Impact fracture toughness of polyethylene materials for injection moulding

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Introduction

This paper presents some results on the toughness of high-density polyethylene (HDPE), when subjected to impact loading at conditions giving semi-brittle fracture with crazing as the main mechanism contributing to the fracture resistance. The test is based on the recent ISO 17281 standard, and the SENB geometry (Fig. 1) is used. This test is supposed to isolate the brittle/crazing fracture in plane strain, and characterises the resistance to unstable crack propagation. The well-known Charpy test (ISO 179), on the other hand, measures a combination of the brittle/crazing toughness (up to maximum force) and the energy absorbed by shear yielding during crack propagation. Furthermore, in ISO 17281 a sharp pre-crack is made with a razorblade, while in ISO 179 the notch is machined with a certain radius, typically 0.25 mm.

The main topic of our study has been to investigate relationships between material parameters (molecular weight and comonomer content) and mechanical response. This paper presents data for a set of HDPE materials having $M_w$ in the range 55-145 kg/mol and nominal density in the range 953-963 kg/m$^3$. In addition to molecular parameters ($M_w$ etc), this paper will focus on effects of processing. Injection-moulded and compression-moulded specimens have quite different fracture toughness values. This is related to the process-induced morphologies. Injection-moulded specimens have higher toughness, but lower stiffness and density. Hence, the higher toughness could be attributed to a higher density of tie-chains and entanglements, due to faster cooling. Fracture mechanics studies are usually based on compression-moulded specimens, due to their rather homogeneous and isotropic morphology, and the possibility to make thick plates (up to 10-12 mm), which are needed to fulfil geometry size criteria of certain fracture mechanics tests. However, injection-moulded specimens are more relevant when there is a need to assess the response of mass-produced products. The inhomogeneity, anisotropy and thickness limitations of injection-moulded specimens are challenges for fracture mechanics tests.

The ultimate aim of fracture mechanics testing is to obtain geometry-independent fracture toughness values. However, due to complicating factors, such values may not be attainable. Complications related to specimen thickness are mentioned in this paper.

Fig. 1 The SENB ('single edge notched bending') test geometry with notation according to ISO 17281 and 13586. The length of the notch including pre-crack (a) is varied according to the specific test. The sharp pre-crack is made with a razorblade. Note that, in fracture mechanics, the machined notch of a SENB specimen usually differs from that of a (ISO 179) Charpy specimen, since the former notch typically is deeper. A typical SENB notch geometry is illustrated on the right.
Materials, experimental procedures and data analysis

Materials: High-density polyethylene (HDPE) materials supplied by Borealis were used in these tests. All the materials are commercial grades with injection moulding as the main application. Table 1 shows the main characteristics of the materials. The grades with the lowest densities are copolymers, with 1-butene as comonomer. Ti-based Ziegler-Natta catalysts were used for all materials.

Table 1 Material parameters and mechanical properties of the polyethylene grades used in this study. The information is based on datasheets and information from Borealis. The mechanical properties were determined with specimens injection-moulded according to ISO 1872-2 and ISO 294-1.

<table>
<thead>
<tr>
<th>PE1</th>
<th>PE2</th>
<th>PE3</th>
<th>PE4</th>
<th>PE5</th>
<th>PE6</th>
</tr>
</thead>
<tbody>
<tr>
<td>MFR[^a] [g/10 min]</td>
<td>31</td>
<td>12</td>
<td>8</td>
<td>4</td>
<td>4</td>
</tr>
<tr>
<td>$M_c[^b]$ [kg/mol]</td>
<td>9</td>
<td>17</td>
<td>19</td>
<td>19</td>
<td>11</td>
</tr>
<tr>
<td>$M_w[^b]$ [kg/mol]</td>
<td>55</td>
<td>74</td>
<td>90</td>
<td>115</td>
<td>115</td>
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<tr>
<td>$M_z[^b]$ [kg/mol]</td>
<td></td>
<td></td>
<td></td>
<td>475</td>
<td>685</td>
</tr>
<tr>
<td>$M_c/M_w$</td>
<td>6.1</td>
<td>4.4</td>
<td>4.7</td>
<td>6.1</td>
<td>10.5</td>
</tr>
<tr>
<td>Density[^c] [kg/m$^3$]</td>
<td>960</td>
<td>961</td>
<td>963</td>
<td>954</td>
<td>953</td>
</tr>
<tr>
<td>Modulus[^d] [MPa]</td>
<td>1050</td>
<td>1150</td>
<td>1200</td>
<td>950</td>
<td>850</td>
</tr>
<tr>
<td>Yield stress[^d] [MPa]</td>
<td>25</td>
<td>26</td>
<td>27</td>
<td>24</td>
<td>22</td>
</tr>
<tr>
<td>Strain at yield[^d] [%]</td>
<td>9</td>
<td>9</td>
<td>8</td>
<td>10</td>
<td>10</td>
</tr>
<tr>
<td>Charpy toughness[^e] at -20°C [kJ/m$^2$]</td>
<td>3</td>
<td>4</td>
<td>6</td>
<td>7</td>
<td>7</td>
</tr>
<tr>
<td>Charpy toughness[^e] at 23°C [kJ/m$^2$]</td>
<td>3</td>
<td>4</td>
<td>7</td>
<td>11</td>
<td></td>
</tr>
<tr>
<td>ESCR[^f] [h]</td>
<td>~1</td>
<td>&lt;10</td>
<td>&lt;10</td>
<td>10</td>
<td>40</td>
</tr>
</tbody>
</table>

