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Tanguy CHOUPIN, Bruno FAYOLLE, Gilles REGNIER, C. PARIS, Jacques CINQUIN, Benoît BRULÉ - Isothermal crystallization kinetic modeling of poly(etherketoneketone) (PEKK) copolymer - Polymer - Vol. 111, p.73–82 - 2017

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Isothermal crystallization kinetic modeling of poly(etherketoneketone) (PEKK) copolymer


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Keywords:
Thermoplastics
Poly(etherketoneketone)
Thermal analysis
Crystallization kinetic
Crystallization induction time
X-ray diffraction

Abstract
Isothermal melt and cold crystallizations of a poly(etherketoneketone) (PEKK) copolymer prepared from diphenyl ether (DPE), terephthalic acid (T) and isophthalic acid (I) with a T/I ratio of 60/40 have been investigated by differential scanning calorimetry, wide-angle X-ray scattering and polarized optical microscopy. For the first time, the two-stage overall crystallization kinetics of PEKK taking into account effects of nucleation and crystal growth has been identified by using a modified Hillier type model. The primary crystallization stage is found to be an instantaneous two dimensional nucleation growth with an Avrami exponent of 2 whereas the secondary stage is found to be an instantaneous one dimensional nucleation growth with an Avrami exponent of 1. The evolution of the crystallization kinetic constants for two-stage crystallizations has been modeled according to the Hoffman and Lauritzen growth theory. Due to low crystallization kinetics, a crystallization induction time has been added to obtain a good fit with experimental data. Based on this modeling, Time-Temperature-Transformation (TTT) diagrams of the relative volume crystallinity have been established for the overall crystallization mechanism and also for the separated primary and secondary crystallization mechanisms providing an original crystallization mapping of the material.

1. Introduction

High performance thermoplastic poly(aryletherketone) (PAEK) matrices are increasingly used in aeronautics for composite structural parts. They have high temperature performances, good impact properties, high chemical and oxidation resistance and as all thermoplastics, they are recyclable and weldable. They are especially suitable for automated composite fiber tape placement and for out of autoclave consolidation. Those two features would help increase the aircraft production rate which is currently a major issue for aircraft manufacturers.

PAEks differ from each other in the ratio of ether/ketone. PEEK has been the most studied since the 1990s [1–8] because of its high temperature performances, and its chemical and oxidation resistance. However, it has a high melting temperature which makes its processing difficult. PEKK copolymers have been studied by few authors in the 1990s [9–14] and they have not aroused great interest until recently [15–18]. They are prepared form diphenyl ether (DPE), terephthalic acid (T) with para phenyl links and isophthalic acid (I) with meta phenyl links and differ from each other with their T/I ratios. Their melting temperatures (Tm) decrease with the T/I ratio without changing significantly their glass transition temperature (Tg) [9]. This is due to a lowering of the macromolecular chains regularity with the T/I ratio which decreases the crystalline phase compactness [9]. PEKK 60/40 has a Tg about 156 °C and a Tm about 300 °C [18], which is lower than the one for PEEK (around 340 °C [2]). This is quite interesting for processing regarding part manufacturing costs and possible degradation at high temperature. At last, PEKK 60/40 crystallizes less and slower than PEEK because ketone linkages are stiffer than ether linkages. The maximum crystallinity varies between 22% and 27% and it does not crystallize for cooling rates above 10 °C/min [18].

Isothermal crystallization mechanisms of PEKK 60/40 have been studied by Gardner et al. [9] and Cortés et al. [18]. After annealing, they observed an additional endotherm during the melting which occurs about 10 °C above the annealing temperature. Such behavior has also been observed for PEEK [2,4,19] and other semi-crystalline polymers as poly(ethyleneterephthalate) [20–22] and is called the
double-melting behavior. This additional endotherm is attributed to the melting of a secondary crystallization associated to the growth of an interlamellar crystalline structure within lamellae [23]. This subsidiary crystallization appears after long annealing or slow cooling rates. Gardner et al. [9] also studied the PEKK 60/40 isothermal crystallization kinetics with a peak-time method (time for which the crystallization exothermic peak is maximum) [9] but did not use a crystallization kinetic model taking into account the two-stage crystallization.

