi(\mathbf{r},T)\phi(\mathbf{r} + \mathbf{y},T) \rangle \), describing correlations between relaxation dynamics, \( \phi(x,t) \), at \( x = \mathbf{r} \) and \( x = \mathbf{r} + \mathbf{y} \), or on the 3-point susceptibility \( \chi_{3}(\mathbf{r},T) = \langle h(\mathbf{r},0)\phi(\mathbf{r} + \mathbf{y},T) \rangle \) relating to the effect of an enthalpy fluctuation at \( \mathbf{r} \) on the relaxation dynamics at \( \mathbf{r} + \mathbf{y} \). (in particular, the second one is directly accessible from experiment, e.g., \( \chi_{3}(0,T) = \phi(0,T)/\phi(T) \). Also within this framework, evidence has been observed of a growing characteristic length on cooling a relaxing liquid.

More recently a further method has been proposed, which considers more closely the features of the energy landscape in a relaxing liquid.\textsuperscript{5,6} In this approach, the cooperativity is derived by fitting the configurational relaxation function, extracted from the data, with a theoretical expression derived within a statistical mechanical framework. The basic idea behind this is that the diffusional motion characterising the rearrangements of units over length scales on the order of the CRR size follows from facilitation by a pre-transitional dynamic state of fluctuational nature.

Also with the latter method, of course, an increase in the cooperativity of a liquid (either polymeric or not) under progressive cooling towards its glass transition has been found; the agreement with Donth’s estimates was good.\textsuperscript{3,8} At the present state of the art, the last circumstance is extremely important, because in the absence of direct measurements of the cooperativity length, a figure of reliability in the analysis is represented by the comparison with the results of independent methods of estimation.

When a certain characteristic length \( L_{h} \) is introduced in a polymeric system through appropriate structural heterogeneities (such as crosslinks, dispersed nano-particles or even the walls of confining nano-pores),\textsuperscript{9,10} it is expected that a change in the relaxation behaviour occurs when the dynamic correlation length approaches that value. On the one hand, this circumstance provides some objective support to the existence of a cooperativity length; on the other, it allows one to probe the relaxation dynamics \textit{via} a proper tailoring of the system.

Elastomers are crosslinked polymer melts having characteristics from the three physical states. They are soft solids which can be shaped due to the presence of crosslinkers, which prevent the polymer melt from flowing; they contain a main component which is a viscous liquid; and the retractive force appearing upon stretching is entropically driven, similar to the case of gases.\textsuperscript{11} The chain mobility in such materials decreases upon cooling down due to the thermal inactivation of conformational changes, and the reduced free volume in the surroundings of the chains.\textsuperscript{12} Therefore, the viscosity increases and the polymer chains freeze into the glassy state where a conformation change is no longer possible, but reorientation of short segments in the polymer backbone or side groups occurs. The high viscosity value at low temperatures in elastomeric networks is the main cause of the slowing down of the dynamics of such systems, together with parameters introducing asymmetry or disorder to the system, e.g., polydispersity, crosslinking, entanglements, looping, dangling ends, supra-molecular interactions.

Polyurea networks with varying degrees of crosslinking have been recently investigated in relation to their mechanical relaxation properties.\textsuperscript{13} In these systems, the presence of hydrogen bonding between urea linkages leads to the formation of nano-segregated hard domains, connected one with the other by flexible chains. By changing the length of the latter, different mechanical constraints on the chains’ configurations were imposed. Considering systems made of soft polyetheramine chains with \( M_{c} \sim 400, \ 2000 \) and \( \ 4000 \ \text{g mol}^{-1} \) segmental molecular weights, it was possible to explore structures turning from an ensemble of hard domains embedded in matrices of increasing rigidity, to a condition in which isolated domains of limited mobility are embedded in an environment of percolated hard domains (\textit{i.e.}, \( M_{c} \sim 400 \ \text{g mol}^{-1} \)). In the less constrained systems, it was possible to separate two distinct segmental relaxations: the faster one associated with the embedding matrix, and the other with a stiffer layer surrounding the hard domains (each with its own glass transition temperature). In the sample with the shortest flexible chain, these processes tended to merge into one and the glass transition temperature of the softer component increased significantly, mainly due to the more restricted mobility of the soft polymer segment pulled from both extremes by the interfacial polymer layer which is well transmitted along the short soft polymer backbone. Such a slowing down of the dynamics can be attributed to the overwhelming effect of the interaction with the confining walls (so called “adsorption”),\textsuperscript{14} over the acceleration expected from pure confinement. This is indeed related to the soft chain pinning to the hard nano-particles.

There are several aspects which deserve to be investigated in this context; one among the others is the interplay between temperature and confinement in the regime of adsorption. In other words, concerning the configurational relaxation, one may question to what extent the enhancement of the constraints can be considered similar to a decrease in temperature. Another aspect regards how things change when the constraints on soft chain ends depletion to some extent.

These two issues represent the main concern of this paper. For the first one, we shall perform dielectric relaxation analyses (necessarily in isothermal conditions), on the systems of ref. 13 with \( M_{c} \sim 2000 \) and \( 4000 \ \text{g mol}^{-1} \) soft chains. For the second, comparisons will be done among the relaxation dynamics thus probed, when using primary or secondary amines with the same \( M_{c} \sim 2000 \ \text{g mol}^{-1} \) in which hydrogen bondings are
not suppressed in the latter, thus allowing a limited mobility to the soft chain ends.

