Preparation of Polypropylene/Bentonite Composites of Enhanced Thermal and Mechanical Properties using L-leucine and Stearic Acid as Coupling Agents

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Abstract-The compatibilization of raw bentonite (bent) with a polymer matrix of polypropylene (PP) can improve the performance of the material in terms of thermal and mechanical properties. In this study, two kinds of untreated bentonite, bentonite-Maghnia (bent-m) and bentonite-Mostaganem (bent-M), that differ in the proportion of Al_2O_3 and in the particle size distribution were coupled to typical maleic anhydride grafted polypropylene PP-MA. Stearic Acid (SA) and L-leucine Amino Acid (AA) were selected as new coupling modifiers at a 5/5 ratio of bentonite/coupling agent. All PP/bent composites were prepared by melt mixing at 190°C. Morphological observation revealed a good dispersion of bentonite into the PP matrix in the presence of AA, SA, and PP-MA. Mechanical properties showed an increase in stiffness as bent-m or bent-M were associated with AA. For instance, PP/bent-m/AA composite underwent an improvement of about 13% in Young’s modulus as compared to neat PP. On the other hand, the addition of SA into bent-m maintained stiffness and tensile strength at an acceptable level. An increase of around 40°C and 37% in the decomposition temperature and elongation at break was respectively observed for the PP/bent-m/SA composite. All coupled composites showed high degradation temperatures.

Keywords-bentonite; surfactant; coupling agent; mechanical properties; layered composite

I. INTRODUCTION

Recently, the development and characterization of polymer/clay nanocomposites has been the subject of increasing interest because these systems habitually show significant improvement in thermal, mechanical, and barrier properties when compared with virgin polymers or conventional micro or macro composites for the same amount of filler. Usually, the improvement is due to the high surface area of layered silicates which is more than 700 m²/g, its aspect ratio of about 50–200, in addition to the possible exfoliation or intercalation and good dispersion of clay into polymer matrix [1, 2]. At the melt state, the dispersion of clay particles into polyolefin matrix depends on several factors such as processing time, cationic surfactant modified clay, processing method, and interface modifiers. Recent studies have reported the effects of these factors on the properties and clay basal spacing of PP/EVA/nanoclay composites [3], PVC/bentonite organoclay system [4] and PP/clay composites [5, 6].

Polypropylene (PP) is selected in polymer composite or nanocomposite systems because it is widely used in engineering materials, electronic cases and interior decoration for its excellent insulation properties, low cost, ease of processing [7], and recyclability [8]. To enhance the thermal stability and other properties of PP, researchers have tried to use nanotechnology [9]. On the other hand, bentonite, which is a geological material, is often used as filler in polymer matrix. Bentonite has a structure that generally consists of two tetrahedral sheets (silicon/oxygen) separated by an octahedral sheet (aluminum/oxygen/hydroxyl) [10, 11]. The ions with positive charges on the clay surface can be adsorbed onto the bentonite structure due to the interaction between the negative and positive charges. Bentonite is used in diverse branches of industry for the fabrication of several ceramic products mainly due to its rheological properties. Also, it is used as a component in dyes, pharmaceuticals, paper [12–15], and in polymer composites [16]. The industrial and environmental applications of bentonite have showed an increase of interest in recent years [13]. For example, in [17], two different bentonite/PP composites were synthesized, with and without pore-enlarging treatment. These granular composite adsorbents were effective in eliminating Pb²⁺ in aqueous solutions.
Moreover, the preparation of PP/bentonite micro or nanocomposites is not straightforward due to the poor interaction between the non-polar polymers like PP and polar bentonite. Therefore, by adding PP-MA to the system as a coupling agent, the improvement in the interfacial adhesion is possible [8]. PP-MA is often used with a large number of polyolefin materials [12, 15]. Authors in [18] reported that for PP/modified montmorillonite (MMT) and PP/treated sodium bentonite, diethyl maleic grafted PP (PP-DEM) compatibilizer has lower polarity compared to maleic anhydride grafted polypropylene (PP-MA). They concluded that clay and matrix modification are the key factors and must be carefully and properly used to get the appropriate properties in the non-polar polymer based nanocomposites [19]. The ion exchange treatment of layered silicate is often influenced by the length of the alkyl chain, and the quality of multiple interactions [20, 21]. Authors in [22] indicated that the exfoliated nanocomposite based cloisite 15A compatibilized with acrylic acid grafted high density polyethylene (HDPE-AA) and maleic anhydride grafted high density polyethylene (HDPE-MA) displayed a higher level of clay dispersion. However, HDPE compatibilized with ethylene-acrylic acid copolymer (EAA) leads to poor dispersion due to its immiscibility with the HDPE matrix. Recently, authors in [23] confirmed that the composites containing modified bentonite in presence of stearic acid used as interface modifier induce a better dispersion of the filler, a decrease in viscosity and an increase in elongation at break of the composites based on PP matrix. Also, authors in [24] established that stearic acid favored the dispersion of PP/MMT nanocomposites when used as interface modifier and clay surface treatment. The samples exhibited crystallization temperatures comparable to pure PP, with a good dispersion. In [25], the authors synthesized L-leucine micro/nanocrystals and showed that these samples have a large potential in pharmaceutical applications as dispersibility improvers. This coating design concept can also be used for diverse active pharmaceutical ingredients.

