EFFECT OF VIRGIN HETERO PHASIC PP COPOLYMER CONTENT ON MOISTURE ABSORPTION, THERMAL AND MECHANICAL PROPERTIES OF RECYCLED POLYETHYLENE WOOD FLOUR COMPOSITES

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ABSTRACT

The effect of virgin heterophasic PP copolymer (vPP) content on moisture absorption, as well as on thermal and mechanical properties of recycled polyethylene/wood flour composites was investigated. The polymer matrix of recycled post-consumed plastic waste (rPE) was composed of a matrix of LDPE and a part of PP. Wood flour of Pinus radiata was used as filler at a constant loading of 45 wt.%. rPE/vPP blends and their composites were manufactured by melt blending, and then by injection molding. The morphology of the blends and composites was analyzed by means of scanning electron microscopy. The addition of vPP improved tensile and flexural moduli and flexural strength of wood plastic composites (WPC). The highest increase of these properties was observed for a WPC sample with a rPE/vPP ratio of 19. WPC made with higher virgin PP content (rPE/vPP ratio = 9 and 5.7) showed lower increase of mechanical properties compared to polymer matrix and WPCrPE samples. The moisture uptake of WPC made of rPE/vPP blend was higher than those of rPE, and their mechanical properties were more adversely affected by immersion in water. TGA results indicate that rPE are thermally more stable than rPE/vPP blends. The incorporation of heterophasic PP copolymer into the recycled polymer matrix of WPC delays the starting of wood flour degradation.

Keywords: recycled LDPE-PP blend, wood plastic composite, mechanical properties, water absorption properties

INTRODUCTION

The recycling of consumed plastics is an attractive alternative to their disposal in landfills, reducing waste streams and their impact on the environment, and the consumption of virgin plastics. Recovery of plastics in Chile was only around 27,000 tons in 2009, 4.3% of the total amount of plastics consumed.1,2 One of the Chilean industries leading in this area, is the aquacultures industry, with 87% recovery of plastics in 2010.3 Most of the time this recycled plastic is reprocessed by down-cycling in order to make new articles of lower quality.

In particular, recycled low density polyethylene alone presents poor mechanical properties, and it is often mixed with other materials to be useful, such as virgin polyolefin and/or additives as fillers.

The presence of propylene homopolymer (PP) in recycled polyethylene matrix is frequently found in plastic waste streams.4 Two of these polymers, although of similar chemical structure, are incompatible in the melting, and the blend exists as two distinct phases with very poor cohesion between them. The incompatibility of polyethylene/PP also contributes to the unsatisfactory mechanical properties of recycled polyethylene-PP (rPE-PP) blends. Therefore, compatibilizers are added during the reprocessing of rPE-PP blends, the most common are elastomeric block and random copolymers and terpolymers, such as ethylene-propylene and ethylene-propylene-diene elastomers.5,6 Graft copolymers have also been used as compatibilizers of rPE-PP blends. For example, by addition of PE-g-(2-methyl-1,3-butadiene) to blends composed of 90 wt.% of rPE and 10 wt.% of recycled PP, a common composition of post-consumer plastic waste, the elongation at break and Charpy impact strength were improved.7 However, heterogeneity of the plastic mixture and the different degree of polymer degradation makes compatibilization of recycled plastics a difficult task. On the other hand, the selection of suitable compatibilizers for industrial applications is limited by economic constraints.

Mechanical properties of rPE-PP blends are improved by the addition of virgin polyolefin resin. Blom et al.8 reported that blends of virgin isotactic PP with a post-consumer plastic, consisting of an rPE-PP blend of (1 part of PP and 2 parts polyethylene), displayed higher impact strength than neat isotactic PP, as well as neat post-consumer resin. This improvement on impact strength was not observed for virgin isotactic PP/high density polyethylene blends.

We have reported the preparation of wood plastic composites (WPC) from recycled plastic composed of a low density polyethylene matrix and a part of PP, and Pinus radiata wood flour.8 Small amounts of virgin homopolymer PP (5, 10 and 15 wt.%) were used to improve the strength and processability of the melting. The addition of virgin homopolymer PP improved mechanical properties and reduced moisture absorption and swelling of the WPC manufactured from recycled plastic/virgin PP. Virgin homopolymer PP in WPC acted also as a reinforcing agent.

