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Molecular mobility interpretation of the dielectric relaxor behavior in fluorinated copolymers and terpolymers

Jean-Fabien Capsal, Eric Dantras, Colette Lacabanne

Abstract

Thermo Stimulated Current and Dynamic Dielectric Spectroscopy have been applied to investigate dielectric relaxation modes of poly(vinylidene-fluoride-trifluoroethylene) copolymer and poly(vinylidene-fluoride-trifluoroethylene-chlorofluoroethylene) terpolymer. The aim of this work is to check the molecular origin of the ferroelectric relaxor behavior of the terpolymer. The combination of data obtained by both dielectric methods allows us to describe the molecular mobility of the amorphous phase and the cooperativity of the order/disorder dipolar transition in the crystalline phase. The introduction of 1,1-chlorofluoroethylene units in the main chain induces an increase of the Cooperative Rearranging Region size associated with less ordered and smaller crystallites. This morphological evolution is responsible of a lack of cooperativity and it explains the dielectric relaxor behavior of the poly(vinylidene-fluoride-trifluoroethylene-chlorofluoroethylene) terpolymer.

1. Introduction

Since the discovery of the piezoelectric behavior of poly(vinylidene-fluoride) – PVDF by Kawai [1] and the understanding of the molecular origin of this electroactivity [2–5], organic ferroelectric materials have attracted the interest of researchers for their potential use in specific applications such as low weight and flexible sensors [6,7], electro-mechanical devices [8,9] and high-K capacitors [10,11]. Many classes of polymers such as odd-polyamides [12], even odd polyamide copolymers [13], have shown piezoelectric and pyroelectric activities. Nevertheless, the poly(vinylidene-fluoride-trifluoroethylene) – P(VDF-TrFE) copolymer [14,15] and the poly(vinylidene-fluoride-trifluoroethylene-chlorofluoroethylene) – P(VDF-TrFE-CFE) terpolymer [16,17]. For electroactive applications, P(VDF-TrFE-CFE) copolymer is the most useful polymer because of the low poling field required to give ferroelectric behavior and high dielectric permittivity, compared with other ferroelectric polymers. Another important point is that this copolymer does not require any mechanical stretching before the poling process. Recently, it has been demonstrated that an electron irradiation of the P(VDF-TrFE) and the random incorporation of the CFE unit in P(VDF-TrFE-CFE) can achieve an electrostrictive strain greater than 5% [18–20] (higher than the P(VDF-TrFE) copolymer). This large electrostrictive behavior has been attributed to the relaxor behavior induced by defects in the crystalline phase caused by the irradiation or introduction of CFE units which is responsible of a high dielectric permittivity at room temperature [21,22]. The introduction of random defects broadens the ferroelectric transition and reduces the ferroelectric–paraelectric transition temperature. However, the molecular origin of the dielectric relaxor behavior of fluorinated terpolymer or electron irradiated copolymer remains obscure.

The main objective of this paper is to investigate the molecular mobility over a wide frequency range by means of thermal and dielectric analysis. Thermo stimulated current analysis of the terpolymer and the copolymer was used for the first time and allows us to give a molecular interpretation of the cooperativity and of the ferroelectric/paraelectric transition, called Curie transition, in fluorinated polymers.

2. Experimental section

2.1. Samples elaboration

Two kinds of fluorinated polymers were used in this study: the poly(vinylidene-fluoride-trifluoroethylene) (P(VDF-TrFE) 70–30 mol%) copolymer and the poly(vinylidene-fluoride-trifluoroethylene-chlorofluoroethylene) (P(VDF-TrFE-CFE) 55.8–35–9.2 mol%) terpolymer. Both of them have been purchased from Piezotech (France). P(VDF-TrFE-CFE) was synthesized by a suspension method with an oxygen-activated initiator [23].

For dielectric experiments, both as-received polymer powders were hot pressed 20 °C above their melting temperature. Films of 100 μm in thickness and 20 mm in diameter were obtained.

2.2. Standard differential scanning calorimetry

Standard differential scanning calorimetry (DSC) measurements were performed using a DSC/TMDSC 2920 setup. The sample temperature was calibrated using the onset of melting of tin (T_m = 231.88 °C),

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indium (T_m = 156.6 °C) and cyclohexane (T_m = 6 °C) with a heating rate of q_h = +5 °C min⁻¹. The heat-flow was calibrated with the heat of fusion of indium (ΔH = 28.45 J g⁻¹) and the baseline was corrected using sapphire. DSC experiments were systematically carried out over a temperature range from the equilibrium state (in order to remove the effect of previous thermal history) T_m = T_m + 20 °C down to the glassy state T_c = T_c - 70 °C with a constant cooling rate q_c = +10 °C min⁻¹, and followed by a linear heating rate q_h = 10 °C min⁻¹. The crystallinity ratios of the copolymer and the terpolymer have been determined. Thanks to the heat of fusion value of the 100% crystalline homopolymer.