\[^a\] MFR = Melt flow rate measured according to ISO 1133 (190°C/2.16kg)
\[^b\] Number-average ($M_c$), weight-average ($M_w$) and z-average ($M_z$) molecular weight calculated from size exclusion chromatography (SEC) data. The $M_z$ value of PE4 was not measured on the actual lot.
\[^c\] These data were measured on the actual lots. Standard: ISO 1183. Circular samples with diameter 5 mm were stamped out of 2 mm thick compression-moulded plates. The cooling rate of the compression moulding process was 15 K/min.
\[^d\] Tensile properties measured according to ISO 527-2. The cross-head speed was 1 mm/min for the tensile modulus data and 50 mm/min at yield.
\[^e\] Charpy impact toughness measured according to ISO 179-1 with notched geometry type 1eA. The actual lots/specimens used in the fracture mechanics tests were also subjected to standard Charpy testing.
\[^f\] Environmental Stress Crack Resistance (ASTM D 1693). Some of these ESCR values were not measured on the actual lots.

Injection moulding of test specimens: Charpy specimens (ISO 179 type 1 bars with dimensions $l = 80$ mm, $w = 10$ mm and $h = 4$ mm) were injection moulded with an Engel ES 330/65 machine. The injection moulding was performed according to ISO 294-1 and ISO 1872-2. The bars are filled along the length direction, and ISO standards specify the melt front speed through the critical cross-section ($h \times w$ in our case). This value is 100 mm/s for PE, and the effective injection time was set according to this (ca. 0.8 s). The temperature of the melt was 210°C, and the mould temperature was 40°C. The packing time was 25 s, and the packing pressure was varied according to the MFR value in order to obtain the correct dimensions, and avoid sink marks, flash and other defects. The packing pressure was 90 MPa for the materials with MFR up to 4, and 70 MPa for the rest. The total cycle time was 45 s.

Compression moulding of test specimens: Specimens were machined from compression-moulded plates, with thickness 4 mm (for making bars with $w = 10$ mm and $h = 4$ mm) and 8 mm (for making bars with $w = 20$ mm and $h = 8$ mm). A Collins laboratory press was used with the following process for the 4 mm thick plates (based on ISO 1872-2):

- Heating for 4 minutes from 23°C to 180°C, while the mould is closing
- Heating for 10 minutes without pressure (180°C)
- Packing for 5 minutes (16.5 MPa, 180°C)
- Cooling for 11 minutes (16.5 MPa, 15 K/min)

For the 8 mm thick plates a pressure of 0.3-0.4 MPa was applied during the second step, and a new step was inserted after this (5 minutes, 1 MPa). Hence, the total cycle time was 35 minutes for the 8 mm thick plates.
Morphology of test specimens: Samples from the bulk, in the vicinity of the notch, of untested specimens were analysed by differential scanning calorimetry (DSC), using a Perkin Elmer Pyris 1 instrument. The samples were heated (10 K/min) to 200°C, cooled (10 K/min) to 20°C and then heated again (10 K/min) to 200°C. The degree of crystallinity was calculated from the normalised integral of the melting endotherm, using the value $\Delta H = 270 \text{ J/g}$ [1] for the enthalpy of fusion of 100% crystalline PE. A density column was used to measure the densities of samples taken from the bulk in the vicinity of the notch of untested specimens. The degree of crystallinity was also calculated from the measured density, using the following values for the density of crystalline and amorphous PE: $\rho_c = 996 \text{ kg/m}^3$ [2] and $\rho_a = 850 \text{ kg/m}^3$ [3]. Thin films (~ 10 $\mu$m) were micromotred from untested specimens, and characterised by optical (transmission) microscopy, using polarised light. The films were micromotred from two planes in the vicinity of the notch: The thickness–length plane ($h-l$ plane in Fig. 1) and the thickness–width ($h-w$) plane.