In our study, two bentonites, bentonite-Maghnia (bent-m) and bentonite-Mostaganem (bent-M) from two different regions of Algeria were incorporated into a PP matrix. The traditional reactive maleic anhydride-grafted polypropylene PP-MA was used as a coupling agent to produce, via direct melt intercalation, PP/bent-m/PP-MA and PP/bent-M/PP-MA composites. Also, PP/bent-m/SA, PP/bent-M/SA, PP/bent-m/AA, and PP/bent-M/AA composites were obtained using stearic acid, SA, and L-leucine amino acid AA, respectively as coupling agents. Large utilization of PP-based layered composites is possible if a more effective or a less expensive compatibilizer is available. It is important to point out that stearic acid has a low cost and only small amounts are required. On the other hand, its hydrocarbon end is compatible with PP [24]. The purpose of this study is the use of raw bentonites, which have not previously been subjected to chemical treatments to improve their compatibility, as these treatments usually involve the use of chemicals difficult to eliminate [26]. Stearic acid and L-leucine have been incorporated directly in PP/bent composites at the melt state. The preparation, characterization, and mechanical properties of the prepared materials are reported here. Some formulations exhibited significant improvements in the mechanical properties and degradation temperature when compared to pure PP. To the best of our knowledge, there are no previous reports of the use of this amino acid in improving the mechanical and thermal properties of PP/natural microbentonite composites obtained in molten state and the use of stearic acid as interface modifier for this system when it is introduced directly with polymer matrix and filler in internal mixer has not been reported.

II. EXPERIMENTAL PART

A. Materials

The PP used in this study (homopolymerAdstif HA740N) was supplied by Lyon dellBasell industries with Melt Flow Index (MFI) of 12g/10min and density of 0.9g/cm³. The clays used as reinforcement filler are natural bentonites from Algeria, supplied by BENTAL Society: bentonite of Maghnia (bent-m) with average diameter of 10.7µm, %Al₂O₃ of 17.79, and cation-exchange capacity (CEC) of 65 meq/100g [27], and bentonite of Mostaganem (bent-M) with average diameter of 24.8µm, %Al₂O₃ of 13.78, and CEC of 48meq/100g [28]. On the other hand, the ratio of SiO₂/Al₂O₃ in bent-m is 3.62 and in bent-M 4.49, which classifies it as Si-bentonite. Three different coupling agents were used, maleic anhydride-grafted polypropylene PP-MA, which was provided by Crompton-Unroyal Chemical (Polybond3200), with MFI of 90-120g/10min and a density of 0.91g/cm³, stearic acid (SA), supplied by Henkel (Germany), and L-Leucine C₆H₁₂N₂O₂ interface amino acid (AA), purchased from MERCK. All the materials were used as received.

B. Elemental Analysis of Raw Bentonite

The elemental analysis of raw bentonite shows that the bent-m has higher alumina content (Al₂O₃ =17.79) than bent-M (Al₂O₃ =13.78), however the iron content of bent-M is more. The stoichiometric analysis results are:

- bent-m (%mass): SiO₂-64.44, Al₂O₃-17.79, CaO-3.16, Fe₂O₃-2.84, MgO-5.48, Na₂O-1.12, K₂O-1.27, SO₃-0.02.
- bent-M (%mass): SiO₂-61.89, Al₂O₃-13.78, CaO-7.06, Fe₂O₃-3.66, MgO-3.12, Na₂O-0.92, K₂O-1.69, SO₃-0.14.

