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A study on the dispersion, preparation, characterization and photo-degradation of polypropylene traced with rare earth oxides

Feliks Bezati, Valérie Massardier, Jean Balcaen, Daniel Froelich

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Site de Plasturgie-IMP/LMM, UMR-CNRS 5223, 6 rue Pierre et Marie Curie, F-01108 Oyonnax, France.

Laboratoire Conception Produit Innovation, Chambéry (LCPI), Institut ARTS et METIERS ParisTech Chambéry, Savoie Technolac, 4 Rue du Lac Majeur, F-73375 Le Bourget du Lac Cedex, France.

Abstract

This research work deals with the effect of rare earth oxides on the PP matrix with respect to the thermal and mechanical properties and to the photo-degradation under UV irradiation exposure. The rare earth oxides are used as tracers for the identification of polymer materials, in order to have an economically efficient recycling and high speed automatic sorting of plastic wastes. The addition of 0.1 wt% of such particles of a micrometric size has a minor effect on the mechanical and thermal properties of the traced materials, as well as on the photo-degradation of the polymer after UV irradiation exposure. For 1 wt% tracer content, before UV irradiation treatment, the melting and crystallization temperatures as well as the thermal stability of the PP matrix are slightly increased, whereas the elongation at break decreases from 10 to 50% for a cross-head speed of 250 mm/min. However, the addition of 1 wt% of CeO₂ improves the photo-degradation resistance of the PP matrix to UV exposure due to the UV light screening effects offered by these particles. The SEM images together with the results obtained from image processing show a homogenous dispersion of tracers in the PP matrix.

1. Introduction

Tracers consist of systems formed by one or several substances dispersed into a material with the aim of adding a selective property to it and improving its identification. In recent works [1,2], an X-ray fluorescence (XRF) detection system based on the use of tracers for the identification of polymer materials was reported by the authors. This system was developed with the objective of having an economical, efficient and automatic sorting of plastic wastes. The addition of X-ray fluorescent tracers in polymer materials during the compounding process, offers many benefits by increasing the efficiency and purity of sorting and providing a solution for the identification of dark plastics.

Tracers must not chemically react with the host material, and cause changes to any of their physical, chemical or mechanical properties beyond acceptable limits. For products which are likely to have long useful lives under normal outdoor environment, such as automotive parts, the tracer particles need to be highly stable under weathering and during plastic processing [3].

The automotive and electrical industries are currently the worst performers concerning the recycling of plastics. This is partly due to the complexity of the waste materials that these sectors produce [4]. For this study, polypropylene (PP), one of the world’s major plastics, abundantly present in polymers to be recycled from the End of Life Vehicles (ELV) and the Waste of Electric and Electronic Equipments (WEEE) was chosen.

XRF is a volume, non-destructive spectroscopic method enabling elemental analysis of materials. Therefore the choice of tracers is directly linked to the elements of the periodic table. Once the toxic and radioactive elements are eliminated, along with the elements present in polymer additives, the low atomic number elements and the elements with limited reserves, the result leads to the selection of rare earth elements. For these selected ones we have chosen the most stable chemical molecules, the oxides [5]. The rare earth oxides (Y₂O₃, CeO₂, Nd₂O₃, Gd₂O₃, Dy₂O₃, Er₂O₃ and Yb₂O₃) used for this study are neither abrasive, nor toxic during their application and use and also stable in the environment [6,7].

The minerals used as fillers in polypropylene are principally talc and calcium carbonate. However, little attention has been paid to...
2. Experimental

2.1. Materials and preparation of traced polypropylene composites

The investigation described in this paper was conducted with a commercial grade of isotactic polypropylene homopolymer (ISPLEN PP 050 G1E) manufactured by Repsol. The original pellets have a melt mass-flow rate of 5.8 g/10 min (2.16 kg at 230 °C) and a density of 0.905 g/cm³. ISPLEN PP 050 G1E is intended for applications that require good impact resistance and is believed to contain low quantities of UV stabilizers and a small amount of heat stabilizer in order to minimize degradation during processing.