Fracture mechanics testing: The tests were based on ISO 17281 using the single edge notched bending (SENB) geometry. The standard conditions were 23°C and an impact speed of 1 m/s, and the tests were performed with a Rosand/Imatek instrumented falling weight impact tester. The notches were milled, and pre-cracks were made with a razorblade. The large compression-moulded specimens (cross-section 20 mm x 8 mm before notching) also had machined side-grooves (Fig. 2), in order to avoid shear lips. The side-grooves were 0.8 mm deep on each side, giving an $h$ value of 6.4 mm at the loaded cross-section. With these materials and specimens, dynamic effects (force oscillations) is a major concern at impact speeds around 1 m/s and above. Hence, mechanical damping (silicone grease) was applied on all fracture mechanics specimens. Values for the dynamic flexural yield stress and the dynamic flexural modulus were also obtained (using un-notched specimens). The impact speed was 0.55 m/s, giving a similar time to yield as the time to fracture initiation in the fracture tests. The yield stress and modulus values were needed in some of the data analyses.

Fig. 2 Testing of a specimen with $a/w = 0.5$. Note that this specimen also has so-called side-grooves (the vertical notches on picture to the right) with opening angle 45° and root radius 0.25 mm.

Calculation of fracture mechanics parameters: ISO 17281 apply to linear-elastic fracture mechanics, characterising theresistance to unstable crack growth by the fracture toughness parameters $K_c$ (the critical stress intensity factor – a force/stress based parameter) or $G_c$ (the critical energy release rate – an energy based parameter). Most PE materials are too tough for a linear-elastic description, even at impact conditions, and an elastic-plastic description must be used. A common parameter for elastic-plastic fracture mechanics is the critical value of the $J$ integral ($J_c$). ($J_c$ is equal to $G_c$ in the linear-elastic limit). There are several methods for obtaining $J_c$. Based on work by Grellmann et al. [4] and Fasce et al. [5], we determined $J_c$ values from data obtained with ISO 17281, using the expression by Rice et al. [6]. This expression is the simplest and most robust; other are sensitive to details in the calculation procedure (e.g. separation of elastic and plastic energy), and require tensile data at similar strain rates. Two other expressions were assessed in the present study, but the discussion of these expressions is beyond the scope of this paper.
Results

DSC, density and microscopy (bars with \( h = 4 \text{ mm} \))

Thermograms from the first heating (Fig. 3 and Table 2) reflect the crystalline structures of the specimens as used in the fracture mechanics test. The temperature range of the first melting endotherm is an indication of the lamella thickness distribution, although factors such as crystal defects and chain constraints also affect the endotherm. As an illustrative reference, the theoretical estimate for the equilibrium melting point of a perfect infinite PE crystal is reported to be in the range 141-146 °C. Furthermore, the normalised integral of the endotherm (enthalpy of fusion, \( \Delta H \)) of the first heating is a measure of the degree of crystallinity (weight fraction). In the second heating scan, compression-moulded and injection-moulded samples should have similar degree of crystallinity, as the history from the original moulding process should be erased to a large extent.

The crystallinities of samples from injection-moulded (IM) specimens vary from 62 to 69 wt% (1st heating), and are correlated with the densities in Table 1. There may also be a molecular weight effect involved. For all samples from IM specimens, a higher crystallinity is recorded in the 2nd heating, due to the lower cooling rate in the DSC than in the IM process. Also, note that for IM specimens the crystallinities recorded during the first heating are correlated with those of the second heating (Fig. 4), indicating that the injection moulding process does not induce totally different microstructures (in the core) from those obtained during slow cooling without flow.

For PE3 it is observed that compression moulding (CM) gives higher crystallinity (1st heating) than IM, again due to the lower cooling rate. For the CM specimen, the crystallinity recorded during the first heating is lower than that of the second. The nominal cooling rate was higher during compression moulding (15 K/min compared to 10 K/min in the DSC pan), but the effective rate at the core was lower, due to the thicker sample. The compression-moulded PE3 specimen had higher density than the injection-moulded specimen, in agreement with the DSC crystallinity trends.