Different studies have been made to incorporate the two-stage crystallization into the crystallization kinetic modeling of polymers, the two most relevant reported are the Velisaris Seferis [5] and the Hillier models [23]. The former postulates that a primary Avrami type crystallization processes [24] occur in parallel independently whereas the second assumes that a primary Avrami type crystallization process occurs followed by a first order crystallization process which grows from the primary crystallization. Hsiao et al. [12] investigated the isothermal crystallization kinetics of a PEKK 70/30 with a modified Hillier model which postulates that the secondary crystallization is not necessarily a first order process and compared it to the Velisaris Seferis model. They concluded that the modified Hillier model has a better physical meaning than the Velisaris Seferis one because it is more suitable to describe the growth of a second crystalline structure within lamellae. In this model, the primary crystallization is expressed by an Avrami equation:

\[ \alpha_1(t) = 1 - \exp(-K_1t^{n_1}) \]  

where \( \alpha_1(t) \) is the relative volume fraction crystallinity at time \( t \), \( w_1 \) is the weight factor between primary and secondary crystallizations, \( K_1 \) is the crystallization rate constant both depending on the annealing temperature and \( n_1 \) the Avrami exponent depending on the nucleation and the growth geometry, for the first mechanism. The secondary crystallization process which occurs once a volume element has been included in the lamellae at time \( 0 \) is expressed by an Avrami equation:

\[ \alpha_2(t) = \int_0^t \alpha_1(\theta) \times \frac{d}{d\theta} (\alpha_2(t-\theta)) d\theta \]  

where \( \alpha_2(t-\theta) \) is the relative volume fraction crystallinity at time \( t \), \( w_2 \) is the weight factor, \( K_2 \) is the crystallization rate constant both depending on the annealing temperature and \( n_2 \) the Avrami exponent depending on the nucleation and the growth geometry, for the second mechanism. The Avrami exponent \( n_2 \) is not necessarily equal to 1 according to Hsiao et al. [12]. The total crystallinity at time \( t \) arising from those two consecutive crystallization processes becomes:

\[ \alpha(t) = w_1\alpha_1(t) + w_2\alpha_2(t) \]  

From Equations (1)–(3):

\[ \alpha(t) = w_1[1 - \exp(-K_1t^{n_1})] + w_2K_2\int_0^t [1 - \exp(-K_1\theta^{n_1})] \times (t - \theta)^{n_2-1} \exp(-K_2(t - \theta)^{n_2}) d\theta \]  

where \( w_1 + w_2 = 1 \), indicating that the relative crystallinity at infinite time is unity which corresponds to the Avrami model hypotheses [24]. Values of weight factors provide the relative importance between both crystallizations. It can be noticed that Equation (4) differs from Hsiao et al. [12] model which could be due to a misprint in their article for the derivative of the secondary crystallization in Equation (2).

This paper aimed to investigate the isothermal crystallization kinetics of PEKK 60/40 with a modified Hillier model for crystallization from the melt and cold crystallization to predict precisely the crystallinity evolution according to processing cycles. Basing on experimental data, parameters associated to the modified Hillier model will be assessed carefully and their physical meanings will be discussed. Finally, Time-Temperature-Transformation (TTT) diagrams of the relative volume crystallinity will be built from the overall crystallization modeling.

2. Experimental

2.1. Material and preparation

PEKK used in this study was provided by Arkema referenced as PEKK 6002. It was prepared from diphenyl ether (DPE), terephthalic acid (T) with para phenyl links and isophthalic acid (I) with meta phenyl links with a T/I ratio of 60/40 (Fig. 1). PEKK 60/40 has a weight-average molecular weight (Mw) of about 60000 g mol\(^{-1}\). PEKK samples were supplied amorphous with no crystalline traces measured by DSC and WAXS. The glass transition and melting temperatures were respectively measured about 155°C and 292°C by DSC with a heating stage at 10°C/min.