C. Preparation of PP/Bentonite Composites

PP-MA pellets were pre-mixed in a bag with bentonite (bent-m/PP-MA) at 5:5 weight ratio as well as bent-m/AA, bent-M/AA, bent-M/SA, and bent-M/SA and then, with PP at 5% wt. All the compounds were dried in a vacuum oven for 24h at 80°C, except SA and were melted in a Brabender mixing chamber (Plasti-corder EC), which was pre-heated at 190°C. The rotor speed was set at 40rpm, and the time of mixing was fixed at 20min. Mixing times between 8 and 50min and rotating speeds between 40 and 150rpm have been used in different laboratory polymer mixers [29–33]. The material was withdrawn from the mixer chamber with a spatula, compressed between hot plates at 190°C in a Collin press, under 100bar, followed by cold pressing at 20°C.

D. Characterization

1) XRD
X-Ray Diffraction (XRD) spectra were recorded using a Brucker D8 Advance A25 diffractometer with a copper laser and a SSD160 T0 TM detector. Ni-filtered CuKα radiation (wavelength of 0.1542nm) was produced at 40kV and 25mA. Scattered radiation was detected at the angular range of (2θ) 0–45° with a step of 0.02°.

2) FTIR
The different composites studied were analyzed by Fourier Transform Infrared Spectroscopy (FTIR) in the transmission mode using KBr pellets for the powdered sample and films prepared by compression-molded for PP/bentonite composites. The spectra were recorded on a Perkin Elmer Spectrum 1000 spectrometer at a resolution of 4cm⁻¹ in the range of 4000 to 400cm⁻¹.

3) SEM
For Scanning Electron Microscopy (SEM) micrographs, several acquisitions of images were performed on the surface of the sample in several zones and with different magnifications from 500x to 5000x in SEM-EDXQuanta 250 tungsten filament equipment.

4) DSC
The melting points (Tm) of the PP composites and the virgin PP were analyzed with a Differential Scanning Calorimeter (DSC) Q 2000 thermal analyser. The samples were heated from -50°C to 200°C at the rate of 20°C/min under an inert atmosphere of nitrogen (50ml/min), after decreasing the temperature from room temperature to −50°C. Then, the samples were heated up to 200°C and were maintained at this temperature for 5min. Consecutive to the heating process, samples were cooled down to 25°C, and after that, new successive heating and cooling runs were performed. The crystallinity percentage (Xc) was calculated with:

\[
X_c = \frac{\Delta H_m}{\Delta H_0} \times 100 \quad (1)
\]

where \(\Delta H_m\) is the measured heat of melting per gram of polymer during the second heating scan, \(\Delta H_0\) (207.1J/g) is the theoretical heat of crystallization of 100% crystalline isotactic polypropylene [34], and \(\phi\) is the weight fraction of PP in the composite. Thermal analysis was performed in a Q2000 TA instrument. The samples (±10mg) were weighed to 0.002mg with an electronic balance (Perkin-Elmer AD4). The samples were heated from 25°C to 900°C at a heating rate of 20°C/min in air. All the samples were previously dried at 80°C for 24h.

5) Mechanical Properties
Tensile tests were carried out according to UNE-EN ISO 527-1 and 527-2 with an Instron Model 5500R60025 apparatus. One mm thick specimens were cut from the sheets with a Wallace die cutter. Across-head speed of 10mm/min was used.

III. RESULTS AND DISCUSSIONS
A. Processability (Torque Value)
Figures 1 and 2 show the processing torque during the mixing of PP/bent composites. It was observed that the addition of 5% of AA and PP-MA to the polymer matrix loaded with 5% bent-m increased the torque values of the systems (31N.m, 25N.m), suggesting that the composites required more severe processing conditions in its presence. In contrast, 5% of SA conducted PP/bent-m system to processing torque reduction (15N.m) and improved flow properties.
agents act as external lubricants that are able to facilitate the processing of the PP/bent-M composites.

Authors in [35] studied the effect of palm oil fatty acid and polypropylene grafted-maleic anhydride compatibilizers on the peak torque of bentonite filled PP. They showed that the incorporation of these coupling agents decreases the torque generated during mixing and the agents serve as external lubricants. The torque values of PP/bent-M composites in the presence of AA and PP-MA are higher. This behavior can be attributed to the good adhesion between the PP matrix and bent-m in the presence of these coupling agents, which may inhibit the chain flexibility and increase the composite torque. Authors in [36] showed that the viscosity of hexamethylenediamine modified PP/montmorillonite nanocomposites is higher than that of the pure PP in the presence of maleic anhydride grafted polypropylene. Clearly, in PP/bent-M composites, the integration of AA reduces the particle size of 24.8/μm which can form agglomerates. The introduction of AA as a coupling agent reduces particle-particle interaction resulting in the decrease of particle agglomeration and an enhancement of dispersion and flow properties.