The peak melting temperature (\( T_m \)) is related to the average lamella thickness, and the upper-end temperature of the melting regime, \( T_u \), is related to the lamellae with the highest melting point (i.e. the thickest lamellae). Fig. 3 shows that \( T_m \) and \( T_u \) values decrease with decreasing (data sheet) density. Hence, adding more short-chain branches (reducing the density) probably reduces the average and maximum lamella thickness. The highest \( T_m \) and \( T_u \) values were observed for the CM specimen. This must be due to the slow cooling, allowing for thick lamellae to develop.

Some micrographs obtained with polarised light are shown in Fig. 5. As expected, the CM specimen is more homogeneous than the IM specimen. The former has a spherulitic morphology with spherulite diameter around 15-20 µm. The spherulites are slightly larger near the centre than near the surface (which touched the mould during compression moulding). A 15-20 µm thick skin layer shows somewhat different polarisation, perhaps due to transcrysallisation. The IM specimen has different morphological zones through the cross-section. This is well-known from the literature. The spherulites near the centre are somewhat smaller than those in the CM specimen. With even higher magnification (not shown in Fig. 5), banded spherulites are observed for the IM specimen. The structure near the surface of the IM specimen is too fine to be resolved by optical microscopy. Sections of the IM specimen cut parallel and perpendicular to the flow direction only differ slightly.

Table 2  DSC and density results for samples cut out from the hulk in the vicinity of the crack of untested specimens (bars). The specimens were injection moulded, except ‘CM’ (compression moulded).

<table>
<thead>
<tr>
<th></th>
<th>PE2</th>
<th>PE3</th>
<th>PE3</th>
<th>PE4</th>
<th>PE5</th>
<th>PE6</th>
</tr>
</thead>
<tbody>
<tr>
<td>First heating</td>
<td>Crystallinity (DSC) [wt%]</td>
<td>69.1</td>
<td>77.0</td>
<td>68.9</td>
<td>63.6</td>
<td>62.3</td>
</tr>
<tr>
<td></td>
<td>Peak melting temp. [°C]</td>
<td>133.5</td>
<td>136.4</td>
<td>134.5</td>
<td>131.4</td>
<td>131.8</td>
</tr>
<tr>
<td></td>
<td>Onset of melting a [°C]</td>
<td>125.8</td>
<td>126.9</td>
<td>125.0</td>
<td>129.0</td>
<td>124.5</td>
</tr>
<tr>
<td>Second heating</td>
<td>Crystallinity (DSC) [wt%]</td>
<td>75.3</td>
<td>73.2</td>
<td>73.5</td>
<td>68.4</td>
<td>65.4</td>
</tr>
<tr>
<td></td>
<td>Peak melting temp. [°C]</td>
<td>132.8</td>
<td>132.1</td>
<td>132.5</td>
<td>131.3</td>
<td>130.0</td>
</tr>
<tr>
<td></td>
<td>Density [kg/m³]</td>
<td>-</td>
<td>962</td>
<td>953</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

\( a \)  The onset of melting is defined as the intersection between the temperature axis and a tangent to the endotherm.
Fig. 3 DSC traces for the first melting. Injection-moulded specimens, except ‘PE3 CM’ which was compression moulded.

Flexural modulus (un-notched bars with \( h = 4 \) mm)

The flexural modulus of elasticity was measured at impact (dynamic) conditions. Data for injection-moulded (IM) specimens and compression-moulded (CM) specimens are shown in Table 3. CM specimens have higher moduli than IM specimens. For quasi-static testing the difference is smaller; for PE6 the modulus of a CM specimen was 27% higher than for an IM specimen.

In our data set for IM specimens, the dynamic moduli of most specimens are lower than moduli from quasi-static tensile testing. This is probably not correct. Quasi-static moduli are obtained for strains between 0.2 and 0.5% (ISO 527). In the dynamic data, a possibly higher slope at such low strains was hidden by force oscillations.

The (pseudo) moduli of IM specimens can be determined with somewhat higher certainty than those of CM specimens. The stress-strain curves of IM specimens are typically linear up to 3% strain, while the curves of CM specimens are only linear up to 2% strain (using the same fitting criteria). The curves of CM specimens also have stronger initial oscillations.

Table 3 Flexural moduli at impact conditions. Bars with cross-section 4 mm (\( h \)) x 10 mm (\( w \)). Data based on three parallels. Quasi-static properties from the data sheets are given for comparison (last row).

