# Thermomechanical Processing Environment and Morphology Development of a Thermotropic Polymer Liquid Crystal

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**ABSTRACT:** We have studied a longitudinal polymer liquid crystal consisting of poly(ethylene terephthalate) (PET) and *p*-hydroxybenzoic acid, namely PET/0.6PHB, where 0.6 is the mole fraction of the second component. The material was injection molded with systematic variations of the melt and mold temperatures and injection flow rate using design of experiments based on a Taguchi orthogonal array. Thermomechanical environment defined by local melt temperatures and shear rates and stresses imposed during processing was estimated by computer simulations of the mold-filling phase. The morphology of the moldings was characterized by optical and scanning electronic microscopy, wide- and small-angle X-ray scattering, and differential scanning calorimetry. An analysis of

#### **INTRODUCTION**

Polymer liquid crystals (PLCs) are high performance polymers and exhibit an advantageous combination of properties, particularly in applications requiring high thermal and mechanical performance, high chemical resistance, and high dimensional stability. However, their high cost has limited the widespread use of these polymers, promoting their blending with thermoplastics engineering polymers.<sup>1–3</sup> variance approach identified the significant processing variables and their contributions to variations of morphological parameters. The processing environment affects strongly the melt viscosity, and there is a strong thermomechanical coupling. The result is a complex multilaminated and hierarchical microstructure, whose morphological features are very sensitive to the processing conditions. Relationships between local thermomechanical variables (rather than global ones) and the morphological parameters are established. © 2009 Wiley Periodicals, Inc. J Appl Polym Sci 115: 2991–3004, 2010

**Key words:** polymer liquid crystal; injection molding; processing; thermomechanical environment; morphology

Longitudinal (or main-chain) PLCs are copolymers comprised of rigid liquid crystalline (LC) sequences connected to flexible sequences by chemical bonds. The rigid mesogenic groups are exclusively in the main chain and are oriented parallel to it (so-called  $\alpha$ -type PLC).<sup>3-6</sup> This segmental nature allows easy orientation of the rigid segments under external applied fields, whether electric, magnetic, or me-chanical.<sup>7–13</sup> In fact, their thermotropic character allows them to be transformed by melt processing techniques, such as injection molding, because of their low viscosity resultant from the easy orientation of the rigid segments in the flowing melt. Because of high relaxation times of PLCs, the orientation induced by the flow cannot easily relax during subsequent cooling, leading to highly oriented structures (self-reinforced polymers) with exceptional mebehavior chanical and highly anisotropic character.14,15 Methods for successful prediction of long-term performance of PLCs from short-term tests have been developed.<sup>16,17</sup>

In a PLC, the rigid phase is inhomogeneously dispersed throughout the material forming a

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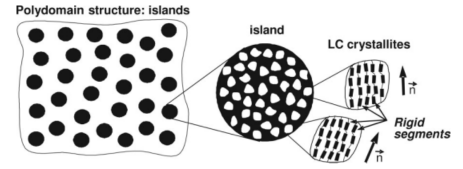


Figure 1 Polydomain hierarchical structure of a PLC (*n* is the orientation director).

hierarchical poly-domain structure<sup>1,18</sup>; Figure 1. The PLC tends to form spherical or ellipsoidal aggregates of LC-rich phase which are called islands.<sup>1,18,19</sup>

The islands constitute of smaller quasispherical shape LC crystallites connected by flexible chain segments. Wide-angle X-ray scattering (WAXS) results show that the LC crystallites (with dimensions of several nm) are mostly composed of highly oriented and packed rigid sequences of the macromolecular chains.<sup>18,19</sup> However, although each LC crystallite is highly oriented, the different crystallites are randomly oriented inside an island, so that no preferred orientation exists inside the islands.<sup>15</sup> The rigid phase is easily oriented during processing forming highly oriented structures. Shear and extensional stress fields have different effects on the morphology developed in PLCs. The former will result in a tumbling flow originating highly oriented sheet-like structures. An extensional flow leads to the development of fiber-like structures.<sup>20</sup>

Injection molding is an attractive technology for the production of large series of parts with a complex geometry. During processing, the polymer melt is subjected to a complex thermomechanical environment that constrains the respective morphological development. In the case of PLCs, with high shear and elongational fields and fast cooling rates, a complex highly heterogeneous laminated microstructure is formed, which is composed of several layers featuring distinct textures and states of molecular orientation.<sup>21-24</sup> Elongational viscosity of several PLC blends shows that the PLC constituent causes channelling of the flexible polymer<sup>25</sup> and thus its orientation along the PLC-this effect is predicted theoretically by an extension of the Flory statisticalmechanical theory of PLCs.<sup>26</sup>

At the micrometric level, a five-layer laminated microstructure is normally formed: two external skins, two intermediate zones, and a central core region. The skin and the intermediate zone can be further discriminated and more sublayers can be identified. The number of identifiable layers depends on the thickness of the molding and the criteria established by the observer. The skin layer exhibits a

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high level of molecular orientation. This feature, combined with the lack of chain entanglements, makes the skin layer easy to delaminate. In the core region, the melt experiences a tumbling flow of LC islands, resulting in molecular orientation along the flow lines. Studies of the morphology development of injection-molded PLCs has been based on the traditional approach of directly relating the global processing conditions (e.g., temperatures, velocities, molding thickness) to the morphological parameters.<sup>23,24,27,28</sup> This does not support the establishment of universally valid processing-morphology relationships, which are molding geometry and injection molded machine dependent. It does not support a predictive effort. Our approach-used in the present work—is different; we believe that these types of relationships must be established locally, relating the local thermomechanical environment to the developed morphology. The processing-morphology relationships are, therefore, of a more general character, independent of selected injection molding machine and molding geometry.

