Influence of the PEO length in gas separation properties of segregating aromatic-aliphatic copoly(ether-imide).

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Summary:
A complete series of aliphatic-aromatic copoly(ether-imide)s has been synthesized in this work. All these copoly(ether-imide)s had the same structure, BPDA-PEOX-ODA, but different lengths of PEO in the final polymer. These copolymers have been thermally treated and characterized by several techniques. A direct relationship between the temperature of treatment, the improvement of phase segregation, and permeability has been demonstrated. The Maxwell model has been applied to predict permeability (for CO₂, CH₄ and N₂) and it has been found that when the segregated PEO can be considered to be amorphous – it is at high temperatures when crystallinity disappears – the model fits reasonably well. This confirms that the aliphatic and aromatic portions of the copolymer behave approximately as a bi-phase of disperse domains within a continuous matrix. Results show that permeability is higher when the PEO chains are longer – when there is no crystallinity, or any kind of internal bonds, distorting the results – while selectivity doesn’t depend on the PEO length in the copolymer. Remarkable are the results for the CO₂/N₂ gas pair, with selectivity-versus-permeability very near to the Robeson’s upper bound at 30 ºC and even in closer proximity to the corresponding trade off line for higher permeation temperatures.

Keywords: Membranes; Polymers; Separations; Selectivity; Copoly(ether-imide); carbon dioxide (CO₂).

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1. Introduction

The liberation to the atmosphere of big amounts of greenhouse gases in a short period can lead to global warming. Over a 60% of the global greenhouse effect is attributable to carbon dioxide (Yamasaki, 2003). There are several sources — such as power plants, steel, cement production plants or chemical industry — where CO$_2$ is generated. Among these, coal-fired power plants are the main emitters of CO$_2$ and are responsible for nearly a 30% of the total CO$_2$ release. It has been estimated that about half of the electricity will be generated by coal-fired power plants at least until 2035 (EIA, 2011). This will cause more CO$_2$ liberated causing a serious intensification of climate change. Hence, it is necessary to develop new technologies to curb as far as possible the emissions of greenhouse gases.

One of the preferred methods to control CO$_2$ emission is its capture and storage (CCS), including separation, transportation and storage stages (EIA, 2011). Around a 70 to 80% of the total costs of CCS is represented by the CO$_2$ capture. The three main options for CO$_2$ capture from power plants are: pre-combustion, oxy-fuel combustion, and post-combustion capture (Figueroa et al., 2008; Ho et al., 2009; Yang et al., 2008). Post-combustion capture is the simplest and most suitable way to perform carbon dioxide capture for newly-built as well as for existing coal-fired power plants (Herzog, 2001).

Post-combustion capture is fundamentally a separation of CO$_2$ from flue gas — generally consisting in N$_2$, CO$_2$, O$_2$ and H$_2$O —, and conventional gas separation technologies, such as chemical absorption, pressure swing adsorption, and membrane separation, could be applied. Of course the separation technique to be used is determined by its effective price (Davison, 2007; Hoffmann et al., 2008; Klmeš et al., 2007). The amount of carbon dioxide emitted and captured are, of course, relevant figures in order to make an adequate decision.

A first instance analysis could lead to conclude that chemical adsorption processes are more interesting because they can give high purity at a relatively low price (Ho et al., 2008; Merkel et al., 2010; Oexmann et al., 2008; Qin et al., 2010). The main disadvantage for chemical absorption is that its cost is still over the target (Figueroa et al., 2008) and further cost reduction is quite complicated (Klara et al., 2009) because the
main cost item corresponds to energy consumption. Very different is the situation of gas separation processes using polymeric membranes that are by nature quite flexible processes. Whenever CO₂/N₂ selectivity exceeds 30, permeability of membrane has more influence on the cost than energy consumption which is very low (Merkel et al., 2010). Moreover, it has been shown that, if the cost of the membrane is low, it is cost-effective to select a material with high selectivity; while if the price of the membrane is high a polymer with a higher CO₂ permeability is more economically effective (Ho et al., 2008; Yang et al., 2011). All these factors concerning the membrane material leave a very wide range of improvement open in terms of operational cost.

It is also convenient, in order to get appropriate membranes for gas separation, to start with a suitable polymeric material, with adequate flow and good mechanical and thermal resistance. Of course when dealing with CO₂ capture, the polymer membrane must have a high affinity for carbon dioxide, so polymers with adequate permeability values are necessary (Bounaceur et al., 2006; Favre, 2007).

Glassy polymers and in particular polyimides are well known for their excellent thermal oxidative stability, good organic solvent resistance and exceptional mechanical properties, along with an extraordinary ability to separate complex mixtures of gases in diverse applications (Bessonov, 1987; Ghosh, 1996; Wilson, 1990). Thus, among all the polymeric membranes, it has been widely demonstrated that the use of aromatic polyimides is one of the best alternatives. Typically these materials have a high selectivity while having a not always high enough permeability (Ayala et al., 2003; Tanaka et al., 1989).

One of the most common approaches to meet these requirements on CO₂ permeability is to increase the affinity of the polymer to CO₂ by using block-copolymers. Specially promising block-copolymers include both hard and soft blocks. The hard block can have a well-packed and rigid structure; as a result it forms a glassy segment of the copolymer chain with usually low free volume. In contrast, the soft block can consist in more flexible chains that can form rubbery segments on the copolymer chain with habitually high free volume. In this way the glassy polymer segments will provide the mechanical support and in the case of polyimides, thermal resistance. The rubbery segments generally form continuous microdomains through which, due to the nature of the
flexible chain structure, gas transport is easy giving a good permeability to the copolymer (Barbari et al., 1988; Li et al., 1997).

The use of polyethylene oxide (PEO) compounds for separating CO\textsubscript{2} from other light gases is known (Hirayama et al., 1999; Kawakami et al., 1982) to give high CO\textsubscript{2} permeability. The interaction between CO\textsubscript{2} and oxyethylene groups has been discussed and used for the development of CO\textsubscript{2} selective membranes by many authors (Car et al., 2008; Lin and Freeman, 2004; Reijerkerk et al., 2010). In view of this, the use of block-copolymers combining aromatic polyimides and PEO, appears to be a successful route (Okamoto et al., 1995; Okamoto, 1993; Suzuki et al., 1998). In effect, these compounds have good permselectivities (permeability-selectivity) results for the couple CO\textsubscript{2}/N\textsubscript{2}. This was attributed mainly to the high solubility-selectivity due to strong interactions between the oxyethylene group and CO\textsubscript{2} (Tena et al., 2012b; Tena et al., 2012c).