The characteristics of the rare earth oxides used as tracers are given in Table 1. The tracer content in the composites is 0.1 and 1%, respectively. All traced materials were prepared under identical mixing and moulding conditions. The injection moulded specimens of 2 mm thickness were produced by using a Battenfeld Unilog B2/350 Plus injection moulding machine. The barrel temperature was fixed at 220 °C for all zones. Before being injected the pellets were extruded twice by using a twin screw extruder CLEXTAL BC21 machine (screw of L/D = 90 and D = 25 mm) for dispersing homogeneously the tracers in the PP matrix. The extrusion temperature, the screw speed and the residence time were 205 °C, 120 rpm and 15 min, respectively. The screw and temperature profiles used in this study are supplied in Fig. 1.

For PP containing 0, 0.1 and 1 wt% of tracers (T), the materials will be designed as PP, PP_T_0.1 and PP_T_1, respectively. For the samples subjected to UV irradiation treatment, they will be referred to as UV-PP, UV-PP_T_0.1 and UV-PP_T_1, respectively.

All samples were kept for at least 1 month at room temperature before undergoing UV exposure in order to ensure that they had reached a near-equilibrium state, following ageing due to post-moulding effects such as secondary crystallization.

2.2. Microscopic observation

The tracers and their dispersion in the PP matrix were observed by Scanning Electron Microscopy (SEM) using a Hitachi S3500N model. The specimens prepared by injection moulding were fractured in liquid nitrogen and sputter-coated with gold before being examined with the microscope at an accelerating voltage of 30 kV. The SEM images were analysed by MATLAB by applying image processing.

2.3. UV irradiation procedure

UV irradiation treatment for the injection moulded specimens were carried out using a QUV accelerated weathering machine (QUV Solar Eye), with a light intensity of 0.68 W/m². The source of UV radiation was fluorescent tubes UVA-340, which have an output matching reasonably close to the solar radiation in the UV range [16]. The surface of the samples was subjected to a cycle of 4 h of UV irradiation at 60 °C and 4 h of condensation at 50 °C. The total exposure time of specimens was 2160 h (3 months).

Table 1
Materials specifications.

<table>
<thead>
<tr>
<th>Materials</th>
<th>Chemical name</th>
<th>Supplier</th>
<th>Particles’ size (μm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PP</td>
<td>Polypropylene</td>
<td>REPSON</td>
<td>2000</td>
</tr>
<tr>
<td>Y₂O₃</td>
<td>Yttrium oxide</td>
<td>RHODIA</td>
<td>2.250</td>
</tr>
<tr>
<td>CeO₂</td>
<td>Cerium oxide</td>
<td>RHODIA</td>
<td>2.250</td>
</tr>
<tr>
<td>Nd₂O₃</td>
<td>Neodymium oxide</td>
<td>AMPERE</td>
<td>3.500</td>
</tr>
<tr>
<td>Gd₂O₃</td>
<td>Gadolinium oxide</td>
<td>RHODIA</td>
<td>1.850</td>
</tr>
<tr>
<td>Dy₂O₃</td>
<td>Dysprosium oxide</td>
<td>RHODIA</td>
<td>2.250</td>
</tr>
<tr>
<td>Er₂O₃</td>
<td>Erbium oxide</td>
<td>AMPERE</td>
<td>8.200</td>
</tr>
<tr>
<td>Yb₂O₃</td>
<td>Ytterbium oxide</td>
<td>AMPERE</td>
<td>1.000</td>
</tr>
</tbody>
</table>

Fig. 1. Screw and temperature profiles for the extrusion processing.
2.4. Infrared spectroscopy

Samples used for Fourier transform infrared spectroscopy (FTIR), were collected from the surface to a depth of 0.5 mm and powdered. Approximately, 1–2 mg of the material powder was mixed and cold pressed with KBr to produce discs. FTIR measurements were obtained in transmission through the discs with a Nicolet Magna-IR 550 spectrometer. The equipment operates with a resolution of 2 cm\(^{-1}\) at a transmission mode using 16 scans.