This work investigated the structure development of an injection-molded thermotropic PLC. The local thermomechanical environment is assessed by computer simulations of the mold-filling phase. The morphology developed by the injection-molded PLC specimens is characterized by several structure-sensitive experimental techniques including the following: scanning electronic microscopy, differential scanning calorimetry, wide-, and small-anlge X-ray scattering. The relationships between the local processing thermomechanical variables and the morphological parameters are then established. A study of the relationships between the developed morphology and the mechanical behavior of injection-molded PLCs will be described in a later article. It is expected that the morphology developed during processing, because of the imposed thermomechanical environment, will affect the mechanical properties of the injection-molded PLCs. This was demonstrated before by molecular dynamics computer simulations of PLCs comprised of amorphous polymeric chains and a rigid LC phase dispersed throughout the flexible matrix in the form of quasispherical islands. Simulation results indicate that both the LC concentration and spatial distribution influence the response of the material.<sup>29,30</sup> Furthermore, the presence of the rigid LC phase affects the predominant molecular deformation mechanisms,<sup>31</sup> and the crack formation and propagation phenomena.<sup>29</sup> On the other hand, experiments tell us that other morphological features—such as the skin-core ratio in injection-molded materials—influence the final macroscopic properties.<sup>32</sup>

# **EXPERIMENTAL**

#### Material and specimen geometry

The material is a longitudinal PLC: a copolyester comprised of 0.6 mole fraction of *p*-hydroxybenzoic acid (PHB) and 0.4 mole fraction of poly(ethylene terephtalate) (PET), referred to as PET/0.6PHB. It was purchased from Unitika, Kyoto, Japan, under the commercial name of Rodrun LC-3000. PET/ 0.6PHB has a density = 1.4 g cm<sup>-3</sup> and a melting temperature =  $200^{\circ}$ C.<sup>33</sup> Above that temperature, we have a smectic E phase coexisting with the isotropic liquid. Upon further heating, we have a 3-phase system (smectic E, smectic B, and isotropic liquid), then a different 2-phase system (smectic B and isotropic liquid) until the clearing temperature  $\approx 420^{\circ}$ C is reached.<sup>33</sup>

The molded specimens are tensile dumbbell-like bars 60-mm long with 20 mm of reference length and uniform rectangular cross-section (4  $\times$  2 mm<sup>2</sup>). The tab gate was located laterally in the middle of the specimen head.

#### Processing and thermomechanical environment

#### Molding programme

The molding program includes systematic variations of three processing variables: melt and mold temperature,  $T_{inj}$  and  $T_{w}$ , respectively, and injection flow rate,  $Q_{inj}$ . These were varied in three levels (1, minimum; 2, medium; and 3, maximum) following a design of experiments (DOE) plan based on a Taguchi orthogonal matrix (L9). This plan ensures that the effects of each processing variable can be analyzed independently of each other. Table I lists the values of the processing variables for each processing run. All the other processing parameters were kept constant. The order of the processing runs was randomized to avoid blocking effects.

The PLC material pellets were dried at 100°C during 12 h before molding. They were then molded in a Klocker-Ferromatic FM 20 injection molding machine. In each run, the weight of the molded samples was statistically controlled.

TABLE I Molding Program Based on a L9 Taguchi Orthogonal Matrix

	-		
Run	$T_{\rm inj}$ (°C)	$T_w$ (°C)	$Q_{\rm inj}$ (cm <sup>3</sup> /s)/ $t_{\rm inj}$ (s)
R1	220	20	4.9/1.00
R2	220	40	18.4/0.45
R3	220	60	27.0/0.10
R4	250	20	27.0/0.10
R5	250	40	4.9/1.00
R6	250	60	18.4/0.45
R7	280	20	18.4/0.45
R8	280	40	27.0/0.10
R9	280	60	4.9/1.00

 $T_{inj}$ , melt temperature;  $T_w$ , mold temperature;  $Q_{inj}$ , injection flow rate;  $t_{inj}$ , injection time.

#### Thermomechanical environment characterization

The thermomechanical environment imposed upon the melt during processing was estimated by computer modeling of the mold-filling stage using a commercial Moldflow package. A thermotropic PLC such as Rodrun LC3000 shows a complex rheological behavior because of its flexible-rigid segmental nature and polydomain structure. The rheological flow curves exhibit distinct behavior depending on the selected temperature,<sup>34</sup> evidencing a strong dependence of the viscosity upon the temperature. In the high shear rates regime ( $10^1$  to  $10^3$  s<sup>-1</sup>), a temperature-dependent shear thinning behavior can be assumed. In this study, the experimental viscosity data reported elsewhere<sup>34</sup> was used and fitted to a power-law equation:

$$\eta = 1.060 \times 10^{14} \dot{\gamma}^{-0.41} e^{-0.199T} \tag{1}$$

with *T* in °C. This simple constitutive model was used in the mold-filling simulations, despite its oversimplification in describing the rheological behavior of the material, which may lead to a less accurate characterization of the thermomechanical environment. The mold-filling simulations allowed the estimation of the through-the-thickness temperature, shear rate, and stress profiles. These profiles were characterized, respectively, by the bulk temperature, *T*<sub>b</sub>, the maximum shear rate,  $\dot{\gamma}_{max}$  and the shear stress at the solid/liquid interface,  $\tau_w$ . A cooling index, *Y*, was defined as<sup>32,35–37</sup>:

$$Y = \frac{T_b - T_t}{T_b - T_w} \tag{2}$$

where  $T_t$  is a transition temperature (a constant value of  $T_t = 200^{\circ}$ C was used). Y characterizes the thermal level of the moldings: a high Y corresponds to a slower cooling process.

# Morphological characterization

# Scanning electronic microscopy

The fracture surfaces of the tensile specimens were observed in a scanning electronic microscope (SEM) Leica Cambridge LS360. The samples were previously sputtered with a thin conductive layer of gold. Two magnifications were used:  $\times 30$  for observing the overall microstructure of the injection moldings and  $\times 5,000$  for observing the intermediate layer. The microstructure of the moldings was evaluated by measuring the thickness of each layer in their multilayered structure. Measurements were made under the lower magnification SEM micrographs using an image analyser software.

# Differential scanning calorimetry

Differential scanning calorimetry (DSC) scans were performed in a PerkinElmer DSC-7 apparatus in heating. Both experiments and calibrations (temperature and heat flow) were performed with indium standard at 20°C/min. Sample cross-sections of 14–20 mg were cut from the central zone of the specimens. One sample per processing run was used. For processing Run 1, the DSC experiment was repeated for three different specimens. Experimental scatter of 4.9, 0.7, and 14.6% was found, respectively, for the enthalpy of melting  $H_m$ , the glass transition temperature  $T_g$ , and the heat capacity change upon melting  $\Delta C_p$  (this error was assumed as equally dispersed within the other runs).