The cooperativity associated with the fast segmental process, and its impact on the structural features of the systems, will be thoroughly discussed by means of the method of ref. 3 and 7. The results will be also compared with the lower bound estimates provided by the 3-point correlation function analysis.5,6 The discussion will be done with reference to structural information extracted by means of large and small angle X-ray scattering (WAXS and SAXS respectively) and by mappings from atomic force microscopy (AFM).

2 Experimental
Sample preparation

The three linear hydrophobic diamino-terminated polyetheramines Jeffamine® D-2000 ($M_n = 2056$ g mol$^{-1}$, $\rho = 991$ kg m$^{-3}$), D-4000 ($M_n = 4000$ g mol$^{-1}$, $\rho = 994$ kg m$^{-3}$) and SD-2001 ($M_n = 2050$ g mol$^{-1}$, $\rho = 978$ kg m$^{-3}$) from Huntsman International LLC, and the triisocyanate crosslinker Basonat HI-100 ($M_w = 504$ g mol$^{-1}$, $\rho = 1174$ kg m$^{-3}$) from BASF SE were used as received. Polyurea (PU) networks are formed by the rapid chemical reaction between a triisocyanate and a diamine to build urea moieties, as shown in Fig. 1. Details of the synthesis of these PU elastomers are well described in the literature.13,15,16 Table 1 compares the main structural characteristics for the three used samples ED-2000, ED-4000 and ESD-2001. The structure of polyurea chains is shown in Fig. 1;

<table>
<thead>
<tr>
<th>Sample</th>
<th>ED-2000</th>
<th>ED-4000</th>
<th>ESD-2001</th>
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<tr>
<td>$M_n$ (g mol$^{-1}$)</td>
<td>2056</td>
<td>4000</td>
<td>2050</td>
</tr>
<tr>
<td>$\rho$ (kg m$^{-3}$)</td>
<td>994</td>
<td>994</td>
<td>978</td>
</tr>
<tr>
<td>$L_s$ (nm)</td>
<td>12.0</td>
<td>24.4</td>
<td>12.0</td>
</tr>
<tr>
<td>$R_s$ (nm)</td>
<td>8.9</td>
<td>18.1</td>
<td>8.9</td>
</tr>
<tr>
<td>$d_{SAXS}$ (nm)</td>
<td>5.23</td>
<td>7.56</td>
<td>—</td>
</tr>
<tr>
<td>$d_{AFM}$ (nm)</td>
<td>5.58</td>
<td>7.94</td>
<td>—</td>
</tr>
</tbody>
</table>

Because of the fast segmental process, the long timescale dynamics of the system is not affected by it.

<table>
<thead>
<tr>
<th>Legend</th>
<th>ED-2000</th>
<th>ED-4000</th>
<th>ESD-2001</th>
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</table>
| a Values obtained by titration. b All-trans conformation of the fully stretched polymer. c Estimated by simulating a ca. 5 repeating unit helical structure. d Values from the stoichiometry used to prepare the samples.

due to strong hydrogen bonding between urea linkages of neighboring chains or neighboring segments in the same chain, the hard segment microphase segregates into so-called “hard domains”. That is the case for samples ED-2000 and ED-4000, whose genitor polymer is a primary amine, but not for the sample ESD-2001 coming from the reaction between a bulky secondary amine-containing polymer and the crosslinker.

Methods

The complex dielectric permittivity $\varepsilon''(\omega) = \varepsilon'(\omega) - i\varepsilon''(\omega)$ was measured over a broad frequency range ($10^2$–$10^9$ Hz) by combining two impedance analyzers: Alpha-A + ZG4 (Novocontrol) for low frequencies and HP4192A for high frequencies. In the latter setup (high frequencies), the cell consists of a parallel plate capacitor placed at the end of a coaxial line. Due to geometry constraints the capacitor consists of two 5 mm diameter gold plated electrodes separated by the polymer as a dielectric. A modified HP16901A coaxial test fixture was used as the sample holder. In order to obtain larger capacitance values at low frequencies, larger samples of 15 mm diameter were employed for the low frequency measurements, with the sample placed between parallel plate electrodes at the end of a homebuilt four-wire holder. Very careful calibration of both systems ensured good matching between the two sets of measurements. In both setups, all the three samples consisted of thin uniform films with a thickness around 0.3–0.4 mm, which were held in a cryostat. Isothermal spectra were recorded at constant temperature with a resolution of 0.1 °C under nitrogen atmosphere.

Damping mode atomic force microscopy (AFM) was carried out using a Multimode 8 scanning force microscope (Bruker). NanoSensors AFM cantilevers for tapping mode (NC1HR) in soft and hard tapping conditions were used at a typical resonant frequency of 300 kHz. Images were simply flattened and analyzed using NanoScope Analysis 1.5 software, and no further image processing was carried out.

Simultaneous small- and wide-angle X-ray scattering (SAXS and WAXS) experiments were performed using a Philips PW 1730 rotating anode (4 kW) in order to obtain direct...
information on the SAXS and WAXS reflections. Cu Kα radiation (λ = 1.5418 Å) filtered by a graphite monochromator and collimated by a 0.8 mm collimator was used. The incident beam was normal to the surface of the film. The scattered X-ray intensity was detected by a Schneider image plate system (700 × 700 pixels, 250 μm resolution). An effective scattering-vector range of 0.1 nm⁻¹ < q < 30 nm⁻¹ is obtained, where q is the scattering wave-vector defined as q = 4πsin(θ)/λ, with a scattering angle of 2θ.