2.2. Differential scanning calorimetry

Isothermal crystallization analyses were carried out in a TA Instruments Q2000 on granules of about 7–8 mg. For cold and melt crystallization, all specimens were first heated at 10°C min\(^{-1}\) from room temperature to 360°C during 5 min. This temperature is above the PEKK 60/40 equilibrium melting temperature (Tm = 340°C [9]) to erase the thermal history and obtain a fully amorphous polymer. For crystallization from the melt, specimens were then cooled at 40°C min\(^{-1}\) to annealing temperatures during 120 min and finally cooled to room temperature at 40°C min\(^{-1}\). During the cooling steps, the polymer does not have enough time to crystallize and the annealing time is long enough to allow the polymer to fully crystallize. For cold crystallization, the polymer is cooled at 40°C min\(^{-1}\) to room temperature and then heated at 80°C min\(^{-1}\) to the isothermal temperature during 120 min and finally cooled at 40°C min\(^{-1}\) to room temperature. For all specimens after crystallization, a heat scan at 10°C min\(^{-1}\) from 360°C was carried out to measure the glass transition, the melting temperature and the melting enthalpy induced by the crystallization cycle.

2.3. Wide-angle X-ray diffraction

The X-ray data were collected by performing wide angle X-ray scattering (WAXS) patterns at room temperature on samples with a thickness of 2 mm thanks to the SAXS/WAXS system from Xenocs. The apparatus consists of an X-ray source using a copper anode K\(_\alpha\) radiation with a wavelength of 1.54 Å delivering a parallel beam. The X-ray patterns were recorded with a MAR300 2D detector.

![Chemical structure of PEKK copolymers.](image-url)
piloted with the MAR300 software. The sample to detector distance was 168 mm. Silver behenate standard reference material was used for sample-to-detector distance calibration. The exposure time was 15 h. The 2D patterns were treated using Fit2D software developed by the European Synchrotron (ESRF). The 2D X-ray diffraction patterns were integrated along the azimuthal direction from isotropic samples.

2.4. Optical microscopy

The optical microscope used was a polarizing Nikon Eclipse LV100 model equipped with a Nikon DS Digital Sight DS-FI1 camera and a hot stage Linkam LTS-420. Observations were carried out with an ocular lens of 30× on PEKK film with a thickness of 5 μm prepared with a Leica RM-2235 microtome. Films were first heated at 10 °C min⁻¹ from room temperature to 360 °C during 5 min to erase the thermal history of the specimen and then cooled down at 40 °C min⁻¹ to 270 °C during 2 h. This annealing temperature corresponds to a low supercooling i.e. to a low nucleation rate allowing the formation of large spherulites.

3. Results and discussion

3.1. Crystalline morphology

WAXS measurements have been carried out on a PEKK 60/40 sample crystallized from the melt and from the glassy state at 230 °C during 2 h (Fig. 2). It has been observed for crystallization from the molten state a diffraction pattern with the (110) peak at 2θ = 23.0° which has already been reported by DuPont and ICI research group [10,11,26–28] and Blundell et al. [13] for PEKK. It has been identified by a second morphology (form II) with a one or two-chain orthorhombic lattice depending on the authors. This subsidiary morphology would appear for high chain stiffness and low molecular mobility. PEKK copolymers have a high ratio of ketone linkages which increases the stiffness of the molecule and allow the apparition of the form II. In fact, ketone linkages have high rotational conformation energy barriers which enhance chain stiffness.

For PEKK 60/40 crystallized from the glassy state, a more important (020) peak has been observed compared to crystallization from the melting state and the (111) peak is non-existent. This is consistent with the fact that cold crystallization induces low chain mobility enhancing the growth of the form II. The disappearing of the (111) peak highlights that PEKK 60/40 crystallized from the glassy state presents only pure form II.