In this article, a commercial heterophasic PP copolymer with 12 wt.% of ethylene was incorporated into the recycled plastic WPC formulation to study how this affects the mechanical and thermal properties, as well as the water uptake behavior of these WPC. In particular, the use of this copolymer would not increase considerably the cost of this kind of WPC of rather low market-value, and could act as reinforcing agent and compatibilizer among recycled polyolefin, interacting with the PP part present in the rPE matrix.8

EXPERIMENTAL

Materials

The post-consumer plastic waste, composed of a matrix of LDPE and a part of PP (MFI = 0.85 g/10 min at 230°C - 2.16 kg), was obtained mainly from packaging of the salmon farming industry, and was used as the main matrix, referred to hereinafter as rPE. The lignocellulosic filler was Pinus radiata D. Don wood flour with less than 2% humidity. Maleic anhydride grafted polypropylene with an acid value of 43 mg KOH/g and MFI of 10,000 g/mol (Liocene PP MA 6542 from Clariant) was used to enhance the wood flour/polymer matrix adhesion. As antioxidants Irgafos 168 and Irganox 1010 from Ciba (BASEF) were used in a 2:1 ratio. A masterbatch of virgin LDPE and 20 wt.% antioxidants was prepared to be added to tested formulations. A commercial heterophasic PP copolymer, PCD 1214, from Petroquim S.A. (Chile) with 12 wt.% of ethylene comonomer content and MFI of 12 g/10 min was employed as reinforcing agent for recycled polyolefin blends and WPC.

Preparation of composites

WPCs and blends were prepared in a co-rotating twin-screw extruder (TSA Industriale, Cernobbio, Italy). The barrel temperature of seven zones varied from 170°C (feeder) to 175°C, 175°C, 175°C, 175°C, 180°C and 185°C (heating zones) and the die temperature was above 190°C. The extruder screw speed was set at 80 rpm and average feed rate was 20 kg/h. More details of sample preparation were described elsewhere.9 The composition of the tested WPCs and blends, is summarized in Table I.

Before molding, pellets of tested samples were dried over night at 100°C. Bars (0.34 x 1.25 x 12.67 cm) and dumbbell tensile specimens were injection molded using Arbarg 420C machine (Lossburg, Germany) for further characterization. The cylinder temperature was maintained constant at 190°C.

In order to determine the compatibility blends of rPE and rPE/vPP blends with different heterophasic PP copolymer content (see Table 1) were previously melt mixed in a batch mixer. The mixing speed, temperature and residence time were 60 rpm, 190°C and 10 min, respectively. Then, these samples were compression molded at 190°C for 4 min, at a compressive load of 15 bars, in order to prepare films (10 x 10 x 1 mm) for their respective SEM characterization.

References

Table I. Sample code and compositions of tested materials.

<table>
<thead>
<tr>
<th>Sample code</th>
<th>Polymer matrix (wt.%)</th>
<th>rPE/vPP ratio</th>
<th>WF (wt.%)</th>
<th>Licocene (wt.%)</th>
<th>Antioxidant masterbatch (wt.%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>rPE</td>
<td>98.0</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>2.0</td>
</tr>
<tr>
<td>rPE/vPP-19</td>
<td>93.2</td>
<td>4.9</td>
<td>19.0</td>
<td>-</td>
<td>1.9</td>
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<tr>
<td>rPE/vPP-9</td>
<td>88.4</td>
<td>9.8</td>
<td>9.0</td>
<td>-</td>
<td>1.8</td>
</tr>
<tr>
<td>rPE/vPP-5.7</td>
<td>83.6</td>
<td>14.7</td>
<td>5.7</td>
<td>-</td>
<td>1.7</td>
</tr>
<tr>
<td>WrPE</td>
<td>53.1</td>
<td>-</td>
<td>45.0</td>
<td>1.0</td>
<td></td>
</tr>
<tr>
<td>WrPE/vPP-19</td>
<td>50.5</td>
<td>2.7</td>
<td>45.0</td>
<td>0.9</td>
<td>1.0</td>
</tr>
<tr>
<td>WrPE/vPP-9</td>
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<td>5.3</td>
<td>45.0</td>
<td>0.9</td>
<td></td>
</tr>
<tr>
<td>WrPE/vPP-5.7</td>
<td>45.2</td>
<td>8.0</td>
<td>45.0</td>
<td>0.9</td>
<td>0.9</td>
</tr>
</tbody>
</table>