Differential scanning calorimetry (DSC) experiments (Fig. S1, ESI†) were carried out on a Netzsch DSC 200 F3 Maia apparatus with heating and cooling-rates of 1 K min⁻¹ under nitrogen atmosphere, using 20 mL aluminum pans with holes.

3 Results

All characteristics for the three elastomeric samples such as the number average molar mass (Mₐ) of the polymer, the corresponding degree of polymerization (DP), the polymer contour length (Lₐ), the polymer end-to-end distance (Rₑ), and the polymer volume fraction (φpolymer) are reported in Table 1. Moreover, since polyurea elastomers are microphase separated materials with soft and hard domains, the hard-to-hard domain distance was evaluated by SAXS (dSAXS) measurements where values of 5.2 and 7.6 nm for the ED-2000 and ED-4000 elastomers were found, respectively, together with the amorphous nature of the polymer backbone as shown from the WAXS profiles (Fig. S2, ESI†). Complementary information was gained from AFM experiment on bulk or on casted elastomeric films (Fig. 1 and Fig. S3, S4, ESI†). AFM images of ED-2000 and ED-4000 show a nanoscale rod-like structure corresponding to the micro-segregation of the hard domains from the soft matrix. The average hard-to-hard distance (dAFM) was evaluated from the Fast Fourier Transform (FFT) of AFM images, and values of 5.6 and 7.9 nm for the ED-2000 and ED-4000 elastomers were obtained, respectively (Fig. S5, ESI†). On the other hand, the sample ESD-2001 showed neither any characteristic peak in SAXS nor in the AFM-FFT evaluation. The reason for the absence of such a characteristic distance is due to the impossibility to establish hydrogen bonds between the urea motifs because of the presence of bulky groups in the secondary amine of the polymer. Thus, the formation of hydrogen bonds in the ED-2000 and ED-4000 elastomeric samples should influence the mechanical properties when compared to the sample ESD-2001 without hydrogen bonds.

Molecular dynamics in the ED-4000, ED-2000 and ESD-2001 polyurea elastomers were investigated by means of broadband dielectric spectroscopy over a wide temperature range. Frequency and temperature dependencies of the dielectric losses for the ED-4000 polyurea elastomer are shown in Fig. 2, and the equivalent representations are given in Fig. S6 (ESI†) for both ED-2000 and ESD-2001 polyurea elastomers. At first glance, all three polymer networks exhibit similar behaviour, i.e., a dielectric spectrum characterized by two relaxation processes, which can be associated with the segmental chain dynamics (i) of the less constrained chain segments embedded in the soft domains (relaxation at higher frequencies or ω₁) and (ii) of the most restricted chain segments directly attached to the hard domains (relaxation at lower frequencies or 1/ω₂), as previously observed for polyurea systems by dielectric spectroscopy and from dynamic mechanical analysis measurements. However, clear differences are also evident when comparing the results obtained from the three samples as shown in the isochronal representation of Fig. 3a. The 1/ω₁-relaxation is predominant in ESD-2001, while for ED-4000 and ED-2000 the 1/ω₂-relaxation and the 1/ω₃-process appear as a pronounced and a broad peak, respectively. However, both relaxation processes can be clearly discerned in the isothermal representations in Fig. 3b and Fig. S7 (ESI†). At temperatures above 40 °C, the shape of the peak related to the 1/ω₃-relaxation process becomes sharper, which can be explained by the progressive rupture of hydrogen-bonding between the urea motifs, and thus by the increase of the dynamics of the chain segments directly attached to the urea motif.

For each temperature, the results were fitted to

$$\varepsilon(\omega) = \sum_k \Delta \varepsilon_k(\omega) + \varepsilon_\infty - i \frac{\sigma_0}{\omega \varepsilon_0} \quad (1)$$

where \(\varepsilon_\infty\) is the high frequency permittivity and \(\sigma_0\) is the dc-conductivity. Both the \(\omega_1\) and the \(\omega_2\)-relaxation process have been characterized by simultaneously fitting the real and imaginary dielectric permittivity to the Havriliak-Negami (HN) function:

$$\Delta \varepsilon_k(\omega) = \frac{\Delta \varepsilon_k}{[1 + (i \omega \tau_k)^{\alpha_k}]^{\beta_k}} \quad (2)$$

where \(\Delta \varepsilon_k\) is the strength of each mode. The parameters \(a_k\) and \(b_k\) (0 < \(a_k\) ≤ 1 and 0 < \(b_k\) ≤ 1) describe the broadness and symmetry of the relaxation spectra, respectively. The frequency
of maximal loss is related to the shape parameters by the following equation:\textsuperscript{20}

\[ f_{\text{max}, k} = \frac{1}{2\pi \tau_{HN, k}} \left[ \sin\left( \frac{\pi a_k}{2 + 2b_k} \right) \right]^{1/a_k} \left[ \sin\left( \frac{\pi b_k a_k}{2 + 2b_k} \right) \right]^{-1/a_k} (3) \]

It should be noted that in all three samples there is a great increase of both \( \varepsilon' \) and \( \varepsilon'' \) at low frequencies, which arises not only from dc-conductivity but also can be related to the interfacial polarization at the internal boundaries between the hard and soft domains. This contribution masks the low frequency segmental relaxation and makes it difficult to unambiguously determine the shape parameters for the low frequency relaxation process or \( a_0 \). To limit its impact on the results, a Debye relaxation could be easily added at low frequencies (Fig. S8 and S9, ESI\textsuperscript{†}) for samples ED-2000 and ESD-2001. However, this approach could not be applied for ED-4000, and the value of the electronic conductivity was determined in the modulus representation, and then fixed in eqn (1) (Fig. S10, ESI\textsuperscript{†}).\textsuperscript{20,21}

The shape parameter values \( a_k \) and \( b_k \) are summarized in Table 2 for the two relaxation processes and for the three samples.