3.2. Nucleation and growth

Fig. 3 shows the crystal growth of a PEKK 60/40 sample crystallized from the melt at 270 °C during 20 min (Fig. 3a), 30 min (Fig. 3b) and 40 min (Fig. 3c). Radial growths of crystalline entities similar to spherulites were observed. During the first step of crystallization, the crystalline entities grow independently to each other with the same size until impingement which is typical to an instantaneous nucleation. The initial number of potential nuclei N₀ has been calculated about 0.0069 μm⁻². It is equal to the number of nuclei at the early beginning of the crystallization divided by the area containing the nuclei which have been both measured with an image processing program. At the end of the crystallization process, the crystalline entities diameters have been measured from 18 μm to 33 μm.

The evolution of the radius of a spherulite during the crystallization process as a function of time at 270 °C from the melting state has been followed in Fig. 4. In order to assess the crystalline growth rate, the spherulite has been chosen to be the most far as possible from the others to collect enough data before impingement. It can be observed three different steps during the crystallization process. The first step is related to the crystallization initiation where the spherulite studied was first observed after 8 min of crystallization with a radius of 2 μm. This time depends on the device resolution. After this initiation step, the spherulite grows constantly until it impinges on another spherulite. The growth rate is independent of the size of the spherulite and calculated from the slope of the line by G = dr/dt. The growth rate of spherulites at 270 °C from the melt has been measured about 0.2 μm min⁻¹. At the end of the crystallization process, the crystal radius has been observed about 16.5 μm. Those results will be compared later with the following DSC study.

3.3. Thermal analysis methods

Isothermal DSC scans of PEKK 60/40 samples at 210 °C and 230 °C cooled at 40 °C min⁻¹ from the melt are reported in Fig. 5a. It can be witnessed that the beginning of the crystallization exothermic peaks are truncated corresponding to a gap between the beginning of the crystallization peak and the baseline (double arrow in Fig. 5a). This phenomenon is due to the instrumental thermal inertia switching from the cooling to the isothermal step. It can be observed that this gap is more important for isothermal at 230 °C than 210 °C due to higher crystallization kinetics at 230 °C. Therefore, the beginning of the crystallization peak has been extrapolated until the base line with a linear curve (dashed line in Fig. 5b) to take into account the overall crystallization process.

As the crystallization kinetics of PEKK 60/40 is relatively low compared to most of semi-crystalline polymers, polyolefins for instance [29], it can be detected a delay between the beginning of

![Fig. 2. X-ray diffraction scans of PEKK 60/40 crystallized at 230 °C during 2 h from the melting state (a) and from the glassy state (b).](image-url)
the isotherm and the crystallization process. This delay has also been observed for other polymers with low crystallization kinetics and modeled as a crystallization induction time \( t_i \) [21,30]. It corresponds to the time to reach a steady state of nucleation i.e. the time to reach enough critical nuclei in the bulk to initiate crystallization. Indeed, the increase in number of critical nuclei which leads into the steady state linear is initially slow [31]. In this work, the induction time \( t_i \) has been defined by the intersection of the baseline and the extrapolation line of the crystallization peak beginning (Fig. 5b). Without taking into account this crystallization induction time, the multilinear regression on Hillier’s model coefficients was not able to give a very good fit of experimental data.

3.4. Isothermal crystallization kinetic modeling

For each isothermal crystallization, the relative volume fraction of crystallinity \( \alpha(t) \) has been calculated by dividing the partial area at time \( t \) by the overall area of the extrapolated crystallization exothermic peak (Fig. 6) and its double logarithm has been plotted in Fig. 7. The induction time as defined previously has been taken into account for the plots of \( \alpha \). Straight line have been observed for the double logarithm plot in Fig. 7 during the beginning of the crystallization which is consistent with the Avrami theory. For every crystallization, the slopes of the lines corresponding to the Avrami exponent have been measured about \( 2.05 \pm 0.01 \) in Fig. 7 (dashed lines). The corresponding Avrami model (Equation (1)) has been plotted in Fig. 6 (dashed lines) with an Avrami exponent \( n \) fixed by 2. It can be observed that the Avrami model fits well with experiments for relative crystallinity below 0.6. According to the Avrami theory, a value of 2 for the Avrami exponent would be associated either to a two dimensional crystallization growth with an instantaneous nucleation or to a one dimensional crystallization growth with a sporadic nucleation. Since the microscopic study has shown previously that the nucleation process is instantaneous (Fig. 3), it can be assumed that the first crystallization mechanism would be a two dimensional disk-like crystallization. However, this assumption does not seem consistent with the microscopic

![Figure 3](image-url) Micrographs of PEKK 60/40 crystallized from the melt at 270 °C during 20min (a), 30min (b) and 40min (c).

