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FRACTURE BEHAVIOUR OF RECYCLABLE ALL-POLYPROPYLENE COMPOSITES COMPOSED OF α AND β MODIFICATIONS

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Abstract

The fracture behavior of all-PP composites was studied under quasi-static loading conditions. Fracture toughness was evaluated by means of different Fracture Mechanics approaches depending on materials' behavior. Composites consolidated at low temperature exhibited popin features and the failure occurs typically by delamination and tape pullout. With increasing consolidation quality – i.e. with increasing processing temperature – the delamination became less pronounced before the specimens break. In composites consolidated at the highest temperature investigated (190°C), the laminate-like structure typical of self-reinforced composites produced according to film-stacking method was lost. Accordingly, composites behave as if they were only α -PP and β -PP matrixes: α -rPP exhibited typical brittle fracture of α -PP, while β -rPP exhibited the stable behavior with fully yielded ligament before crack propagation commonly observed for β -PP. In general, stress-strain behavior changed from stable to unstable and fracture toughness strongly decreased as consolidation quality increased. Based on these results and previous findings, it can be concluded that the properties of self-reinforced PP composites can be tailored for a given application through the quality of consolidation.

Keywords: polypropylene; self reinforced composites; fracture behavior; toughness

1. Introduction

Polypropylene (PP) is apparently the major polymeric construction material of the future in view of its impressive growth figures of the past years. However, PP as such has to be reinforced to meet the high demands on stiffness and strength in engineering applications and glass fibers are the major reinforcing elements used in these materials. Unfortunately, in view of recyclability, glass fibers are components which still cause environmental problems, both in mechanical recycling and thermal recycling (incineration).

An important contribution to environment preservation and energy saving may come from our ability to recover, recycle and/or reuse materials. Consumer consciousness, waste management regulations and environmental legislation are all pushing the manufacturers of raw-materials and end-products to carefully consider the environmental impact of their products at all stages of the life cycle, including ultimate disposal. This scenario obviously involves also composite materials, increasingly used in several industrial sectors. In fact, there is a marked interest to improve the methods for recycling and reusing the existing composites, or to develop new materials intrinsically more suitable to be recycled and reused. In this framework, a great deal of efforts has been expended for the development of the so called 'single-polymer' composite materials [1].

The concept of 'single polymer' composites was first described by Capiati and Porter [2] using oriented polyethylene filaments and polyethylene powder with different melting points. Following this work, a large number of different techniques have been reported in the literature, for combining an oriented fiber with a separate phase to form the matrix. The so-called ''all-PP'' composites are designed to compete with traditional thermoplastic composites such as glass fiber reinforced PP (GF-PP). In comparison to the classical composites, self-reinforced composites have some advantages in addition to the good mechanical properties: the enhanced recyclability and low density which are achieved by using the same polymer for both fiber and matrix phase of the composite [3-10]. Unlike GF-PP, all-PP composites can be entirely melted down at the end of the product life for recycling into PP feedstock. As a consequence, all-PP composites are gaining acceptance in automotive applications.

Further promising possibility to widen the processing window is to exploit the polymorphismrelated difference between the melting temperature of β -(matrix) and α -phases (reinforcement) of PPs. Note that the β -PP has a markedly lower melting temperature that the α form [11-18]. Therefore the β -PP can fulfill the role of matrix, while the stretched α -PP should work as reinforcement. The resulting composite is really a PP homocomposite – the matrix and the reinforcement differ from one another only in their crystalline modifications. This was confirmed in previous works [10, 19-20].

Consolidation degree was proved to influence mechanical behavior [21]. For assurance of the reliability of these materials in structural applications, it is of paramount interest to study and understand the fracture resistance of composites. In addition, the study may be useful also in the optimum design of structural materials.

Traditionally toughness has been characterized by the Izod or Charpy impact energy. It has long been recognized that the impact energy is a very complicated strain rate function of the plastic and fracture work with generally the plastic work dominating. The Izod and Charpy tests have lost favor in mechanical engineering because they cannot be used directly in design, but they still have use for comparing the toughness of a particular polymer composite system. Design of structural parts, their connecting and assembly may be based on fracture mechanical approaches. In their desire to characterize toughness of polymer composites more exactly, many researchers have turned to Fracture Mechanics.

In this work, the fracture behavior of all-PP composites was studied under quasistatic conditions and toughness was evaluated by means of Fracture Mechanics approaches.