<table>
<thead>
<tr>
<th></th>
<th>( a_k )</th>
<th>( b_k )</th>
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<tbody>
<tr>
<td>ED-4000</td>
<td>0.70–0.75</td>
<td>0.34–0.60</td>
</tr>
<tr>
<td>ED-2000</td>
<td>0.40–0.50</td>
<td>0.6</td>
</tr>
<tr>
<td>ESD-2001</td>
<td>0.65–0.70</td>
<td>0.60</td>
</tr>
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</table>

The obtained relaxation frequencies of both relaxation processes as function of temperature for the three samples are gathered in Fig. 4a, which follow the Vogel–Fulcher–Tamman law (VFT)

\[ f_k = f_0 \exp\left\{ -\frac{B_k}{T - T_{0,k}} \right\} (4) \]

![Table 2 Relaxation shape parameters, \( a_k \) and \( b_k \), for both the \( \alpha \) and \( \alpha' \)-processes, and for the three elastomeric samples](image)

![Fig. 3](image) (a) Temperature dependence of the real \( (\varepsilon') \) and imaginary \( (\varepsilon'') \) components of the complex dielectric permittivity at 10 Hz, for the three elastomeric samples. (b) Complex dielectric permittivity vs. frequency on a logarithmic scale for the ED-4000 sample at different temperatures. In both, full and empty symbols correspond to the real and imaginary component, respectively.

![Fig. 4](image) (a) Arrhenius plot of the frequency of the maximum loss for the \( \alpha \)- (filled symbols) and the \( \alpha' \)-process (empty symbols) – as deduced from the fits to eqn (1) of the three elastomeric samples, together with the corresponding fits to the VFT law. (b) Angell’s plot of the relaxation times for the \( \alpha \)- (filled symbols) and the \( \alpha' \)-process (empty symbols) for the three elastomeric samples, together with the corresponding fits to the VFT law.
where the pre-exponential factor $f_{0,bs}$ the so-called Vogel temperature $T_{0,bs}$, and the exponential factor $B_\varepsilon$ (related to the fragility and to the apparent activation energy) are the fitting parameters. The dielectric glass transition temperature $T_g$ is calculated when the corresponding structural relaxation time is $\tau_{\text{max}} = 100$ s. Although in calorimetric measurements a single glass transition is observed, dielectric measurements clearly show two segmental relaxation processes associated with a dielectric glass transition. The best VFT fits are plotted in Fig. 4a, and the corresponding fitting parameters together with the obtained $T_g$ values are reported in Table 3, and compared to those obtained by dynamic mechanical analysis (DMA) and by differential scanning calorimetry (DSC, Fig. S1, ESI†) measurements.$^{13,15,16}$ It should be pointed out that, in DMA measurements, the storage and loss moduli are obtained at a given frequency as a function of temperature and the VFT behaviour is assumed in the fitting procedure. Thus, it is noteworthy that despite the difference between the methods, values are in reasonable agreement.

In Fig. 4a, the $\alpha$-relaxation process looks similar for the three samples at high temperatures, and the mobility is slightly but noticeably higher for the sample ED-4000 due to its higher segmental molecular weight. However, such a difference is enhanced as the temperature decreases. On the other hand, the slow $\alpha'$-relaxation process behaviour clearly differs between the samples with hydrogen bonding (ED-4000 and ED-2000) and the sample without (ESD-2001), which has a larger and more thermally activated characteristic frequency.

Poly(propylene oxide), PPO, is one of the few polymers which show two characteristic relaxation modes in dielectric spectroscopy: one related to the segmental relaxation mode (at high frequencies), and the second connected to the end-to-end distance characteristic of amorphous polymers (at lower frequencies).$^{22}$ The latter is present when the repeating monomer unit has no plane of symmetry perpendicular to the contour of the polymer backbone.$^{23}$ In a previous paper, two mechanically active modes were observed for both hydrogen bonded samples, i.e., ED-2000 and ED-4000, which correspond to the segmental dynamics of the $\alpha$- and $\alpha'$-relaxation processes.$^{15}$ Thus, in this work, we could assign the low frequency peak to the normal mode for the ESD-2001 sample, but unambiguously to the $\alpha'$-relaxation process for the segregated systems ED-2000 and ED-4000, where the normal mode is suppressed due to pinning of the PPO chains by means of hydrogen bonding. This matches the picture of nano-segregated hard domains dispersed in a soft matrix, with the faster $\alpha$-relaxation process related to the mobility of chain segments embedded in the soft domains and the slow $\alpha'$-relaxation process arising from the restricted mobility of those chain segments in the vicinity of the hard domains. In this way, hydrogen-bonds further decrease the mobility of the restricted chain segments, thus the rupture of the physical interaction between them is expected to result in similar mobility for the three samples at temperatures higher than 70 °C. The longer segmental molecular weight in sample ED-4000 results in a higher mobility for the chain segments in the soft domain, corresponding to the $\alpha$-relaxation process, as seen in Fig. 4a. Thus, the mobility at high temperatures is similar for both the elastomeric samples ED-2000 and ESD-2001 with comparable segmental molecular weight, and increasingly differs on cooling due to the constraints imposed on the dynamical correlation length when hydrogen-bonding is formed for the sample ED-2000 which at the same time decreases $T_g$.