![Figure 4](image-url) Spherulite radius vs time during crystallization at 270 °C from the melting state.
observations where growing crystalline entities behave like spherulites. This phenomenon has already been observed for Poly(ethylene Oxide) by Banks and Sharplees [32] and they concluded that the Avrami exponent is not necessarily satisfactory to provide information on crystallization mechanisms [33].

For values of relative crystallinity above 0.6, crystallization kinetics become slower than the Avrami model which corresponds to a bending of the double logarithm plots in Fig. 7. This phenomenon is typical of the formation of a secondary crystallization process within the primary crystallization with a slower crystallization kinetic [5,12]. The modified Hillier model previously presented (Equation (4)) has been used to take into account this secondary crystallization and fit with experiments during the overall crystallization process. The best fit of the Hillier model determined with a multi linear regression fitting has been found for a secondary crystallization Avrami exponent $n_2 = 1$. It is associated to a one dimensional crystallization growth with an instantaneous nucleation. This result was also reported for polymethylenes by Hillier [23] attributed to a first-order crystallization law within the crystalline lamellae. The same approach has been done for cold crystallization and same Avrami exponents have been found. Values of $n_1$ and $n_2$ are lower than those reported for PEEK [4] and PEKK 70/30 [12] which can be explained by a different nucleation and growth process. Results of the fitting of the modified Hillier model with $n_1 = 2$ and $n_2 = 1$ for $K_1$, $K_2$ and $w_1$ are discussed below.

Following the general approach reported in literature for nucleation and crystallization growth, crystallization rate constants ($K_1$ and $K_2$) can be expressed as a function of the initial number of potential nuclei ($N_0$) and the crystal growth rate ($G$) depending on the dimensionality of crystallization and the nucleation mode. For the primary crystallization which has been shown to have a two dimensional crystallization growth with an instantaneous nucleation, the crystallization rate constant can be written:
\[ K_1 = \pi N_{01} G_1^2 \]  
(5)

with \( N_{01} \) the initial number of potential nuclei and \( G_1 \) the crystallization growth rate for the primary crystallization.

For the secondary crystallization which has been shown to have a one-dimensional crystallization growth with an instantaneous nucleation, the crystallization rate constant can be written:

\[ K_2 = N_{02} G_2 \]  
(6)

with \( N_{02} \) the initial number of potential nuclei and \( G_2 \) the crystallization growth rate for the secondary crystallization.

According to the Hoffmann and Lauritzen theory [34–36], the crystal growth can be expressed by:

\[ G_i(T) = G_0i \exp \left( -\frac{U^*}{R(T - T_{\infty})} \right) \exp \left( -\frac{K_{gi}}{T \Delta T} \right) \]  
(7)

where \( G_0i \) is a pre-exponential factor independent of the temperature. The first exponential term contains the contribution of the diffusion process to the growth rate, where \( U^* \) is the activation energy of the molecular transferring through the melt crystal interface \( (U^* = 6300 \text{ J mol}^{-1}) [35]) \), \( T_{\infty} \) is the temperature below which diffusion stops \( (T_{\infty} = T_g - 30 \text{ K}) \) and \( R \) is the gas constant. The second exponential term is the contribution of the nucleation process, where \( K_{gi} \) is the activation energy of nucleation for a crystal with a critical size, \( \Delta T \) is the degree of supercooling \( (\Delta T = T_m - T) \) with \( T_m = 340 \text{ K} \) and \( f \) is a correction coefficient for the temperature dependence of the melting enthalpy \( (f = 2T/(T_m0 - T)) \). The index \( i \) for \( G_0 \), \( G_0i \) and \( K_{gi} \) is equal to 1 for primary crystallization and 2 for secondary crystallization.