2.5. Differential scanning calorimetry (DSC) characterization

Differential scanning calorimetry characterization (DSC Q10, TA Instruments) was performed to investigate the crystallization and the melting behaviours of the traced material. Thin slices of samples about 0.5 mm thick and 5–10 mg weight were cut from the surfaces of the specimens for the measurement. Thermograms were recorded in three consecutive runs: (1) a first heating, from room temperature to 210 °C, followed by (2) cooling, from 210 to 25 °C after holding the sample at 210 °C for 5 min, and finally (3) a second heating, from 25 to 210 °C. All experiments were performed at heating/cooling rate of 10 °C/min under argon atmosphere to avoid thermal degradation. The degree of crystallization, \(\chi_C\), was calculated considering a melting enthalpy of 209 J/g for a 100% crystalline polypropylene [17].

2.6. Thermal stability analysis

A TA Instruments Q500 Thermogravimetric Analyzer was used to characterize the thermal stability of specimens before and after UV irradiation. Measurements were conducted under an argon flow rate of 90 mL/min at a heating rate of 20 °C/min. The scanning temperature ranged from 25 to 500 °C. Similar to the DSC characterizations, thin slices of samples about 0.5 mm thick and 10–15 mg weight were cut from specimens for the measurement. The temperature at 5% weight loss (\(T_{5}\)) and the temperature of maximum decomposition rate (\(T_{\text{max}}\)) were calculated for each traced PP sample.

2.7. Mechanical characterization

Mechanical testing was carried out on specimens, prepared according to ISO 527-1:1993, using an Instron 4469 uniaxial tensile testing machine. Yield stress and elongation at break were measured at ambient temperature for a cross-head speed of 250 mm/min and the values reported represent an average of six measurements.

3. Results and discussions

3.1. Characterization of the dispersion of tracers

The homogenous dispersion of the filler particles in the PP matrix is of prime importance since it may strongly affect their detection and the mechanical properties [18,19]. Therefore, a quantitative characterization of the dispersion status of the
tracers is important for understanding their distribution in the PP matrix. Image analysis may be a powerful tool for describing quantitatively the dispersion of tracers, but this technique is subtle and delicate to use. The quality of results is highly dependent on the user’s knowledge of this technique. In particular, the setting parameters are able to create artefacts or erase a part of the information and by this way bias the results. Consequently, the processed images should show a high contrast between the objects (rare earth oxides) and their environment (PP matrix). The image processing must be subject to an easy programmable criterion uniformly applicable to all images in question.

In our case, the objective is to extract the rare earth oxides particles from a polymer matrix by SEM. The use of back scattered electrons simplifies the identification of tracers since the atomic number of rare earth oxides is higher than the elements composing the polymer matrix. This difference in the atomic weight contrast makes them appear like white dots on a black background.

In order to have a global view of the distribution of tracers, the samples have to be fractured so as to visualize the particles in material’s depth. This consideration causes two main problems. First, the slice of polymer material at ambient temperature makes them melt in front of the blade and second, as rare earth oxides are rigid particles, they break in multiple parts under blade pressure by modifying their geometrical properties.

To avoid these problems so as to conserve charge and matrix geometry, the cryofracture method was chosen for sample preparation. However, this method creates a relief in the SEM images which may interfere with the detection of rare earth oxides. Likewise, as non-specific interactions between charge and matrix are known, the fracture will propagate preferentially from charge to charge by probably inducing an over evaluation of charge population.

The SEM image of PP matrix containing 1% of gadolinium oxide is given in Fig. 2-A. Even if the image shows a good contrast between charge and matrix, the relief creates illumination variation which makes the charge extraction from the matrix complicated.