# Wide- and small-angle X-ray scattering

The crystalline phase morphology of the moldings was studied by X-ray synchrotron radiation source (transmission mode), generated at the Soft Condensed Matter beam line (A2 polymer), HASYLAB, at DESY, Hamburg, Germany (Gc with the wavelength of  $\lambda = 0.15$  nm). The injection-molded specimens were placed vertically (flow direction in the vertical) and perpendicularly to the incident X-ray beam. The experimental set-up allowed for the simultaneous acquisition of both wide-angle (WAXS) and small-angle X-ray (SAXS) bidimensional scattering patterns. For both techniques only one specimen per processing run was used.

The WAXS patterns were recorded on a phosphorus-based image plate after the exposure time of 60 s. The specimen-to-detector distance was 22 cm. The  $\theta$ -2 $\theta$  scattering scans for the equatorial reflection and the azimuthal angle scans for the mesogenic group reflection (highest intensity reflection arc) were obtained from the 2D WAXS patterns. Both these curves were corrected for background scattering and beam-intensity and normalized with respect to the specimen thickness and accumulation time. In the  $\theta$ -2 $\theta$  scans the scattering angle coordinate was calibrated using a semi-crystalline PET sample.

The level of orientation (or second-order orientation function), *s*, was calculated from the angular intensity distribution  $I(\phi)$ , using the Herman's orientation function as discussed by Gedde and Wiberg<sup>15</sup>:

$$s = \frac{1}{2} \left( 3 \left\langle \cos^2 \phi \right\rangle - 1 \right) \tag{3}$$

where is the angle between the axis of the average mesogen direction (director) and the preferred direction of the nematic phase. The angle brackets mean the averaging over the azimuthal angle, i.e.,

$$\left\langle \cos^2 \phi \right\rangle = \frac{\int_0^{\pi/2} I(\phi) \cos^2 \phi \sin \phi \, d\phi}{\int_0^{\pi/2} I(\phi) \, \sin \phi \, d\phi} \tag{4}$$

where  $I(\phi)$  is the scattered intensity of the mesogenic reflection at the azimuthal angle.

The parameter *s* can also be obtained by other routes, for instance by proton and deuterium nuclear magnetic resonance as described by Arrighi et al.<sup>38</sup> It follows from the definition of the angle that s = 1 for a system comprising molecules perfectly aligned in the flow direction. As already noted, given a highly layered configuration of injection-molded PLC, the use of a single global parameter to characterize the orientation state may be misleading. Tentatively, we might use a global through-the-thickness value of *s* as a measure of the mean orientation averaged over the thickness of the sample.

The SAXS reflections were recorded in a MARCCD detector, from which bidimensional patterns were obtained. The specimen-to-detector distance was 202 cm. The same exposure time was used as in the WAXS analysis. From these 2D patterns the intensity-scattering vector scans (in the meridional/flow direction) were calculated. Again, and as before, the intensity values were corrected from background scattering and beam intensity, and normalized with respect to the specimen thickness and exposure time. In the SAXS experiments, the scattering vector coordinate was calibrated by a rat tendon tail standard.

# RESULTS

# Thermomechanical environment characterization

Table II lists the computed values of the variables selected to characterize the thermomechanical environment associated with the experimental programme. The processing conditions resulted in a considerable variation of the shear stress and shear rate, also due to the high dependence of the

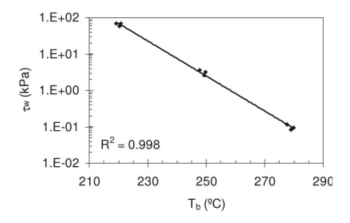
Computed Thermomechanical Variables								
Run	$T_b$ (°C)	$\tau_w$ (kPa)	$\dot{\gamma}_{max}~(s^{-1})$					
R1	219.3	70.0	$3.82 \times 10^{2}$					
R2	220.3	57.4	$8.69 \times 10^{2}$					
R3	220.8	67.7	$3.98 \times 10^{3}$					
R4	249.9	3.20	$3.97 \times 10^{3}$					
R5	247.7	3.68	$3.81 \times 10^{2}$					
R6	249.3	2.55	$8.62 \times 10^{2}$					
R7	279.1	$8.37 \times 10^{-2}$	$8.62 \times 10^{2}$					
R8	279.9	$9.46 \times 10^{-2}$	$3.98 \times 10^3$					
R9	277.6	$1.18  imes 10^{-1}$	$3.80 \times 10^{2}$					
Variation <sup>a</sup> (%)	27.6	$8.35 \times 10^{4}$	$9.95 \times 10^3$					

**TABLE II** 

 $T_{b}$ , bulk temperature;  $\dot{\gamma}_{max}$ , maximum shear rate;  $\tau_{w}$ , shear stress at the solid/melt interface.

<sup>a</sup> Variation = [(maximum value)-(minimum value)]/ (minimum value)  $\times$  100.

rheological behavior on the temperature and shearing levels. The values of  $T_b$  are only dependent upon the melt temperature, both temperatures being very similar. The bulk temperature increases linearly with the melt temperature. The highest variations were found in the shear stress, which is influenced mostly by the melt temperature. This is a consequence of the strong temperature dependence of the melt viscosity. The shear rate shows a two-decade



**Figure 2** Relationship between the wall shear stress,  $\tau_{w}$ , and the bulk temperature,  $T_b$ , for all the moldings evidencing a strong thermo-mechanical coupling (the solid line is the fitting of the data with  $\tau_w=2.55\times 10^{12}e^{-0.111T_b}$  equation ( $R^2$  is the coefficient of correlation).

amplitude of variation and is affected only by the injection flow rate: an increment on the injection flow rate induces a strong increase on the shear rate, as would be expected. In this shear rate range  $(10^2)$ to  $10^3 \text{ s}^{-1}$ ), the temperature and shear rate dependence of the viscosity were assumed to be well described by a power law model with an Arrhenius thermal dependence.

