Study of Thermo-Mechanical Properties of Polypropylene and Low Density Polyethylene Composite Material

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ABSTRACT

The goal of this research paper is preparation and characterization of the mechanical and thermal properties of polypropylene (PP) and linear low density polyethylene (LLDPE) base polymer that blend filled with talc and magnesium carbonates.

Key words: Polypropylene, Low density polyethylene, Magnesium carbonate, Talc, Thermo-mechanical properties.

INTRODUCTION

From ancient periods, the quest for new materials has been developed. PP applications are very useful in engineering materials for the automotive, electrical appliances and packaging manufacturing due to its excellent properties such as rigidity and stiffness. Oil resistance and thermal stability is usually confined due to its low impact strength and Young's modulus, particularly at low and high-temperature loading conditions. These PP impediments can be considerably enhanced by blending PP with other polymers. LLDPE is also similar polymer like PP. It is commonly used in automotive and packaging industries. The blending of two or more polymers is a cheaper and more efficient option, not only for the development of polymers with new properties but also for recycling of materials. It is possible to achieve improvement in properties such as modulus, yield stress, and tensile strength by an appropriate combination of polymer components. PP and LLDPE are deemed to be compatible in the liquid state.

PVC and ENR blend prepared using Brabender plasticorder by the melt flow behavior of ENR decreases the Brabender torque, increases the melt flow index (MFI), and decreases the melt viscosity of PVC in the blends 1. Species of wood flour added with PP and one or two broad particle-size distributions. On increasing wood flour content, flexural and tensile modulus, density,
heat deflection temperature, and notched impact energy increased, and some of the properties are decreased. Viscoelastic properties of a 60/70 penetration grade bitumen improved. Stability tests performed combining oscillatory flow and microscopy results disclose that blends with the higher polymer proportion (3%) are susceptible of phase separation after 24 hr of storage at 165°C, but 1% blends are stable for at least 4 days. A general evaluation of the results indicates that the performance of this bitumen as a binder for road pavement is particularly improved when 1% of recycled EVA or virgin EVA is added.

The influence of long-chain branching on the foaming behavior of polypropylene (PP) is investigated. Different branching contents are achieved by blending a linear PP (L-PP) and a long-chain branched PP (LCB-PP), it improve the optimal foaming process of PPs with a chemical blowing agent, and that additions of the linear material up to 50 wt% to the LCB-PP improve the optimal foaming process of PPs with a chemical blowing agent.

Thermal, mechanical and fire retardant properties of silane cross linked extra low-density polyethylene (XLPE) containing ethylene-vinyl acetate (EVA) copolymer using incorporation of EVA into XLPE gel content increased and curing time decreased. Mechanical properties of XLPE/EVA blends improve by increasing EVA content up to 15 wt%. Study the isotactic polypropylene and ethylene vinyl acetate blends prepared with five different talc concentrations, 3, 6, 9, 12 and 15 wt%, with addition to i-PP/EVA (88/12) to produce ternary composites and find the different mechanical properties those are effected by heat treatments. Prepared composite panels using virgin and recycled high-density polyethylene (VHDPE and RHDPE) and synthesize five types of natural fibers by melt compounding and compression molding. Fiber characteristics and the influences of fiber type and loading rate on HDPE crystallization behavior and composite mechanical properties were investigated. Evaluated the effect different filled cross linked thermoplastic elastomers of styrene butadiene rubber (SBR)/high density polyethylene blends (HDPE) blends and prepared using silica, HAF-carbon black, china clay and TiO₂. The reinforcement ability of the filler was increased in the order of silica>HAF-black>clay>TiO₂. Investigated the effect of prepared rice husks filled SBR/LLDPE (50/50) blends with a compatibilizer, maleic anhydride (MAH) using a Brabender plasticorder. Increasing MAH concentrations in SBR/LLDPE blends resulted in an increase in the tensile strength, elongation at break and hardness.