In order to optimize the necessary properties, a good balance between the hard and soft block parts of the copolymer is needed. Previous studies from our group reported that when the copolymer has a percentage of PEO around 50\% it has the best properties attending to the necessary mechanical resistance that should be compromised by too high PEO contents (Tena et al., 2012a). Moreover, the crystallinity of polyether segments is known to increase when the length of PEO increases leading to a decrease in gas sorption because the crystalline phase acts as a non-sorbing and impermeable barrier reducing the permeability severely. Therefore, in addition to the percentage of PEO, the length of its chain has also an important effect on the permeation properties of the copolymer.

For this reason, we propose here a complete analysis of the influence of PEO length on the properties of separation for a system where both the hard part, in this case the aromatic polyimide BPDA-ODA, and the proportion between soft and hard segments are common in all copolymers to be studied. The length of the PEO chain will be varied from short lengths where all the polyethylene oxide is in amorphous state to percentages as high as to lead to crystallinities over a 40\% of the total PEO in the copolymer. Some copolymers have been also synthesized including hexamethylene diisocyanate in order to extend the polyether chain while reducing the increase of crystallinity but improving the phase segregation after thermal treatment. The influence of thermal treatments at
different temperatures on PEO segregation and permselectivity will be also analyzed here.

The properties of these copolymers have been analyzed by standard techniques of characterization (DSC, TGA, TMA, mechanical properties and densitometry) and by SAXS. The properties of permeation have been evaluated at temperatures from 30 °C to 50 °C, a temperature high enough so that all samples had the polyethylene oxide chains in an amorphous state. Finally, we have used the Maxwell’s model to predict the permeability to different gases.

2. **Experimental**

2.1. **Chemicals**

3,3′,4,4′-biphenyltetracarboxylic dianhydride (BPDA), and 4,4′-oxydianiline (ODA) were purchased from Sigma-Aldrich Co. These products were purified by sublimation at high vacuum just before being used. Bis(2-aminopropyl) poly(ethylene oxide) with nominal molecular weight of 900 and 2000 g/mol were kindly donated by Huntsman (Holland) (PEO-900 and PEO-2000 from here on). α,ω-Diamine-poly(ethylene oxide) with nominal molecular weight of 6000 g mol\(^{-1}\) was kindly donated by Kawaken Fine Chemicals Co., Ltd. (Tokyo, Japan), (PEO-6000 from here on). The polyethers were dried at 70 °C in vacuum for 5 hours and stored in a desiccator at vacuum until use. Hexamethylenediisocyanate (HDI) was purchased from Sigma-Aldrich and purified by distillation at vacuum before use. Anhydrous N-methylpyrrolidinone (NMP), to be used as polymerization solvent, was purchased from Sigma-Aldrich Co.

![Figure 1. Chemical structure of the monomers.](image)

2.2. **Synthesis of copoly(ether-imide)s**

The polymerization reaction was carried out in a three-neck flask, under N\(_2\) atmosphere and mechanical stirring, following this procedure: PEO (x mmol) was dissolved in anhydrous NMP, and when necessary, a certain amount of HDI (y mmol) added in order to obtain a PEO oligomer of approximately 6000 g/mol of final molecular weight.
Polyether chains included in the synthesized copolymers were: PEO900, PEO900+HDI, PEO2000, PEO2000+HDI, PEO900+PEO2000 (50/50 w/w), and PEO6000. After some minutes of stirring at room temperature, the aromatic diamine (ODA) (z mmol) in a weight ratio 2:1 (polyether chain : aromatic diamine), was added to the solution to give a final concentration of 20 mL solvent for 10 meq of amino group. When the solid aromatic diamine was dissolved the precise stoichiometric amount of the BPDA aromatic dianhydride was added at once. The reaction was stirred at room temperature for 12 hours.

From here on the copolymers to be studied will be named by adding cPI to the name of the PEO entering the composition, i.e.: cPI-PEO900, cPI-PEO900+HDI, cPI-PEO2000, cPI-PEO2000+HDI, cPI-PEO900+PEO2000 and cPI-PEO6000.

2.3. Preparation of the copolymers dense films

The resultant viscous copolyamic acid solution was diluted with NMP to attain an appropriate viscosity for casting, filtered through a nominal #1 fritted glass funnel, degassed, and cast onto a levelled glass plate. The resulting film was covered with a conical funnel to avoid too fast evaporation of the solvent, dried at 80 ºC overnight, and treated at 120 ºC for 6 hours in a vacuum oven. For the pure aromatic homopolymer (BPDA-ODA), the procedure was similar but the final heat treatment was carried out at 200 ºC in a vacuum oven for 24 hours. Films with 40-60 µm in thickness were obtained. The additional thermal treatments (see Figure 2) for the copolymeric films were carried out under inert atmosphere.

2.4. Characterization Methods

Attenuated total internal reflectance-Fourier transform infrared analyses (ATR-FTIR) were performed at room temperature using a PerkinElmer Spectrum One infrared spectrometer equipped with an ATR accessory. IR spectra were the average of 16 scans recorded in the spectral range of 4000–650 cm\(^{-1}\) with a 4 cm\(^{-1}\) resolution.
A Thermal Analysis Q500 instrument was used for thermogravimetric analysis (TGA). Disc samples cut from films with weights between 5 and 15 mg were tested. When running dynamic scans, it was done in Hi-Resolution mode, where the heating rate is automatically adjusted in response to changes in the rate of weight loss, which results in improved resolution, with an initial heating rate of 10ºC/min under a flux of nitrogen.

Differential scanning calorimetry (DSC) analyses were carried out in a Mettler Toledo (DSC 822e) calorimeter equipped with a liquid nitrogen accessory. Disc samples cut from films weighting 5–15 mg were sealed in aluminium pans. Samples were heated with the following cyclic method in order to monitor the changes in thermal properties with thermal treatment: from 25 ºC, the sample was heated at 10ºC/min to a target temperature; once reached, the sample was cooled at the maximum cooling rate accessible for the instrument to -90 ºC, held at this temperature for 15 min to equilibrate and reheated at 10 ºC/min to the next target temperature. The procedure was followed until the last treatment temperature was reached and a final run from -90 ºC to 80 ºC was performed. In this way, in each heating run, the thermal properties for the copolymers after “instantaneous” treatment to the previously reached temperature were obtained.

Thermomechanical (TMA) tests were performed in a Rheometric Scientific instrument model DMTA V. Rectangular test pieces of 3 mm width and 20 mm length were cut from films. A distance of 10 mm was set between fixation clamps. Runs were carried out from ambient temperature at 10 ºC/min with a static stress of 3 MPa.

The densities (ρ) of the dense membrane films were determined using a CP225D Sartorius balance, provided with an immersion density kit.

