Influence of Cation and Anion Type on the Formation of the Electroactive β-Phase and Thermal and Dynamic Mechanical Properties of Poly(vinylidene fluoride)/Ionic Liquids Blends

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Abstract

Films based on poly(vinylidene fluoride) (PVDF) blended with ionic liquids (ILs) comprising different cations and anions were developed to investigate the IL influence on the resulting PVDF crystalline phase. Blends with 25 wt % IL content were produced by solvent casting followed by solvent evaporation at 210 °C in an air oven. Five different ILs containing the same cation 1-ethyl-3-methylimidazolium [Emim] and five ILs containing the same anion bis(trifluoromethylsulfonyl)imide [TFSI] were selected. The formation of the different phases and the resulting thermal and dynamic mechanical properties were studied by Fourier transform infrared spectroscopy, differential scanning calorimetry, and dynamic mechanical analysis. The incorporation of [Emim]-based ILs successfully directs the PVDF crystallization from the nonpolar α-phase toward the electroactive and highly polar β-phase. On the contrary, blends containing [TFSI] as a
common anion yield a mixture of α and β phases. Overall, the induced β-phase ranges between 14 and 95% depending on the incorporated IL type. Interestingly, 1-ethyl-3-methylimidazolium chloride [Emim][Cl] is the most effective IL among the studied ones to significantly enhance the β-phase content, showing also a marked nucleating effect. The results suggest that ions favorably interact with PVDF chains and are located occupying the amorphous interlamellar PVDF regions. Furthermore, the studied ILs act as plasticizers, yielding lower glass-transition temperatures of the amorphous phase and decreasing mechanical storage modulus at room temperature.

**Keywords:** PVDF; ionic liquids; anion; cation; crystallization.
1. Introduction

Poly(vinylidene fluoride) (PVDF) is a semicrystalline polymer showing very interesting multifunctional properties for sensor and actuator applications, among others. [1,2]. The most common polymorph is the α-phase, where CF2 groups are established in trans–gauche–trans conformations, which result in a nonpolar chain configuration as C–F bonds are arranged in an alternating fashion along the chain. From the other possible four polymorphs (β, γ, δ, and ε), the β-phase is the most technologically relevant for applications, where C–F bonds are normal to the backbone chain and the obtained all-trans conformation leads to a large piezoelectric and ferroelectric response [2]. A wide range of applications such as capacitors, battery separator, or sensors have been proposed so far for β-PVDF [3–5]. The α phase is the most common form of PVDF as it is obtained during crystallization from the melt. As a result, many works have been focused on the development of β-phase containing PVDF. The techniques reported so far to obtain the β-phase PVDF include, but are not limited to, filler addition [6], blending [7], poling [8], stretching [9], or by nanoconfinement [10].

Although such techniques have proven efficient toward the development of electroactive PVDF, these approaches suffer from challenges associated with homogeneous filler dispersion (composite preparation), phase separation (polymer blending), or expensive and tedious processes (poling, stretching, and nanoconfinement), which limit their processability at industrial level. An efficient and straightforward approach to circumvent the above-mentioned issues may be the incorporation of organic compounds in a PVDF matrix by simple mixing processes. In that framework, room-temperature ionic liquids (ILs) emerge as a perfect choice as they can be easily processed at room temperature by simply incorporating selected ILs into dissolved PVDF and have been proven to be highly compatible with many polymer matrices [11–13]. ILs are salts whose melting point is below 100 °C and are composed of a small anion and an asymmetric heterocyclic cation [14]. Moreover, ILs are generally considered as environmentally benign [15], present a negligible vapor pressure, and their high thermal stability and ionic conductivity may serve to upgrade the multifunctional properties of PVDF [16]. The influence of the incorporation of ILs into the PVDF polymer matrix, such as 1-hexadecyl-3-methylimidazolium bromide [C16mim][Br] [17], or butyl methylimidazolium hexafluorophosphate [Bmim][PF6] [18], on the physical and chemical properties of the PVDF has been studied. It has been reported that the
incorporation of ILs into the PVDF polymer matrix induces an extended conformation of PVDF into its polar β-phase and that PVDF/IL blends show excellent mechanical characteristics, with significantly increased ductility, good optical transmittance, and higher electrical conductivity [18]. However, to the best of our knowledge, no studies have systematically reported on the influence of the ILs with different functional groups on the PVDF crystallization process toward the formation of the β-phase. In this sense, strong dipolar interactions are expected between the heterocyclic cation of ILs and the CF2 moiety of PVDF. Similarly, the nature of the anion could also influence the resulting interactions between the IL and the polymer. PVDF/IL blends in the form of films with IL with different cation chain sizes have been evaluated showing that the characteristics of the IL strongly influence the physicochemical characteristics of the composites, including the increase of the alternating current conductivity [19].