<table>
<thead>
<tr>
<th>Sample No.</th>
<th>PP (Wt %)</th>
<th>LLDPE (Wt %)</th>
<th>Talc (Wt %)</th>
<th>MgCO₃ (Wt %)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>70</td>
<td>30</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>B</td>
<td>60</td>
<td>20</td>
<td>20</td>
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<tr>
<td>C</td>
<td>60</td>
<td>20</td>
<td>-</td>
<td>20</td>
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<tr>
<td>D</td>
<td>40</td>
<td>20</td>
<td>20</td>
<td>20</td>
</tr>
<tr>
<td>E</td>
<td>60</td>
<td>20</td>
<td>10</td>
<td>10</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Sample</th>
<th>Thermo-Mechanical Properties of PP/LLDPE/Talc/MgCO₃ composition</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>A</td>
</tr>
<tr>
<td>HDT (°C)</td>
<td>119.6</td>
</tr>
<tr>
<td>Impact resistance (KJ/m²)</td>
<td>1.181</td>
</tr>
<tr>
<td>Charpy Impact resistance (KJ/m²)</td>
<td>1.190</td>
</tr>
<tr>
<td>Tensile strength (Kg/cm²)</td>
<td>68.07</td>
</tr>
<tr>
<td>Hardness</td>
<td>57.6</td>
</tr>
</tbody>
</table>
Synthesize the equal amount of PP/LLDPE and maleic anhydride grafted polypropylene blend systems. The SEM and DSC results of the PP/LLDPE/WP blend composites showed that WP located itself more in the LLDPE phase. The TGA results show that the MAPP/LLDPE blend and composites are more thermally stable than the PP/LLDPE blend and composites. Focused on the investigation of the changes on thermal, mechanical and morphological properties of PP-LDPE/DAP (90/10/0.06 wt. %) blend and PP-LDPE/DAP (90/10/0.0 wt. %) blend when different ratios. With increase in temperature up to 150°C the observation of TG/DTG/DTA curves revealed that the thermal stabilities of composites increased usually by increasing amounts of CaCO3 and the blends are incompatible. Evaluated the melt blending of thermoplastic polyurethane (TPU) with polyolefins (PO’s) which reduced the cost and improve mechanical and chemical properties. In addition to that it gives excellent performance in all aspects. In this research work, impact strength of untreated, alkali treated, potassium permanganate treated, benzoyl chloride treated and acrylic acid treated increased with increase in fiber loading up to 60% and then showed a decline for all untreated and chemically treated areca fiber reinforced epoxy composites.

Talc addition with PP improves the mechanical properties (stiffness, flexural strength, modulus etc) and heat deflection temperature. In this study, the improvements of mechanical properties have shown with increasing suitable percentage (by weight) of talc for new applications, and cellulose acetate/Fe2TiO5 nanocomposites have been synthesized successfully. Nanocomposites indicate a ferromagnetic paramagnetic behavior, as evidenced by using vibrating sample magnetometer (VSM) at room temperature. A mononuclear complex of [has been prepared by the reaction of 1-thenoyl-4,4,4-trifluoracetone in a 2:1 molar ratio. Thermogravimetric (TG) studies of 3 - 6 show a great thermal stability up to 400 °C for polymers, wherein the nickel oxide is formed at higher temperature. Experimental studied for formulations of PP and LLDPE viz. 90/10, 80/20, 70/30, 60/40 and 50/50 wt % were prepared for injection moulding machine. From the work they concluded that It optimum composition of PP/LLDPE blends provided the best mechanical, thermal properties and a moderate rheological property is 70/30. Investigated the effect of incorporation of talc and calcium carbonate PP/LLDPE blend that led to increase the hardness, MFI and melt density while decreased the impact resistance of blend. The results help to fix the optimum composition of (PP/LLDPE/Talc/MgCO3) wt% at which blend will show excellent thermo-mechanical properties.

**EXPERIMENTAL METHODOLOGY**

The preparation and characterization of Thermo-mechanical properties of polymer composite consist of number of sequential steps, which are as follow:

- Polymer pallets (PP-LLDPE) will be crushed in the powdered form using pulverizer.
- Talcum powder and Magnesium carbonate (powder form) along with PP-LLDPE (powder form) will be mixed uniformly in desired proportion using an agitator.
- This raw material will be fed in Compression Molding Machine (minimum 3Kg) at 250°C temperature and 50 bar pressure.
- The final product will be in the form of sheets which will be cooled at room temperature and then will be used further for various thermal and mechanical tests.