Characterization

Mechanical testing

Tensile and flexural tests were performed on a Karg Industrietechnik machine (Krailling, Germany) according to ASTM D638/97 and D790/97 standards, respectively. Prior testing, specimens were conditioned for 40h at 23°C and 50% relative humidity. The flexural measurements were performed under a three-point bending configuration. The crosshead speed was set at 1.27 and 50 mm/min for the flexural and tensile tests, respectively. The Young’s modulus, strength, and elongation percentage at break were obtained from the stress-strain curves. At least 18 individual measurements were carried out for each formulation and mean values and standard deviations of tensile and flexural properties were reported. Collected data were evaluated with a one-way analysis of variance at the 95% confidence level. Differences among the mean values of variables were analyzed with Duncan’s multiple-range test. The factors and levels used in the experiments are summarized in Table II. The statistical analyses were made with Statgraphics Centurion XV software, Statistical Graphics, United States.

Table II. Factor and level used in the experimental design.

<table>
<thead>
<tr>
<th>Factor</th>
<th>Level</th>
<th>I</th>
<th>II</th>
<th>III</th>
<th>IV</th>
</tr>
</thead>
<tbody>
<tr>
<td>rPE/vPP ratio</td>
<td></td>
<td>19.0</td>
<td>9.0</td>
<td>5.7</td>
<td>Without vPP</td>
</tr>
<tr>
<td>Wood flour</td>
<td>yes</td>
<td>no</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Water immersion</td>
<td>before</td>
<td>after</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

Morphological analysis of fractured surfaces

The fractured surfaces of the mechanical test specimens and films obtained by compression molded (sputtered with a gold coating of ca. 50 nm) were observed at different magnifications, ranging from 100 to 3000 with a field emission scanning electron microscope (JEOL JSM 6380 LV, Tokyo, Japan) operated at 20 and 5 kV.

Moisture uptake

Specimens type bars (0.34 x 1.25 x 12.67 cm) obtained by injection molding were used in long-term immersion tests. Water absorption was determined according to ASTM D570-98. Ten specimens of each formulation were selected and dried for 24h at 50°C. The specimens were placed in distilled water and maintained at 23°C for 98 days. For each measurement, specimens were taken from the water and surface moisture was removed with absorbing paper. The sample weight was measured with 0.001 g precision at different times during time immersion. Specimens were considered substantial when the increase in weight was less than 1% in three consecutive measurements. The percentage of water absorption was calculated according to the following expression:

Water absorption (%) = \( \frac{\text{wet weight (g)} - \text{ooven-dry weight (g)}}{\text{ooven-dry weight (g)}} \times 100 \)

Thermal analysis

Thermal stability of blends and WPCs was evaluated using a thermogravimetric analyzer NETZSCH 209 F3 TGA (Tarsus, Selb, Germany). Thermogravimetric (TGA) curves were recorded at 10°C/min, in a range between 30 and 640°C, under nitrogen atmosphere (20 mL/min).

RESULTS AND DISCUSSION

Exploratory studies to evaluate the compatibility of rPE/vPP blends with different heterophasic PP copolymer content were performed. Cryogenically fractured surfaces were prepared by means of compression-molding of samples and observed by SEM (see Figure 1). Micrographs of the fractured surface of rPE revealed a co-continuous morphology with droplets spread out in the rPE matrix. The dispersed droplets corresponded to recycled PP, the minority phase. rPE/vPP blends also displayed phase separation, but the microstructural phase separation of rPE/vPP blends with rPE/vPP ratio of 19 was finer and of better homogeneity than neat rPE. In addition, the size of virgin PP domains were more uniform for blends with 4.9 wt.% of PP content (sample: rPE/vPP-19). Similar morphology has been reported by other authors for rPE-PP blends of 80/20 and 90/10 (PE/PP) with circa 5 wt.% of a compatibilizing agents, such as ethylene-propylene-diene monomer and ethylene-propylene monomer. In general, elastomers used as compatibilizer for polyolein PE-rich blends have a high content of ethylene monomer.

The domain size of virgin PP, the dispersed component, was larger for those rPE/vPP blends with rPE/vPP ratio = 5.7, indicating a decrease in the compatibility by increasing content of virgin PP in the blend. This may be partly due to the self-association, which is more favorable than hetero-association when the content of virgin heterophasic PP copolymer is higher than 4.9 wt.%. It is important to note that there was no difference in the morphology of compression-molded and injection-molded specimens.