2. Experimental

2.1. Materials and sample preparation

A plain woven fabric (Stradom S. A., Czestochowa, Poland) composed of highly stretched split PP tapes with a nominal weight of 180 g/m² (approximately 180 μ m in thickness) was used as reinforcement. The reinforcing tape has a tensile strength of 465±32 MPa (measured on a single tape). Two kinds of PP were used as matrix materials: a random PP copolymer (TIPPLEN R351F TVK Nyrt., Tiszaújváros, Hungary), and β form of the same copolymer. Thermal properties of matrices and PP fabric are depicted in Table 1. The melting temperature of the β -modification was clearly below that of the corresponding α -version, as expected. The matrix films (9 layers) and the reinforcing woven fabrics (8 plies) were laminated according to the film-stacking method. Since the properties of the fabrics showed some directional anisotropy, they were assembled adopting a cross-ply lay-up to make the resulting sheets orthotropic. Self-reinforced PP composite sheets with a thickness of 2.5 mm and a nominal reinforcement (i.e. α -PP fabric) content of 50 wt% were produced by compression moulding of a film-stacked package at 4 different processing temperatures (147, 162, 177 and 190°C). They were selected 5-45°C above the relevant matrix melting temperature. For the latter, the DSC melting peak was considered. The consolidation process took place as follows: after heating up the moulds, the film-stacked package was inserted and held for 30 s without pressure and for 90 s under a pressure of 7 MPa, and then it was cooled down to 50°C with a cooling rate of 7.5°C/min and demoulded. It is noteworthy that the holding time at processing temperature was kept as short and low, respectively, as possible to prevent shrinkage (relaxation) of the fibers.

2.2. Mechanical tests and data reduction

Fracture characterization was carried out on mode I double edge-notched tensile specimens (DENT) cut from 3 mm thick plaques (nominal width *W* was 23 mm and nominal length *S* was 100 mm), at a crosshead speed of 10 mm/min. Sharp notches were introduced by scalpel-sliding a razor blade having an on-edge tip radius of 0.13 μ m.

Different Fracture Mechanics approaches were applied depending on materials' behavior. In cases where there is not a significant crack growth resistance the value of the *J*-integral at initiation, J_{Ic} , is a good measure of toughness. The *J*-integral is conventionally defined for non-linear elastic materials as a path independent line integral. In fact, the single-specimen *J* formulation has been extensively used in the past to characterize ductile fracture in polymers [22-23]. Although ASTM E813 and ASTM E1152 apply only to ductile fracture, more recent standards, allow *J* integral of testing materials that fail by cleavage. The J_c parameter [22, 24] is applicable to characterize quasi-brittle failure behavior (load-displacement curves with sharp load drop at the point of fracture) of specimens with a crack to depth ratio close to 0.5. J_c was evaluated at the instability load point by calculating the fracture energy required to produce cleavage behavior of pre-cracked specimens having a crack depth to width ratio of $0.45 \le a/W \le 0.55$ as:

$$J_c = \frac{\eta U_{tot}}{B(W-a)} \tag{1}$$

where U_{tot} is the overall fracture energy, *i.e.* the total area under the load-deflection curve, *B* the thickness of tested specimens, and η a geometry factor that for DENT specimens is expressed as [25]

$$\eta = -0.06 + 5.99 \left(\frac{a}{W}\right) - 7.42 \left(\frac{a}{W}\right)^2 + 3.29 \left(\frac{a}{W}\right)^3$$
(2)

The *J*-integral approach is a natural extension of Linear Elastic Fracture Mechanics and works best for not too ductile fractures. A methodology that works best for very ductile fracture that has been used for polymer composites is the Essential Work of Fracture (EWF) approach. This approach was first proposed for plane stress ductile metal fractures [26] and later applied to polymers [27]. The aim of the EWF approach is to separate the work performed in the fracture process zone, W_e , from the total work of fracture, W_f , that in ductile polymers is often dominated by the work of plastic deformation, W_p .

$$W_f = W_e + W_p \tag{3}$$

The work performed in the fracture process zone, termed the essential work of fracture, is a quasi-material property only dependent on the specimen thickness. The EWF method makes use of the fact that the essential work and the plastic work scale differently:

$$W_f = w_e l B + \beta w_p l^2 B \tag{4}$$

where *B* is the plate thickness, β is the shape factor, and w_e and w_p are the specific essential work of fracture and the specific non-essential work of fracture, respectively. By dividing W_f by the ligament area *lB*, one obtains the specific total work w_f that can be expressed as:

$$w_f = w_e + \beta w_p l \tag{5}$$

If the entire specimen ligament deforms plastically before fracture initiation, then the specific essential work can be found by testing different ligament lengths and extrapolating the specific total work of fracture to zero ligament length. In recent years the EWF method has been extensively applied to polymers and there is a draft standard of the European Structural Integrity Society (ESIS) [28]. The EWF method delivers a single fracture parameter that is representative of crack propagation.

It is noteworthy that J_{Ic} and w_e can be compared since they should be identical or very similar, as was indeed corroborated by several authors [29-32]. The EWF approach can also be used for plane strain fracture J_{Ic} either obtained from slow strain rate tests [33-34] or impact tests [35].

Broken samples were studied by optical microscopy and lateral views of tested samples were examined by scanning electron microscopy (SEM) after they had been coated by a thin layer of gold.