SAXS measurements were performed at the beamline BM16 at the European Synchrotron Radiation Facility (Grenoble, France). Wave length of the X-ray beam was 0.980 Å. Detector calibration was done with silver behenate (AgC_{22}H_{43}O_{2}). Disc samples cut from films were placed in a Linkam hot stage and heated at 10 ºC/min while the SAXS spectra were recorded. Calibration of temperature gave a difference of
approximately 7 °C between the temperature reading at the hot stage display and the real temperature at the sample.

2.5. **Gas Permeation and Selectivity**

The permeability, $P$, for several gases (namely: $O_2$, $N_2$, $CH_4$ and $CO_2$) was determined by using a permeator with constant volume and variable pressure which uses the “time-lag” operation method. The measurements were carried out at 3 bar at temperatures set from 30 to 50 °C. The equations used and the mode of operation of the equipment, as well as an outline of it, has been described previously (Tena et al., 2010).

3. **Results and Discussion**

3.1 **Copoly(ether-imide)s imidization**

Poly(ethylene oxide) chains are prone to oxidation (Scheirs et al., 1991), and therefore, a great care was taken to carry out the imidization. The films were heated at 120 °C for 6 h. After 6 h heating at 120 °C, the solvent was almost completely removed as proved by thermogravimetric analysis. All the copolymers resulted to be insoluble in DMAc (dimethylacetamide), NMP, Hexane, Toluene, THF (tetrahydrofuran), and $CH_2Cl_2$ (dichloromethane) after this process.

Infrared spectra were recorded to check for the progress of imidization. As for other described copoly(ether imide)s (Marcos-Fernández et al., 2010; Tena et al., 2012b; Tena et al., 2012c), it was possible to observe the IR bands centered around 1774, 1713, 1372 and 738 cm$^{-1}$, typical for imidized samples, and the peaks appearing around 2800 cm$^{-1}$, due to the aliphatic polyether segments. There were no noticeable differences between the copolymers with different polyether lengths, or between samples containing PEO and the samples with PEO+HDI.

For similar copoly(ether-imide)s to the ones presented here, Hangzheng et al. (Hangzheng et al., 2010) used IR spectra to prove the imidization at 200 °C. Even though the complete imidization of our copoly(ether imide)s at a temperature as low as 120 °C is actually remarkable, we must mention that this full imidization at relatively
low temperatures has already been found by us for copoly(ether-imide)s based on PEO segments (Marcos-Fernández et al., 2010; Tena et al., 2012b) and PPO segments (paper in preparation). Also, two works of Okamoto et al., studying similar copoly(ether-imide)s to the ones presented here, stated that a thermal treatment at 170 °C completely imidize the precursor poly(amic acid)s to polyimides (Okamoto et al., 1995) although they did not show any IR spectra to prove the imidization state of the copolymers.

3.2 Thermal Stability

Thermogravimetric analysis was performed to evaluate the thermal stability of the synthesized copolymers treated at 120 °C during 6 hours (Figure 3).

![Figure 3. TGA curves in dynamic conditions for the synthesized copolymers.](image_url)

The behaviour is similar for the different samples. The initial loss from ambient temperature to 100 °C, can be attributed to the adsorbed water in the sample. A second step approximately from 100 to 250 °C is due to residual solvent in the film, with a weight loss of 1 to 1.7%. The third step is thought to be due to the contribution of the poly(ethylene oxide) included in the copolymer composition, and it is therefore assigned to the loss of polyether block sequences plus the hexamethylenediisocianate in the case of the copolymers where the polyether chain was extended. The final stage of weight loss is due to the thermal decomposition of the remaining aromatic polyimide segments. For all copolymers, the residual carbon content at 800 °C was 25-35 %.

The mass loss in this third step for the different copolyimides is close to the percentage of polyether blocks in the feed (see Table 1), proving that it is related to the degradation of this part of the copolymer, as already found previously in other similar copolymers (Marcos-Fernández et al., 2010; Tena et al., 2012b; Tena et al., 2012c). From Figure 3, it is clear that the polyether extended copolymers cPI-PEO900+HDI and cPI-PEO2000+HDI start to degrade at lower temperatures (circa 250 °C) than the other copolymers (circa 350 °C). This is due to the lower stability of the urea groups formed by the reaction of the HDI. This relatively low thermal stability of the urea groups is well known in the field of polyurethanes.
Table 1.- Results obtained by TGA for the prepared copolymers.

Thus, TGA analysis confirmed that the aliphatic-chain thermal stability is much lower than the thermal stability of the aromatic polyimide segments, and therefore a selective degradation of this part of the chain can be performed in these copolymers.

3.3 Calorimetric Studies

The samples were heated in a DSC instrument with a cyclic method in order to monitor the changes in the thermal properties of the films with the thermal treatment. When the polyether moiety is able to crystallize, the melting enthalpy can be used to evaluate the percentages of amorphous-crystalline PEO in the copolymers (Marcos-Fernández et al., 2010). No transition related to the aromatic polyimide hard segments was detected by DSC in any case.

A curve with the developed crystallinity for PEO vs. temperature thermal treatment can be built for all copolymers. The melting enthalpy of PEO was taken from literature as 8.67 kJ/mol (Van Krevelen, 1990). In Figure 4 crystallinity is shown for all the copolymers thermally treated at different temperatures.

No crystallinity was observed for samples cPI-PEO900 and cPI-PEO900+2000. For cPI-PEO900, the polyether chain is too short to develop crystallinity when the chain ends are linked to the polyimide segments, whereas for cPI-PEO900+2000, the PEO900 chains hamper the crystallization of the longer PEO2000 chains. In the case of sample cPI-PEO2000 the crystallinity found was almost negligible, a maximum of 0.9 % after the thermal treatment at 300 °C, whereas for cPI-PEO6000, crystallinity increases with thermal treatment from almost inexistent in the original sample, heated at 120 °C, to a 32 % after the thermal treatment at 300 °C. The maximum increase in crystallinity, up to approximately 250 °C, happens simultaneously with the release of the residual solvent.
as shown by TGA. From these data it is clear that lengths over 2000 g mol$^{-1}$ are necessary to develop crystallinity and longer chains develop higher crystallinities.