![Figure 1: SEM images of fractured surface of neat rPE and rPE/vPP blend samples prepared by compression molding: (a) rPE, (b) rPE/vPP-19, (c) rPE/vPP-9 and rPE/vPP-5.7. (Scale bar: 5 µm)](image_url)
without polymer matrix coating. As expected, an increasing content of virgin PP in the WPC displayed a lower amount of holes in fractured surfaces. The virgin PP in the composite favors the diffusion of the exposed PP chain of the coupling agent into the polyolefin matrix phase of mainly recycled and virgin PP chains, which contributes to the improvement of interfacial adhesion. There was no visible morphological difference of surfaced fracture of the WPC specimens before and after a long term immersion test (data not shown).

Figure 2: SEM images of fractured surface of WPCs after 2352 h of immersion in water: (a) WrPE, (b) WrPE/vPP-19, (c) WrPE/vPP-9 and (d) WrPE/vPP-5.7. (Scale bar: 20 µm).

Table III: Thermal properties of neat rPE, rPE/vPP blends and their WPCs.

<table>
<thead>
<tr>
<th>Sample</th>
<th>$T_{5%}$</th>
<th>$T_{\text{max}}$</th>
<th>$W_{L_{\text{Tmax}}}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>rPE</td>
<td>425</td>
<td>469</td>
<td>66.3</td>
</tr>
<tr>
<td>rPE/vPP-19</td>
<td>421</td>
<td>465</td>
<td>62.7</td>
</tr>
<tr>
<td>rPE/vPP-9</td>
<td>420</td>
<td>461</td>
<td>55.8</td>
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<tr>
<td>rPE/vPP-5.7</td>
<td>393</td>
<td>462</td>
<td>70.0</td>
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</table>

<table>
<thead>
<tr>
<th>Sample</th>
<th>$T_{\text{onset-1}}$</th>
<th>$T_{\text{max-1}}$</th>
<th>$W_{L_{\text{Tmax-1}}}$</th>
<th>$T_{\text{onset-2}}$</th>
<th>$T_{\text{max-2}}$</th>
<th>$W_{L_{\text{Tmax-2}}}$</th>
<th>$T_{\text{onset-3}}$</th>
<th>$T_{\text{max-3}}$</th>
<th>$W_{L_{\text{Tmax-3}}}$</th>
</tr>
</thead>
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<tr>
<td>WrPE</td>
<td>31</td>
<td>72</td>
<td>1.9</td>
<td>199</td>
<td>362</td>
<td>26.6</td>
<td>383</td>
<td>466</td>
<td>71.3</td>
</tr>
<tr>
<td>WrPE/vPP-19</td>
<td>32</td>
<td>71</td>
<td>1.7</td>
<td>203</td>
<td>352</td>
<td>28.4</td>
<td>385</td>
<td>466</td>
<td>71.2</td>
</tr>
<tr>
<td>WrPE/vPP-9</td>
<td>30</td>
<td>64</td>
<td>2.3</td>
<td>210</td>
<td>349</td>
<td>28.2</td>
<td>383</td>
<td>463</td>
<td>73.3</td>
</tr>
<tr>
<td>WrPE/vPP-5.7</td>
<td>31</td>
<td>68</td>
<td>2.2</td>
<td>213</td>
<td>349</td>
<td>26.5</td>
<td>385</td>
<td>464</td>
<td>69.2</td>
</tr>
</tbody>
</table>

$T_{\text{5\%}}$ Temperature at which 5% weight loss occurs in rPE and rPE/vPP blends samples.

$T_{\text{max}}$ Temperature at which maximum rate of decomposition occurs.

$W_{L_{\text{Tmax}}}$ Weight loss associated at maximum temperature of each decomposition step.

$T_{\text{onset}}$ The onset temperature corresponding to each decomposition step of WPC.

The weight loss during heating of WPCs occurred in three steps, as shown in Figure 3. Small weight loss (around 3%) between 30 - 120°C is attributed to the evaporation of water from wood fibers. After 200°C the weight loss rate quickly increased and two degradation steps were distinguished, the first one corresponds to the decomposition of wood fiber components such cellulose, hemicellulose, lignin and wood extractives. The maximum rate of decomposition temperature of the second mass-loss step ($T_{\text{onset-2}}$) was around 350°C. The third degradation step between 383 - 493°C is due to the decomposition of main chains of polyolefin matrix.