For this reason, it is necessary to obtain an image in which each pixel attributed to a charge will get the value “1” and each pixel attributed to the matrix will get the value “0”. One way to obtain this binarisation is by clipping. Threshold value is made by minimising environmental residue while maximising charge visibility. The choice of this value is a very important parameter and is taken by comparing the original SEM image to the proceeded one.

Fig. 2-B shows the thresholding image for a filter of 37% of its total dynamic. In the dark part of the image it can be observed a loss of “white dots”, compared to the original SEM image, which may bias the distribution results.

One of the charge properties not used in the previous thresholding process is their small size compared to the relief. This property can be used by applying a classic image processing named “unsharpening”, consisting in blurring a copy of the image, mostly with a convolution by a Gaussian function, and subtract it from the original image with the aim of only saving the fine details. As it can be seen in Fig. 2-C, the relief is mitigated while the charge remains as white as before the processing.

After the “unsharpening” process the image of Fig. 2-C was also proceeded to a clipping and the obtained image is shown in Fig. 2-D. A point to point comparison between the original image and the processed one shows that quality has been improved. Thus the charge surface on the image passed from 0.26% for the simple

![Fig. 4. Cubic and hexagonal close packing arrangement representation.](image-url)

![Fig. 5. Number average distribution of particles’ diameter for the PP\_Gd\(_2\)O\(_3\)_1 sample (A) Surface average distribution of particles’ diameter for the PP\_Gd\(_2\)O\(_3\)_1 sample (B).](image-url)
clipping to 0.38% for the unsharped thresholding by increasing to 36% the charge gain.

Once the image has been binarised, the coordinates of the constitutive points of the charge can be extracted. From this data, multiple parameters like gravity centre position of each particle and their size can be obtained. The gravity centre position can be used for characterizing the dispersion of the charge in the matrix. The mean distance from a particle to all the others can be affected by boundary effects and the minimal distance to the first neighbours is extremely sensitive to object fragmentation. In order to minimize boundary and local effects the mean distance to the first ten neighbouring particles was chosen as a representative value.

Fig. 3 presents the curve of spatial dispersion of the first ten neighbouring particles distances for the PP_Gd2O3_1 sample. The noisy spectrum (raw curve) corresponds to the number of counted particles. This number is a consequence of the sampling and the scale chosen for the image. The image sampling should be chosen between 1/10 and 1/100 of the inter-particle distance in order to have a reasonable sampling noise on each measure. Thus, as the image has a characteristic scale of 1000 × 1000 pixel, the number of particles will be about 100–1000 by image (in our case, 1421 objects in a 2560 × 1920 pixels image), and as they are represented on a scale of 1000 step, most of them are visible individually.

For comparing the results given by image processing, Fig. 3 shows the mean distance to the ten proximal neighbouring particles in a cubic and hexagonal close packing arrangement of the same surface and the same number of particles as in the case of the experimental data. As it can be seen in Fig. 4 the hexagonal network is more isotropic at low scale than the cubic one and this is the reason why the mean distance appears shorter.

Fig. 5-A, B gives the results of the particles diameter dispersion in number and in surface, respectively. As it can be seen, the distribution of the particles diameter in number follows a decreasing exponential curve, which can be explained by the fact that during the extrusion process particles are subject to brittle in smaller pieces. However, the distribution of particles diameter in surface is represented by a linearly decreasing curve showing the importance of each population.

The results obtained for the distance of the first ten neighbouring spheres (d calculated) calculated for the SEM data and the theoretical distance for a cubic (dth_cubic) and hexagonal (dth_hexagonal) close packing arrangement, together with average diameter in number (DNumber) and average diameter in surface (DSurface) are given in Fig. 6.

For characterizing the rare earth oxides dispersion in the PP matrix, the mean distance of the first ten neighbouring spheres calculated for the SEM data (d calculated) was compared with the theoretical distance for a cubic (dth_cubic) or hexagonal (dth_hexagonal) close packing arrangement. As d calculated is ranged between the cubic and hexagonal theoretical values this implies that the dispersion isotropy is quite good and homogenous. Moreover the SEM image of Fig. 2-A leads to the same conclusion.