The samples with extended polyether, cPI-PEO900+HDI and cPI-PEO2000+HDI, have a very characteristic behavior that differs from that shown by cPI-PEO900 and cPI-PEO2000. When the polyether chain is extended with HDI, although all the copolymers contain very similar weights of polyether, the molar ratio of the aromatic reactants over the aliphatic ones increases for the copolymers with extended polyether chains, leading to longer aromatic polyimide blocks. This, in turn, leads to an increase in the incompatibility of the blocks and therefore to a better phase separation, as indicated by the higher crystallinity achieved for polyether extended copolymers. In fact, for cPI-PEO900+HDI, crystallinity appears at 200 °C whilst the parent copolymer cPI-PEO900 remains amorphous at all thermal treatments, and for cPI-PEO2000+HDI, crystallinity is significant from the beginning, and increases further with thermal treatment, with values that are very similar to those for the cPI-PEO6000.

However, when treated at high temperatures, a different behavior is found for polyether extended copolymers. cPI-PEO900+HDI starts to lose crystallinity above 230 °C and it completely disappears at 260 °C, whereas cPI-PEO2000+HDI reaches a pseudo plateau between 200 and 240 °C and above this temperature crystallinity still has a small increase. Although the trend is just the reverse for these two copolymers, the origin of the behavior is the same. Above 230-240 °C the urea groups connecting the extended polyether chains start to deteriorate, as shown by TGA, and when this occurs, polyether chains with free chain ends are produced. The chains with low polyether molecular weight, 900 g mol$^{-1}$, are unable to crystallize and plasticize the PEO phase, decreasing crystallinity until its disappearance, whereas chains with higher molecular weight, 2000 g mol$^{-1}$, have less restrictions to crystallize and crystallinity increases (it is worth to point out that the starting bis(2-aminopropyl)poly(ethylene oxide) of 2000 g mol$^{-1}$ is highly crystalline).

It is worth noting that here we have been referring to maximal cristallinity that would appear at low enough operation temperatures. It is well known that gases do not easily pass through the crystal structures (Casado-Coterillo et al., 2012), and for this reason it is important to know the actual crystallinity of our samples at the temperature of
permeation measurement (from 30 to 50 °C). For all the copolymers, except for cPI-PEO6000, the melting point measured by DSC is below 30 °C after thermal treatment at any temperature, and therefore, PEO chains will be amorphous. For copolymer PEO6000, the melting point reaches a value of 43 °C after thermal treatment at 300 °C (approximately 40 °C after treatment at 250 °C), thus, when measuring permeation properties at 50 °C, PEO chains will be amorphous, but at 30 °C, part of these PEO chains could still be crystallized.

In figure 5 we can see the changes in the polyether T_g for the different treatment temperatures. Only minor variations are seen, and T_g remains almost unchanged with thermal treatment except for copolymers PEO900+HDI and PEO2000+HDI.

| Figure 5. T_g of the aliphatic part of copolymers after “instantaneous” treatment for the copolymers treated at 160 °C. |

For these two HDI extended copolymers, when urea groups start to degrade for treatments above 220 °C, T_g increases. This could be interpreted as an increase in the mixing of the rigid polyimide segments in the polyether phase. This increase in T_g is very noticeable for copolymer PEO900+HDI, and coincident with the decrease in crystallinity of the PEO phase. For copolymer PEO2000+HDI the change coincides also with the small increase in crystallinity produced by urea groups degradation, but because of the ability of the PEO chains of 2000 g mol^{-1} to crystallize, the mixing of the aromatic polyimide segments in the PEO phase is very limited and the increase in T_g is very small. If we compare the T_g values after treatment at 200 °C, where the T_g is practically constant for all the copolymers and the copolymers with extended polyether chain have not started to degrade, it follows the expected trend for the copolymers with un-extended polyether chain, the longer the PEO chain, the lower the T_g:

(PEO6000) < (PEO2000) < (PEO900)

In the case of the copolymers with extended polyether chain, their T_g are well below the values for the copolymers with the parent polyethers, PEO900 and PEO2000, as a consequence of the better phase separation achieved. For PEO2000+HDI the value is
similar to the value for PEO6000, and for PEO900+HDI is around 6 °C higher, probably due to the restrictions imposed by hydrogen bonds between the urea groups, present in a higher concentration in PEO900+HDI compared to PEO2000+HDI. Yet, $T_g$ for PEO900+HDI is almost 10 °C lower than for copolymers with PEO2000.

Thus, from these results, it can be affirmed that for all the copolymers studied, at a temperature of permeation of 50 °C, the polyether segments were totally amorphous and well above their $T_g$.

3.4. Thermomechanical Analysis

Thermomechanical analysis was carried out in order to detect the glass transition temperature of the aromatic polyimide hard segments, which could not be detected by DSC. It was assumed that a good estimation of $T_g$ corresponds to the temperature when strain is 10 times that of the sample at 100 °C. This criterion could be accepted at least for comparative purposes (Marcos-Fernández et al., 2010). The so-obtained results for samples treated at 160 °C for 2 hours in order to develop a phase separated structure are shown in Figure 6.

Figure 6. $T_g$ for the aromatic polyimide segments as a function of the PEO length for the copolymers treated at 160 °C.

All samples have similar content in PEO, but because of the stoichiometry, the longer the polyether chain, the longer the aromatic polyimide would be, and therefore, the higher the $T_g$ of the aromatic polyimide segments, as shown in Figure 6. The order is:

$$(c\text{PI-PEO900}) < (c\text{PI-PEO900+2000}) < (c\text{PI-PEO2000}) < (c\text{PI-PEO6000})$$

In addition, when PEO chains are shorter, the mixing of PEO segments in the polyimide phase is more likely, and can contribute further to decrease the polyimide $T_g$ by plasticization. This must be the cause of the difference in polyimide $T_g$ for the chain extended polyethers. For cPI-PEO900+HDI, cPI-PEO2000+HDI, cPI-PEO6000, with theoretically similar polyimide segments length, $T_g$ of polyimide decreases when PEO
length decreases (HDI content increases). In these copolymers, although the longer polyimide segments lead to a better phase-separated polyether phase (and bigger domains, as it will be shown latter in SAXS), apparently, the polar urea groups in the polyether chains also produce some mixing in the polyimide phase lowering the $T_g$. This decrease is of the same order than the increase due to the longer polyimide chains for copolymer cPI-PEO900+HDI, and the $T_g$ value is approximately the same as for the copolymer with the parent polyether, cPI-PEO900, whereas for copolymer cPI-PEO2000+HDI is slightly higher and $T_g$ value is slightly lower than for the copolymer with the parent polyether, cPI-PEO2000.

For all the copolymers, the $T_g$ of the aromatic polyimide is well above ambient temperature, although lower than the corresponding $T_g$ for a pure aromatic polyimide homopolymer (270 °C (Okamoto et al., 1995), 287.3 °C (Zhang et al., 1999), 305 °C (Marcos-Fernández et al., 2010)), due to the much lower polymerization degree and, consequently, lower length of the aromatic polyimide segments in the copolymer as compared to the corresponding homopolymer.