TGA curves of WPC samples in nitrogen atmosphere showed slight differences in the onset degradation temperatures corresponding to first ($T_{\text{onset-1}}$) and third ($T_{\text{onset-3}}$) decomposition steps; this did not change systematically by variation of virgin PP content into polymer matrix. However, the second onset degradation temperature ($T_{\text{onset-2}}$) was shifted to higher temperatures with increasing PP content in WPC samples. The differences in $T_{\text{onset-2}}$ of WPC indicate that virgin PP loading delays the starting of wood fibers degradations. The apparent thermal stability of wood fibers can be attributed to stronger entanglement between PP chains of coupling agent (MAPP) and PP of polymer matrix around the fiber, which results in a diffusion delay of the gaseous degradation products out of the composite.

Mechanical properties of neat rPE, rPE/vPP blends and their WPCs are shown in Figure 4. The incorporation of wood fiber (45 wt.%) into polymer matrix improved tensile and flexural properties of the studied materials, except for the elongation at breaking. Reinforcing effect of wood fibers is related to their intrinsic properties such as tensile strength, stiffness, chemical inert and dimensional stability. The improvement of tensile and flexural strengths as well as moduli of recycled PE by addition of wood fiber has been previously reported. The major issue in obtaining WPC with superior mechanical properties is an adequate interfacial adhesion between hydrophilic fiber and hydrophobic polymer matrix. In a previous work it was reported, that maleic anhydride modified PP with higher grafting capacity as coupling agent provided significant improvement in mechanical properties of rPE/wood flour composite. 

Thermal stability of neat rPE, rPE/vPP blends and their WPCs was analyzed by TGA. Figure 3 shows the TGA and derivative of the TG (DTG) sample curves in the temperature range of 20 - 640°C. Temperature at which maximum rate of decomposition occurs ($T_{\text{max}}$) and weight loss associated at this temperature ($W_{L_{\text{Tmax}}}$), as well as the onset temperature ($T_{\text{onset}}$) corresponding to each decomposition step of WPCs and temperature at which 5% of weight loss occurs for rPE and rPP/vPP blends, are summarized in Table III. The TGA curves reveal a one-step decomposition pattern for neat rPE and rPE/vPP blends. The neat rPE remains stable from room temperature up to 150°C, with no change of weight of the sample. After a temperature of 150°C the rate of decomposition was slow up to 415°C, with a weight loss of about 3%. Then, the sample started to decompose fast and T$_{\text{max}}$ was 469°C. A weight loss of 66% was recorded at this T$_{\text{max}}$. In an inert atmosphere, the thermal degradation of polyethylene is associated with random scission of main polymer chains as well as branching polymers. Both processes occur simultaneously resulting in a single mass loss step. 

The degradation of rPE/vPP blends followed a similar trend as neat rPE. T$_{\text{onset}}$ decreased with the increasing of virgin PP content from 4.9 to 14.7 wt.%. A similar behavior is observed for T$_{\text{max}}$. The results indicate that the fragmentation of rPE chains is slightly accelerated by the presence of virgin PP, especially blends with rPE/vPP ratio $= 5.7. This behavior is probably due to the higher susceptibility of PP to ultraviolet and oxidative degradation than PE.
The effect of rPE/vPP ratio on tensile and flexural properties before and after immersion in water of rPE/vPP blends and their WPCs was also analyzed. Before immersion, there was a statistical significant difference among values of tensile and flexural strengths, and flexion modulus for both sample groups: rPE/vPP blends and WPCs made with different rPE/vPP ratio. However, WPC sample made with rPE/vPP ratio = 19 showed the highest increase of tensile and flexural strengths and moduli in relation to polymer matrix (rPE) and WrPE sample. An increase of about 70% in tensile and flexural moduli and flexural strength of all WPC samples in relation to polymer matrix was obtained. We can conclude that the addition of small amounts of virgin PP (rPE/vPP = 19) to the rPE matrix of WPC improves mechanical properties due to heterophasic PP copolymer acting as a reinforcing agent. PP is a stronger but less flexible material than PE, because of a rigid shortly methyl group attached to every second carbon atom of the polymer main chain, which restricts rotation of the chain. Furthermore, virgin PP chain favors diffusion of the exposed PP chain of the coupling agent into the polyolefin matrix phase. Both factors, in turn, contributed to better mechanical properties of the WrPE/vPP-19 sample.