In a previous work, the mixture of PVDF and ionic liquid 1-ethyl-3-methylimidazolium [Emim] bis(trifluoromethylsulfonyl)imide [TFSI] was evaluated, analyzing the dependence of the melting temperatures and crystallization on IL concentration within the polymer matrix. It was observed that the presence of IL induced the formation of the β phase of PVDF [20]. In the present work, we have extended the study to a wide series of ILs with different anions and cations to analyze the influence of the interaction of the anion and cation with the polymer chains in the crystallization of the PVDF. Accordingly, in this work, blends of PVDF with 10 different ILs were prepared to study the effect of the anion (by keeping the cation unchanged) and cation (by keeping the anion unchanged) on the conformation and crystalline structure of PVDF. The associated thermal transitions and dynamic mechanical properties are also discussed.

2. Experimental section

2.1. Materials

PVDF (Solef 6020, $M_w = 700\,000$ g mol$^{-1}$) was purchased from Solvay, and N,N-dimethylformamide (DMF) (99.5%) was obtained from Merck. The ILs used in this work, comprising different cations and anions, were obtained from IoLiTec and are indicated in Table 1. The effect of the anion was evaluated using the following ILs: 1-ethyl-3-methylimidazolium hexafluorophosphate [Emim][PF$_6$], 1-ethyl-3-methylimidazolium chloride [Emim][Cl], 1-ethyl-3-methylimidazolium hydrogen sulfate [Emim][HSO$_4$], 1-
ethyl-3-methylimidazolium ethylsulfate [Emim][C₂SO₄], and 1-ethyl-3-methylimidazolium bis(trifluoromethylsulfonyl)imide [Emim][TFSI], 1-Hexyl-3-methylimidazolium bis(trifluoromethylsulfonyl)imide [Hmim][TFSI], 1-octyl-3-methylimidazolium bis(trifluoromethanesulfonyle)imide [Dmim][TFSI], N,N,N-trimethyl-N-(2-hydroxyethyl)ammonium bis(trifluoromethylsulfonyl)imide [N1112OH][TFSI], trihexyl(tetradecyl)phosphonium bis(trifluoromethylsulfonyl)imide [P66614][TFSI], and 1-butyl-1-methylpyrrolidinium bis(trifluoromethanesulfonyle)imide [C₄C₁Pyrr][TFSI] were used to study the effect of the cation on the properties of the blends. The selection of the different cations and anions is based on the different structural and dimensional characteristics, as presented in Table 1, modulating therefore the interactions with the polymer chain.

**Table 1** - Structure and Melting Temperature of the ILs Used for the Preparation of the IL–PVDF Blends.

<table>
<thead>
<tr>
<th>IL</th>
<th>Cation</th>
<th>Anion</th>
</tr>
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<tbody>
<tr>
<td>[Emim][Cl]</td>
<td></td>
<td>Cl⁻</td>
</tr>
<tr>
<td>[Emim][PF₆]</td>
<td></td>
<td>[PF₆]²⁻</td>
</tr>
<tr>
<td>[Emim][HSO₄]</td>
<td></td>
<td>[HSO₄]⁻</td>
</tr>
<tr>
<td>[Emim][C₂SO₄]</td>
<td></td>
<td>[C₂SO₄]⁻</td>
</tr>
<tr>
<td>[Emim][TFSI]</td>
<td></td>
<td>[TFSI]⁻</td>
</tr>
<tr>
<td>[Hmim][TFSI]</td>
<td></td>
<td>[TFSI]⁻</td>
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<tr>
<td>[Dmim][TFSI]</td>
<td></td>
<td>[TFSI]⁻</td>
</tr>
<tr>
<td>[C₄C₁Pyrr][TFSI]</td>
<td></td>
<td>[TFSI]⁻</td>
</tr>
</tbody>
</table>
2.2. Sample preparation

PVDF was dissolved in DMF in a proportion of 15/85 wt % under magnetic stirring at room temperature. After complete polymer dissolution, the selected ILs (25 wt %) were added to the solution. The IL content of 25 wt % was selected as it has been demonstrated that this amount is high enough to allow substantial nucleation of the electroactive phase by different ILs, allowing a suitable comparison among them [21,22]. PVDF/IL films with a thickness of approximately 50 μm were obtained after spreading the solution at room temperature on a glass substrate followed by solvent evaporation at 210 °C (above melting temperature of the polymer for complete and rapid solvent evaporation) in an air oven (P-Selecta), following the general procedure described in ref [23].