In this experimental study, five different talc and magnesium carbonate concentrations were added to PP/LLDPE (70/30) to produce composites make up a total of 3kg. The raw material was first crushed into fine powder form using pulverizer. Talc and MgCO3 were provided already in fine powder form by LOBA CHEMIE Pvt. Ltd. LLDPE and PP were in granules forms. PP and LLDPE were crushed separately in the pulverizer. Then the powder form PP/LLDPE/Talc/MgCO3 were mixed uniformly using an agitator. This prepared sample (3Kg) was then fed into compression molding. The machine was preheated to 250°C. Then the upper plate was made to apply pressure of 50 bars on mould, and finally sheets were taken out. It took 03hrs to prepare a single sheet. The processed samples were allowed to cool at room temperature for 48 hours and 50 ± 5% humidity. Then different mechanical and thermal tests were carried out. PP/LLDPE filled with talc and
magnesium carbonate to produce PP/LLDPE/Talc/ 
MgCO$_3$ composite is shown in table 1 and prepared 
sample are enlisted in figure 1-5

Analytical Standardization (ASTM)

The thermal properties of the blends were 
evaluating through heat deflection temperature 
(HDT) test. The test was carried out on a Ceast 
tester machine according to the ASTM D648. The 
tests specimens were molded in a size of 12.7 mm 
(width), 3.2 mm (thickness). Prior to the test all 
specimens were kept at room temperature at least 
48 h and 50 ± 5 % humidity. Izod impact strength 
values of the blends were evaluated with a Resil 
impact test instrument according to the ASTM D256 
test procedure at room temperature. Izod impact tests 
specimens were molded in a size of 12.7 mm (width), 
3.2 mm (thickness) and notched width 10.20mm. 
Prior to the test all specimens were kept at room 
temperature at least 48 h and 50 ± 5 % humidity. Izod 
impact strength values of the blends were evaluated 
with a Resil impact test instrument according to the 
ASTM D256 test procedure at room temperature. 
Izod impact tests specimens were molded in a size 
of 12.7 mm (width), 3.2 mm (thickness) and notched 
width 10.20mm. Prior to the test all specimens were 
kept at room temperature at least 48 h and 50 ± 5 % 
humidity. The hardness test of the composition was 
carried out on shore D testing machine according to 
the ASTM D 2240 and ISO 868. The tensile strength 
test of the blends was carried out on a universal 
testing machine according to the ASTM D638. A 
computer was connected to the load cell and data 
acquisition program recorded the force measured by 
the load cell. Test specimens were molded in a size 
of 12.7 mm (width) and 3.2 mm (thickness). Prior to 
the test all specimens were kept at room temperature 
at least 48 h and 50 ± 5 % humidity.
RESULTS AND DISCUSSIONS

Heat Deflection Temperature (HDT)

The heat deflection temperature (HDT) test is used to investigate the thermal properties of PP/LLDPE/Talc/MgCO3 composition. It is the temperature at which a polymer or plastic sample deforms under a specified load. The HDT of PP/LLDPE/Talc/MgCO3 compositions are shown in Table 2.

The heat deflection temperature of PP/LLDPE blend without any filler was 119.6°C. Addition of 20 wt% of talc to PP/LLDPE blend decreased the heat deflection temperature to 108.6°C. Addition of only 20 wt% of MgCO3 to PP/LLDPE blend increased the HDT to 113.1°C. Addition of 20 wt% of talc and 20 wt% of MgCO3 to PP/LLDPE blend has decreased the heat deflection temperature to 112.2°C. It is clearly seen that the addition of 10 wt % of talc and 10 wt% of MgCO3 to PP/LLDPE has increased the heat deflection temperature to 121.6°C. The increase in heat deflection temperature of composite may relate to the interaction between PP/LLDPE and Talc/MgCO3 and this restrict the mobility and deformability of PP/LLDPE, especially for lower contents. Table 2 indicates that the optimum composition percentage of PP/LLDPE/Talc/MgCO3 provides the best impact strength temperature is (60/20/10/10) wt %.