In order to quantify the change in the dynamics when approaching $T_g$, we can make use of the fragility metric, which describes to what extent dynamics differ from the basic Arrhenius behavior. Following the formalism introduced by Angell,$^{24–28}$ the fragility index is defined by the slope value at $T_g$ for a log$_{10}\tau$ vs. $T_g/T$ representation (Fig. 4b). The minimum $m$ value is assigned to conventional Arrhenius behaviour $\tau = \tau_0 \exp[\varepsilon/(k_B T)]$, which corresponds to $m = 16$ for the pre-factor $\tau_0 = 10^{-14}$ s. According to this metric, glass-forming materials can be classified as “fragile” ($m > 50$) and “strong” ($m < 30$). In the present case, all materials show two segmental relaxation processes, whose dynamics freeze at different $T_g$, and thus a fragility value $m$ can be obtained for each relaxation mode. Fits to the VFT law (Fig. 4b) allow one to calculate the fragility index $m$ as:

$$m = \frac{B_\varepsilon}{\ln(10)T_g} \left[ \frac{d\log_{10}\tau_g}{dT_g/T} \right]_{T=T_g}$$

(5)

The $m$ values obtained for ED-4000, ED-2000 and ESD-2001 are given in Table 3. For all three materials, values slightly above 100 for the fast $\alpha$-relaxation process were obtained.

| Table 3 | The Vogel–Fulcher–Tamman (VFT) fitting parameter, $B_\varepsilon$, $\tau_0$, and $T_0$, and the $T_g$ values obtained by broadband dielectric spectroscopy, dynamic mechanical analysis (DMA) and differential scanning calorimetry (DSC) |
|----------|----------|----------|----------|----------|
| ED-4000  | $\alpha$  | $\alpha'$ | $\alpha$  | $\alpha'$ |
| $B_\varepsilon$ | 945 ± 18 | 2477 ± 87 | 944 ± 12 | 3081 ± 70 |
| $\tau_0$ (s) | $10^{-12.4}$ | $10^{-12.0}$ | $10^{-12.2}$ | $10^{-12.7}$ |
| $T_0$ (K) | 175 ± 1 | 137 ± 2 | 180 ± 1 | 120 ± 2 |
| $T_g$ (K) | 204 ± 1 | 213 ± 7 | 208 ± 1 | 211 ± 5 |
| $T_g$ (DMA) | 201 ± 1 | 222 ± 1 | 199 ± 1 | 208 ± 2 |
| $T_g$ (DSC) | 206 ± 1 | — | 213 ± 1 | — |
| $m$-index | 103 ± 9 | 39 ± 6 | 102 ± 10 | 34 ± 4 |

The normal mode is suppressed due to pinning of the PPO chains by means of hydrogen bonding. This matches the picture of nano-segregated hard domains dispersed in a soft matrix, with the faster $\alpha$-relaxation process related to the mobility of chain segments embedded in the soft domains and the slow $\alpha'$-relaxation process arising from the restricted mobility of those chain segments in the vicinity of the hard domains. In this way, hydrogen-bonds further decrease the mobility of the restricted chain segments, thus the rupture of the physical interaction between them is expected to result in similar mobility for the three samples at temperatures higher than 70 °C. The longer segmental molecular weight in sample ED-4000 results in a higher mobility for the chain segments in the soft domain, corresponding to the $\alpha$-relaxation process, as seen in Fig. 4a. Thus, the mobility at high temperatures is similar for both the elastomeric samples ED-2000 and ESD-2001 with comparable segmental molecular weight, and increasingly differs on cooling due to the constraints imposed on the dynamical correlation length when hydrogen-bonding is formed for the sample ED-2000 which at the same time decreases $T_g$.

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(5)

The $m$ values obtained for ED-4000, ED-2000 and ESD-2001 are given in Table 3. For all three materials, values slightly above 100 for the fast $\alpha$-relaxation process were obtained.