2.3. Characterization techniques

Fourier transform infrared (FTIR) measurements were carried out with a Jasco 4100 spectrometer in the attenuated total reflection (ATR) mode from 4000 to 400 cm\(^{-1}\). FTIR spectra were collected after 64 scans with a resolution of 4 cm\(^{-1}\). Differential scanning calorimetry (DSC) was carried out with a DSC Mettler Toledo 823e for low-temperature measurements conducted in the region of glass transition and a DSC 8000 of PerkinElmer for scans in the crystallization and melting region, in both cases under flowing nitrogen (N\(_2\)) atmosphere. The mass of the sample was between 5 and 10 mg. Dynamic mechanical analysis (DMA) was carried out in a PerkinElmer DMA 8000 apparatus in tensile mode. The storage modulus and loss tangent were measured as a function of temperature at a frequency of 1 Hz from −100 to 125 °C at a heating rate of 2 °C min\(^{-1}\). The dimension of each sample was around 10 × 4 × 0.050 mm\(^3\). For each sample, the FTIR spectra and DSC and DMA curves were measured in triplicate.
3. Results and discussion

3.1. FTIR characterization

Both blend constituents, ILs and PVDF, have the ability to crystallize, with the melting temperature of the ILs below the melting temperature of PVDF. At temperatures above the melting temperature of PVDF, a homogeneous mixture of IL and PVDF liquid chains is expected. Thus, on cooling, nucleation and growth of PVDF crystals take place in an environment in which chain segments interact with IL ions and, on the other hand, an environment in which the conformational mobility of PVDF macromolecules is higher than in pure PVDF, which is attributed to the interaction of the ILs with the PVDF polymer chain (cation–polymer and anion–polymer interactions) [24]. Both these factors are expected to influence the crystallinity and the crystal phase of the polymer. Therefore, here we attempt to analyze the effect of different ILs on the crystalline structure of PVDF blends.

To obtain a homogeneous mixture of PVDF and IL, the samples were obtained after dissolving PVDF and the required IL in DMF, followed by a solvent evaporation process at high temperatures (210 °C) [23] to promote the crystallization process through cooling from the molten blend.

The effect of the different ILs on the PVDF chemical properties was evaluated by FTIR-ATR analysis through the identification and quantification of the different crystal phases of PVDF. Figure 1a displays the FTIR spectra of both PVDF film, [Emim][C₂SO₄] IL, and the corresponding blend. Neat PVDF displays sharp bands at 615, 763, 796, and 976 cm⁻¹, which correspond to the α-phase [25]. It is interesting to note that no conformational changes are obtained upon the processing of PVDF powder into films by solvent casting followed by drying at 210 °C (as supported by the FTIR spectra in Figure S1, where both samples present virtually identical spectra). The PVDF/[Emim][C₂SO₄] blend presents a weak absorption band at 1275 cm⁻¹ and a stronger one at 840 cm⁻¹, both ascribed to the β-phase [25]. The absorption bands at 1071, 1176, and 1402 cm⁻¹ are also assigned to β PVDF [24,26]. Remarkably, with the IL incorporation, the intensity of the absorption bands characteristic of the α-phase significantly decreases. The absence of a shoulder absorption band at 833 cm⁻¹ and also the absence of peaks at 1234 and 811 cm⁻¹, characteristic of the γ phase, indicate the PVDF/[Emim][C₂SO₄] blend crystallization into the polymorph β phase [27,28]. This is in contrast to the previously reported works where a transformation from α to a mixture of β and γ crystal phases has been reported.
by mixing PVDF with 1-hexadecyl-3-methylimidazolium bromide [C16mim][Br] [17], or 1-butyl-3-methylimidazolium hexafluorophosphate [Bmim][PF₆] [18]. In marked contrast to the directing effect of [Emim][C₂SO₄], Figure 1b shows that the incorporation of [Emim][PF₆] into PVDF does not promote modifications in the crystal phase of PVDF films as the obtained spectra are a mere superposition of the α-containing PVDF and the IL due to the small size of the anion. This fact reveals the relevance of the selection of ILs with an appropriate anion–cation pair.