### 3.2. Stability under UV irradiation/condensation cycles

The main products of polypropylene degradation, carbonyls, are easily observed in the wavelength range [1700–1800 cm⁻¹] by
giving a fairly broad peak as shown in Fig. 7 [20]. In addition, for a more quantitative characterization of the PP molecular degradation, the carbonyl index, corresponding to the areas under the carbonyl peak divided by the area of a reference peak was calculated, the carbonyl index, corresponding to the areas under the absorption band (in the range [1700–1800 cm⁻¹]) between the exposed samples and the unexposed unfilled PP, it is clearly visible that it increases after the UV irradiation treatment. From the FTIR

![Table 2](Image)

<table>
<thead>
<tr>
<th>Sample</th>
<th>ΔH_f (J/g gab)</th>
<th>T_m (°C)</th>
<th>ΔH_c (J/g gab)</th>
<th>T_c (°C)</th>
<th>χ_c (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PP</td>
<td>110 ± 3</td>
<td>167.0 ± 0.5</td>
<td>100 ± 3</td>
<td>105.0 ± 0.5</td>
<td>53 ± 1</td>
</tr>
<tr>
<td>PP_Y2O3_0.1</td>
<td>113 ± 3</td>
<td>166.5 ± 0.5</td>
<td>103 ± 3</td>
<td>106.9 ± 0.5</td>
<td>54 ± 1</td>
</tr>
<tr>
<td>PP_CeO2_1</td>
<td>110 ± 3</td>
<td>166.1 ± 0.5</td>
<td>100 ± 3</td>
<td>108.1 ± 0.5</td>
<td>52 ± 1</td>
</tr>
<tr>
<td>PP_NdO3_0.1</td>
<td>108 ± 3</td>
<td>167.1 ± 0.5</td>
<td>99 ± 3</td>
<td>106.3 ± 0.5</td>
<td>52 ± 1</td>
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<tr>
<td>PP_GdO3_1</td>
<td>109 ± 3</td>
<td>166.1 ± 0.5</td>
<td>99 ± 3</td>
<td>108.4 ± 0.5</td>
<td>52 ± 1</td>
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<tr>
<td>PP_Dy2O3_0.1</td>
<td>109 ± 3</td>
<td>167.0 ± 0.5</td>
<td>99 ± 3</td>
<td>105.1 ± 0.5</td>
<td>52 ± 1</td>
</tr>
<tr>
<td>PP_Er2O3_0.1</td>
<td>107 ± 3</td>
<td>166.6 ± 0.5</td>
<td>96 ± 3</td>
<td>105.5 ± 0.5</td>
<td>51 ± 1</td>
</tr>
<tr>
<td>PP_Yb2O3_0.1</td>
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<td>168.3 ± 0.5</td>
<td>101 ± 3</td>
<td>109.5 ± 0.5</td>
<td>53 ± 1</td>
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<td>168.8 ± 0.5</td>
<td>99 ± 3</td>
<td>106.1 ± 0.5</td>
<td>51 ± 1</td>
</tr>
</tbody>
</table>

![Fig. 9. Second heating run DSC thermograms of the PP samples of 1 wt% tracer content exposed to UV irradiation compared to the unexposed unfilled PP.](Image)

![Fig. 10. TGA curves of the PP samples of 1 wt% tracer, before (A) and after (B) UV irradiation exposure.](Image)

![Table 3](Image)