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<tbody>
<tr>
<td>ED-4000</td>
<td>$\alpha$</td>
<td>$\alpha'$</td>
<td>$\alpha$</td>
<td>$\alpha'$</td>
</tr>
<tr>
<td>$B_\varepsilon$</td>
<td>945 ± 18</td>
<td>2477 ± 87</td>
<td>944 ± 12</td>
<td>3081 ± 70</td>
</tr>
<tr>
<td>$\tau_0$ (s)</td>
<td>$10^{-12.4}$</td>
<td>$10^{-12.0}$</td>
<td>$10^{-12.2}$</td>
<td>$10^{-12.7}$</td>
</tr>
<tr>
<td>$T_0$ (K)</td>
<td>175 ± 1</td>
<td>137 ± 2</td>
<td>180 ± 1</td>
<td>120 ± 2</td>
</tr>
<tr>
<td>$T_g$ (K)</td>
<td>204 ± 1</td>
<td>213 ± 7</td>
<td>208 ± 1</td>
<td>211 ± 5</td>
</tr>
<tr>
<td>$T_g$ (DMA)</td>
<td>201 ± 1</td>
<td>222 ± 1</td>
<td>199 ± 1</td>
<td>208 ± 2</td>
</tr>
<tr>
<td>$T_g$ (DSC)</td>
<td>206 ± 1</td>
<td>—</td>
<td>213 ± 1</td>
<td>—</td>
</tr>
<tr>
<td>$m$-index</td>
<td>103 ± 9</td>
<td>39 ± 6</td>
<td>102 ± 10</td>
<td>34 ± 4</td>
</tr>
</tbody>
</table>
However, interesting differences can be noticed in the case of the slow $\alpha'$-relaxation process. The fragility in this case is greatly affected by the hydrogen-bonds between the urea motifs, taking values of around 35 for ED-2000 and ED-4000, and almost double for EDS-2001, which accounts for the restricted dynamics of the chain segments near the hard domains.

4 Information from the shape parameters of the $\alpha$-mode

The picture provided by the VFT analysis in the previous section can be refined if the relaxation frequency profiles are also taken into account. The conductivity component characterizing our systems, however, is too large to let the shape parameters of the $\alpha'$-process be assigned without ambiguity. On the contrary, the $\alpha$-process contribution is sufficiently well defined in order to allow for a reliable analysis of the cooperativity. Before doing this, we give a brief outline of the theoretical scheme that underlies our data treatment; further and more thorough details can be found in ref. 7.

Basic description of segmental relaxation

The $\alpha$ dynamic regime in polymeric melts refers to a condition whereby the diffusional motion of segments is promoted by activated processes. The energy threshold $\zeta$ of a rearranging unit, however, is not constant in time, as it varies continually due to local configurational changes; rather, each unit explores a distribution $p(\zeta)$ of such barriers. In a stationary state, a constant excess chemical potential $\Delta\mu(T) > 0$ is then established, which describes the deviation of the liquid from equilibrium at the temperature $T$.

Configurational transitions take place by facilitation from precursory (or pre-transitional) excited states in which the rearranging units move about fixed sites, probing the $\zeta$-space that is locally accessible. Such non-diffusive dynamics is characterized by a certain energy per unit which has been gained by fluctuation; when this energy is so large that the probability of the excited state gets lower than that associated with a completely unconstrained mobility state – with ideal specific configurational entropy $s_c$ – then a small number $z$ of units may acquire a diffusional component in their overall motion (in other words, the acquisition of a diffusional component in the motion is a spontaneous process). The local constraints that have contributed until then to hinder large-scale configurational changes in the surroundings deplete; more units start diffusing, further disentangling the neighboring ones in a sort of a cascade process. Facilitation, as just described, eventually leads to the formation of a high mobility spot – the Cooperatively Rearranging Region (CRR) – surrounded by glassy domains (possibly, local conformations characterized by high rearrangement barriers are pushed away from the spots’ bodies to bank up at some liquid/glass interface). The number of units forming such mobility domains is on the order of

$$N_\alpha \approx z(\lambda + 1)$$  \hspace{1cm} (6)

where $\lambda$ (in general $> z$) is the Lagrange multiplier associated with the condition that $\Delta\mu$ is held constant when extremizing the thermodynamic potential for the derivation of $p(\zeta)$. More intuitively, the $z$ units may initiate a large-scale rearrangement provided other $\lambda$ units are dynamically correlated with each of them.

The nature of the CRRs is fluctuational; they form and are subsequently re-absorbed in the quasi-glassy liquid, so their total number is a non-conserved quantity. This means that a chemical potential can be associated with them and that it must be zero, implying in turn that the relation

$$\lambda \approx [1 - w(\zeta^*)] \frac{\zeta^*}{T_{S_c}}$$  \hspace{1cm} (7)

holds, where $w \equiv w(\zeta)$ is the probability that a unit has an energy not lower than $\zeta$, and $\zeta^*$ is the average barrier height that each unit of a CRR has to overcome in order that facilitation initiates, leading eventually to a large scale configurational change. Note that, in proximity to $T_g$, $\zeta$ is in fact close to the average energy per unit needed to initiate the large scale rearrangement from a non-diffusive pre-transitional dynamic state. This relation is an important indirect link between $\zeta^*$ and $N_\alpha$.

Concerning the CRR shape, the scheme proposed in ref. 7 does not provide any prescription; thus, besides the common picture of compact structures,3,4 the possibility of branched domains can also be envisaged.29

Data analysis

Information about $\lambda$ and $z$ is carried by the relaxation function $\varphi_{\exp}(t)$ associated with the $\alpha$-process. $\varphi_{\exp}(t)$ can be obtained from the Fourier cosine transform of the HN frequency loss profile in the complex permittivity $\varepsilon(\omega)$. The obtained expression can be then fitted with the function

$$\varphi_{\exp}(t) = \int_0^\infty d\zeta p(\zeta) \exp\left\{-t\nu^* e^{-z\Delta\mu(\zeta)/k_B T}\right\}$$  \hspace{1cm} (8)

where $\nu^*$ is an attempt rate and $\Delta\mu(\zeta) = -k_B T \ln w(\zeta)$ is the excess chemical potential associated with a barrier height $\zeta$ (its $p$-weighted average yields the chemical potential $\Delta\mu$ mentioned at the beginning of the preceding subsection). Of course, $\lambda$ enters $p(\zeta)$ as a parameter and in fact, not too far from $T_g$, $p(\zeta) \approx \exp\{-\lambda \Delta\mu(\zeta)/k_B T\}$.