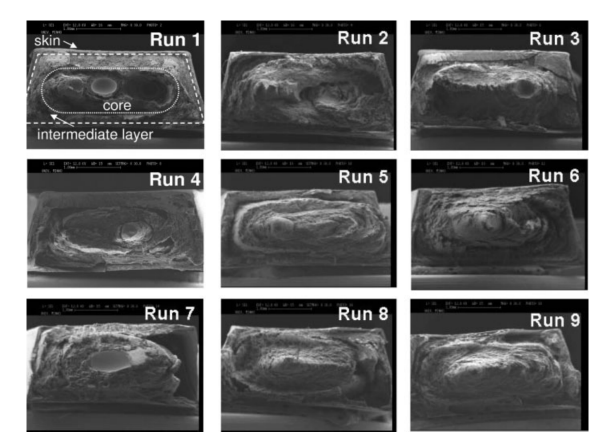


Figure 3 Effect of the processing variables (see Table I) on the microstructure of the moldings: SEM micrographs of the fracture surfaces.

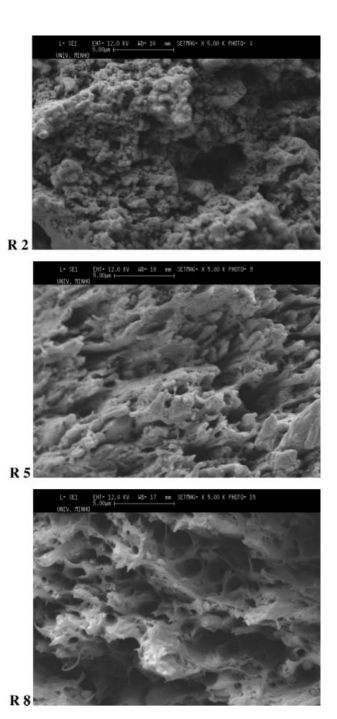
A strong thermal and mechanical coupling is evidenced during processing, as depicted in Figure 2, and already anticipated from the data in Table II. In fact, the data can be divided in three groups accordingly to  $T_b$  and  $\tau_w$  ranges. This thermomechanical coupling strongly determines and limits the morphology development in injection molding.<sup>35–37</sup>

# Morphological characterization by SEM

Figure 3 depicts the SEM micrographs of the fracture surfaces of all the tensile specimens. The processing conditions induce distinct patterns in the fracture surfaces.

The overall trend is the formation of a five-layer laminate composed of a very thin external layer, a large intermediate zone and an internal core, as illustrated in Figure 3 (Run 1). The thinner skin layer is developed during the filling stage as the hot material enters in contact with the cold mold wall, and its thickness is determined by the heat transfer conditions.<sup>23</sup> In the core region the melt experiences a tumbling flow of the poly-domains. This results in molecular orientation along the flow lines of this tumbling flow.<sup>19,23</sup> The fracture surfaces follow this imposed orientation, giving rise to parabolic patterns seen in the cores of the fracture specimens. In the intermediate layer, the molecular chains can be highly oriented in the flow direction. This layer can be therefore easily delaminated, creating visible holes (e.g., Run 7 in Fig. 3). A magnified view of these intermediate zones is presented in Figure 4 for selected processing runs.

For processing condition R2 it is possible to observe some spherical domains that were not deformed during the mechanical drawing. These aggregates are more evident for the low-melt temperature moldings (Runs 1 to 3), suggesting a weak coupling between these aggregates. Conversely, condition R8 shows a highly deformed structure, typical of samples molded at the highest melt temperature (Runs 7 to 9). In this case, the stress is better transferred to the aggregates which are able to deform, resulting in high local plasticity. For condition R5, some fibrillar-like structures can be observed (similarly also in Run 6). This type of morphology appears as intermediate with respect to those referred to above. The thicknesses of the layers were measured from the SEM micrographs. The skin layer is very small (c.a. 11% of the total molding thickness) and the measurements were not very accurate. For that reason, only two layers were considered: a highly oriented external layer (corresponding to the sum of the thickness of the skin and intermediate layers,  $S_i$ ; see Fig. 3) and a central core layer,  $C_a$ .



**Figure 4** Magnified (×5,000) SEM micrographs of the intermediate layer for selected moldings. (R2):  $T_{inj} = 220^{\circ}$ C;  $T_w = 40^{\circ}$ C,  $Q_{inj} = 18.4 \text{ cm}^3/\text{s}$ ; (R5):  $T_{inj} = 250^{\circ}$ C;  $T_w = 40^{\circ}$ C,  $Q_{inj} = 4.9 \text{ cm}^3/\text{s}$ ; (R8):  $T_{inj} = 280^{\circ}$ C;  $T_w = 40^{\circ}$ C,  $Q_{inj} = 27.0 \text{ cm}^3/\text{s}$ .

# Differential scanning calorimetry

The DSC scans for all the moldings are shown in Figure 5. The degree of crystallinity of the moldings is relatively low, as expected because the room temperature PET/0.6PHB solid contains 4 phases: PET crystals, PHB-rich islands, isotropic PET-rich glass and PHB-rich glass.<sup>33</sup> A very small and broad crystalline peak is observed centred approximately at

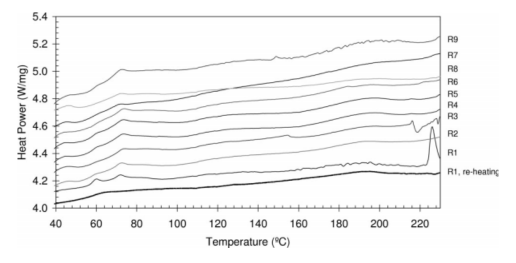
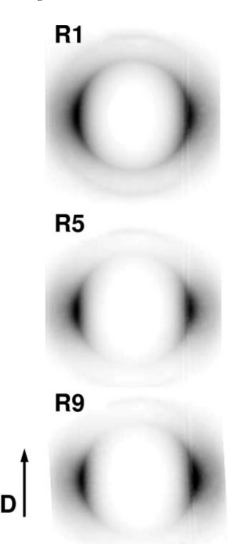


Figure 5 DSC scans for all the processing runs (see Table I for processing conditions and run identification).