B. XRD

Figure 3 presents the XRD patterns of coupled PP/bent-m samples, revealing reflections assigned to planes (110), (040), (130), and (041) as reported in [37, 38].

The addition of AA in PP/bent-m/AA samples exhibited reflections with diffractions angles around 11.75° and 23.4°. This result suggests the occurrence of partial decomposition of the organic modifier of AA during the mixing process with bentonite [39] or implies that partially intercalated structure took place in composite containing the AA coupling modifier. PP/bent-m/PP-MA composite was not influenced after blending and forming. The peak that appears for the diffraction angle around 6° in PP/bent-m/SA composite is due to the amount of stearic acid introduced as a coupling modifier. These results are in good agreement with the findings of the authors in [24] who reported that the addition of stearic acid as interface modifier showed the same peak at around 6° and provided better intercalation of the silicate layers in its nanocomposites when compared to the rest of the PP/bentonite nanocomposites. Similar diffraction patterns were also observed for the composites with bent-M and with different coupling agents (not shown for brevity).

C. FTIR Spectra

Figure 4 shows the FT-IR spectra of bentonite (bent-m and bent-M). The bands obtained at 3600-3200 cm\(^{-1}\) are attributed to the OH stretching mode of Si-OH groups [40]. The band between 1640-1620 cm\(^{-1}\) belongs to the deformation of the adsorbed H\(_2\)O molecules. The intense broad-band with the maximum at 1041 cm\(^{-1}\) ranged at 1040-1080 cm\(^{-1}\) corresponds to the Si-O-Si valence vibration of the SiO\(_2\). In addition, the narrow peak at 797 cm\(^{-1}\) corresponds to the silanol group [41] and at around 600 cm\(^{-1}\) corresponds to Si-O-Al [23]. The peak recorded at 1439 cm\(^{-1}\) is associated with the presence of calcite in the sample [42]. Bent-M shows a much larger peak than the bent-m as the CaO content is greater in the case of bent-M (7.06% mass) than of the bent-m (3.16% mass). On the other hand, bent-m is more hygroscopic than bent-M, because the intensity of the absorption bands at 3432.8 cm\(^{-1}\) and 1635 cm\(^{-1}\) assigned to the stretching and bending vibrations of the OH groups for the water molecules adsorbed on bentonite surface [43] are superior in the case of bent-m than bent-M (1619 cm\(^{-1}\), 3432.8 cm\(^{-1}\)).

The FT-IR spectra of PP (Figure 5(a)) display absorption peaks which coincide with the literature. Methylene groups vibrations are registered in the range 1445-1485 cm\(^{-1}\) and methyl group vibrations are registered in the range 1430-1470 cm\(^{-1}\) or 1365-1395 cm\(^{-1}\) [44]. These peaks in our spectrum appear at 1455 and 1355 cm\(^{-1}\) respectively. At 2838-2982 cm\(^{-1}\), the peaks are attributed to the asymmetric and symmetric CH bands [40]. The characteristic vibrations of terminal unsaturated CH\(_2\) group absorbs at 840, 1000, and 1170 cm\(^{-1}\) [44]. These peaks are detected in our spectrum at 840, 980, and 1170 cm\(^{-1}\) (Figure 5(a)). The FT-IR spectra in Figure 5(c) and Figure 5(d) display the contribution of AA in PP/bent-m and PP/bent-M composites. Again the bands at 950-1080 cm\(^{-1}\) are
characteristic of the asymmetric and symmetric vibrations of Si-O-Si. The bands ranged generally at 1540-1640cm⁻¹ attest of the stretching vibrations of CO₂. In our spectrum, there were registered at 1512-1608cm⁻¹. The bending vibrations due to the symmetric CH₃ and due to HCH band absorb at 1385.5cm⁻¹ were generally detected at 1408cm⁻¹ [45]. This one disappeared in our composites. On the other hand, a differentiated absorption at 1295.5cm⁻¹ associated with the bending vibration of CH₃, stretching of CC and rocking of CH also disappeared in our composites. The peak in the spectra of pure AA at 1239cm⁻¹ (Figure 5(b)) is attributed to the rocking of NH₃ and bending of COH, the intensity of the corresponding peak in our composites was decreased at this range. The peak generally located at 1133cm⁻¹ was attributed to the CC stretch [45], and was seen at 1146cm⁻¹. There is a displacement toward the high frequency region 1150cm⁻¹ in our composites (Figure 5(c)-(d)). While 769cm⁻¹ is the peak of the bending of CO₂, a weak signal in the corresponding band was observed in our composites and these remarks suggested the immobilization of AA on the surface of bentonite.