The information provided by the analysis of $\varphi_{\exp}(t)$ is related directly to the precursory state, since the parameters $z$ and $\lambda$ characterize the local dynamics immediately preceding the large scale rearrangements.7 The attempt rate in eqn (8) is expected to depend on $T$. For the present purposes, it suffices to treat it as a fitting parameter.

Results and discussion

Fig. 5 shows that in all samples the rearrangement threshold eventually increases when the proper $T_g$ is approached on cooling. The absorbas report the difference $T - T_g$, and steeper growths are found where configurational restrictions are weaker; this means that increasing $\zeta^*$ by enhancing the constraints is not equivalent to cooling a less constrained system, though it
Comparison of ED-2000 with ED-4000 shows that the energy barrier $\zeta^*$ at fixed $T - T_g$ is larger for shorter soft chains. Concurrently, both the central relaxation time and the peak width (in the $-50^\circ C \leq T \leq 30^\circ C$ interval) increase upon shortening the soft chain length. This conforms to constraining conditions in the adsorption regime of nano-pore confinement, that is, when the relaxing regions are strongly coupled with the walls of the confining volume; indeed, this coupling is represented here by the soft chain pinning to the hard domains.

A possible reason for the increase of $\zeta^*$ is the emergence – or increasing significance – of additional mechanisms, not directly related to facilitation, through which the energy of the pre-transitional state can return to the heat bath. In this case, the characteristic time facilitation takes to start from the excited state would compete with the lifetime of the precursory state itself, which is shortened by the presence of these additional paths of energy loss. Larger initial energies of the pre-transitional dynamic state – and also cooperativities, after eqn (6) and (7) – would thus be needed in order that the energy of the precursory state still remains high enough to render facilitation thermodynamically advantageous (i.e., spontaneous) should it happen to initiate. Incidentally, the emergence of new paths of energy regression could be hypothesized also in the tentative explanation of the $\zeta$ increase at higher temperatures noticed above, that is found on heating above temperatures specific to each of the samples. Of course, a reliable description of the heat transfer among different degrees of freedom (i.e., vibrational and configurational) would be required to support this hypothesis, but we must defer this issue to future work.

The overall slowing down of the ED-2000 dynamics compared to ED-4000 is, of course, associated with a shortening of the soft chains, since the way constraints are introduced and the nature of the soft/hard interface are the same for both. Such a situation seems to be counterintuitive, because a restriction of the available volume would be expected to also limit the mean size of the CRRs and, together with it, the activation energy for the configurational changes. On the other hand, cooperativity is not just a consequence of a “drag”effect characterizing the diffusional motion of rearranging units. Rather, what we observe is in fact the precursory state itself, which is shortened by the presence of these additional constraints in the adsorption regime of nano-pore confinement,30 that is, when the relaxing regions are strongly coupled with the walls of the confining volume; indeed, this coupling is represented here by the soft chain pinning to the hard domains.

For a comparison, we also report in Fig. 6 the results found by means of the 4-point correlation function approach.5,13,29,31 In that framework, indeed, a lower limit to the number of correlated units is estimated by the equation

$$N_{corr} = \frac{k_B}{\Delta C_p} \left[ \frac{\beta(T)}{e} \right]^2 \left( \frac{d \ln \tau_\alpha}{d \ln T} \right)^2,$$

where $e$ is the Euler number, while $\beta$ and $\tau_\alpha$ parametrize the Kohlrausch-Williams-Watts (KWW) function, exp$[-(t/\tau_\alpha)^\beta])$, that reproduces our $\varphi_{exp}(t)$ at each temperature in the relevant
time interval. The agreement between \( N_s \) and \( N_{\text{corr}} \) has to be considered good, also from a quantitative viewpoint.

The cooperativity in ED-2000 increases on cooling at a lower rate than in ED-4000. In this respect, the behaviour shown in Fig. 6 does not differ qualitatively from that of Fig. 5 – except in that no upwards bending is found on heating at high temperatures. Considering the general tendency of cooperativity to grow on cooling, this difference in slope may be intuitively interpreted as a saturation effect of geometric nature, since the CRR growth is contrasted by the presence of chain pinning points and the soft domains in ED-2000 are smaller. Unfortunately, \( \lambda \)-relaxation patterns at temperatures closer to \( T_c \) cannot be extracted in the present conditions, and a crossing between the ED-2000 and ED-4000 lines can only be guessed.