2.3. Thermostimulated currents

Complex Thermo Stimulated Currents [24] (TSC) thermograms were carried out using a TSC/RMA Analyser. For complex experiments, the sample was polarized by an electrostatic field E = 3 kV mm⁻¹ over a temperature range from the polarization temperature (T_p = 40 °C for the terpolymer and T_p = 110 °C for the copolymer) down to the freezing temperature T_c (LNT temperature) with a constant cooling rate. Then, the field was turned off and the depolarization current was recorded with a controlled heating rate (q_h = +7 °C min⁻¹); the equivalent frequency of the TSC thermogram was f = 10⁻² - 10⁻³ Hz. Elementary TSC thermograms were obtained with a polarization window of 5 °C. The field was removed and the sample cooled to a temperature T_c = T_c - 30 °C. The depolarization current was recorded with a constant heating rate q_h. The series of elementary thermograms was generated by shifting the polarization window by 5 °C toward higher temperature. ΔH and ΔS uncertainties, extracted by this method, have been estimated near 10 and 20% respectively.

2.4. Dynamic dielectric spectroscopy

Dynamic Dielectric Spectroscopy (DDS) experiments were performed using a BDS400 set up covering a frequency range of 10⁻² - 3.10⁶ Hz, with 10 points per decade. Experiments were carried out in a temperature range from -100 °C to 130 °C. Dielectric isothermal spectras were measured every 2 °C. During each frequency scan, the temperature was kept constant to ±0.2 °C. The real ε′ and imaginary ε″ parts of the relative complex permittivity ε_r were measured as a function of frequency F at a given temperature T.

3. Results and discussion

3.1. Physical structure

Differential scanning calorimetry thermograms of the terpolymer and the copolymer are shown in Fig. 1. This experimental technique allows us to perform a quantitative and comparative study of the thermal transitions in semi-crystalline polymers. The heat capacity steps associated with the glass transition are weak. The glass transition temperature (T_g) and the freezing temperature (T_c) are reported as function of temperature for various frequencies ranging from 10 Hz to 1 kHz. Two relaxations are pointed out. The low temperature relaxation called α is localized near T_g = -15 °C at 10 Hz. This relaxation has a Vogel–Tammann–F Fulcher (VTF) behavior which has been associated with the dielectric manifestation of the glass transition. It is in agreement with previously published results [25]. The high temperature relaxation located near T_c = 15 °C at 10 Hz has been attributed to the dielectric manifestation of the Curie transition according to DSC results. The thermal evolution of the dielectric manifestation of the Curie process is more complex than the one of the α relaxation. For frequencies ranging from 10 Hz to 15 kHz, this relaxation is quasi-isothermal process; it is consistent with a first order Curie transition. At f_m = 15 kHz, a merging between the α and Curie relaxations takes place. Above f_m, this merging gives rise to a unique relaxation mode with a thermal evolution similar to the α mode. The relaxor behavior of the terpolymer seems to be governed by the α process.

The real and imaginary parts of ε_r of P(VDF-TrFE-CFE) are shown on Fig. 3-a/b. In both cases, two relaxations are found similar to the terpolymer. The VTF α process is located near T_g = -25 °C at f = 10 Hz and the Curie peak is located at 110 °C according to DSC measurements: P(VDF-TrFE) is a ferroelectric polymer with an isothermal first order Curie transition.

The dielectric strength Δε and the dielectric energy losses of the α relaxation in the terpolymer are higher than those of the copolymer. The high values of ε′ and ε″ of the terpolymer above f_m are attributed to the superposition of the α and Curie processes.

3.2. Dynamic dielectric relaxations

In order to understand the dielectric relaxor behavior of P(VDF-TrFE-CFE), dynamic dielectric spectroscopy and thermally stimulated currents have been used to characterize the molecular mobility over a wide frequency range. In Fig. 2-a/b the evolution of the imaginary (ε″) and real (ε′) parts of the dielectric permittivity of P(VDF-TrFE-CFE) are reported as function of temperature for various frequencies ranging from 10 Hz to 1 kHz. Two relaxations are pointed out. The low temperature relaxation called α is localized near T_g = -15 °C at 10 Hz. This relaxation has a Vogel–Tammann–Fulcher (VTF) behavior which has been associated with the dielectric manifestation of the glass transition. It is in agreement with previously published results [25]. The high temperature relaxation located near T_c = 15 °C at 10 Hz has been attributed to the dielectric manifestation of the Curie transition according to DSC results. The thermal evolution of the dielectric manifestation of the Curie process is more complex than the one of the α relaxation. For frequencies ranging from 10 Hz to 15 kHz, this relaxation is quasi-isothermal process; it is consistent with a first order Curie transition. At f_m = 15 kHz, a merging between the α and Curie relaxations takes place. Above f_m, this merging gives rise to a unique relaxation mode with a thermal evolution similar to the α mode. The relaxor behavior of the terpolymer seems to be governed by the α process.