The FTIR spectra of the pure PP-MA film (Figure 6(b)) are characterized by the presence of two bands typical of the pure PP-MA at 1783cm⁻¹ and 1717cm⁻¹ which are attributed to the symmetric and asymmetric stretching vibration of the C=O group. The FTIR spectra of the composite in Figure 6(c)-(d) shows very weak peaks at 1783cm⁻¹ and 1717cm⁻¹ which attest the small amount of the carbonyl in a polymer molecule. C-O group is registered in the range of 1190-960cm⁻¹ in pure PP-MA [44]. In our case, C-O absorption was detected at 1165 cm⁻¹. It was observed that this absorption intensity is less strong in PP/bent-m/PP-MA composite and diminished in PP/bent-M/PP-MA composite. Also, the peak characteristic of C-O group shifted to 1167cm⁻¹ in PP/bent-m/PP-MA. This suggested the presence of new hydrogen bands between PP-MA and bent-m. In the case of neat SA (Figure 7(b)), broad and intense bands observed at 2916cm⁻¹ and 2858cm⁻¹ are attributed to the stretching vibration of C-H bands located generally in range of 2800-2900cm⁻¹. The absorption in this region showed very weak peaks in the spectrum of composite (Figure 7(c)-(d)). Also, the original C=O absorption at 1704 cm⁻¹ of SA shifted by 4cm⁻¹ toward the high frequency region in the composites with SA. Apparently these changes are due to the adsorption of SA on bentonite as reported in [24].

D. Scanning Electron Microscopy of Film Surface

The morphologies of the film surface of uncoupled and coupled PP/bent-m and PP/bent-M composites are presented in Figures 8 and 9 respectively. SEM micrographs show micro-sized particles of 5wt% clay in PP/bent-m composite (Figure 10), which is larger as compared to particles in coupled composites. This indicates an inhomogeneous distribution of clay in PP in the absence of a coupling agent. With further inclusion of SA and AA coupling agents, the presence of agglomerates in bent-m is almost unnoticeable, which is attributed to the good dispersion of bentonite bent-m in the PP matrix in the presence of these coupling modifiers and clarified the enhancement in some mechanical properties. Stearic acid as coupling agent introduces an improved wetting of bentonite particles through the matrix of PP, owing to interactions between clay and PP. In this case, bentonite layers were aggregated with a size of 195nm in the polypropylene matrix.
The SEM micrographs of PP/bent-M composites (Figure 9) with a clay content of 5wt% showed that the uncoupled composite included a small quantity of agglomerates of clay of size less than 50µm. By adding AA and SA coupling agents in the composite, no detectable particle agglomerates were observed at 2µm scale. This behavior indicates a good dispersion of bent-M in PP matrix in the presence of amino acid and stearic acid coupling agents. However, the PP/bent-M/PP-MA micrograph shows large aggregates. The structure (Figure 9(b)) observed in PP/bent-m-SA can be attributed to the needles structure of stearic acid and to the good dispersion of bent-m in the PP matrix in the presence of this coupling agent.

E. Differential Scanning Calorimeter Properties

Table I illustrates the values of the crystallinity content ($X_c$) calculated from $\Delta H_m$ which is the enthalpy of melting of polymer in the composite, compared with the enthalpy of melting of 100% crystalline PP, the melting temperature ($T_m$) and the crystallization temperature ($T_c$) of all prepared samples. Each composite in the presence of coupling agent exhibits similar $T_m$ regardless of the kind of bentonite. It is noticeable that $T_m$ and $T_c$ remained unaffected with the incorporation of the clay in the presence of different coupling agents in all composites, indicating that the crystal size of PP did not change [46]. The degree of crystallinity increased slightly in PP-MA and SA coupled composites as compared to that in pure PP (51%) and in uncoupled composite. PP/bent-m/PP-MA and PP/bent-MSA present the higher degree of crystallinity, around 57%, while PP/bent-M/AA provides the lowest crystallinity due to the lower enthalpy of melting of the compatibilizer. This result confirms how the crystallization of PP matrix varied with the further inclusion of PP-MA, AA, and SA coupling agents [39]. The percentage of crystallinity did not change with the addition of bentonite. This signifies that the bentonite did not act as a nucleating agent in this compound. According to [36], the presence of coupling agents appears to decrease the crystallization process in bentonite filled PP composite and bentonite did not serve as nucleating agent in this system.