Focusing now on ESD-2001, Fig. 6 shows that its cooperativity line crosses that of ED-2000 at \( T \approx T_{cr} \approx -40^\circ \text{C} \). The length of the soft chains is the same in both systems, but in ESD-2001 their ends are not pinned to fixed positions; rather, some limited mobility – though somewhat contrasted by the crosslinking units and the other chains they drag – is allowed. At \( T > T_{cr} \), the cooperativity in the less constrained system is smaller, conforming to what has been found above comparing ED-2000 with ED-4000. On the other hand, a dichotomic behaviour emerges at \( T < T_{cr} \); upon releasing configurational constraints with respect to ED-2000, we find that \( N_s \) decreases in ED-4000 and increases in ESD-2001. Clearly, qualitative arguments simply based on just the effectiveness of the constraints with respect to ED-2000, we find that \( \lambda \) decreases a little over the explored \( T \)-interval (filled circles); on the other hand, \( z \) increases dramatically on cooling (open circles). At \( T \approx T_{cr} \), a relatively smaller number of units (\( z \)) initiate the facilitation process in ED-2000, but each of them needs a comparatively large number of correlated units (\( \lambda \)) to be able explore the \( \zeta \)-space before the large scale transition starts; the opposite is true in ESD-2001. Thus, although at this temperature the cooperativities are almost the same, the stronger mechanical constraints of ED-2000 tend to restrict the size (\( z \)) of the spots where the configurational transitions may initiate. Note that in these circumstances, the mean barrier height is also larger (cf. Fig. 5).

The effect of the rigid crosslink units in ESD-2001 is twofold: on the one hand, they contribute to hinder the configurational degrees of freedom of the soft chains, though not as much as in ED-2000, and on the other hand, they couple mechanically all soft chains attached. Since our analysis probes the precursory state, we have direct evidence of a mechanical coupling in the absence of significant diffusing motion. When \( T > T_{cr} \), the precursory state involves regions of so limited an extension that the pinned ends of the soft chains are only marginally involved. In these conditions, ESD-2001 really appears to be less constrained compared to ED-2000, and the cooperativity turns out to be lower, approaching that of ED-4000. For lower temperatures, instead, the cooperativity analysis certainly reveals a change of regime, in which the crosslinking units are now involved; however, the cooperativity estimate may be expected not to have the same accuracy as in the \( T > T_{cr} \) case.

5 Concluding remarks

The arrest of crystallization is probably the most elementary and widely known example of how structure may affect the primary relaxation dynamics in polymeric systems.\(^{32}\) In that case, for semi-crystalline materials as well as confined systems, e.g., crosslinked networks, elastomeric nano-composites, filled nano-porous materials, after the crystalline structure has evolved in a more or less controlled manner, the amorphous domains invariably reach a state in which the configurational degrees of freedom are frozen. In the systems considered here instead, polyurea materials with segregated hard domains form a continuous amorphous matrix in which the interface could influence the dynamics of the low glass transition domains; the configurational constraints are established at the outset and do not necessarily force the amorphous regions to vitrify.

Appropriate experimental methods, together with an exhaustive relaxation analysis, allowed for a detailed description of the interplay between the structure and the fast contribution to the primary relaxation of the amorphous domains. Concerning instead the slow dynamics of the semi-rigid layers surrounding the hard domains, one has to resort to different probes for the reasons illustrated previously. This may help to extract, by direct analysis in terms of both the cooperativity and the rearrangement threshold energy \( \zeta^* \), valuable information concerning, e.g., the thickness \( d_{\text{p}} \) of these interfacial layers. Indeed, the information extracted by the relaxation analysis adopted in the present case, depends only weakly on the probe, as it was shown in ref. 7 and 8;
this is supported also by the good agreement between the
ED-2000 cooperativities worked out here and in ref. 13 by dynamic
mechanical spectroscopy.

A reasonable upper bound to \( d_a \) can be estimated in the
case of the ED-2000 sample on the basis of the fast mode
cooperativity (Table S1 (ESI†), which considers low-\( T \) extra-
polated cooperativities, and Table S2 (ESI†), which refers to
the actual data of Fig. 6). With reference to Fig. 1c, simulations
yield a diameter of the hard core (blue colored circles in the
figure) ca. 2.1 nm in the all-\textit{trans} conformation of the hexam-
ethylene chains from the crosslinker moiety. On the other
hand, SAXS and Fourier transform AFM analyses yield a 5.2
to 5.6 nm, respectively, typical distance between the centers
of these hard domains. Assuming that the cooperativity of the fast
mode is close to saturation at the lowest probed temperature,
one finds a diameter of about 2.5 nm for an approximately
spherical CRR. This implies a surrounding layer of restricted
mobility of thickness 0.4 < \( d_a \) < 0.5 nm, which is compatible
with the estimate of about 0.2 nm derived by independent
considerations.\(^3\) Such a value (0.4 < \( d_a \) < 0.5 nm) would
be at most two monomers in the \textit{trans} conformation.

The possibility of probing the dielectric response at temperatures
closer to the fast mode \( T_g \), would in particular allowed to discuss the effect of
hydrogen bonding on cooperativity.\(^3\) When dealing with con-
fine mechanisms, in particular, geometric considerations alone cannot
in general account for all aspects of the process. In particular,
as touched upon in the discussion on fast relaxation, the role of the
threshold energy \( T_\ast \) seems to be important, and indeed a step fur-
ther in the understanding of the cooperative dynamics
would also need a satisfactory description of heat transfer among
the different degrees of freedom. How efficient is the transfer of the
pre-transitional energy towards the configurational modes,
compared to other regression paths, is indeed central for a more
complete understanding of confinement problems in general, just
leaving aside the still lively problem of the super-\textit{arrhenian}
behavior of fragile liquids.

Conflicts of interest

There are no conflicts to declare.

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