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3.3. Fine structure of dielectric relaxations

The complex thermostimulated current (TSC) and the elementary TSC thermograms of P(VDF-TrFE-CFE) and P(VDF-TrFE) are reported in Fig. 4-a and b/c respectively. For the terpolymer, two relaxation modes are shown. The $\alpha$ and Curie modes are located at $T_{\alpha} = -15^\circ C$ and $T_{c} = 25^\circ C$ respectively; these values are consistent with low frequencies data from DSC and DDS. The $\alpha$ mode is attributed to the dielectric manifestation of the glass transition. For the P(VDF-TrFE), $\alpha$ and Curie relaxations are centered around $T_{\alpha} = -24^\circ C$ and $T_{c} = 103^\circ C$ respectively. The $T_{\alpha}$ value is slightly different from a previous work [29] determined near $-20^\circ C$. This difference has been associated with the influence of the Curie mode at higher temperature. We note that the Curie mode of the terpolymer is broader than the one of the copolymer. We assume that each process can be analyzed as a distribution of relaxation times; we extract from the elementary processes the activation enthalpies and the whole polarization of the dipolar relaxation [30–33]. Fig. 5 reports the activation enthalpies $\Delta H$ versus temperature.

The evolution of the $\alpha$ process of the copolymer and the terpolymer is very similar. In both cases, the activation enthalpy values depart from the null activation entropy line [34]. The delocalized mobility behavior of the dipolar entities involved in this process is characteristic of the relaxation associated with the glass transition. The maximal activation enthalpy of the terpolymer $\alpha$ mode is higher than for the copolymer. As previously shown by DSC, the CFE units strongly affect the crystalline regions. As a consequence, the Curie transition activation enthalpies of the terpolymer and copolymer are very different. For the terpolymer, the activation enthalpies are close to those of the $\alpha$ relaxation and are not influenced by the peak temperature. The delocalization of the Curie process is of the same order than for the $\alpha$ process. In other words, CFE units give rise to less ordered crystalline regions. For the copolymer, the evolution of the activation enthalpies associated with the Curie transition clearly indicates a more delocalized mobility due to long range cooperativity.

The polarization $P_0$ of each elementary processes, proportional to the number of dipoles, is shown in Fig. 6-a and b for the terpolymer and the copolymer respectively. For both polymers, two peaks are distinguishable. They are associated with the maximum of $\alpha$ and Curie relaxations in TSC. $P_0(\alpha, \text{copo})$, i.e. $P_0$ for the copolymer $\alpha$ mode, is 6 times lower than for the terpolymer, while $P_0(\text{Curie, copo})$ is nearly 30 times higher. This opposite behavior of $P_0(\text{Curie})$ and $P_0(\alpha)$ is attributed to the crystallinity ratio decrease due to CFE units. The activation entropy versus activation enthalpy for the $\alpha$ and Curie relaxations of these polymers are reported in Fig. 7.

3.4. $\alpha$ relaxation mode

For both polymers, the $\alpha$ relaxation is characterized by a compensation law between activation entropy and activation enthalpy. This behavior is associated with the $\alpha$ relaxation cooperativity. The activation enthalpy values for the $\alpha$ relaxation range from 100 kJ mol$^{-1}$ to 280 kJ mol$^{-1}$. These values are in good agreement with activation parameters of a dipolar relaxation associated with the glass transition [35–37]. The slopes of the terpolymer and copolymer plots are similar, indicating that the CFE units weakly influence the molecular dynamics in the amorphous phase. The activation enthalpy range for...
the copolymer is 20% lower than for the terpolymer due to the lower crystallinity ratio of the terpolymer. It leads a less constrained amorphous phase with higher cooperative rearranging regions (CRR) size in the model introduced by Adam and Gibbs [38] and adapted by Donth [39,40]. The decrease of CRR [41–43] size with the decrease of the amorphous phase content report in this study is analogous with previously published data on the evolution of the CRR in ceramic/polymer nanocomposites [44,45]: the densification of the composite with the increase of the inorganic phase volume fraction tends to reduce the activation enthalpy of the α process and consequently the size of the CRR.