Finally, the primary crystallization rate constant can be written:

\[ K_1(T) = K_{01} \exp \left( -\frac{2U^*}{R(T - T_{\infty})} \right) \exp \left( -\frac{2K_{g1}}{T \Delta T} \right) \]  
(8)

And the secondary crystallization rate constant can be written:

\[ K_2(T) = K_{02} \exp \left( -\frac{U^*}{R(T - T_{\infty})} \right) \exp \left( -\frac{K_{g2}}{T \Delta T} \right) \]  
(9)

with the pre-exponential factors independent of temperature \( K_{01} = \pi N_{01} G_{01}^2 \) and \( K_{02} = N_{02} G_{02} \).

\( K_1 \) and \( K_2 \) have been plotted respectively in Fig. 8a and Fig. 8b and fitted with respectively Equations (8) and (9) for crystallization from the melt and cold crystallization. As expected, we observed that both plots have typical bell shapes regarding the Hoffman and Lauritzen growth theory [35]. Crystallization kinetics are the most important around 230 °C and very low close to the \( T_g \) and \( T_m \) corresponding to low diffusion and nucleation kinetics respectively. Interestingly, it appears that primary crystallization rate constants for cold crystallization are higher than those for crystallization from the melt. However, it is the opposite for secondary crystallization rate constants. This has been attributed to a lower final degree of crystallinity (i.e. \( \Delta H_c \)) for cold crystallization than crystallization from the melt (Table 1). Values of \( K_{gi} \) are similar to those reported in the literature for other PAEKs [37,38]. It is noteworthy that crystallization rate constants \( (K_1 \) and \( K_2 \) for isothermal crystallization at 200 °C, 260 °C and 270 °C (unfilled marks in Fig. 8) have not been taken into account for the fitting because of a lower final crystallinity compared for the other temperatures (Table 1) involving higher rate constants. Indeed, for those temperatures, crystallization kinetics are very low which makes difficult the integration of the end of the DSC crystallization peak which truncates the overall area of the crystallization peak and underestimates the crystallization enthalpy.

Since, the crystallization rate constant \( K_1 \) is linked to \( G_1 \) with Equation (5), we propose to crosscheck \( K_1 \) value with the previous microscopic study assuming that the spherulite growth rate is equal to the crystalline lamellae growth rate. Thanks to this study, the growth rate \( G \) at 270 °C from the melt and the initial number of potential nuclei \( N_0 \) were measured about 0.2 \( \mu \text{m min}^{-1} \) and 0.0069 \( \mu \text{m}^2 \) respectively. By assuming that the growth rate is mainly associated to the primary crystallization process, the crystallization rate constant \( \ln(K_1) \) were calculated about –7.04 min^{-1}.  

![Fig. 8. Logarithmic plot of the primary (a) and the secondary crystallization (b) rate constants vs temperature for isothermal crystallization from the melt (■) and cold crystallization (▲) with the Hoffman and Lauritzen model fitting with a solid line for crystallization from the melt and dashed line for cold crystallization. Unfilled markers have not been taken into account for the fitting.](image-url)
with Equation (5). This value is consistent with the \( \ln(K_1) \) value calculated with the Hillier model at 270 °C from the melt about \(-6.8\) min\(^{-1}\) (Fig. 8). It is noteworthy that the \( \ln(K_1) \) value calculated with the microscope study is slightly lower than the one calculated with the Hillier model and fits better with the Hoffman and Lauritzen modeling (solid line in Fig. 8). In fact, the issue of the truncation of the end of the DSC crystallization peak related previously (Table 1) does not exist for the spherulites growth method. However, this method needs high temperature crystallizations allowing the growth of large spherulites to be able to measure accurately their size during crystallization which becomes difficult for temperatures below 260 °C.