To search for further suitable ILs as directing agents toward the β crystal phase, PVDF has been blended with a series of ILs having [Emim] as a common cation. Figure 2 displays the obtained results, where the most relevant PVDF fingerprint regions have been enlarged for the sake of comprehension. First, Figure 2a is focused on the 900–700 cm⁻¹ region, where the bands at 763, 840, and 870 cm⁻¹ are assigned to the α-phase, β-phase, and amorphous PVDF, respectively. As an overall trend, all of the blends showed a decreased intensity (or even vanishing) of the 763 cm⁻¹ absorption band together with an increase of the 840 cm⁻¹ band, suggesting that the presence of the α-phase is markedly reduced upon the introduction of such ILs. Moreover, it is also observed that the amorphous 870 cm⁻¹ band of CF₂–CH₂ bending vibration tends to increase, which is indicative of the presence of larger amorphous region within the blend. This may be due to the fact that, apart from the directing effect of ILs toward the β-phase, their incorporation also brings a plasticizing effect (absorption of ions that migrate between the macromolecular chains, thus allowing the plastic part to lose stiffness) [29], which limits the organization of PVDF macromolecules into crystalline regions [12,13]. Interestingly, it should be noted that the amorphous PVDF band becomes wider and is
shifted to a higher-energy region by 4–6 cm⁻¹, indicating that >CF₂ groups effectively interact with the IL [30]. This interaction denotes the good miscibility between PVDF and ILs, where IL may be found occupying amorphous interlamellar regions. The results in Figure 2b agree with the directing effect of ILs as an increase of the 510 cm⁻¹ band (corresponding to β-phase) at the expense of the α-phase 485 cm⁻¹ band is also observed. Finally, Figure 2c demonstrates that the PVDF/[Emim][TFSI] sample shows an absorption band around 1230 cm⁻¹, indicative of the presence of γ-PVDF [28]. For all of the other samples, no trace of γ-PVDF is observed.

![FTIR spectra of PVDF/IL blends](image)

**Fig. 2.** FTIR spectra of PVDF/IL blends having [Emim] as a common cation in the: a) 900-700 cm⁻¹; b) 600-400 cm⁻¹ and c) 1300-1100 cm⁻¹ regions.

Furthermore, it has been observed that ILs having [Emim] as a common cation have a similar effect on the crystalline phase of PVDF, independently of the anion.

Blends having an IL with a common anion have also been obtained to investigate the effect of the cation on the resulting crystalline phase. Accordingly, Figure 3 shows the FTIR spectra corresponding to the PVDF/IL blends with [TFSI] as a common anion. Therefore, those results allow to understand the effect of different cations on the crystalline phase of PVDF. Both PVDF/[Dmim][TFSI] and PVDF/[N1112OH][TFSI] blends present the α-phase as indicated by the bands located at 763 and 485 cm⁻¹, while
the 840 cm\(^{-1}\) band indicative of the \(\beta\)-phase is achieved for all of the blends (despite the notable intensity differences). Similarly, blends show the \(\beta\)-phase absorption band at 510 cm\(^{-1}\) (corresponding to \(\beta\)-phase) together with a displacement toward higher-energy region of the 870 cm\(^{-1}\) band, suggesting that ILs interact with the PVDF chains. Interestingly, the blends showing \(\alpha\)-phase (PVDF/[Dmim][TFSI] and PVDF/[N1112OH][TFSI]) do not present a notable shift on the amorphous 870 cm\(^{-1}\) band and also the intensity of the 840 cm\(^{-1}\) band (\(\beta\)-phase) is very low. Therefore, contrarily to [Emim]-containing ILs, the cationic ions in [C4C1Pyrr] and [P66614] are able to interact with the \(\text{>CF}_2\) moieties in PVDF to locate \(\text{>CF}_2\) along one side of the main chain, similarly to previously obtained PVDF samples blended with imidazolium-ion-based ionic liquids [18].

Overall, while most of the blends having [Emim] as a common cation yield the electroactive \(\beta\)-PVDF phase, blends having [TFSI] as a common anion yield a mixture of \(\alpha\), \(\beta\), and \(\gamma\) phases. In any case, it is interesting to note that most samples with different IL types result in the crystallization of the polymer just in the electroactive and highly polar \(\beta\) phase, which is in contrast to previously reported works where a mixture of both \(\beta\) and \(\gamma\) phases was obtained [17,18,30].

![Fig. 3. FTIR spectra of PVDF/IL blends having [TFSI] as a common anion in the: a) 900-700 cm\(^{-1}\); b) 600-400 cm\(^{-1}\) and c) 1300-1100 cm\(^{-1}\) regions.](image-url)
It is observed that the chain structure (i.e., the anion and cation type) affects the ability of the ILs to act as directing agents for PVDF phase crystallization, highlighting the relevance of properly selecting the anion to be able to achieve the electroactive $\beta$-phase. The quantification of the $\beta$-phase could shed light on the efficiency of different ILs to give the all-trans chain conformation of PVDF as a result of the electrostatic PVDF–IL interactions through ion–dipole interactions [31]. When no coexistence of the $\beta$ and $\gamma$ crystal phases exists, the extent of the polar $\beta$ crystalline phase ($F(\beta)$) can be computed as assuming that IR absorption follows the Lambert–Beer law [32]:

$$F(\beta) = \frac{A_\beta}{1.26 \cdot A_\alpha + A_\beta}$$

(1)

where $A_\alpha$ and $A_\beta$ are the absorbances at 763 and 840 cm$^{-1}$, respectively. The obtained results are reported in Table 2. It is observed that depending on the IL, $F(\beta)$ could reach up to a maximum value of 95% for the PVDF/[Emim][Cl] blend, indicating that such IL plays the most effective role (among the studied ILs) as directing agent for the nucleation of the electroactive $\beta$-phase during the solvent evaporation process. This may be due to the presence of a large electrostatic interaction between the negatively charged chlorine anion (Cl$^-$) and the positively charged PVDF’s dipolar moment, boosting the development of the $\beta$-phase more effectively [33]. These results prove that by selecting the proper IL, an efficient way to obtain large $\beta$-phase fractions could be achieved through this method as other approaches such as uniaxial stretching render materials with an $\alpha$-phase content that can be higher than 20% (depending on the applied stretching ratio and temperature) [2,34].

**Table 2.** $\beta$ Crystalline PVDF Phase Obtained Depending on the IL Used for an IL Content of 25 wt %.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Neat PVDF</th>
<th>[Emim][C$_2$SO$_4$]</th>
<th>[Emim][Cl]</th>
<th>[Emim][HSO$_4$]</th>
<th>[Emim][TFSI]</th>
<th>[Emim][PF$_6$]</th>
</tr>
</thead>
<tbody>
<tr>
<td>$F(\beta)$</td>
<td>33 %</td>
<td>80 %</td>
<td>95 %</td>
<td>82 %</td>
<td>83 %</td>
<td>0 %</td>
</tr>
<tr>
<td>Sample</td>
<td>[C4ClPyr$</td>
<td>$][TFSI]</td>
<td>[Dmim][TFSI]</td>
<td>[Hmim][TFSI]</td>
<td>[N1112OH][TFSI]</td>
<td>[P66614][TFSI]</td>
</tr>
<tr>
<td>$F(\beta)$</td>
<td>85 %</td>
<td>43 %</td>
<td>86 %</td>
<td>36 %</td>
<td>87 %</td>
<td>83 %</td>
</tr>
</tbody>
</table>
3.2. DSC results

The effect of ILs on the crystallinity of PVDF was also evaluated by DSC measurements. Accordingly, DSC cooling scans of melted PVDF/IL blends having different anions and cations are depicted in Figure 4a,b, respectively. The achieved bell-shaped endothermic peak upon cooling from the melt arises from the arrangement of PVDF chains into well-ordered regions. The peak crystallization temperatures ($T_c$) for all of the samples together with their crystallization enthalpy ($\Delta H_c$) are listed in Table 3. It is observed that upon the addition of ILs, the $T_c$ of PVDF decreases from 134 °C to a minimum value of 113 °C for the PVDF/[Emim][PF$_6$] as a result of the incorporation of the ILs [18,35].

![Fig. 4. DSC cooling scans of PVDF/IL blends having different a) anions and b) cations. Normalized heat flow (heat flow divided by the sample mass and the cooling rate) is represented.](image-url)

Table 3. Crystallization Temperature ($T_c$) and Enthalpy ($\Delta H_c$) upon Cooling from the Melt and Melting Temperature ($T_m$) for the Different PVDF/IL Blends (The effect of the anion is shown on rows 2 to 6, while rows 6 to 10 displays the effect of the cation).

<table>
<thead>
<tr>
<th>Arrow</th>
<th>Blend</th>
<th>$T_c$ ± 1 °C</th>
<th>$\Delta H_c$ ± 2% (J·g$^{-1}$)</th>
<th>$T_m$ ± 1 °C</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>PVDF</td>
<td>134</td>
<td>47</td>
<td>169</td>
</tr>
<tr>
<td>2</td>
<td>PVDF/[Emim][Cl]</td>
<td>127</td>
<td>63</td>
<td>165</td>
</tr>
<tr>
<td>3</td>
<td>PVDF/[Emim][HSO$_4$]</td>
<td>139</td>
<td>86</td>
<td>172</td>
</tr>
<tr>
<td>4</td>
<td>PVDF/[Emim][PF$_6$]</td>
<td>113</td>
<td>48</td>
<td>158</td>
</tr>
<tr>
<td>5</td>
<td>PVDF/[Emim][SO$_4$]</td>
<td>124</td>
<td>57</td>
<td>164</td>
</tr>
<tr>
<td>6</td>
<td>PVDF/[Emim][TFSI]</td>
<td>120</td>
<td>47</td>
<td>162</td>
</tr>
</tbody>
</table>
PVDF presents a well-defined single endothermic peak associated with melting at 168 °C (Figure 5a), which matches with the melting of the α-phase previously reported [2]. 