190°C and ranging from approximately 150– 220°C (we recall the melting temperature resulting from a combination of results obtained by several techniques =  $200^{\circ}C^{33}$  whereas that temperature from DSC alone is  $212^{\circ}C^{10}$ ). The calculated enthalpies of fusion are very small, of the order of 2–4 J/g, whereas the experimental errors are significant. In Ref. 1 the value of  $H_m = 3$  J/g is reported, in the middle of our range. The low  $H_m$  value is typical for melting PLCs because the transition on heating does not result in a single isotropic phase but in a fairly well-ordered LC-containing phase.

Two glass transitions can be detected in all systems at 58 and 78°C. These processes are identified here by the endothermic peaks resulting from a structural relaxation process that occurred at room temperature before the DSC experimental run. By dynamic mechanical analysis a loss peak was found at 65°C (at 3 Hz) in PET/0.6PHB, ca. 12°C below the  $T_{\sigma}$  of the pure PET.<sup>33</sup> Also in dielectric relaxation spectroscopy a  $\alpha$  relaxation is identified for PET/ 0.6PHB at 60°C (maximum of the dielectric loss at 1 Hz), and it is attributed to the glass transition of the PET-rich domains.<sup>39</sup> A weak process, labelled  $\alpha'$ , was found at approximately 40°C and was ascribed segmental dynamics within the PHB-rich to domains; apparently this mechanism was not visible in the DSC results of the present work. The presence of the  $\alpha$ -relaxation in the PLC material was also confirmed by thermally stimulated depolarization, where a depolarization peak was observed at 65°C, and again attributed to the glass transition within the PET-rich phase.<sup>40</sup>

No significant differences were found in the two glass transition detected from the range of sample prepared. Second run DSC experiments were performed right after the first ones (see first lower curve in Fig. 6), to investigate the thermal properties of the samples after erasing the processing history. As



**Figure 6** WAXS patterns for different processing conditions. (R1):  $T_{inj} = 220^{\circ}$ C;  $T_w = 20^{\circ}$ C,  $Q_{inj} = 4.9 \text{ cm}^3/\text{s}$ ; (R5):  $T_{inj} = 250^{\circ}$ C;  $T_w = 40^{\circ}$ C,  $Q_{inj} = 4.9 \text{ cm}^3/\text{s}$ ; (R9):  $T_{inj} = 280^{\circ}$ C;  $T_w = 60^{\circ}$ C,  $Q_{inj} = 4.9 \text{ cm}^3/\text{s}$ . FD is the flow direction.

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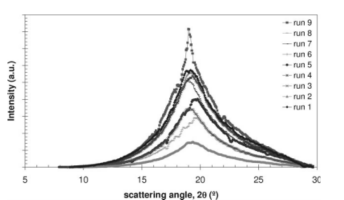


Figure 7 WAXS equatorial intensity scans for all the processing runs.

expected, the two endothermic peaks were no longer detected, as the thermal history was erased. Moreover, the high temperature glass transition process almost vanished, and only the glass transition at  $\approx$  58°C was detected as a step in the heat flow curve. This result is consistent with the findings observed for PET/0.6PHB discussed before. The high glass transition process should be then assigned to the segmental mobility of a confined PET-rich phase, induced during the processing. The developed shear stresses may create PET-rich amorphous regions that are geometrically confined within oriented crystalline structures. The difference in the dynamics for conformational mobility of bulk-like and confined amorphous PET was already quantified<sup>41</sup> from a practical point of view; apparently amorphous PET entrapped within crystalline structures exhibits a higher  $T_g$  than the one found in pure amorphous PET. Interestingly, the confinement effect due to the processing studied in this work seems to be similar in both skin and core of the injected parts. Samples taken from these two regions were also tested by DSC and, in the first run, the same position in the temperature axis and similar relative intensities of the two glass transitions were observed. Another possibility for the assignment of the two glass transitions is to consider the existence of two kinds of glassy states: an isotropic one and a more ordered glass resulting from the freezing of the mesophase structure. It was proposed before that the ordered phase should be characterized by a lower  $T_{\sigma}$ with respect to the amorphous one.42 Under this picture, we could conclude from the double glass transition events observed in the DSC results that after injection molding the two glassy states could coexist, as the fast cooling in the mold could prevent the complete organization of the chains. In the second DSC run just the lower  $T_g$  is observed indicating that the previous cooling was slow enough to allow the formation of the mesophase and the formation of a ordered glassy structure.

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# Wide-angle X-ray scattering

WAXS patterns of selected moldings are presented in Figure 6.

All the patterns show an intense equatorial reflection, reflecting the presence of the PHB mesogenic groups. The angular dimension of this reflection and its spread are both dependent on the processing conditions. For this set of moldings the effect of the thermal level on the developed morphology is already visible: as the thermal level increases (from runs R1 to R9), the mesogen reflection becomes narrower and the angular scattering is reduced. As such, a higher level of orientation of specimen R9 is anticipated (as compared to condition R1).

Figure 7 represents the WAXS equatorial scattering scan for all the moldings. The mesogenic group reflection is centred at  $2\theta = 19-20^{\circ}$ .<sup>6</sup>

The effect of processing is noticeable. The diffraction peak gets more intense and broad (compare conditions R2 and R9). The peak position is the same, within the experimental error. The high increase of intensity of the reflection peak reflects an increase of magnitude of the relative order of the mesogenic groups (higher LC order). The increase of the peak width is related to a decrease of the mean LC crystallite size inside an island.

Figure 8 shows the WAXS azimuthal scans for the mesogen group reflection. As expected, the level of molecular orientation is higher in the flow direction (FD) for all the molding conditions. The different thermomechanical environments imposed on the melt during processing result in significant differences in the level of the molecular orientation (mainly in the FD).

#### Small-angle X-ray scattering

The SAXS patterns of selected moldings are presented in Figure 9.

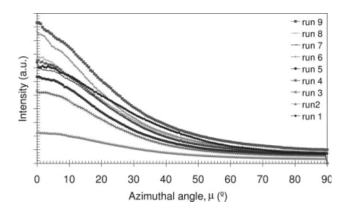
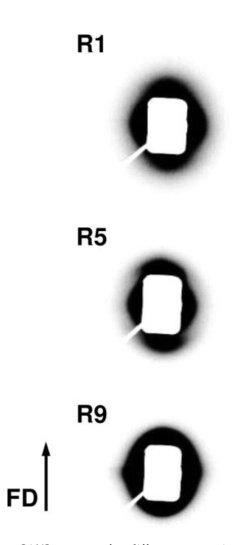


Figure 8 WAXS azimuthal intensity scans for the mesogen reflection for all the processing runs.