<table>
<thead>
<tr>
<th>Sample</th>
<th>ΔH_f (J/g gab)</th>
<th>T_m (°C)</th>
<th>ΔH_c (J/g gab)</th>
<th>T_c (°C)</th>
<th>χ_c (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>UV-PP</td>
<td>97 ± 3</td>
<td>137.6 ± 0.5</td>
<td>89 ± 3</td>
<td>100.3 ± 0.5</td>
<td>47 ± 1</td>
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<tr>
<td>UV-PP_Y2O3_0.1</td>
<td>98 ± 3</td>
<td>139.8 ± 0.5</td>
<td>86 ± 3</td>
<td>100.8 ± 0.5</td>
<td>47 ± 1</td>
</tr>
<tr>
<td>UV-PP_CeO2_1</td>
<td>93 ± 3</td>
<td>144.8 ± 0.5</td>
<td>98 ± 3</td>
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<td>45 ± 1</td>
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<td>UV-PP_NdO3_0.1</td>
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<td>139.1 ± 0.5</td>
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<td>100.0 ± 0.5</td>
<td>48 ± 1</td>
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<tr>
<td>UV-PP_GdO3_1</td>
<td>107 ± 3</td>
<td>167.2 ± 0.5</td>
<td>102 ± 3</td>
<td>108.3 ± 0.5</td>
<td>53 ± 1</td>
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<td>102 ± 3</td>
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<td>100.4 ± 0.5</td>
<td>49 ± 1</td>
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<td>139.6 ± 0.5</td>
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<td>51 ± 1</td>
</tr>
<tr>
<td>UV-PP_Yb2O3_0.1</td>
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<td>138.4 ± 0.5</td>
<td>92 ± 3</td>
<td>102.1 ± 0.5</td>
<td>44 ± 1</td>
</tr>
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<td>UV-PP_Y2O3_1</td>
<td>96 ± 3</td>
<td>138.2 ± 0.5</td>
<td>97 ± 3</td>
<td>101.6 ± 0.5</td>
<td>46 ± 1</td>
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<tr>
<td>UV-PP_CeO2_1</td>
<td>93 ± 3</td>
<td>164.8 ± 0.5</td>
<td>99 ± 3</td>
<td>107.2 ± 0.5</td>
<td>44 ± 1</td>
</tr>
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<td>UV-PP_NdO3_1</td>
<td>95 ± 3</td>
<td>141.1 ± 0.5</td>
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<td>46 ± 1</td>
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<td>UV-PP_GdO3_1</td>
<td>99 ± 3</td>
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<td>UV-PP_Er2O3_1</td>
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<td>UV-PP_Yb2O3_1</td>
<td>104 ± 3</td>
<td>141.5 ± 0.5</td>
<td>91 ± 3</td>
<td>100.3 ± 0.5</td>
<td>50 ± 1</td>
</tr>
</tbody>
</table>

![literature for PP [20–23], as 840, 1166, 1455 and 2720 cm⁻¹. The peak at 1455 cm⁻¹ is overlapped with several others [24], whereas 840 and 1166 cm⁻¹ are sensitive to PP crystallinity [25]. Consequently, the peak at 2720 cm⁻¹, associated with CH3 stretching and CH bending, was chosen as the most appropriate [20,26]. By comparing in Fig. 7, the area evolution of the carbonyl peak absorption band (in the range [1700–1800 cm⁻¹]) between the exposed samples and the unexposed unfilled PP, it is clearly visible that it increases after the UV irradiation treatment. From the FTIR
The results obtained for the carbonyl index show that the carbonyl peak intensities decrease slightly with tracer content, except the CeO$_2$ for which the carbonyl index decreases drastically compared to the UV-PP. Cerium oxide is well known to be an UV absorber\cite{13}, and these results show that it plays an important role in PP stabilization by delaying the photo-degradation process. The CeO$_2$ particles act as optical filters by absorbing part of the UV irradiation and releasing the excess energy as heat, thus the UV intensity which can promote the oxidation of the PP chains is reduced. The same explication can be given for the Y$_2$O$_3$ particles which absorb the beginning of the UV irradiation at 300 cm$^{-1}$ and this is the reason why its carbonyl index has a more important diminution compared to the other tracers.

By comparing the un filled PP sample with the tracer filled PP samples after the UV irradiation treatment, no new peaks are observed, suggesting that their photo-degradation mechanisms are identical. Similar photo-stabilization effect of metal oxides has also been reported by Zhao et al., Tessier et al. and Sato et al. [12,13,26].