$T_m$ is lowered upon the addition of [Emim][TFSI], [Emim][Cl], and [Emim][PF6] to a minimum of 158 °C. It should be taken into account that the melting temperature of both α and β PVDF phases is reported to be found within the range of 167–172 °C [2]. Therefore, although samples containing ILs show a $T_m$ below the 167–172 °C range, their crystal structure still corresponds to α or β phase, as the lowering of the melting temperature is due to the freezing point depression and the presence of specific interactions between PVDF macromolecules and free IL chains as shown in many miscible amorphous–crystalline (PVDF–IL in this specific case) blends [36–38].

**Fig. 5.** DSC heating scans showing a) the glass transition and b) the melting of PVDF/IL blends. Normalized heat flow (heat flow divided by the sample mass and the heating rate) is represented.

The thermodynamic equilibrium criteria lead to the fact that when a solute is added to a liquid, its melting temperature decreases. This is known as the freezing point depression. Thus, if in the liquid mixture, above the melting temperature of PVDF, the ionic liquid dissolves homogeneously in the liquid PVDF, then the melting temperature of PVDF should decrease, as it is observed in the mixtures with [Emim][TFSI], [Emim][Cl], and [Emim][PF6]. The inset in Figure 4a shows a schematic representation of the phase diagram of a blend that is miscible in the liquid phase but not in the solid phase. By

<table>
<thead>
<tr>
<th></th>
<th>PVDF/[C4C1pyrr][TFSI]</th>
<th>PVDF/[Hmim][TFSI]</th>
<th>PVDF/[N1112OH][TFSI]</th>
<th>PVDF/[P66614][TFSI]</th>
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<tr>
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<td>173</td>
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cooling of a blend rich in PVDF, as it is the case of our blends, PVDF crystals form in equilibrium with a liquid blend, which becomes richer in the IL. This equilibrium takes place at a temperature lower than that of pure PVDF. Depression of both crystallization and melting temperatures (see Figure 5b) is a strong indication of the formation of PVDF crystals from a homogeneous liquid phase containing IL. Actually, in the case of polymeric systems, other kinetic factors must be considered. After IL incorporation, PVDF chains require longer migrating distances to reach the crystal growth front within the formed miscible blend, which in turn yields slower crystallization kinetics. As a consequence of such delayed crystallization ability, lower crystallization temperatures are obtained. Nevertheless, the relatively small IL content and the observed decrease of the melting temperature leads us to think that, in the present case, kinetic effects are less influential than thermodynamic ones.

If, on the contrary, the liquid PVDF and IL phases separate, the ionic liquid does not act as a solute of the PVDF and, therefore, the freeze-point depression does not occur and crystallization takes place in a pure PVDF phase. Furthermore, as the temperature of crystallization depends on the process of nucleation of the crystals and crystallization occurs more easily on surfaces already formed, such as impurities or interfaces, when phase separation occurs, the formed interfaces can act as nucleating agents and facilitate crystallization, leading to an increase of the crystallization temperature. The blends containing [Emim][HSO₄] (Figure 4a) and [P66614][TFSI] (figure 4b) show an increase of approximately 5 °C on their crystallization temperature, and the melting temperature \( T_m \) increases by 5 °C as well (Figure 5b). These results suggest that the liquid phase is immiscible and, in addition, an efficient nucleating effect from the ionic liquids as the crystallization is accelerated [39]. It is also observed in Table 3 that the crystallization enthalpy per gram of PVDF in the blend increases from 47 J g⁻¹ up to a maximum of 86 J g⁻¹ for the blend containing [Emim][HSO₄]. This increase indicates that a notably larger fraction of PVDF is able to crystallize during the cooling process when such IL is added. The addition of other ILs does not have such a marked nucleating effect. Therefore, it can be concluded that the role of most of the ILs is to act as a directing agent toward the \( \beta \)-phase, while [Emim][HSO₄] also acts as an effective nucleating element to enhance the extent of the crystalline fraction (and not only its conformation).