**Figure 9** SAXS patterns for different processing conditions. (R1):  $T_{inj} = 220^{\circ}$ C;  $T_w = 20^{\circ}$ C,  $Q_{inj} = 4.9 \text{ cm}^3/\text{s}$ ; (R5):  $T_{inj} = 250^{\circ}$ C;  $T_w = 40^{\circ}$ C,  $Q_{inj} = 4.9 \text{ cm}^3/\text{s}$ ; (R9):  $T_{inj} = 280^{\circ}$ C;  $T_w = 60^{\circ}$ C,  $Q_{inj} = 4.9 \text{ cm}^3/\text{s}$ .

The scattering pattern is typical of fibrillar morphology, with a high-long spacing and different levels of perfection. The SAXS patterns also show void scattering (voids are oriented parallel to the FD) dependent upon the processing conditions as well. Figure 10 depicts the correspondent meridional intensity scans for all the processing runs.

The local maximum indicates some periodicity of the morphology. However, the position of this maximum is not affected by the processing conditions, showing a constant long spacing in the range from 270 to 290 nm. This value should correspond to the size of the LC-rich island. However, the perfection of this regular structure seems to be very dependent upon the thermomechanical environment imposed during processing.

The processing runs with high melt temperature (R7, R8, and R9) show more intense and welldefined peaks becasue of the development of more ordered structures. Conversely, the conditions set at low melt temperature (R1, R2, and R3) are the ones presenting broader and less intense peaks, and therefore a more heterogeneous structure.

# Quantitative analysis

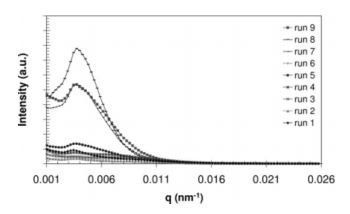
Table III summarizes quantitatively the morphological characterization of the moldings, in the most important measurable parameters. Processing has a pronounced effect on the layered microstructure of the moldings.

The highest variations are found in the values of (in descending order): heat capacity, intermediate skin/core layers thickness, average level of orientation. The glass transition and melting temperatures,  $T_g$  and  $T_m$ , are only slightly affected by processing. Figure 11 compares the effect of the processing conditions on the development of the morphological parameters. Run 1 shows the lowest values of *s*,  $\Delta C_p$ , but relatively high values of the skin thickness  $S_i$ . No apparent relationship between all the morphological parameters is evident.

The variations described earlier are averaged through the molding thickness. It is expected that the morphology of each layer will also be sensitive to the applied local thermomechanical environment. A more detailed through-the-thickness characterization (layer by layer) would elucidate this point. Nevertheless, there are significant variations of the morphology induced by processing, which forecasts a distinct mechanical behavior of the moldings.

#### PROCESSING EFFECTS ON THE MORPHOLOGY (ANOVA)

The morphology of the moldings was found to be very sensitive to the processing conditions. Even though establishing direct links between the processing parameters (machine set), and the developed morphology of the moldings is not a satisfactory route, <sup>32,35,36</sup> it is important to analyze what are the



**Figure 10** SAXS intensity scans for the mesogen reflection for all the processing runs.

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Results of the Morphological Characterization									
Run	DSC		WAXS		SEM				
	$T_g$ (°C)	$\Delta C_p (J/g^{\circ}C)$	$T_m$ (°C)	S	I <sub>max</sub> (a.u.)	$S_i ({ m m}/{ m m})$	$C_a (m/m)$		
R1	67.6	0.200	191.8	0.217	0.182	0.581	0.419		
R2	61.6	0.248	191.5	0.264	0.045	0.405	0.595		
R3	61.9	0.226	192.8	0.325	0.097	0.541	0.459		
R4	65.8	0.246	192.9	0.323	0.157	0.630	0.370		
R5	63.8	0.248	192.9	0.316	0.119	0.436	0.564		
R6	61.6	0.238	195.4	0.325	0.159	0.510	0.490		
R7	59.1	0.330	195.3	0.326	0.146	0.566	0.434		
R8	63.8	0.253	190.9	0.304	0.082	0.529	0.471		
R9	63.5	0.228	194.4	0.317	0.195	0.505	0.495		
Variation (%)	14.4	65.0	2.3	50.5	333.3	55.5	60.8		

TABLE III Results of the Morphological Characterization

 $T_{g}$ , glass transition temperature;  $\Delta C_p$ , heat capacity;  $T_m$ , melting temperature; s, level of orientation;  $I_{max}$ , maximum WAXS equatorial intensity;  $S_i$ , external layers thickness;  $C_a$ , core layer thickness.

contributions of the different processing parameters to the morphology development. Furthermore, this has been the general approach followed in the literature,<sup>23,24,27,28</sup> and it will be—initially—adopted in this discussion. The results were treated by analysis of variance (ANOVA). The relevant processing parameters were identified for a significance level of 0.9. The effect of processing on the microstructure of the moldings is shown in Figures 12–15.

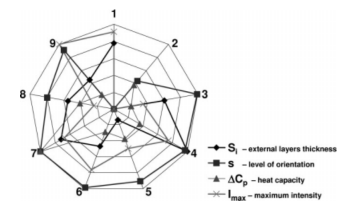
Figure 12 represents the variations of the thickness of the external layer  $S_i$  with the processing conditions. Only  $T_w$  and  $Q_{inj}$  are significant factors. The mold temperature has the highest influence with the contribution of 71%.