3.5. **Density**

The densities ($\rho$) of the dense membrane films were determined according to the equation:

$$\rho = \rho_o \frac{W_{air}}{W_{air} - W_{liq}}$$

(1)

where $\rho$ is the density of the film, $W_{air}$ and $W_{liq}$ are the weights of the film in the air and immersed in an auxiliary liquid (in these measurements, isooctane was used because this polymer is reported to be insoluble in isooctane (Brandrup, 1999)) and $\rho_o$ is the density of the auxiliary liquid (isooctane). The density for all copolymers is lower than that for the pure aromatic polyimide, BPDA-ODA, which was 1.40 g mL$^{-1}$, and follows this trend:
\[\rho(\text{cPI-PEO900}) = 1.308 > \rho(\text{cPI-PEO900+2000}) = 1.303 > \rho(\text{cPI-PEO2000}) = 1.294 > \]
\[\rho(\text{cPI-PEO6000}) = 1.281 > \rho(\text{cPI-PEO2000+HDI}) = 1.276 > \rho(\text{cPI-PEO900+HDI}) = 1.266\]

It is clear that the longer the polyether chain, that is, the better the phase separation, the lower the density should be but, actually, the differences within the copolymers are very small as shown in Figure 7. Copolymers with extended polyether chains, with long polyether chains and well developed phase separation – as well as cPI-PEO6000 – should give low densities. These copolymers should give big free volumes and thus gas should pass through the membrane quite easily if there were no cristallinity (cPI-PEO6000) and/or hydrogen bonding (HDI extended copolymers).

Figure 7. Density as a function of the polymer length.

Note that density decreases when \( T_{g}^{AI} \) decreases. The lower densities and lower \( T_{g}^{AI} \) appear due to the ability of the polar groups placed throughout the polymer chains to increase the intermolecular forces increasing the free volume along with the mobility of the PEO chains.

3.6. Small-Angle X-ray Scattering

From the curves of scattered intensity \( I \) versus the scattering vector \( (q = 4\pi(\sin\theta)/\lambda, \lambda = \text{wave length}, 2\theta = \text{scattering angle}) \), two parameters can be evaluated: the relative invariant, \( Q' \), as the integral below the curve \( Iq^{2} \) vs. \( q \), which is related to the extent of the phase separation; and the maximum on the scattering curve, \( q_{\text{max}} \), related to the size scale, \( L \), of the separated phases, calculated also from the curve \( Iq^{2} \) vs. \( q \). The equations used and the mode of operation of the equipment, as well as an outline of it has been explained in previous studies (Tena et al., 2012b).

The scattering profile is very different for the polyether extended copolymers when compared with the copolymers with the parent polyethers. For copolymers cPI-PE900, cPI-PEO2000, cPI-PEO900+2000 and cPI-PEO6000, a single broad peak centered at 0.4-0.6 nm\(^{-1}\) appears, as seen in Figure 8 for two selected copolymers. However, for the
polyether extended copolymers, cPI-PEO900+HDI and cPI-PEO2000+HDI, the maximum in the scattered intensity is located close to the beam stop (see inset in Figure 8 were these peaks appear clearer).

Figure 8. Intensity (logarithmic scale) vs q curves for copolymers cPI-PEO900, cPI-PEO2000, cPI-PEO900+HDI and cPI-PEO2000+HDI. In the insert, the Iq^2 vs q curves for the same copolymers are presented.

In Figure 9, the relative invariant (Q’) is plotted versus treatment temperature. Copolymers with the extended polyethers were heated up to 250°C (before degradation of the urea groups), whereas the others until 300°C. For copolymer cPI-PEO-900, Q’ does not change significantly with temperature, with only a small increase above approximately 165°C. For this copolymer, with the shortest polyether chain and the shortest polyimide segments of those synthesized here, the segments are partially mixed, as seen in the DSC results, and the phase separated structure almost doesn’t improve with thermal treatment. For copolymers cPI-PEO2000, cPI-PEO900+2000 and cPI-PEO6000, at a certain temperature (approximately 180, 170 and 150°C respectively), Q’ increases noticeably up to the final heating temperature, showing a notorious improvement on their phase separated structure. The same improvement is found for the polyether extended copolymers cPI-PEO900+HDI and cPI-PEO2000+HDI, starting in both cases at approximately 180°C up to the final heating temperature. This improvement is coincident with the increase in PEO crystallinity seen by DSC and, as already mentioned, it is also coincident with the release of the residual solvent in the films.

Figure 9. Changes in the relative invariant for the synthesized copolymers treated at 120°C.

When the morphology of the phase separated structure is unknown, as in our copolymers, a calculation of the spacing length, L, from the curve I vs. q should be preferred (Baltá-Calleja and Vonk, 1989). However, we are more interested on the changes of the spacing scale with temperature than on the absolute value for this length.
scale of the segregated phases, thus, $Iq^2$ vs. q curve was used to calculate L attending to the higher accuracy in determining the maximum on this curve.

As shown in Figure 10, changes in L take place almost simultaneously to changes in Q’. Big differences are found referring to L between the copolymers with the extended polyether chains and those with the parent polyethers. For copolymers cPI-PEO900, cPI-PEO2000, cPI-PEO900+2000 and cPI-PEO6000, L is initially similar for all of them, circa 8 nm, and on heating, grows to a value around 12 nm (i.e. an approximate 50% increment), except for cPI-PEO6000, for which the value reached is almost 19 nm (around 300% increment). This longer L value means bigger PEO domains that are able to crystallize extensively as seen by DSC. Therefore, there is not a trend between L value and polyether length, and only for polyether PEO6000, the L value is clearly above the others. Apart from the strong increase on L for copolymer cPI-PEO6000, it is worth to comment that for copolymer cPI-PEO900, although Q’ has a very small increase with temperature, L value changes as much as for copolymers cPI-PEO2000 and cPI-PEO900+PEO2000.

Copolymers with extended polyether chains have a very high L value from the start, around 40 nm for cPI-PEO900+HDI and 50 nm for cPI-PEO2000+HDI, and when Q’ increases, the increase in L is very modest, around 8% for both. This large L corresponds to very large PEO domains, and that is the reason why cPI-PEO2000+HDI can reach also very high values of PEO crystallinity, of the order of cPI-PEO6000.

![Figure 10](image-url) Evolution of L as a function of temperature for the synthesized copolymers.