As miscible blends display a single glass-transition \( T_g \) intermediate between those of pure blend constituents and noncompatible blends typically have a double \( T_g \) region, DSC
emerges as one of the most useful methods to study the miscibility in polymer blends [40,41]. DSC heating scans displaying glass transition are shown in Figure 5a. It is observed that depending on the incorporated IL, a single or a double glass transition is observed (marked by the arrows in Figure 5a). For instance, the PVDF/[Emim][HSO₄] blend presents a well-defined step in the heat capacity at −98 °C and a broader transition in the temperature range of −50 to −10 °C. The first transition is ascribed to the glass transition of the [Emim][HSO₄], while the second one corresponds to the glass transition of a blend phase rich in PVDF [42]. PVDF/[P66614][TFSI] blend also displays two separated glass-transition regions, indicating that the blend is not miscible, while PVDF/[Emim][PF₆] shows indication of two broad glass transitions that are ascribed to the partial miscibility between the two components and no signs of IL melting is observed. On the contrary, PVDF/[Emim][TFSI] blend displays a single and broad transition, suggesting a good miscibility between the ILs and the amorphous regions of PVDF, as previously observed in FTIR results. The fact that the vitrification behaviors of the different blends are much different from each other indicates a very different interaction between the PVDF and the anion of the IL.

The heating scans of Figure 5 show melting peaks that are clearly widened toward the low-temperature side of the peak, as expected to occur in a mixture of the polymer with a low-molecular-weight substance [43]. However, except in the case of the PVDF mixture with [P66614][TFSI], the thermogram does not show a melting peak associated with the melting of IL crystals or the eutectic point. This melting peak in PVDF/[P66614][TFSI] appears at very low temperatures, around −70 °C. In all of the other blends, at temperatures below 0 °C, the glass transition appears and the vitrification of the IL prevents the appearance of the eutectic point. The temperature of the endothermic peak corresponds thus to the melting of the PVDF crystals and are listed in Table 3.

Although DSC does not allow distinguishing between α and β phases due to their very similar $T_m$ (167–172 °C[2]), the degree of crystallinity ($X_c$ %) obtained during the cooling process can be extracted from the DSC melting enthalpy (Figure 5b) according to the following equation [2]:

$$X_c(\%) = \frac{\Delta H_s}{(x\Delta H_{\alpha} + y\Delta H_{\beta})W_m}$$  \hspace{1cm} (2)
where $\Delta H_s$ is the melting enthalpy of the sample, $W_m$ is the PVDF matrix weight fraction, and $\Delta H_\alpha$ and $\Delta H_\beta$ are the heats of fusion for 100% crystalline $\alpha$-PVDF and $\beta$-PVDF, respectively (set at 93.07 and 104.6 J g$^{-1}$, respectively) [2].

When accounting for the $X_c$ of PVDF/IL blends, one should be aware that the mixing enthalpy may also contribute to the measured enthalpy. In any case, since melting enthalpy is much larger than the mixing enthalpy between polymer and IL, in this work, $X_c$ has been estimated according to Equation 2. Overall, in comparison to neat PVDF, blends show a slight $X_c$ depression from 55 to nearly 45% as a result of the restriction suffered by PVDF macromolecules to organize themselves into ordered regions during the cooling from the melt, which also denotes a good miscibility between both blend constituents. [18].

3.3. PVDF segmental relaxation

FTIR and DSC results indicate that miscibility in PVDF/IL blends is quite dependent on the IL structure within the explored composition range. In this sense, dynamic mechanical analyses (DMA) have been carried out to investigate the effect of different IL on the chain segmental relaxation of PVDF. Figure 6 displays the obtained dynamic mechanical storage modulus ($E'$) and the mechanical damping factor ($\tan \delta = E''/E'$; also known as loss factor) for the processed PVDF/IL blends over the temperature range of −100 to 125 °C. It is observed in Figure 6a,c that in the glassy region, at temperatures below −50 °C, all of the blends present similar values of $E'$. The exception is the blend [Emim][PF$_6$] in which $E'$ in the glassy state is around half of the other blends. In the main dynamic mechanical relaxation of the amorphous phase, $E'$ drops up to a decade upon IL incorporation with respect to pure PVDF. In other words, neat PVDF can keep its structural integrity once the $T_g$ has been overcome, suggesting an increased chain stiffness in comparison to IL-containing samples[44]. Similar behavior has also been observed for IL-blended zwitterionic copolymers, where the IL incorporation results in a reduced electrostatic interaction between ion pairs within the copolymers, increasing the chain mobility and therefore lowering the rubbery plateau [45].
As observed in Figure 6b, neat PVDF is characterized by two main relaxations centered at −35 and 73 °C, attributed to the $\beta_{DMA}$ relaxation arising from the cooperative segmental motions in the amorphous region and to the $\alpha_{DMA}$ relaxation of the imperfections present within the crystalline phases, respectively [46]. This second relaxation achieved at temperatures above $T_g$ involves $180^\circ$ flip motions of the chain sections within the crystalline lamellae and has been also found in other semicrystalline polymers such as isotactic polypropylene, polyethylene, or poly(ethylene oxide) [47,48]. Additionally, the shoulder in the tan $\delta$ signal corresponding to neat PVDF that can be observed in Figure 6b at temperatures around 23 °C has been ascribed to the chain movements at the amorphous–crystal interface [49].