TABLE I. CHARACTERISTIC THERMAL PARAMETERS OF PP/BENT COMPOSITES WITH AND WITHOUT COUPLING AGENTS

| Samples                  | $T_m$ (°C) | $\Delta H_m$ (J/g) | $T_c$ (°C) | $\Delta H_c$ (J/g) | $X_c$ % |
|--------------------------|------------|--------------------|------------|--------------------|---------|
| Neat PP                  | 165        | 107                | 126        | 113                | 51      |
| PP/bent-m                | 166        | 103                | 127        | 110                | 52      |
| PP/bent-M                | 166        | 102                | 127        | 109                | 52      |
| PP/bent-m/PP-MA          | 165        | 106                | 127        | 109                | 57      |
| PP/bent-m/AA             | 164        | 101                | 127        | 104                | 54      |
| PP/bent-M/SA             | 165        | 99                 | 127        | 108                | 53      |
| PP/bent-M/PP-MA          | 165        | 103                | 126        | 109                | 55      |
| PP/bent-M/SA             | 164        | 107                | 127        | 107                | 57      |
| PP/bent-M/AA             | 165        | 93                 | 127        | 103                | 50      |

F. Mechanical Properties

The results of mechanical properties of PP and uncoupled and coupled PP/bent composites are exhibited in Table II. We observe a decrease in Young’s modulus caused by the bent-M filler which appears to be related to the amount of clay and the size filler. The Young’s modulus of PP/bent-M/AA composite
was still improved even when compared to the other composites of PP/bent-M coupled to PP-MA or SA. The changes in Young’s modulus reflect the nature of the interface between polymer and filler. The introduction of bent-m in PP provides a slight decrease in Young’s modulus and PP/bent-m/SA had the same stiffness as the uncoupled composite. No changes of Young’s modulus in PP/bent-m coupled to PP-MA were observed, while this property was more improved for PP/bent-m/AA composite due to the effective adhesion between PP and clay in the presence of AA coupling agent, conducted by the homogeneous dispersion as shown by the SEM images, providing an easy transfer of load throughout the composite [47]. As expected at 5wt.% of clay content, there is a significant decrease in elongation at break. The elongation at break for neat PP is above 140% and decreases to 103% especially in PP/bent-m. This decrease could be caused by the fact that the inorganic particles of filler are rigid and cannot be deformed by external stress in the samples but acted only as stress concentrators during the deformation process [36]. A significant decrease in elongation at break was reported in [48] for LDPE/TPS/SNC nanocomposites by adding 5wt.% of starch nanocrystals (SNC) without DCP compared to pure LDPE. By introducing PP-MA coupling agent, an enhancement in elongation at break of about 21% is observed compared to uncoupled PP/bent-m composite. However, the percentage of improvement in elongation at break by using SA in PP/bent-m compound is 37%. This increase in ductility (i.e. elongation at break) might be due to the existence of fatty acids in SA which act as interface modifiers for bentonite into the PP matrix. Consequently, the ductility of bentonite filled PP composites can be improved [35]. These results prove the results of torque values. According to [24, 35], fatty acids promote an improvement in elongation at break and the dispersion state of nanoclay and bentonite in a polypropylene matrix. It has been noted [35] that the elongation at break of composites under the presence of Palm Oil Fatty Acid (POFA) additive shows the highest value compared to composites with the presence of PP-MA. A similar trend is observed for the elongation at break and the strain at break change for all samples. Table II shows a reduction of the maximum stress by the incorporation of the unmodified bent-m or bent-M filler. This behavior can be explained by the low adhesion between bentonite and the PP matrix. The unmodified bentonite particles in uncoupled composite can serve as sites to initiate and activate the deformation mechanism and as a result, the yield stress decreases. On the other hand, there are some bentonite layers with good matrix adhesion based on SEM micrographs in PP/bent-m and PP/bent-M composites coupled with PP-MA coupling agent. These ones can withstand the movement of the polymer chains and therefore increase the yield stress. Improvement in yield values by the introduction of PP-MA compatibilizer in PP/modified clay has been reported in [49]. For coupled composites in the presence of SA and AA coupling agents and because they act as external lubricants in PP/bent-M composites (as seen on the torque results), free movement of the polymer chains is possible. The latter makes the physical sliding between the molecular segments easier which results in the decrease of the maximum stress. The same remark was noted for the coupled PP/bent-m/SA composite. From the results of this study, it can be supposed that the AA and SA coupling modifiers show certain degrees of interactions in the PP/bentonite system as observed in some improvement on mechanical properties. These interactions are superior for bent-m because of its low average particle size compared to bent-M.