The primary crystallization weight factor \( (w_1) \) for crystallization from the melt and cold crystallization has been plotted as a function of temperature in Fig. 9. It has been modeled with a linear equation:

\[
w_1(T) = a_1T + b_1
\]

where \( a_1 \) and \( b_1 \) are two constants independent of the temperature. We observed that \( w_1 \) is closed to 1 for low temperatures and then decreases to values lower than 0.5 for temperatures above 260 °C. This shows that the formation of the secondary crystallization is promoted for low degree of supercooling. This observation has also been reported by few authors [18,39] with the increase of a first endothermic peak which occurs 10 °C above the annealing temperature attributed to the melting of the secondary crystallization. In fact, for high annealing temperatures, the diffusion process is very important with a high mobility of the chains allowing the enhancement of the crystalline phase by the growth of a second crystalline structure within lamellae.

### 3.5. Induction time modeling

The crystallization induction time due to nucleation initiation has to be considered in terms of processing time and thus in crystallization models. Following the general approach reported in literature for nucleation and crystallization growth [34,35] and other works [21,30], the temperature dependence of the induction time \( (t_i) \) is the same as the one of lamellae growth and therefore follows the Hoffman and Lauritzen model:

\[
t_i(T) = K_{t0} \exp\left(\frac{E_{t1}}{R(T-T_g)}\right) \exp\left(\frac{E_{t2}}{R(T_{m0}-T)}\right)
\]

where \( E_{t1} \) and \( E_{t2} \) are the activation energies for nucleation respectively below the equilibrium melting temperature \( T_{m0} \) and above the glass transition temperature \( T_g \).

The induction time \( (t_i) \) measured as reported in Fig. 5 has been plotted in Fig. 10 and fitted with Equation (11) for crystallization from the melt and cold crystallization. Activation energy values \( E_{t1} \) and \( E_{t2} \) range between 2.5 and 4.8 kJ mol\(^{-1}\) are low and consistent with literature [21,30]. Contrary to later modeling, induction times measured at 200 °C, 260 °C and 270 °C have been taken into account to fit the model because they depend only on the beginning of the crystallization. The induction time at 270 °C is about 9 min.

### Table 1

Comparison of the isothermal crystallization enthalpy \( \Delta H_c \) for crystallization from the melt and cold crystallization.

<table>
<thead>
<tr>
<th>Temperature (°C)</th>
<th>( \Delta H_c ) melt (J g(^{-1}))</th>
<th>( \Delta H_c ) cold (J g(^{-1}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>200</td>
<td>18.4</td>
<td>17.1</td>
</tr>
<tr>
<td>210</td>
<td>21.5</td>
<td>20.2</td>
</tr>
<tr>
<td>220</td>
<td>23.3</td>
<td>19.6</td>
</tr>
<tr>
<td>230</td>
<td>23.2</td>
<td>21.1</td>
</tr>
<tr>
<td>240</td>
<td>23.9</td>
<td>21.0</td>
</tr>
<tr>
<td>250</td>
<td>20.0</td>
<td>20.1</td>
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<tr>
<td>260</td>
<td>17.1</td>
<td>16.0</td>
</tr>
<tr>
<td>270</td>
<td>15.8</td>
<td>15.7</td>
</tr>
</tbody>
</table>

Fig. 9. Weight factor of the primary crystallization vs temperature for isothermal crystallization from the melt (●) and cold crystallization (▲) with the Hoffman and Lauritzen model fitting with a solid line for crystallization from the melt and dashed line for cold crystallization. Unfilled markers have not been taken into account for the fitting.

Fig. 10. Induction time vs temperature for isothermal crystallization from the melt (●) and cold crystallization (▲) with the Hoffman and Lauritzen model fitting with a solid line for crystallization from the melt and dashed line for cold crystallization.
which is consistent with the crystallization initiation time measured with the microscopic study in Fig. 4. It has been observed that for temperatures closed to the $T_g$, induction times for crystallization from the melt and cold crystallization are similar whereas for temperatures near $T_m$, the induction time of cold crystallization is lower than crystallization from the melt. This has been explained by the formation of nuclei during the heating from the glassy state to the annealing temperature for cold crystallization. For annealing temperatures closed to $T_g$, the nucleation mechanism is very fast thus no difference has been observed for induction time for both crystallization.