<table>
<thead>
<tr>
<th></th>
<th>PE1</th>
<th>PE2</th>
<th>PE3</th>
<th>PE4</th>
<th>PE5</th>
<th>PE6</th>
</tr>
</thead>
<tbody>
<tr>
<td>E, IM bars [GPa]</td>
<td>–</td>
<td>1.04 ± .03</td>
<td>0.99 ± .04</td>
<td>0.87 ± .06</td>
<td>0.86 ± .03</td>
<td>0.89 ± .06</td>
</tr>
<tr>
<td>E, CM bars [GPa]</td>
<td>1.37 ± .09</td>
<td>1.62 ± .04</td>
<td>1.50 ± .07</td>
<td>1.32 ± .02</td>
<td>1.18 ± .04</td>
<td>1.36 ± .02</td>
</tr>
<tr>
<td>E_{quasi-static}, IM bars [GPa]</td>
<td>1.05</td>
<td>1.15</td>
<td>1.20</td>
<td>0.95</td>
<td>0.85</td>
<td>0.90</td>
</tr>
</tbody>
</table>

Fig. 4 Crystallinities calculated from the 1st and 2nd DSC heating scans. IM = Injection moulded. CM = Compression moulded. The figure is based on data in Table 2.
Overview (specimen surface at bottom, centre of specimen indicated by dash-dot line)

Position near the centre

PE3 CM, E = 1.5 GPa

Position near the surface

PE3 IM, E = 1.0 GPa,

**Fig. 5** Optical microscopy of microtomed sections from 4 mm thick compression-moulded (left) and injection-moulded (right) specimens. The sections of the injection-moulded specimen were cut parallel to the flow direction. Values of the dynamic moduli (E) are given in the bottom line.
Fracture mechanics results (bars with $w = 10\,\text{mm}$ and $h = 4\,\text{mm}$)

In this section we will mainly focus on injection-moulded (IM) vs. compression-moulded (CM) specimens, but some other main trends will also be summarised. A fracture surface with some of the main features is shown in Fig. 6. The size of the craze zone increases with increasing molecular weight and decreasing density. When injection-moulded, the toughest material (PE6) does not fracture in a semi-brittle manner, as the other materials.

IM and CM specimens of PE3 are compared in Fig. 7 and Fig. 8. The CM specimen absorbs less energy to peak force (less pronounced craze zone), and also seems to absorb less energy during crack propagation (no clear shear lips).

![Fracture surface with main features](image1)

**Fig. 6** Left: Optical microscopy of fracture surface of IM PE3. The crack tip (notch) made with a razorblade is indicated by the red arrows. Right: Fractured specimen (fracture surfaces inside the red circle). Also note the strain gauge that was used on some specimens.

![Representative fracture surfaces](image2)

**Fig. 7** Representative fracture surfaces for injection-moulded (IM) and compression-moulded (CM) specimens of PE3 ($h = 4\,\text{mm}$ and $w = 10\,\text{mm}$).
Fracture toughness values are shown in Table 4. Note that IM specimens have higher $G_C$ and $J_C$ values, but lower $K_C$ values (i.e. lower maximum force). The difference between IM and CM specimens is larger for larger for PE5 than for PE3. Charpy (ISO 179) data for PE6 showed even larger differences. At 23°C the IM specimen had a factor 3.5 higher Charpy toughness, while at -20°C the factor was 2.0.

With the linear-elastic approach ($G_C$ and $K_C$), the criteria given in the standard could not always be satisfied with these specimen dimensions and the chosen notch depths and test conditions (impact speed and temperature).

**Table 4** Results for linear-elastic fracture toughness ($K_C$ and $G_C$), elastic-plastic fracture toughness ($J_C$). The impact speed was 1 m/s and the temperature was 23 °C. Only fracture mechanics parameters in bold are valid with regard to the criteria of the standards/protocols.