3.5. Curie relaxation mode

The relaxation associated with the Curie transition has been assigned to dipolar reorientations in the crystalline phase of these semi-crystalline polymers. It has been shown that the CFE units strongly influence the crystalline phase by acting as defects. The compensation diagram of the copolymer and terpolymer is reported in Fig. 7. Activation enthalpies of the Curie relaxation are higher than for the α mode which is consistent...
3.6. Ferroelectric behavior

Fig. 7. Activation entropies \( \Delta S \) versus enthalpies \( \Delta H \) extracted from elementary TSC thermograms for the P(VDF-TrFE-CFE) terpolymer and the P(VDF-TrFE) copolymer. Lines are given as guides to the eye.

with the fact that the Curie mode is attributed to an order–disorder transition in a crystalline environment with higher potential barriers.

The decrease of the activation entropy for the Curie mode regarding the \( \alpha \) one, reflected by the relative position of the two compensation lines, is consistent with the decrease of the number of accessible sites in the crystalline phase, regarding the amorphous phase. The narrow temperature range in which this relaxation takes place (\( \Delta T = 4 \) °C) reflects the homogeneity of the crystalline phase.

The Curie relaxation of the terpolymer is quite different. No compensation law is found and the activation enthalpies and entropies are close to each other in the temperature range in which this relaxation takes place (\( \Delta T = 4 \) °C) reflecting the homogeneity of the crystalline phase.

The Curie relaxation loses its cooperative behavior. Since the range of cooperativity is linked to the range of order, this is consistent with the small crystallite size near \( \sim 10 \) nm — induced by defects reported by Ang et al. [46] for the terpolymer in comparison with the submicron crystallite size (\( \leq 0.5 \) \( \mu \)m) of the copolymer. The temperature range of the Curie relaxation is 15 °C. The CFE units tend to create defects in the crystalline phase creating a less ordered crystalline structure with heterogeneous morphologies.

The typical size of the terpolymer crystallites close to the CRR size of the amorphous phase and the non-cooperative behavior of the Curie transition tend to indicate that the molecular dynamics at the glass transition highly affect the dynamic response of Curie mode. At low frequency, the \( \alpha \) relaxation is far from the isothermal Curie relaxation. As the frequency increases, the temperature location of the \( \alpha \) relaxation gets closer to the Curie relaxation until a merging takes place. At \( f_m \), the dielectric relaxor dynamic is activated by the relaxation associated with the vitreous transition.

4. Conclusion

The understanding of the molecular origin of the dielectric relaxor behavior of fluorinated terpolymers is crucial because of their potential applications in electro-mechanical transduction systems. The molecular dynamics, the cooperativity of the dipolar entities involved in the Curie transition and the thermal properties have been characterized by DSC and dielectric experiments. A comparative study of a fluorinated terpolymer and a copolymer allows us to quantify the role of the CFE units on the relaxor like behavior of the P(VDF-TrFE-CFE) terpolymer. CFE units tend to reduce the size of crystallites, cohesive energy, and the rate of crystallinity. The molecular dynamics of the relaxation associated with the glass transition are quite similar in the terpolymer and the copolymer. Moreover, the decrease of the crystallinity with CFE units results in larger CRR size as defined from the relaxation associated with \( T_g \).

The most important point is the loss of the cooperative behavior of the Curie transition in the ferroelectric relaxor terpolymer. The activation enthalpies of the Curie relaxation of the terpolymer remain close to those involved in the relaxation associated with \( T_g \). This result is consistent with the fact that the crystallite size is of the same order than the CRR size which is not sufficient for cooperativity. Above the merging frequency \( f_m \), the \( \alpha \) process associated with \( T_g \) governs the dielectric behavior of the relaxor.

References


Fig. 8. Dielectric displacement versus applied electric field of the P(VDF-TrFE) copolymer (black line) and the P(VDF-TrFE-CFE) terpolymer (gray line) measured at room temperature and under a sinusoidal signal with a period of 18 s.

The \( D(E) \) curve of the P(VDF-TrFE-CFE) terpolymer is a mix between a High-K \( D(E) \) curve of a normal dielectric material (large maximum polarization of 35 mC m\(^{-2}\)) and a poorly ferroelectric hysteresis cycle characterized by a low remnant polarization of 4 mC m\(^{-2}\). As the electric field is increased, dipoles in the crystalline phase are oriented. Most of the electrical energy injected in the terpolymer is recovered when the electric field is turned off. The cooperativity of the dipolar mobility requires long range order. Consequently, the lack of cooperativity observed in the terpolymer may be assigned to the reduction of the crystalline region inducing a loss of ferroelectric behavior.


