Chemical Composition Analysis of Polyolefins by Multiple Detection GPC-IR5

W. W. Yau, A. Ortín, P. del Hierro, Polymer Char, Valencia, Spain

Introduction

Polyolefin (PO) is the largest volume industrial polymer in the world for making a wide range of commercial products that touch nearly every aspect of our daily lives, such as automobile parts, pipes, packaging films, household bottles, baby diapers, and so on. Polyolefin as a group includes high-density and low-density polyethylene (HDPE, LDPE), polypropylene (PP), EP rubber, and linear low density (LLDPE) copolymers of ethylene with alpha olefins (propylene, 1-butene, 1-hexene, 1-octene).

Though chemically simple, being made up of only carbon and hydrogen atoms, PO products derive their wide range of enduse properties from their semicrystalline structure. The ability to incorporate co-monomers in PE to create short chain branches (SCB) makes it possible to control the polymer crystallinity and crystalline morphology, and thus control the rigidity and flexibility of PO products. The controlling factor also depends on the SCB variations across the molar mass distribution (MMD).

In dilute solution, the effect of SCB causes a reduction in polymer coil size with an increase of methyl groups off the backbone. Such structural changes can be studied by high temperature size exclusion chromatography (SEC, also known as gel permeation chromatography, or GPC), equipped with a multiwavelength infrared detector (IR5 MCT)¹, a light scattering (LS) detector, and a viscosity detector in the triple-detector GPC configuration (TD-GPC)². These techniques allow the analysis of SCB variation across MMD as it is illustrated by using the homopolymer PP and PE samples described in Table I together with their blend.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Mw (g/mol)</th>
<th>Mn (g/mol)</th>
<th>Mw/Mn</th>
<th>IV (dL/g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Broad PP</td>
<td>530,000</td>
<td>141,200</td>
<td>3.75</td>
<td>2.44</td>
</tr>
<tr>
<td>Broad HDPE</td>
<td>51,900</td>
<td>20,600</td>
<td>2.52</td>
<td>1.01</td>
</tr>
<tr>
<td>Blend (50/50)</td>
<td>291,000</td>
<td>36,000</td>
<td>8.09</td>
<td>1.73</td>
</tr>
</tbody>
</table>

Experimental Conditions

- Eluent: TCB (stabilized with 300 ppm BHT).
- Columns: 3 × PLgel 10 µm, 300 × 7.5 mm.
- Flow rate: 1.0 mL/min.
- Injection volume: 200 µL.
- Sample concentration: 2 mg/mL.
- Temperature: dissolution 160 °C; columns: 145 °C; detectors 160 °C; MALLS 150 °C.
- Chromatographic system: GPC-IR5 by Polymer Char.
- Detectors: IR5 MCT infrared detector by Polymer Char. 4-capillary viscometer by Polymer Char. Multiple angle laser light scattering HELEOS™ 8+ by Wyatt Technology.
- Data handling: GPC One by Polymer Char.

Results and Discussion

1. Ethylene-propylene Composition Analysis by GPC-IR5 MCT:

The MMD obtained by GPC-LS for the three samples in Table I are shown in Figure 1, where the IR5 MCT detector total CH channel is used to record the sample concentration across the GPC elution curve.

For studying chemical composition variation across MMD, the combination of detector signals from the IR5 MCT methylene (CH₂) and methyl (CH₃) channels, shown in the bottom part of Figure 2, is used. Since the PP part in the blend sample is of higher MM than the PE part, a higher methyl signal is seen in the higher MM end of the sample, as expected.

The trend of changing ethylene percent, seen in Figure 2, can be predicted a priori from the co-adding of the original PE and PP GPC elution curves. This ethylene to propylene transition also can be determined from the difference between the CH₂ and CH₃ signals of the IR5 MCT detector. The validity of this analytical approach is supported by the agreement seen in the Figure between the predicted and the IR5 MCT results.
2. Ethylene-propylene Composition Analysis by TD-GPC:

In the triple detector GPC, the ratio of LS signal over sample concentration at each GPC elution volume gives a measure of the polymer weight-average MM ($M_w$). Similarly, the ratio of the viscosity detector signal over sample concentration gives the polymer intrinsic viscosity (IV). The plot of Log IV versus Log $M_w$ by LS is called the Mark-Houwink (MH) plot, which is an often used tool to reveal polymer branching and conformational structures.

Both light scattering and viscosity detector signals are more responsive to higher MM molecules. This is the reason that the LS and viscosity curves of the blend sample are skewed more to the higher MM region than the IR concentration curve shown in Figure 3.

The corresponding MH plots for the three samples are shown in Figure 4. The MH line of the PP sample is seen to have a shift to higher MM from that of the PE sample. This is because of the extra methyl group off the chain backbone that leads to higher MM and lower IV for the PP sample. The MH plot of the blend sample in Figure 4 is seen to go through a transition from the PE line to the PP line as the MM increases. The experimental data agrees with the predicted curve very well. This agreement supports the use of TD-GPC for studying polymer blends and block copolymers of EP, PE, EO (ethylene/ octene), and so on.

Conclusions

The micro-structural difference of Polyethylene and Polypropylene as well as the compositional variation in EP can be detected by GPC-IR5 MCT and TD-GPC, as it has been validated by using a test sample of known composition created by a 50-50 PE-PP blend. This result of SCB variation across MMD provides the additional polymer structural information that complements perfectly the CRYS TAF (crystallization analysis fractionation), TREF (temperature rising elution fractionation), and CEF (crystallization elution fractionation) results which give SCB distribution across the crystallization temperatures and to the cross-fractionation technique.

References