<table>
<thead>
<tr>
<th></th>
<th>$K_C$ [MPa m$^{0.5}$]</th>
<th>$G_C$ [kJ/m$^2$]</th>
<th>$J_C$ [kJ/m$^2$]</th>
</tr>
</thead>
<tbody>
<tr>
<td>PE2 IM</td>
<td>1.38 ±0.1</td>
<td>2.05</td>
<td>1.5 ±0.3</td>
</tr>
<tr>
<td>PE3 IM</td>
<td>1.55 ±0.1</td>
<td>2.4</td>
<td>1.9 ±0.5</td>
</tr>
<tr>
<td>PE3 CM</td>
<td>1.60 ±0.1</td>
<td>1.6</td>
<td>1.5 ±0.1</td>
</tr>
<tr>
<td>PE4 IM</td>
<td>1.7 ±0.1</td>
<td>2.8</td>
<td>3.6 ±0.2</td>
</tr>
<tr>
<td>PE5 IM</td>
<td>1.8 ±0.1</td>
<td></td>
<td>3.6 ±0.3</td>
</tr>
<tr>
<td>PE5 CM</td>
<td>1.93 ±0.03</td>
<td></td>
<td>2.1 ±0.3</td>
</tr>
<tr>
<td>PE6 IM</td>
<td></td>
<td></td>
<td>7 ±1</td>
</tr>
</tbody>
</table>
Fracture mechanics results for large compression-moulded bars
($w = 20 \text{ mm and } h = 6.4 \text{ mm (8 mm minus side-grooves)}$)

The most common validity criteria for elastic-plastic fracture mechanics tests were satisfied when using compression-moulded bars with this cross-section. Also, the side-grooves prevented the formation of shear lips (yielding), thereby isolating a semi-brittle plane-strain fracture with crazing as the main mechanism contributing to the fracture resistance. All the criteria of ISO 17281 (linear-elastic fracture mechanics) were still only satisfied for PE1, PE2 and PE3 (with our rather conservative yield stress values). Typical load curves for PE1 (the least tough material) and PE6 (the toughest material) are shown in Fig. 9. Note that PE6 exhibits a time to fracture which is a factor 2 higher than that of PE1.

![Graph showing force vs. time for PE1 and PE6](image)

Fig. 9 Recorded force vs. time for PE1 (upper diagram) and PE6. Data for three tested specimens are shown (red, blue and green curves).
The main trends for this set of materials are the same as reported earlier. The main effects of molecular weight and density are seen in Fig. 10.

For the materials PE3 and PE5, compression-moulded specimens were made in two ‘sizes’. The large specimens (with $w = 20 \text{ mm}$ and $h = 6.4 \text{ mm}$) gave higher toughness values for all three toughness parameters ($K_C$, $G_C$ and $J_C$). The values were a factor 1.3-1.4 higher.

Finally, some comments on the reproducibility and statistics:

- The reproducibility of the force vs. time curves is good, taking into account the impact speed and the rather low forces involved, see Fig. 9.
- The least tough material (PE1) had the lowest standard deviation of the $J_C$ value.
- The average coefficient of variation (CV) for all six materials was 7%.
- The $J_C$ values generally had somewhat higher standard deviation and CV than the $K_C$ values. The average CV of the $K_C$ values was 4%.

![Fracture toughness vs. weight-average molecular weight for large compression-moulded bars.](image)

**Fig. 10** Fracture toughness vs. weight-average molecular weight for large compression-moulded bars.

**Discussion**

**General**

Based on the literature from the last 20 years, we can say that the mechanisms resisting the initiation and propagation of crack growth in HDPE (crazing, yielding), and their relationships with MWD (molecular weight distribution) and CCD (comonomer concentration, type and distribution) via morphology, are qualitatively understood. Fracture mechanics is an important tool in analysing structure-property relationships. Some important features of fracture mechanics are 1) the isolation of fracture mechanisms and 2) the concept of geometry-independent parameters. However, in order to analyse the performance of a set of materials, a rigorous fracture mechanics approach needs to be complemented with high-quality data characterising the material (MWD, CCD), the morphology, and the fracture surfaces. This is especially important when analysing subtle effects of catalyst system or multimodality. Recent in-situ studies of crack tip deformation have also revealed interesting features.

Quantitative impact fracture mechanics is not straight forward. The fracture mechanisms can sometimes not be decoupled, thermal effects may affect the tests results, and the size criteria in the literature
are perhaps not suitable for tough polymers. Furthermore, dynamic oscillations in the recorded force are a problem at high impact speeds.

A geometry-independent toughness value is the ideal. However, validity criteria, especially with regard to specimen size, are seldom mentioned in the scientific literature on impact fracture mechanics of polymers. When discussing and comparing the responses of different materials, all guidelines and criteria may not need to be taken literally, i.e. some ‘technically invalid’ results may still be interesting, showing qualitatively correct trends.

Fracture toughness vs. molecular weight and comonomer content

The main trend for this set of materials is that the toughness increases with increasing weight-average molecular weight ($M_w$). This is known from the literature. It is attributed to increasing density of interlamellar tie-chains and chain entanglements, which give a higher craze fibril strength, and, hence, a higher energy for initiation of fracture. An increase in the size of the craze zone with increasing $M_w$ is seen on the fracture surfaces.