3.6. Overall crystallization kinetic modeling

From modeling of kinetic constants ($K_1$ and $K_2$), weight factors ($w_1$ and $w_2$) and induction times ($t_i$) depending on temperature, the evolution of the relative crystallinity depending on time can be plotted with the modified Hillier model (Equation (4)) for the overall crystallization process as well as for the separated primary and secondary crystallizations. Fig. 11 shows the plot of the modified Hillier model compared to experiments for the overall crystallization, the primary and secondary crystallizations at 210 °C, 230 °C and 250 °C for crystallization from the melt. At the beginning of the crystallization, the curves of overall crystallization and primary crystallization are confounded which shows that the secondary crystallization is negligible during the early stage of crystallization. Then, the secondary crystallization occurs with a lower crystallization kinetics corresponding to the enhancement of the first crystallization with the growth of a second crystalline structure within lamellae. As it was observed for the crystallization weight factor, the proportion of the secondary crystallization process increases with the annealing temperature which shows that this mechanism is facilitated for high temperatures corresponding to high diffusion mechanisms.

As the relative crystallinity of PEKK 60/40 depending on time can be determined for any temperatures with the modified Hillier model, TTT diagrams of the relative volume fraction crystallinity for the overall crystallization (Fig. 12a) and also for the separated primary (Fig. 12b) and secondary crystallization (Fig. 12c) have been built allowing the determination of the crystallinity of the material for any isothermal crystallization process as well as corresponding crystallinity ratios of each crystallization mechanism. As expected, crystallization kinetics are the most important for temperatures around 230 °C and become very slow close to the glass transition and the melting temperature. For instance, Fig. 12a shows that around 230 °C, it takes approximately 25 min to reach 95% of the relative crystallinity whereas the last 5% takes more than 45 min to be done. This is consistent with the fact that the first crystallization mechanism is fast compared to the secondary crystallization. As it was shown previously, the secondary crystallization process is promoted for high annealing temperatures (Fig. 12c) which is the opposite for the primary crystallization (Fig. 12b).

4. Conclusion

 Isothermal crystallization kinetics of PEKK 60/40 for crystallization from the melt and cold crystallization were characterized by using a DSC. It was observed a crystallization induction time related to nucleation initiation delaying the crystallization process due to the low crystallization kinetics of PEKK 60/40. This induction time was modeled with a Hoffman and Lauritzen and taken into account in the crystallization kinetics modeling. It was observed lower induction times for cold crystallization at low degree of supercooling probably induced by the formation of nuclei during the heating from the glassy state to the annealing temperature. Isothermal crystallization kinetics of PEKK 60/40 were modeled with a modified Hillier model related to the growth of a first crystallization mechanism followed by a slower secondary process attributed to
the enhancement of the crystalline structure with the growth of a second crystalline structure within lamellae. From the fitting of the Hillier model with isothermal crystallization experiments, the Avrami exponent of the first crystallization mechanism was found to be equal to 2 related to a two dimensional growth with an instantaneous nucleation confirmed by microscopic observations whereas the Avrami exponent of the secondary crystallization was found to be equal to 1 related to a one dimensional growth with an instantaneous nucleation. Primary and secondary crystallizations rate of the modified Hillier model were modeled by the Hoffman and Lauritzen growth equation showing a very fast crystallization kinetics around 230 °C. Cold crystallization process seems to be faster than melt crystallization due to a lower final degree of crystallinity. According to the linear modeling of the primary crystallization weight factor, the secondary mechanism was found to be enhanced for low degree of supercoolings due to a high diffusion process. Thanks to a full modeling of crystallization kinetics, TTT diagrams of the relative volume fraction crystallinity of PEKK 60/40 for the overall crystallization and also for the corresponding primary and secondary crystallization mechanisms have been plotted providing a useful tool for PEKK 60/40 processing.

**Acknowledgments**

The authors wish to thank Arkema company for providing PEKK materials and Vincent Michel from PIMM laboratory for the WAXS measurements.

**References**


Fig. 12. TTT diagram of the relative volume fraction crystallinity for the overall crystallization (a), the primary crystallization (b) and the secondary crystallization (c) for crystallization from the melt.