Regarding the remaining tracers (Nd$_2$O$_3$, Gd$_2$O$_3$, Dy$_2$O$_3$, Er$_2$O$_3$ and Yb$_2$O$_3$) their effect on the photo-degradation of PP doesn’t seem to play an important role.

3.3. Effect of tracers on thermal and crystallization behaviours

The thermal and crystallization data obtained from DSC for the un filled PP and the traced polymer composites before UV irradiation exposure are summarized in Table 2. As it can be reflected by $\Delta H_f$, $T_m$, $\Delta H_c$, and $T_c$, the melting and crystallization behaviours of the PP matrix are slightly affected by the tracer addition before UV irradiation treatment. Regarding the PP crystallinity ($\chi_c$) the values are so close for considering a variation with the tracer addition. Even if it was expected that the rare earth oxides will act as nucleating agents \cite{8-10} in polymer crystallization by increasing the crystallinity, their micrometric size seems to prevent them for having a real impact. In fact the average particle size of the rare earth oxides studied by Liu et al.\cite{8,9} was in the range $[0.05-0.6 \mu m]$, in contrary to the particles used in this study that was superior to 1 $\mu m$.

By comparing the data in Tables 2 and 3 and the DSC thermograms displayed in Fig. 9, it can be seen that UV irradiation exposure has a real impact on both melting and crystallization behaviours of the PP matrix. The melting temperature of the un filled PP dropped from 166.7 to 137.6 $^\circ$C after 3 months of UV irradiation. This decrease of about 30 $^\circ$C, is observed for almost all the traced samples, except the one containing CeO$_2$ at 1 wt% (PP_CeO$_2$-1). The chain scission caused by photo-degradation can release the entanglement of molecules and produce more freed segments, thus the recrystallised material (Fig. 9) contains molecules that are smaller and defective due to groups like carbonyls.
Consequently, the reduction of molecular weight leads to the decrease of the melting temperature.

3.4. Effect of tracers on thermal stability

The addition of rare earth oxides in the PP matrix could improve the thermal stability and onset degradation [10]. The TGA measurement results before and after the UV irradiation treatment, are shown in Fig. 10 and Table 4. From the TGA curves of Fig. 10, it is obvious that the thermal decomposition for all the samples is a one-step process. PP thermally degrades to volatile products above 250 °C through a radical chain reaction process propagated by carbon centred radicals due to the lack of oxygen. The formed radicals initiate and propagate the subsequent radical chain reactions [26,27].

In Fig. 10-A, it can be seen that the TGA curves of the samples containing the tracers are nearly overlapping with the un filled PP. By comparing the results before the UV irradiation exposure in Table 4, the temperatures of 5% weight loss ($T_{5\%}$) and of maximum decomposition rate ($T_{max}$) for the traced materials appear to be superior to 1−5 °C than the un filled PP. This means that rare earth oxides could provide thermal stability to the PP matrix, even though better results could be obtained by increasing their content [10].

Fig. 10-B shows that the majority of TGA curves of the UV irradiation exposed traced samples are close to the un filled UV-PP, except the UV-PP_CeO2_1 sample for which the TGA curve is closer to the unexposed un filled PP. Moreover, by comparing the results of $T_{5\%}$ for the UV irradiation exposed samples with the unexposed un filled PP sample, it appears that it drops dramatically of about 125 °C. On the contrary, for the samples containing the CeO2 this diminution is of about 70 °C and 25 °C for the 0.1 and 1 wt% tracer content, respectively. The positive thermal stability provided by cerium oxide can be explained by the protective effect upon UV irradiation due to the UV screening properties of CeO2 particles.

3.5. Effect of tracers on mechanical properties

In previous work [2], it has been shown that the addition of 1 wt % of Nd2O3 and Gd2O3 leads to a slight increase of the Young's modulus and a slight decrease of the yield stress. Regarding the tensile tests, the most important effect of these rare earth oxides was observed on the elongation at break. Fig. 11 presents the results of elongation at break for all the tracers used in this study before UV irradiation exposure.