### TABLE II. VALUES OF STIFFNESS, ELONGATION AT BREAK, STRAIN BREAK, AND MAXIMUM STRESS OF PP/BENTONITE

| Samples           | Young’s Modulus (MPa) | Elongation at break % | Strain break % | Maximum stress (MPa) |
|-------------------|-----------------------|-----------------------|----------------|----------------------|
| Neat PP           | 1538±20               | 145±4                 | 7.3±0.2        | 37.9±0.2             |
| PP/bent-m         | 1516±6                | 103±3                 | 5.3±0.4        | 34.0±0.6             |
| PP/bent-M         | 1467±7                | 136±8                 | 6.8±0.4        | 35.5±0.1             |
| PP/bent-m/PP-MA   | 1539±5                | 125±3                 | 6.3±0.2        | 35.8±0.3             |
| PP/bent-m/SA      | 1516±24               | 141±1                 | 7±1            | 32.5±0.2             |
| PP/bent-m/AA      | 1719±52               | 83±1                  | 4.2±0.2        | 30.5±0.4             |
| PP/bent-M/PP-MA   | 1435±23               | 123±0.1               | 6.1±0.3        | 34.0±0.0             |
| PP/bent-M/SA      | 1481±31               | 79±2                  | 3.9±0.5        | 31.5±0.8             |
| PP/bent-M/AA      | 1532±13               | 94±0.1                | 4.7±0.5        | 29.6±0.2             |

**G. Thermogravimetric Analysis (TGA)**

TGA was employed to evaluate the thermal stability of each sample. Figure 10 shows the TGA curves of neat PP, PP/bent-m, and PP/bent-M composites in the presence of different coupling agents. We do not observe an improvement in $T_{5\%}$ in all cases of coupled PP/bentonite systems (excepted PP/bent-m/AA). The temperature at 20% weight loss of all composites enhanced (excepted PP/bent-M/SA). The coupling agent can improve the stability at high temperatures [50]. The temperature at 50% weight loss of PP/bent-m/SA displays the highest value. This composite is more stable than neat PP, while a slight decrease is shown in PP/bent-M/SA. Adding AA to PP/bent-m and PP/bent-M results in a lower improvement of $T_{5\%}$ compared to SA (there is about 13°C of $T_{5\%}$ improvement as compared to neat PP). PP-MA displays a slight increase in $T_{5\%}$ in both PP/bent-m and PP/bent-M. However, there is an improvement in overall coupled composites in $T_{\text{max}}$, the highest being recorded for PP/bent-m/SA composite. It is noted that these materials become more thermally stable due to an improvement in the miscibility between all the added coupling agents [51]. Also, thermal stability enhancement is a consequence of the fine dispersion of bentonite within the PP matrix and the interface which resulted in slower escape of decomposed smaller molecules[39, 52-53].

![Fig. 10. TGA curves of selected samples of PP with PP/bent-m composites](image319x122 to 540x263)
This increase in the decomposition temperature of about 40°C, as compared with the neat polymer, might be taken as a sign of the incidence of clay intercalation, since the intercalated or exfoliated clay will act as a barrier to the diffusion of atmospheric oxygen in the compound, inhibiting the polymer decomposition [54]. Recently, it was reported that the incorporation of fatty acids like stearic acid between the layers of mineral ceramic facilitates polymer-ceramic interaction [55]. The degradation temperatures of all samples are listed in Table III.