3. Results and discussion

Figure 1 shows typical DENT load-displacement curves along with typical failure behavior for all composites. Composites consolidated at lower temperature (147°C) exhibited pop-in features, i.e. crack propagation occurred by the stick-slip mechanism. Each load drop resulted in a peak linked with the crack progress. The failure occurred typically by delamination and tape pullout and load remained constant during test till it dropped to zero gradually. With increasing consolidation quality (with increasing processing temperature) the delamination became less pronounced when the specimens broke (note that the delamination process absorbs high energy but easy delamination is linked to poor tensile and flexural mechanical properties). Composites consolidated at 162°C and 177°C exhibited non-linear quasi-brittle behavior with abrupt load drop. A different phenomenon was observed in composites consolidated at 190°C. Laminate-like structure typical of self-reinforced composites produced according to film-stacking method was lost. Accordingly, composites behaved as if they were only α -PP and β -PP matrixes: α -rPP consolidated at 190°C exhibited typical brittle fracture of α -PP, while β -rPP consolidated at 190°C exhibited stable behavior with fully yielded ligament before crack propagation commonly observed for β -PP. It should be recalled that with increasing hot pressing temperature the consolidation improved and a significant

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transcrystalline layer formed between the reinforcement and the matrix [36]. It seems that at 190°C partial or total melting of reinforcing tapes occurred, so the reinforcement and the matrix are not only adhesively but also cohesively bonded.

It is noteworthy that composites consolidated at the lowest temperature $(147^{\circ}C)$ absorb considerable higher energy than the better consolidated at intermediate temperatures (167 and 177°C).

Optical micrographs give extra evidence of failure mechanisms. Some examples are shown in Figure 2. In poorly consolidated samples, debonding of tapes is clearly seen as white lines, and stretching of tapes can be easily observed. Moreover, lateral SEM inspection of failed samples confirms massive delamination of fabric tapes (Figure 3). In α -rPP based composites consolidated at 190°C, on the other hand, no signs of plastic deformation are found, while β -rPP based composites consolidated at 190°C exhibited fully yielded ligament. Based on load-displacement curves and observed failure surfaces, toughness was evaluated using both Fracture Mechanics approaches described in experimental section. In Figure 4, load-displacement curves showing similarity along with work of fracture graphs are shown for composites in which toughness was estimated by the EWF approach. Toughness values are listed in Table 2 along with their deviations. It is clearly seen that those values decrease with increasing processing temperature (consolidation) (Figure 5) and no significant differences can be found between the two different matrices investigated. From the above discussion it can be concluded that in poor consolidated composites PP tapes debond from the PP matrix, so crack advances passing through the PP matrix leaving PP fibers practically intact and available for load carrying. This toughening mechanism, in which the fibers completely bridge the crack faces, preventing the composite from undergoing catastrophic failure, is known as a fully bridging situation [37]. After a certain degree of crack face displacement, fibers no longer bridge the crack faces and load starts to fall, while fibers are continuously pulled out from the matrix, allowing the composites to reach a higher level of final displacement. The extension of this mechanism depends on the degree of adhesion between matrix and fabric: the lower the adhesion, the higher the degree of crack bridge and pull out. Hence, the superior performance of poor consolidated composites is based on the high degree of PP fibrillation, giving rise to improved energy absorption capability with sustained crack growth stability through crack surface bridging and large drawability.

4. Conclusions

The fracture behavior of all-PP composites was studied under quasi-static loading conditions and toughness was evaluated by means of different Fracture Mechanics approaches depending on the materials' response.

It was found that stress-strain behavior changed from stable to unstable and fracture toughness strongly decreased as consolidation quality increased.

The superior performance of poor consolidated composites is based on the high degree of PP fibrillation, giving rise to improved energy absorption capability with sustained crack growth stability through crack surface bridging and large drawability.

Based on the results of this investigation and previous results on similar composites [21, 36], it can be concluded that the properties of the composites can be tailored for a given application through the quality of consolidation.

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Figure captions

Figure 1. Load-displacement curves and macroscopic failure behavior of all-PP composites

Figure 2. Light microscopy photographs of some typical failures

Figure 3. SEM micrograph of typical failure of a poorly consolidated composite

Figure 4. Load-displacement curves and work of fracture plots for composites which exhibited stable behavior

Figure 5. Toughness of the composites as a function of processing temperature.

of the composites ..



Load-displacement curves and macroscopic failure behavior of all-PP composites 191x168mm (500 x 500 DPI)

5 mm





SEM micrograph of typical failure of a poorly consolidated composite 180x140mm (500 x 500 DPI)



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Load-displacement curves and work of fracture plots for composites which exhibited stable behavior 206x241mm (500 x 500 DPI)



Toughness of the composites as a function of processing temperature 105x88mm (500 x 500 DPI) **Table 1.** Thermal properties of materials determined by differential scanningcalorimeter (DSC) [10].

Material	Melting temperature [°C]	
α–rPP	142.5	
β–rPP	131.3	
α-PP fabric	172.4	

Table 2. Toughness of self-reinforced PP composites

Composite	$w_e [\text{kJ/m}^2]$	$J_c [\mathrm{kJ/m^2}]$
α–rPP 147	904.1 ± 115.1	-
α–rPP 162	-	536.6 ± 46.5
α–rPP 177		299.4 ± 39.5
α–rPP 190		6.8 ± 1.0
β–rPP 147	975.8 ± 129.9	-
β–rPP 162	-	547.8 ± 46.7
β–rPP 177	-	413.3 ± 104.3
β–rPP 190	5.25 ± 0.9	-

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