From SAXS data it can be concluded that thermal treatment improves the phase separated structure except for cPI-PEO900, where the segments are mixed in a substantial extent. For this copolymer, however, a significant change in long spacing takes place with thermal treatment. The size scale of the phase separated structure after thermal treatment is similar for copolymers with parent copolyethers PEO900 and PEO2000, and is much bigger for copolymer based on polyether PEO6000. For copolymers with extended polyether chains, the size scale of the phase separated structure is much bigger than for copolymers with parent polyethers, and even than for copolymer cPI-PEO6000, with similar PEO content, similar total PEO chain length and
polyimide segment length, thus, chain extension of relatively short polyether chains is a very effective way to modify the phase separated structure of the copolymers.

3.7. Permeability properties

3.7.1. Behaviour with the thermal treatment

As seen for other aliphatic-aromatic copoly(ether imide)s (Tena et al., 2012c), the results show an improvement in permeation properties after thermal treatment while selectivity remains substantially constant, so that there is a direct relationship between: treatment temperature, phase segregation and permeation properties as can be seen in Figure 11 for the sample cPI-PEO900+2000.

Figure 11. Evolution of the permeability (a) and selectivity (b) as a function of the treatment temperature for cPI-PEO900+2000.

In the case of samples with extended polyether chain, the picture is somewhat different. Figure 12 shows the comparative behavior of sample cPI-PEO900+2000 with samples cPI-PEO900+HDI and cPI-PEO2000+HDI. As already seen, segregation increases with heat treatment, but by TGA and DSC it was clear that at certain temperature, urea groups decompose. When polyether extended copolymers are heated at 200°C for 2 hours in an oven, urea groups would decompose, producing some mixing of segments and some loss of polyether chains that lead to the observed decrease in CO₂ permeability.

Figure 12. CO₂ permeability as a function of the thermal treatment.

3.7.2. Behaviour with the measurement temperature

The movement of gas molecules to penetrate the polymer membranes depends on the thermally activated motion of polymer chains. Not only diffusivity but also solubility is related to the interaction between the polymer chain and the place left to gas
condensation. In this sense it is very important to know the properties of the polymers that have a strong influence on the process of separation, such as the free volume. When a polymer is in the crystalline state, the gases hardly penetrate across this structure because the chain movements are very limited giving quite low permeability. The crystals would operate as a non-sorption, impermeable barrier and reduce the permeability severely. On the other hand, when the melting temperature, $T_m$, is surpassed, the polymer is amorphous, making chains more mobile and thus favoring the permeation of gases through the samples. Therefore, in samples that present or may present crystallinity at work temperature, it is advisable to conduct a study of their behavior as a function of measurement temperature.

Taking into account the temperature dependence for the diffusion and sorption coefficients, the temperature effect on gas permeability follows an Arrhenius law (Kesting and Fritzsche, 1993):

$$
(2)
$$

where $E_P$ is the activation energy of permeation. In Figure 13 the Arrhenius’ increase of permeability with the measurement temperature is shown for several gases and the cPI-PEO6000 sample. Several intermediate permeabilities at temperatures from 30 °C to 50 °C are shown in the Figure. The activation energy of permeation is shown in Figure 14 as a function of the PEO length. It seems clear that cPI-PEO-6000 has lower activation energy than expected. This could be attributable to its residual crystallinity.

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Figure 13. CO$_2$ permeability as a function of permeation temperature for cPI-PEO6000 membranes treated at 160°C.

Figure 14. Activation energy as a function of the molecular weight of PEO (a) and for the longest PEO copolymers (b). In the (a) figure only cPI-PEO2000+HDI, among the copolymers with the longest PEO chains, is shown.

3.7.3. Permselective properties as a function of the PEO length
Some authors have recently assumed that there should be a maximum length over which permeability decreases (Hangzheng et al., 2010; Yave et al., 2010). As mentioned above, the amount of crystalline PEO in these samples is important especially for the longest PEO chains (especially cPI-PEO6000). Could this crystallinity be the cause of such an apparent decrease in permeability for copolyetherimides containing very long PEO chains?

In order to analyze the influence of crystallinity in an eventual decrease of permeability for long PEO chains. It is worth to note that at 50 °C all residual crystallinity would be eliminated for all the copolymers studied guaranteeing that all they are in amorphous state.

Figure 15 shows the results obtained, at the lowest (30 °C) and the highest (50 °C) permeation temperature, for the permeability of CO₂ (the trend is similar for all gases tested). The results at 30 °C show that there is a maximum length over which permeability decreases as seen in literature. However, the results at 50 °C show that there is a monotonous increase of permeability for increasing PEO lengths. This is clear for samples containing only pure PEO. For samples with HDI in its structure permeabilities are lower than for the sample cPI-PEO6000. For the polymers with chain extender (HDI) the restriction of chain motion due to possible hydrogen bonds in between urea groups in the soft segments would produce a reduction in the permeability values. The existence of these hydrogen bonds that restrict polyether chain mobility has already been proved in aromatic isocyanate chain extended polyethers (Marcos-Fernández et al., 1997).

Therefore it seems clear that when permeation is measured at a high enough temperature to have only amorphous PEO longer chains improves the passage of gases. This indicates that for each length of PEO it is necessary to find an optimum working temperature to get all PEO in amorphous state, in order to obtain the maximum
performance. Referring to selectivities, it seems clear that they can be considered almost constant; what means that for these block-copolymers, the aromatic part determines the selectivity of the sample (as mentioned in this study the proportion of PEO is almost constant). This leads to a clear possibility of controlling permeability without damaging selectivity by simply increasing the length of PEO chains. On the other hand by varying the aromatic part, higher selectivity could, of course, be obtained. Naturally, for the copolymers that are already amorphous at 30 °C, permeability increases with the permeation temperature while selectivity decreases only very slightly.

3.7.4. Permselective properties as a function of the PEO percentage

For all samples the content of PEO is quite similar, ranging from 42 to 45 %, (between 42 and 49 % considering the HDI+PEO content— see Table 1). Consequently the effect of the polyether proportion should be very small in permeability. Here results for both the permeation temperatures of 30 and 50 °C are considered too.

Figure 16a shows the evolution of permeability as a function of the percentage of total polyether in the final polymer. It seems clear that the amount of total polyether or PEO doesn’t play a decisive role for this narrow range of percentages, and therefore it is the length of the PEO chains the key parameter.

In the case of samples with chain extender (HDI), that have the highest percentages of polyether in the sample, it can be seen that they don’t show the best permeabilities. The reduction of mobility of the chains causes lower permeability (Marcos-Fernández et al., 1997) and activation energies as seen in Figure 14.