 $S_i$  shows a minimum value for intermediate  $T_w$ . At the lowest  $T_{w}$ , the high cooling rates contribute to thicker  $S_i$  (faster cooling). For the highest  $T_w$ , the expected lower melt viscosity induces a higher level of flow-induced molecular orientation, resulting in a thicker oriented layer due to the low relaxation capabilities of PLCs.<sup>20</sup> This also applies to the variations of  $S_i$  with  $Q_{inj}$ ; as  $Q_{inj}$  is increased, the melt viscosity decreases and the flow-induced molecular orientation increases, resulting in thicker  $S_i$ . The formation of the external layer appears to be dictated by the imposed cooling and shear levels. The negligible effect of  $T_{inj}$  may also be due to this balance. As  $T_{inj}$  increases the cooling rate decreases (lower  $S_i$ ), but simultaneously the flow-induced orientation increases (higher  $S_i$ ).  $S_i$  is therefore insensitive to variations of T<sub>inj</sub> within the studied range. As expected, the effect of the processing conditions on the thickness of the core layer,  $C_a$ , is opposed to that of  $S_i$ .

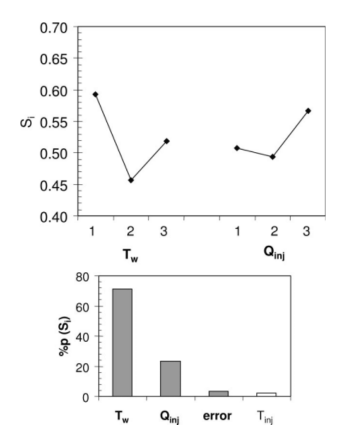
Figure 13 depicts the evolution of the orientation parameter *s* with the processing variables. Generally, *s* increases with the increase of all the studied processing variables. The contribution of  $T_{inj}$  is the highest (c.a. 47 %). Again, the increase of the thermal

level and imposed shear rate contributes to a higher molecular orientation in the moldings.

Despite the higher melt and mold temperatures, the flow-induced orientation is not allowed to relax due to the high relaxation times of PLCs. Higher Q<sub>inj</sub> will decrease the melt viscosity and will enhance the orientability of the melt. The final level of molecular orientation seems to be a compromise between the orientability of the melt (related to its viscosity) and the characteristic relaxation times of the PLC, as is the case for semicrystalline polymers.<sup>37</sup> Furthermore, the high value of the contribution of the pooled error anticipates an eventual important interaction between the studied processing variables, as already suggested.43 In fact, the effects of  $T_{inj}$  and  $Q_{inj}$  on the orientation levels are sometimes opposed. In general, low injection rates lead to high levels of molecular orientation,<sup>27,42-44</sup> but this is not always the case.<sup>42</sup> Higher melt temperatures result in higher levels of molecular orientation, but exceptions also exist here.36 This reveals



**Figure 11** Effect of processing conditions on the morphological parameters (the values are normalized between 0 and 1 for comparison purposes).



**Figure 12** Effect of the processing variables on the thickness of the external layers,  $S_i$  ( $T_{inj}$ , melt temperature;  $T_{uv}$ , mold temperature;  $Q_{inj}$ , injection flow rate; % p, percentage of contribution; error, pooled error, 1-2-3 are, respectively, the low, medium, and high level sets of the processing variables). Lines in the graph are for guidance.

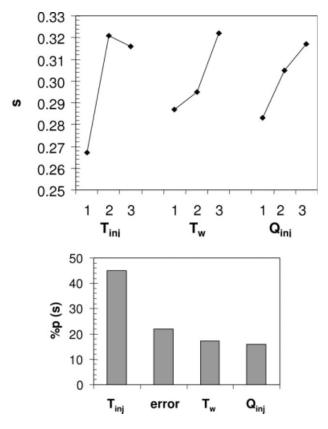
some interactions between the processing parameters, their effects being a function of their relative levels. The direct link between the overall processing parameters and the morphology is therefore not a satisfactory approach.

Figure 14 represents the maximum intensity of the WAXS peak,  $I_{max}$ , as a function of the processing variables.  $I_{max}$  is proportional to the order of the LC phase: a more ordered structure gives a higher maximum intensity of the WAXS reflection.

 $I_{\text{max}}$  increases with the increase of  $T_{\text{inj}}$ , especially for the low values of this processing parameter and it also decreases with the increment on  $Q_{\text{inj}}$ , in particular for lower  $Q_{\text{inj}}$  values. The dependences of  $I_{\text{max}}$  upon  $T_w$  show a minimum value for intermediate  $T_w$ . The most important processing parameter is  $T_w$ , with the contribution of 57%, followed by  $Q_{\text{inj}}$ and  $T_{\text{inj}}$  with 27 and 12.5 %, respectively.  $I_{\text{max}}$  seems to be determined by both the cooling conditions and shear levels applied during the molding phases. A higher thermal level results in a higher level of molecular orientation as a consequence of the lower melt viscosity. Incrementing  $Q_{\text{inj}}$  leads also to a higher level of orientation, but also to a thicker skin layer resulting in a less ordered structure. The effect of  $T_w$  on the extent of order depends upon the balance between the imposed cooling and shear levels. For low  $T_w$ , the shear level is higher and for high  $T_w$ the cooling rate is lower; both these conditions leading to a higher order state. For intermediate  $T_w$  values, these opposed trends lead to low order state. Nonetheless, these relationships are not easy to establish when an average through-the-thickness  $I_{max}$ value is considered.

# PROCESSING THERMOMECHANICAL ENVIRONMENT AND MORPHOLOGY RELATIONSHIPS

Clearly, the morphology development in injection molding is determined by the thermomechanical environment imposed on the melt during the processing stages. In fact, the morphology evolves inside the molding tool and is controlled locally by the thermal and mechanical fields applied to the material. Again, as already argued, it would therefore be desirable to establish relations between the local thermomechanical environment and the local morphology development.<sup>32,35–37</sup>



**Figure 13** Variation of the level of orientation, *S*, with the processing variables ( $T_{inj}$ , melt temperature;  $T_w$ , mold temperature;  $Q_{inj}$ , injection flow rate; %*p*, percentage of contribution; error, pooled error, 1-2-3 are, respectively, the low, medium, and high level sets of the processing variables). Lines in the graph are for guidance.

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**Figure 14** Effect of the processing variables on the maximum intensity,  $I_{max}$ , of the WAXS peak ( $T_{inj}$ , melt temperature;  $T_{w}$ , mold temperature;  $Q_{inj}$ , injection flow rate; % p, percentage of contribution; error, pooled error, 1-2-3 are, respectively, the low, medium, and high level sets of the processing variables).