With respect to the amorphous region, there is a correlation between the presence of a single broad glass transition in the DSC thermograms and a broad tan $\delta$ peak in the $\beta_{DMA}$ of the samples, in which there is an homogeneous mixture of IL molecules and amorphous PVDF chains in the amorphous phase of the blend. Thus, Figure 6b shows in this temperature range a single very wide tan $\delta$ peak in the PVDF/[Emim][TFSI] mixture,
which covers practically the entire interval between −80 and +20 °C (red arrow in Figure 6b), confirming the miscibility of this mixture. In the opposite case, the PVDF/[Emim][HSO₄] mixture clearly shows a tan δ peak close to −100 °C, which is associated with the glass transition of the IL, while between −60 and −10 °C, a complex relaxation process appears, which is associated with the glass transition of the phase of the mixture richest in PVDF chains (blue arrows in Figure 6b), correlating with the two clearly separated glass transitions observed in the DSC thermograms (Figure 5a).

Overall, the tan δ value slightly increases for the blends, suggesting an increased presence of additional free volume upon blending with ILs. The rather marked decrease in E’ for the PVDF/[Emim][PF₆] blend at around 53 °C is accompanied by a lowering in the relaxation temperatures corresponding to the chain movements at the amorphous–crystal interface (from 23 to −10 °C) and within the crystalline phases (from 73 to 57 °C). This effect is ascribed to the plasticizing effect of [Emim][PF₆], which, apart from introducing additional free volume within the blend, limits the intermolecular interactions between PVDF chains. As a result, lower energy (temperature) is required to reach a given relaxation. The plastizicing effect of the IL on the glass transition of PVDF supports the mixture of PVDF chain segments and IL molecules in the amorphous phase. Conversely, despite to the plasticizing effect of ILs, the other blends show increased transition temperatures compared to neat PVDF. This may be ascribed to the presence of a more packed chain structure, as the PVDF crystalline phase shifts from the α- to β-phase. Indeed, β-PVDF shows a more packed crystalline structure in regard to the α-PVDF (with a density of 1.99 and 1.92 g cm⁻³ for β- and α-phases, respectively) [50].

Since IL molecules cannot fit into the crystalline structure of the PVDF crystallites, when PVDF chains crystallize from the homogeneous liquid mixture with the low-molecular-weight IL, IL molecules are displaced to the amorphous interlamellae and interspherulite spaces. In fact, the DMA data in the region of the βDMA relaxation and the DSC data in the glass-transition region demonstrate that in some mixtures, an homogeneous blend with the polymer is formed, while in other mixtures, two separated amorphous phases coexist, as in the case of PVDF/[Emim][HSO₄].

Overall, those results agree with FTIR and DSC findings, where a concomitant plasticizing and directing (toward the β-phase) effect of ionic liquids has been observed.
4. Conclusions

Different ILs comprising different cations and anions were incorporated into the PVDF polymer matrix. Ten different ILs containing 1-ethyl-3-methylimidazolium [Emim] as a fixed cation (five of them) and ILs containing bis(trifluoromethylsulfonyl)imide [TFSI] as a fixed anion (another five) were incorporated into PVDF at a concentration of 25 wt % through solvent casting followed by solvent evaporation at 210 °C in an air oven. The effect of IL on the conformational, thermal, and dynamic mechanical behaviors of PVDF was evaluated. It is shown that while most of the blends having [Emim] as a common cation crystallize in the electroactive $\beta$-PVDF phase, blends having [TFSI] as a common anion crystallize in a mixture of $\alpha$, $\beta$, and $\gamma$ phases.

As a general trend, IL incorporation delays the PVDF crystallization and decreases both the glass-transition and melting temperatures of neat PVDF. Conversely, [Emim][HSO$_4$] and [P66614][TFSI] increase the crystallization and melting temperature of PVDF, suggesting a nucleating role of such ILs. Overall, a concomitant plasticizing and directing (toward the $\beta$-phase) effect of ionic liquids has been observed. Experimental findings here reported shed light on the relevance of properly selecting the optimum cation–anion balance to promote the formation of the electroactive and highly polar PVDF $\beta$-phase, which play a relevant role in applications such as sensors and actuators, biomedicine, and energy storage systems.

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Supporting Information

**Fig. S1.** FTIR spectra for the as-received PVDF powder and processed PVDF film after dissolving the powder in DMF and subsequent drying at 220 °C and cooling down to room temperature.