A higher comonomer content (lower bulk density) is also known to increase the density of tie-chains, at least in a certain range. Higher toughness for grades with lower bulk density is observed in our study.

The two grades with similar $M_n$ and density (PE4 and PE5) have almost the same fracture toughness (within the uncertainty). The bimodal grade (PE5) has higher $M_z$ (Table 1), and should have a better comonomer distribution (more comonomers in the high MW fraction). On the other hand, the lower $M_n$ value may have a negative effect. Overall, it was expected that the bimodal grade would have a higher toughness. We will discuss this further in another publication, where also other mechanical parameters will be compared for these two materials.

Finally, note that the tensile modulus of the material may affect the apparent impact toughness indirectly via an adiabatic micomechanism. According to Leevers [7], adiabatic effects will reduce the toughness with increasing modulus ($G_c$ proportional to $E^{-1/3}$).

Fracture toughness of injection-moulded (IM) vs. compression-moulded (CM) specimens

Fracture mechanics tests in the literature are usually performed with specimens machined from CM plates. This gives isotropic and homogeneous specimens (surface effects can be minimised by machining off the surfaces of the original plate). IM specimens, on the other hand, always have a certain degree of anisotropy (processing-induced molecular orientation), and a skin-core variation in morphology (caused by a variation in cooling rate, pressure and shear stress during solidification). Another difference between CM and IM specimens is the cooling rate, which is much higher for the latter. For an injection moulding grade of HPDE, the morphology obtained by injection moulding is the most relevant to probe by fracture mechanics testing. However, when testing IM specimens, possible effects of inhomogeneities and anisotropy should be assessed.

In our data set, IM specimens are tougher\(^1\) than CM specimens. The main explanation for the higher toughness of IM specimens is probably the higher density of tie-molecules and chain entanglements, due to the higher cooling rate. Flow-induced orientation in the IM specimens may also play a role [8, 9]. In our tests, the fracture plane is perpendicular to the flow direction in the IM process, i.e. the loading is parallel to the flow direction. However, it is difficult to separate the orientation effect from the cooling rate effect without also testing specimens with loading perpendicular to the flow direction. This could be done by machining out specimens from a plate with width at least the length of the specimens. Schrauwen et al. [8] reported that IM specimens with thickness above 1 mm and different injection temperatures, loaded parallel and perpendicular to the flow direction, had similar Izod impact toughness values. The toughness of a 1 mm thick CM plate was similar to that of a 1 mm thick IM specimen with low orientation level loaded perpendicular to the flow direction.

The IM specimens have lower flexural modulus (Table 3). This must be due to the lower crystallinity in the core (Table 2), and probably also lower crystallinity out towards the surface. We are saying probably, because the shearing near the surface, and the pressure during crystallization further into the specimens, increase the nucleation and crystallization rates. Since the IM specimens have lower flexural modulus, the anisotropy effect (orientation in the flow direction) is overshadowed by the crystallinity effect.

The yield stress may have a separate effect on the toughness. A lower yield stress will give a larger damage zone, and also affect the relative amounts of crazing and shear yielding. CM and IM specimens have

\(^1\) IM specimens have higher $J_C$ and $G_C$ values. However, the $K_C$ values are slightly lower. Different trends for parameters based on energy and force (stress field) are not uncommon in the literature.
different yield stress. Above the $\gamma$ relaxation the yield stress should reflect the degree of crystallinity, which
is higher for CM specimens. In our tests with un-notched specimens in three-point bending at impact
conditions, CM specimens have a higher stress than IM specimens at the highest strains in the test, but a
local maximum or a clear yield plateau was not reached in these tests. Results for two common yield criteria
based on the deviation from elastic behaviour were not conclusive regarding CM vs IM specimens. The first
criterion was based on stiffness reduction to 95%. The other criterion was based on 'strain offset', i.e. the
intersection between the stress curve and a line parallel to the elastic part of the curve, but translated (offset)
to a higher strain. Strain offset values of 0.002 and 0.005 were tried. Testing with a high-speed tensile tester
is planned.

The standard deviation of the fracture toughness is typically lower for CM specimens than for IM
specimens. Optimising the IM process to obtain a more stable processing window, or controlling the IM
process via pressure sensors in the mould, could reduce the shot-to-shot variation of the IM specimens.
Removing the skin layer of the IM specimens could maybe also reduce the scatter. However, the
inhomogeneous and anisotropic structure of IM specimens is unavoidable.