![Figure 16. CO₂ permeability (a) and selectivities (b) for some gas pairs as a function of the PEO percentage for HDI extended copolymers (treated at 160 °C) and for the rest of copolymers (treated at 250 °C).](image)

It has been carried out an analysis of the influence of the percentage of polyether on the selectivity for different gas pairs. Figure 16b shows the selectivity for the synthesized copolymers and also for the homopolymer BPDA-ODA and for the pure PEO at 30 °C. It can be seen that all samples show very close selectivities regardless of the percentage
of polyether. A comparison of selectivity with those of the pure aromatic polyimide, and pure PEO shows that for the couples O₂/N₂ and CO₂/CH₄ selectivities are between those of both BPDA-ODA and pure PEO. In the case of couples CO₂/N₂ and CH₄/N₂, the samples showed exceptional results, surpassing the selectivities of BPDA-ODA and pure PEO.

3.7.5. Robeson’s plot

After confirming that the working temperature and the length of the PEO chains, for similar percentages of polyether, play key roles in permeation properties, these permselective properties are themselves considered here on. A visual way of representing the permeability values obtained for different gases are the Robeson’s representations (Robeson, 1991, 2008). In these representations the distance to the so called upper bond gives us an idea of the permeation properties of the samples.

In addition to our copolymers, we have represented the values for pure PEO at 35 ºC (Lin and Freeman, 2004) and for BPDA-ODA (synthesized in our lab) at 30 ºC. The representative points for our copolymers should be in between these two values. As it’s well known, the Robeson’s upper bounds depend on the measurement temperature (Rowe et al., 2010). This is also taken into account when comparing data at 30 and at 50 ºC.

Figure 17. Robeson’s plot for O₂/N₂. Full circles correspond to a temperature of permeation of 30 ºC; while hollow ones were measured at 50 ºC.

Figure 18. Robeson’s plot for CO₂/CH₄. Full circles correspond to a temperature of permeation of 30 ºC; while hollow ones were measured at 50 ºC.

Figure 19. Robeson’s plot for CO₂/N₂. Full circles correspond to a temperature of permeation of 30 ºC; while hollow ones were measured at 50 ºC.

Figure 20. Robeson’s plot for CH₄/N₂. Full circles correspond to a temperature of permeation of 30 ºC; while hollow ones were measured at 50 ºC. A pseudo Robeson upper bound has been drawn by us from the best results in literature.
In these Figures, the crosses correspond to BPDA-ODA and the stars to pure PEO. The full circles represent the measurements at 30 °C and the hollow circles represent the measurements at 50 °C. The best permeabilities at 30 °C were given by the cPI-PEO2000 sample; whereas for 50 °C, the best results are shown by cPI-PEO6000. As already mentioned, the high permeability for cPI-PEO6000 is due to the elimination of all crystallinity in this copolymer at 50 °C. Therefore the high length of cPI-PEO6000 leads to higher permeability.

For couples O₂/N₂ and CO₂/CH₄ (Figure 17 and 18), the results are far from their respective upper bound. For the couple CO₂/N₂ the polymers showed an excellent performance (see Figure 19). Whereas when the measuring temperature was 30 °C the polymers showed values close to the Robeson’s limit, when the measurement temperature was 50 °C results clearly are even closer to the upper bound for this temperature. For the couple CH₄/N₂ the polymers showed a reverse selectivity getting reasonably good figures. For this couple, there is not a Robeson’s plot thus a tentative limit (drawn by us) is shown in Figure 20 (Tena et al., 2012c).

3.7.6. Permeability prediction

The permeability for the samples BPDA-ODA-PEO with different PEO lengths in the polymer may be predicted by the Maxwell equation as exposed below (Maxwell, 1954):

\[
P_{\text{eff}} = \phi_D P_D + (1 - \phi_D) P_C
\]  

(3)

where \( P_{\text{eff}} \) is the effective permeability, \( P_C \) and \( P_D \) are the permeability of the continuous phase and the dispersed phase, respectively, and \( \phi_D \) is the volume fraction of the dispersed phase in the block copolymer. In this type of polymers, we should take into consideration that the roles of continuous or disperse phases could be played by the hard (aromatic) or by the soft (aliphatic) segments of the copolymer. We will consider the aliphatic (PEO containing) phase as the continuous phase, because it has already been proven, for this type of system, that when the amount of PEO is higher than 20 %, the disperse phase is the aromatic part (Hangzheng et al., 2010). Under these assumptions we built the Maxwell prediction curve. The permeability of pure PEO in the amorphous
state is used as the permeability of the soft segments (continuous phase) (Lin and Freeman, 2004) and the permeability of BPDA-ODA to the hard segments (disperse phase).

Figure 21. Predicted values for CO$_2$ and N$_2$ permeability (a) and CO$_2$/N$_2$ selectivity (b) using Maxwell equation along with the corresponding experimental results.

Figure 21 shows the prediction of permeability and selectivity prediction for the CO$_2$/N$_2$ gas pair according to the Maxwell model. The model shows a good fit for permeability, although samples with high crystallinity showed a slightly poorer fit, and the polymers with longer PEO have the highest crystallinity. On the other hand, the phase segregation was better for longer PEO chains, thus the model should be better adapted. Referring to selectivity, the fit was acceptable for all samples. This model brings new evidence of the importance of crystallinity in the samples, so for that to get the best fit, it is necessary that the samples are in amorphous state and phase segregation be as high as possible.

4. Conclusions

A series of copoly(ether-imide)s with approximately 45% weight of polyether in the final copolymer presenting good gas separation properties have been prepared. These copolymers have been synthesized by the reaction between an aromatic dianhydride (BPDA), an aromatic diamine (ODA) and a diamine terminated poly(ethylene oxide) having a different molecular weight (900, 2000 and 6000 g / mole), or diisocyanate (HDI) extended poly(ethylene oxide) 900 and 2000 with the aim of obtaining a greater length of PEO (approximately 6000 g/mol) but with less tendency to crystallization.

TGA analysis confirmed the weight amount of PEO in the copolymers and showed that the polyether moiety can be eliminated selectively by thermal treatment. Thermal treatment decreased the Tg value for the polyether phase in the completely amorphous copolymers or increased its crystallinity for the copolymers with polyether crystallization capability except for copolymer with the shortest PEO, cPI-PEO900 for which no change in thermal properties was observed. For the chain extended polyethers, the surprising low Tg value and the higher crystallinity respect to the parent polyethers 900 and 2000 proved that chain extension is an effective way to increase the phase
separation in these copolymers. TMA data complemented DSC data and a $T_g$ well above ambient temperature for the aromatic polyimide phase was measured. This $T_g$ increased with the increase of PEO length due to the increase in the aromatic polyimide segment length, but for the copolymers with chain extended polyethers, the value was lower than expected, probably due to some mixing of soft segments in the hard segment phase promoted by the urea groups.