Figures 15 and 16 present the variation of the assessed morphological parameters with thermomechanical variables locally computed from the moldfilling simulations. The shown dependences were chosen taking into consideration the results from previous Sections (*Processing effects – ANOVA results*) and *Thermomechanical environment characterization*) and an equation that best fits the experimental data (as evaluated by the coefficient of multiple correlation,  $R^2$ , and the percentage residuals). The low values of  $R^2$  of the fittings may indicate that other thermomechanical variables, interactions or more complex relationships need to be considered. The values of the residuals must be analysed taking into account the experimental scatter. The fitted simple equations, and the respective graphical representation, aim only at presenting the main trends of variations, evidencing the most relevant thermomechanical parameters, and their effects on the morphology development.

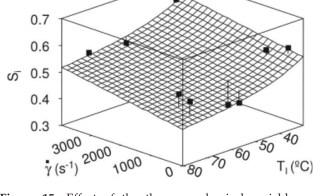
Figure 15 shows the variations of the outer skin ratio,  $S_i$ , with the maximum shear rate and temperature at the mold/polymer interface,  $T_i$ .

The latter is an interface temperature defined as an averaged temperature of the polymer (subscript p) and mold (subscript w) weighted by their thermal effusivities, b:

$$T_{i} = \frac{b_{w}T_{w} + b_{p}T_{p}}{b_{w} + b_{b}p}$$
(5)

Where  $T_m = T_p$ ,  $b = (\rho k c_p)^{0.5}$  ( $\rho$  is the density, *K* is the thermal conductivity, and  $c_p$  is the heat capacity). In this case,  $T_i$  is mainly governed by  $T_w$ .

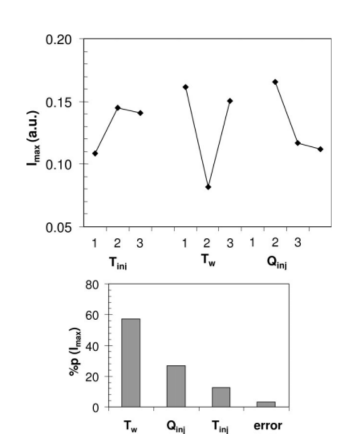
The main processing parameters influencing  $S_i$  are  $T_w$  and  $Q_{inj}$  (Fig. 13).  $\dot{\gamma}_{max}$  is mainly determined by  $Q_{inj}$  (Table II). A higher  $\dot{\gamma}_{max}$  leads to a thicker oriented layer. During mold-filling, the flow-induced orientation increases with  $\dot{\gamma}_{max}$ . This molecular orientation cannot relax due to the long relaxation times of PLCs, thus resulting in a thicker oriented layer. Conversely, a higher  $T_i$  results in less severe cooling conditions and in a thinner external layer. The thickness of this layer seems to be related to the



**Figure 15** Effect of the thermomechanical variables on the external skin layer ratio,  $S_i$  ( $\dot{\gamma}_{max}$  - maximum shear rate,  $T_i$  – interface temperature (mold/polymer),  $R^2 = 0.55$ ; average residuals of 8.0 %; maximum residual of 23.0%).

 $\begin{array}{c} 0.5 \\ 0.4 \\ 0.3 \\ 0.2 \\ 0.1 \\ 0 \\ 0.0^6 \\ 0.0^4 \\ 0.0^2 \\ 0 \\ 0.1 \\ 0 \\ 0.0^{-15} \\ 0.2 \\ 0 \\ 0.35 \\$ 

**Figure 16** Effect of the thermomechanical variables on the level of orientation of the moldings, s ( $\tau_{vv}$  shear stress at the solid/liquid interface; Y, cooling index,  $R^2 = 0.71$ , average residuals of 5.6 %; maximum residual of 11.0%).



shear and thermal levels – as in semi-crystalline polymers.<sup>35,37</sup>

Figure 16 shows the dependence of the orientation parameter *s* on the local shear stress and cooling index. *s* is affected by all processing parameters (Fig. 13). It increases with both *Y* and  $\tau_w$ . The dependence on *Y* is stronger than on  $\tau_w$ .

Y is mainly determined by  $T_{inj}$ . At high  $T_{inj}$  the melt viscosity is lower allowing a better melt orientability, thus, resulting in a higher flow-induced orientation. Despite the higher thermal level of the moldings, the molecular relaxation process is constrained by the high relaxation times of the rigid PLC molecules, and the flow-induced orientation is essentially preserved. The orientation parameter increases slightly with  $\tau_w$  because of the strong coupling between the thermal and stress levels. In fact, the highest  $\tau_w$  are obtained for the lowest Y, which tends to reduce s. On the other hand,  $\tau_w$  increases also with the flow rate, Qini, contributing to an increase of the flow-induced orientation and of the final level of orientation. In Figure 16, the data points are located along a narrow area. This is due to the high coupling between the thermal and mechanical fields in the injection molding process, as mentioned earlier (Fig. 2).

# CONCLUSIONS

We have established certain relationships between the local thermomechanical environment and the morphology developed in a thermotropic PLC. The main conclusions can be summarized as follows:

- A strong thermal and mechanical coupling is evidenced by the moldings.
- The imposed thermomechanical environment results in the development of a complex multilaminated and hierarchical microstructure whose thicknesses and morphological features are very sensitive to the processing conditions.
- The skin thickness is mainly determined by the mold temperature and injection flow rate, acting in opposite ways. The thickness increases with increasing melt flow shear rate and a reduction of the mold/melt interface temperature. The thickness of this layer seems to be related to the shear and thermal levels—as in semi-crystalline polymers.
- The molecular orientation increases with the increase of all processing parameters, but most significantly with the melt temperature. It is maximized for the highest thermal levels and shear stresses evidencing the preponderant effect of the flow-induced molecular orientation and high relaxation times of PLC materials.

• The order parameter (as reflected by the maximum intensity on WAXS scans) is mainly influenced by the mold temperature and injection flow rate. It seems to be determined by both the cooling conditions and shear levels applied during the molding phases.

The mechanical properties of the PLC injection moldings are expected to be determined by the morphology developed upon the constraints imposed by the processing thermomechanical environment. This will be reported later, where dependence of mechanical properties upon morphological parameters ( $S_i$ , s, and  $I_{max}$ ) will be demonstrated.

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