**Effect of specimen size**

Our data, as well as data from Imperial College London on some of the same materials, shows that the large
compression-moulded specimens give higher toughness values than the small compression-moulded
specimens. In linear-elastic fracture mechanics, a lower or equal toughness value is expected when
increasing the specimen thickness ($h$ in Fig. 1), see Fig. 11. Williams [10] also reported a decrease in the
apparent impact toughness of an HDPE when increasing the specimen thickness ($h$) in the range 6-12 mm.

The higher apparent toughness of the large specimens in our study is in line with the thermal
decohesion model [7, [11,. According to this model, $G_c$ should be a factor $2^{1/3}$ (1.26) higher for specimens
with twice the size and the same shape. (According to ref. [7], $G_c$ is proportional to $w^{1/3}$). Our ratios for $G_c$
and $J_c$ are somewhat higher. It would be interesting to study size effects further, in particular to study the
effects of $h$ and $w$ independently. Size criteria for fracture mechanics of polymer materials is a field where
further work is needed, but this is beyond the scope of the present study.

In the scientific literature on impact fracture mechanics of PE, the specimens are typically made from
8-12 mm thick compression-moulded plates, i.e. $h = 8-12$ mm. However, size criteria are usually not
discussed in the scientific literature. From a practical point of view, injection-moulded (IM) 4 mm thick
(Charpy) specimens are interesting, because they can be made fast for extensive screening and routine
testing, and because their morphology is the relevant one for IM grades. Grellmann et al. [4] used such small
specimens, and also focused on size criteria [12, 13, 14].

![Plane stress, Plane strain](image)

**Fig. 11**  a) Plastic zone around crack tip, and its variation through the specimen thickness ($h$), according to
linear-elastic fracture mechanics [15]. b) The resulting effect of specimen thickness ($h$) on apparent
toughness. Note the plane-strain lower-bound toughness plateau for thick specimens.
In order to study effect of specimen thickness and processing further, we propose trying to weld together specimens, e.g. welding three specimens with cross-section 10 mm x 4 mm together, making a specimen with $h = 10$ mm and $w = 12$ mm. Since the most common relative pre-crack length tests is $a/w = 0.5$ (used for $K_C$ and $J_C$), welding together three specimens would give us a pre-crack ending halfway between two weld lines.

Could differences in morphology contribute to the size effect? The cooling rate is lower at the centre of the 8 mm thick plate. Slower cooling would mean higher crystallinity and lower tie-chain density. The direct effect of this (if significant) would probably rather be the opposite, i.e. lower toughness for the thicker specimen.

Finally, it could be mentioned that for thin injection-moulded products the plane-stress toughness is a relevant value, although the plane-strain toughness is the ‘lower-bound’ (conservative) toughness. Different methods can be used to obtain the plane-stress toughness. Preliminary work by the authors indicates that the so-called reversed Charpy test [16, 17] may not be suitable for the least tough HDPE materials in this study.

**Conclusions**

Values for the fracture toughness of six injection moulding grades of HDPE were obtained. We have determined critical parameters for the onset of unstable crack growth in plane strain at impact conditions (1 m/s) and 23 °C. The testing and data analysis was based on the ISO 17281 standard (linear-elastic fracture mechanics) using the single edge notched bending (SENB) geometry. In addition, values for the critical $J$ integral (elastic-plastic fracture mechanics) were calculated.

- With compression-moulded side-grooved specimens machined from 8 mm thick plates, the toughness values for all materials were valid in terms of criteria for elastic-plastic fracture mechanics ($J$ integral). The formation of shear lips was prevented, and the craze zone length was quite uniform over the fracture surface cross-section.
- The fracture toughness increased with increasing $M_w$ and increasing comonomer content.
- Injection-moulded specimens had higher fracture toughness than compression-moulded specimens. In both cases, the main morphological factor determining the fracture toughness was believed to be the density of tie-chains and chain entanglements. A higher ‘network’ density increases the strength of craze micro-fibrils.
- Two (compression-moulded) specimen sizes were tested, and the largest specimens gave the highest toughness value. This could be explained by the thermal decohesion model by Leevers et al. Further work is needed to assess size effects, in particular to study the effects of specimen height and width (SENB geometry) independently. This could also teach us more about how to interpret the results for injection-moulded specimens, with limited (wall) thickness.
References


[2] Clark ES, Unit cell information on some important polymers, Ch. 30 in Physical Properties of Polymers Handbook, Mark JE (Ed.), American Institute of Physics, 1996


[11] Leevers PS, private communication