Predictably, the obtained results reveal that the elongation at break drops upon increasing the filler fraction [28−30]. However, at 0.1 wt% tracer content, the elongation at break slightly decreases from 5 to 25% with a standard deviation of about 5−15%. The obtained values are close to the standard deviation of measurements and hence it can be considered that the influence of rare earth oxides in mechanical properties is minor at 0.1 wt%.

As it can be seen in Fig. 12, the ductility of the tensile specimens was reduced considerably after UV irradiation treatment. Indeed, the majority of the samples, except UV-PP_CeO2_0.1 and UV-PP_CeO2_1, were brittle and it was difficult to carry out the tensile tests. Fig. 13 presents the results of elongation at break obtained for the samples submitted to 10 days UV irradiation treatment. The tensile tests results of UV-PP_CeO2_0.1 and UV-PP_CeO2_1 for 10 days and 3 months UV irradiation exposure are displayed in Fig. 14.

Fig. 13 shows that elongation at break decreases significantly from about 100% to 20% for the samples exposed to UV irradiation. However, for the majority of samples containing the rare earth oxides at 0.1 and 1 wt% the values of elongation at break are similar to the un filled exposed PP. The unique exception of the traced samples are UV-PP_CeO2_0.1 and UV-PP_CeO2_1 for which elongation at break and yield stress (Fig. 14) increase with CeO2 particle content.

Based on the mechanical tests results, it can be summarized that Y2O3, Nd2O3, Gd2O3, Dy2O3, Er2O3 and Yb2O3 of micrometric mean particle size, have a minor effect on the photo-degradation of the PP matrix at 0.1 wt%. On the contrary the addition of 1 wt% of CeO2 can act as an effective UV light screen for reducing the degree of photo-degradation of the PP matrix.

4. Conclusions

Results from this work reveal that the addition of rare earth oxides of micrometric particle size at 0.1 wt% content in the PP matrix has a minor effect on the mechanical and thermal properties of the traced materials as well as on the photo-degradation of the polymer after UV irradiation exposure.

However, the addition of 1 wt% of rare earth oxides particles into the PP matrix decreases the elongation at break by 10−50% before UV irradiation treatment for a cross-head speed of 250 mm/min. For the same concentration, the melting and crystallization behaviours are slightly increased, whereas the thermal stability is improved, increasing the temperatures of 5% weight loss and maximum decomposition rate by 1−5 °C.

Regarding the UV irradiation treatment, the results of FTIR, DSC and ATR analysis together with mechanical characterization, pointed out that the addition of CeO2 particles in the PP matrix, which is believed to contain low quantities of UV stabilizers, can lead to significant improvements on the photo-degradation resistance of PP to UV exposure. Indeed, the CeO2 particles act as UV filters by absorbing part of the UV irradiation and thus reducing the UV intensity which can promote the oxidation of the PP chains. For the remaining tracers, the results obtained show the same behaviour on the photo-degradation as in the case of the un filled PP matrix.

The dispersion effect of tracers in the PP matrix was quantitatively characterized by applying image processing to the SEM images with the help of MATLAB. The distance to the first ten neighbouring spheres calculated for the SEM data was compared.
with the theoretical distance for a cubic and a hexagonal close packing arrangement of the same surface and the same number of particles as in the case of the experimental data. The SEM images and the results obtained from image processing show a homogeneous dispersion of tracers in the PP matrix.

In order to study the effect of rare earth oxides in the PP matrix, a polypropylene homopolymer free of fillers like talc and calcium carbonate and which is believed to contain only low quantities of UV stabilizers was chosen in this study. As it was shown, CeO₂ particles can act as UV absorbers by protecting the polymer matrix from photo-degradation. Complementary work can be carried out with other grades of PP, more adapted to outdoor environment applications, for studying probable antagonistic effects between the CeO₂ particles and the UV stabilizers already contained in the polymer matrix.

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References