Phase separation of blend morphology is another factor to be taken into account in the analysis of mechanical properties of studied samples. Weak mechanical properties are associated to higher domain sizes of minor phase and poor interfacial adhesion between polymer constituents of the blends. Phase separation observed for rPE/vPP blends by SEM could explain why those samples of WPC with higher virgin PP content (WrPE/vPP-9 and WrPE/vPP-5.7) show lower increase on their mechanical properties.

The mechanical properties of samples after and before immersion displayed similar trends: WPC showed higher tensile and flexural moduli and strengths than polymer matrices (rPE and rPE/vPP blends). The mechanical properties of rPE and rPE/vPP blends almost were not affected by immersion, whereas mechanical properties of WPC samples were strongly affected by immersion. In general, the values of tensile and flexural properties of WPC samples significantly decreased after immersion. Thus, tensile modulus and strength decreased around 35% and 15% after immersion for WPC, respectively. Similar behavior has been reported by other authors. They postulate that water molecules absorbed by the wood flour not only change the structure and properties of fibers, but also the interface between the polymer matrix and the fibers. Furthermore, water uptake of WPC samples contributes to a loss of compatibilization between fibers and matrix, weakening the interface adhesion. As a result the stress transfer and the load-bearing ability of the composites were deteriorated. The loss in mechanical properties caused by water uptake was higher in WPC samples made with rPE/vPP blends as polymer matrix, due to their higher water uptake during long-term immersion tests (see Figure 5).
Water uptake of neat rPE, rPE/vPP blends and their WPCs as a function of the square root of immersion time is illustrated in Figure 5. For all samples, water uptake increased with immersion time up to a saturation point, from which the content of water in the composites remained constant. The water absorption of WPC samples is higher than those of rPE and rPE/vPP blends. The hydrophilic nature of wood flour is required for water uptake in WPCs.

Fibers of Pinus radiate are rich in cellulose and hemicellulose (around 60 wt.%), which have free OH-groups that come in contact with water through hydrogen bonding. The water-fiber interaction results in water uptake and weight gain in WPC samples. In addition, microgaps at the fiber and polyolefin matrix interface can contribute to water uptake of the composites because of increased porosity.22

The hydrophobic polymeric matrix (rPE or rPE/vPP blends) had no significant water uptake. rPE/vPP blends absorbed less water compared to rPE sample. The lowest water uptake was observed for the rPE/vPP-19 sample, which exhibits better interfacial adhesion between the rPE matrix and virgin PP, as discussed above. However, composites based on the rPE/vPP blends absorbed more water than the rPPE sample. The possible reason is the presence of interfacial voids between rPE and virgin PP, which allow water to reach the wood fibers. The swelling by water absorption of WPC can lead to microcracking with considerable decreasing of mechanical properties, as has been discussed above.25

CONCLUSION

The incorporation of a virgin heterophasic PP copolymer into the rPE matrix improves the mechanical properties and thermal stability of the composite. WPC made of rPE/vPP blend showed higher values of tensile and flexural moduli and flexural strength than WPC made of rPE. However, the improvement of mechanical properties depends on the rPE/vPP ratio. Composite manufactured with rPE/vPP ratio = 19 showed a higher increase of tensile and flexural strength and moduli with respect to neat rPE and composite made of rPE. Heterophasic PP copolymer acts as a reinforcing agent and favors diffusion of the exposed PP chain of the coupling agent anchored to wood flour into the polyolefin matrix phase. WPC manufactured with rPE/vPP ratio = 9 and 5.7 displayed a slight increase of their mechanical properties. This increment is associated to poor interfacial adhesion between polymer constituents of the blends, factor that is opposed to the reinforcement effects of virgin PP.

TGA results shows that the addition of virgin PP to the polyolefin matrix of WPC delays the starting of wood flour degradation. The apparent thermal stability of wood fibers is attributed to a stronger entanglement between PP chains of coupling agent (MAPP) and PP of polymer matrix around the fiber.

The immersion of WPCs in water adversely affects their mechanical properties, especially of those WPCs manufactured with a rPE/vPP polymer matrix. The presence of interfacial voids between rPE and virgin PP favors water entering into and swelling of the wood fibers.

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REFERENCES