Izod & Charpy Impact Strength

The Izod impact test is used to investigate the mechanical properties of PP/LLDPE/Talc/MgCO3 composition. The impact properties of PP/LLDPE/Talc/MgCO3 composition are summarized in Table 2. The impact strength of PP/LLDPE blend without any filler was 1.181KJ/m². It is clearly seen that the addition of only 20 wt% of talc to PP/LLDPE blend increased the impact resistance to highest value of 2.804KJ/m². Addition of only 20 wt% of MgCO3 to blend decreased the impact resistance to 2.120KJ/m². The addition 20 wt % of talc and 20 wt% of MgCO3 to PP/LLDPE blend has decreased the impact resistance to 1.648KJ/m². It is clearly seen that the addition of 10 wt % of talc and 10 wt% of MgCO3 to PP/LLDPE blend has increased the impact resistance to 2.623KJ/m². The increase in impact resistance of composite may relate to the interaction between PP/LLDPE and Talc/MgCO3 and this restricts the mobility and deformability of the PP/LLDPE, especially for lower contents.

The Charpy impact test is used to investigate the mechanical properties of PP/LLDPE/Talc/
MgCO₃ composition. The impact properties of PP/LLDPE/Talc/MgCO₃ composition are summarized in Table 2. The impact resistance of PP/LLDPE blend without any filler was 1.190 KJ/m². It is clearly seen that the addition of only 20 wt% of talc to PP/LLDPE blend increased the impact resistance to highest value of 2.872 KJ/m². Addition of only 20 wt% of MgCO₃ to PP/LLDPE blend decreased the impact resistance to 1.817 KJ/m². The addition of 20 wt% of talc and 20 wt% of MgCO₃ to PP/LLDPE blend further decreased the impact resistance to 1.590 KJ/m². It is clearly seen that the addition of 10 wt% of talc and 10 wt% of MgCO₃ to PP/LLDPE blend increased the impact resistance to 1.842 KJ/m².

**Hardness and Tensile Strength**

The hardness of PP/LLDPE blend without any filler was 57.6. Addition of only 20 wt% talc has increased the hardness to 70.7. Addition of only 20 wt% of MgCO₃ to PP/LLDPE blend has decreased the hardness to 61.4. Addition of 20 wt% of talc and 20 wt% of MgCO₃ to PP/LLDPE blend has decreased hardness to lowest value of 50.7. But it can also be seen that the addition of 10 wt% of talc and 10 wt% of MgCO₃ to PP/LLDPE blend has increased the hardness to 69.3. The increase in hardness of composite may relate to the interaction between PP/LLDPE and talc/MgCO₃ and this restricts the mobility and deformability of the PP/LLDPE, especially for lower contents.

The tensile strength of PP/LLDPE blend without any filler was 68.07 Kgf/cm². It can also be seen that the addition of only 20 wt% talc to PP/LLDPE blend has increased the hardness to highest value of 79.98 Kgf/cm². It can also be seen that the addition of 20 wt% of MgCO₃ to PP/LLDPE blend has decreased the tensile strength to 68.37 Kgf/cm². Addition of 20 wt% of talc and 20 wt% of MgCO₃ to PP/LLDPE blend has decreased tensile strength to lowest value of 32.75 Kgf/cm². Addition of 10 wt% of talc and 10 wt% of MgCO₃ to PP/LLDPE has decreased the tensile strength to 57.75 Kgf/cm². The decrease in tensile strength of composite may relate to the interaction between PP/LLDPE and talc/MgCO₃ and this restricts the mobility and deformability of the PP/LLDPE, especially for lower contents.

**CONCLUSIONS**

The compositions of PP/LLDPE (70/30) with different percentage of talc and magnesium carbonate fillers were prepared. The work obtained incorporate talc and magnesium carbonate as fillers on PP/LLDPE blend led to increasing the heat deflection temperature (HDT), impact resistance, and hardness. While decreased the tensile strength of PP/LLDPE blend. It is inferred that the optimum composition provided the good thermal properties is PP/LLDPE/Talc/MgCO₃ (60/20/10/10) wt%. It is also concluded that the optimum composition provided the good mechanical properties is PP/LLDPE/Talc/MgCO₃ (60/20/20/00) wt%. As filler talc & magnesium carbonate enhanced the thermo-mechanical properties of PP/LLDPE blend but only at lower wt. % of talc & magnesium carbonate. Magnesium carbonate as a filler proved to be costly, so it is not cost effective as talc. Also due to the presence of carbonate ions it can shows rust problems. On the other hand silica, alumina, titanium dioxide can be used at the place of magnesium carbonate which can increase thermo-mechanical properties of PP/LLDPE in future.

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