SAXS analysis confirmed the existence of a phase separated morphology in these copolymers that improved when they were thermally treated at increasing temperatures below the temperature of the onset of decomposition. After initial treatment temperatures that released the remaining solvent in the imidized copolymers, a fast improvement in the phase separation was produced at higher temperatures. The size scale of the phase separated structure was initially similar for the copolymers based on the parent polyethers, around 8 nm, and increased simultaneously with the improvement in the phase separation up to a value of 12 to 19 nm. For the copolymer with chain extended polyethers, long spacing was much higher, 40 to 50 nm, with almost no change with treatment temperature. The large size of the phases would explain the DSC results for these copolymers.

Gas permeability for the synthesized copolymers was directly related to the phase separated structure present in the material, with an increase in permeability when treatment temperature, and phase separation, was increased. For copolymers with chain extended polyethers, the degradation of the urea groups led to an increase of segregation but with considerable internal bonding that led to a decrease in permeability. At the same time, selectivity was almost unaffected and remained constant at any treatment temperature. In this way, we could conclude that chain enlarging with HDI is not efficient in increasing permeability.

Temperature is important in the case of polymers that show a high content of crystallized polyether. Thus studies of the behavior of the copolymers as a function of working temperature have been carried out. The amount of crystalline PEO is zero or negligible for these samples when the temperature is 50 ºC. For this reason, in this case, we have analyzed the results of permeability and selectivity for different pairs as a function of the length of PEO at two temperatures of work, a standard 30 ºC and a
sufficiently high so that all the PEO is in amorphous state, 50 °C. Thus, it was observed that the greater the length of PEO the higher permeability was observed, because the PEO is in purest state. For samples with HDI results were lower than for the sample with PEO6000 owing to the hydrogen bridges forming urea structures. The results of selectivity for all PEO lengths were similar.

Analyzing the results in terms of percentage of PEO in the samples, the highest results were found for 43.5 % for the measurement at a temperature of 30 °C, which corresponds to the sample PEO2000, and to values close to 45 % when as the temperature is 50 °C, which coincides with the sample PEO6000. This again highlights the importance of crystallinity, when all the poly(ethylene oxide) is in amorphous state, the best values are for the longest PEO, the 6000 in this case.

Regarding the Robeson’s plots, the best results were found for the CO₂/N₂ gas couple. At 30 °C the results are quite close to the Robeson upper bound, and in the case of the measures at 50 °C the permselectivity closely approaches the limit for this temperature. In conclusion it seems clear that this type of polymer is suitable for separations where CO₂ is involved, and especially for the separation CO₂/N₂. This separation has a great importance in the greenhouse effect and makes these membranes good candidates for their application at industrial level. An interesting application of CO₂/N₂ separation is in the purification of coal gas. In this process the working temperatures are high, so this type of materials is presented as an attractive alternative.

We have compared the results with those calculated by using the Maxwell’s model. In this case also the percentage of amorphous PEO should be used and crystallinity has to be taken into account for low permeation temperatures. The samples with a low or negligible crystallinity showed a good fit. The need to take into account the truly amorphous portion of PEO makes necessary to optimize both the amount of poly(ethylene oxide) in the sample and the operating temperature for a specific length and amount of PEO.

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6. References.


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Figure 1. Chemical structure of the monomers.

**Dianhydride**

![BPDA](image1)

**Aliphatic Diamine**

![PEO X Diamine Terminated](image2)

Where \( X = 900 \) and \( 2000 \)

**Aromatic Diamine**

![ODA](image3)

\( \alpha,\omega \)-Diamine-poly(ethylene oxide) PEO6000

**Chain Extender**

\( O=C=\left(CH_2\right)_6N=C=O \)

HDI
Figure 2. Thermal treatments.
Figure 3. TGA curves in dynamic conditions for the synthesized copolymers.
Figure 4. Crystallinity of the aliphatic part of the copolymers. The copolymers that do not appear don’t have any crystallinity.
Figure 5. $T_g$ of the aliphatic part of copolymers after “instantaneous” treatment for the copolymers treated at 160 °C.
Figure 6. $T_g$ for the aromatic polyimide segments as a function of the PEO length for the copolymers treated at 160 $^\circ$C.
Figure 7. Density as a function of the polymer length.
Figure 8. Intensity (logarithmic scale) vs q curves for copolymers cPI-PEO900, cPI-PEO2000, cPI-PEO900+HDI and cPI-PEO2000+HDI. In the insert, the $Iq^2$ vs q curves for the same copolymers are presented.
Figure 9. Changes in the relative invariant for the synthesized copolymers treated at 120°C.
Figure 10. Evolution of $L$ as a function of temperature for the synthesized copolymers.
Figure 11. Evolution of the permeability (a) and selectivity (b) measured at 30ºC as a function of the treatment temperature for cPI-PEO900+2000.
Figure 12. CO$_2$ permeability measured at 30°C as a function of the thermal treatment.
Figure 13. CO₂ permeability as a function of permeation temperature for cPI-PEO6000 membranes treated at 160°C.
Figure 14. Activation energy as a function of the molecular weight of PEO (a) and for the longest PEO copolymers (b). In the (a) figure only cPI-PEO2000+HDI, among the copolymers with the longest PEO chains, is shown.
Figure 15. CO$_2$ permeability (a) and selectivities (b) for some gas pairs as a function of the PEO length, for HDI extended copolymers (treated at 160 ºC) and for the rest of copolymers (treated at 250 ºC).
Figure 16. CO₂ permeability (a) and selectivities (b) for some gas pairs as a function of the PEO percentage for HDI extended copolymers (treated at 160 ºC) and for the rest of copolymers (treated at 250 ºC).
Figure 17. Robeson's plot for $O_2/N_2$. Full circles correspond to a temperature of permeation of 30 °C; while hollow ones were measured at 50 °C.
Figure 18. Robeson’s plot for CO$_2$/CH$_4$. Full circles correspond to a temperature of permeation of 30 ºC; while hollow ones were measured at 50 ºC.
Figure 19. Robeson’s plot for CO$_2$/N$_2$. Full circles correspond to a temperature of permeation of 30 °C; while hollow ones were measured at 50 °C.
Figure 20. Robeson’s plot for CH$_4$/N$_2$. Full circles correspond to a temperature of permeation of 30 °C; while hollow ones were measured at 50 °C. A pseudo Robeson upper bound has been drawn by us from the best results in literature.
Figure 21. Predicted values for CO$_2$ and N$_2$ permeability (a) and CO$_2$/N$_2$ selectivity (b) using Maxwell equation along with the corresponding experimental results.