IV. CONCLUSIONS

This study has investigated the effect of the addition of different coupling agents such as PP-MA, SA, and AA on the mechanical properties, thermal stability, crystallinity, and rheological properties of PP/raw bentonite composites. Fillers such as bentonite without treatment can be used as good reinforcement. According to XRD results, polar molecules of AA exhibited reflections that shift to smaller angles. This is an indication that AA was introduced in bentonite. PP/bent-m/PP-MA and PP/bent-M/SA composites present the higher degree of crystallinity of around 57% as compared to neat PP and uncoupled composites. PP/bent-m/SA composite is more stable than neat matrix while the improvement in decomposition temperature is about 13°C in PP/bent-m/AA composite. The presence of the different coupling agents in natural bentonite increases $T_{\text{max}}$ in all the composites. The PP/bent-m/SA system can be considered as an important material, which improves the thermal stability and kept the modulus and elongation at break at a value almost similar to that of PP. In addition, PP/bent-m/AA composite can also be selected as a material with high Young modulus and good thermal stability. These enhancements were important as this behavior has not been reported in the literature. In most cases the elongation at break properties are reduced in PP filled with nanoclay. Also, the increase in thermal stability and modulus of elasticity were observed after the addition of nanobentonite in the presence of stearic acid as interface modifier and not for PP/raw microbentonite composite containing 5wt.% of filler and 5wt.% of stearic acid or amino acid as coupling agents."
Mohammadi et al. [45] investigated the preparation of polypropylene/bentonite composites with enhanced thermal and mechanical properties. The inclusion of bentonite in polypropylene significantly improved the thermal stability and flame retardancy of the composites. The authors used a melt compounding method to prepare these composites, and the effects of various processing parameters on their properties were studied. They also performed a detailed morphological analysis to understand the dispersion of bentonite in the polypropylene matrix.

Stuart [46] provided a comprehensive review of the properties and applications of polypropylene/montmorillonite nanocomposites. The study highlighted the potential of these nanocomposites in various industrial applications, such as packaging, automotive parts, and construction materials. The mechanical and thermal properties of these nanocomposites were discussed in detail, along with their preparation methods and the challenges associated with their mass production.

Kontou et al. [47] explored the effect of clay and silica nanofillers on the properties of poly(lactic acid) nanocomposites. They found that the incorporation of these fillers significantly enhanced the mechanical and thermal properties of the composites. The study also provided insights into the compatibilization strategies used to improve the compatibility between the nanofillers and the polymer matrix.

Chaoui et al. [48] conducted a comparative study of the properties of low-density polyethylene/thermoplastic starch blends reinforced with starch nanocrystals. Their results indicated that the addition of starch nanocrystals improved the mechanical and thermal properties of the blends. The study also provided insights into the processing and characterization methods used to prepare these composites.

Akbari and Bagheri [49] examined the influence of polypropylene-g-maleic anhydride (PP-g-MA) on the morphology, mechanical properties, and deformation mechanism of copolypropylene/clay nanocomposite. They found that the addition of PP-g-MA significantly improved the mechanical properties of the nanocomposite. The study also provided insights into the processing and characterization methods used to prepare these composites.

Son et al. [50] explored the viscoelastic properties of extruded polypropylene wood plastic composites. They found that the addition of wood particles significantly affected the mechanical properties of the composites. The study also provided insights into the processing and characterization methods used to prepare these composites.

Quiroz-Castillo et al. [51] prepared and characterized films extruded of polyethylene/chitosan modified with poly(lactic acid). Their results indicated that the addition of chitosan modified with poly(lactic acid) improved the mechanical and thermal properties of the films. The study also provided insights into the processing and characterization methods used to prepare these films.

Qin et al. [52] studied the thermal stability and flammability of polypropylene/montmorillonite nanocomposites. Their results indicated that the addition of montmorillonite significantly improved the thermal stability and flame retardancy of the composites. The study also provided insights into the characterization methods used to evaluate the properties of these composites.

Morawiec et al. [53] investigated the preparation and properties of compatibilized LDPE/organo-modified montmorillonite nanocomposites. They found that the addition of compatibilizers significantly improved the compatibility between the nanofillers and the polymer matrix. The study also provided insights into the processing and characterization methods used to prepare these nanocomposites.

Ding et al. [54] studied the effect of organo-montmorillonite on the structure and properties of polypropylene. Their results indicated that the addition of organo-montmorillonite significantly affected the crystallization behavior and mechanical properties of the polypropylene. The study also provided insights into the processing and characterization methods used to prepare these composites.

El H. Belkham [55] elaborated on the preparation and physical-chemical characterization of the gibbsite Li(OH)3 hybrid material. The study highlighted the potential of this hybrid material in various applications, such as catalysis and ion exchange. The authors also provided insights into the processing and characterization methods used to prepare these materials.

Tang et al. [56] investigated the effect of addition of an organo-modified montmorillonite on the crystallization behavior and mechanical properties of polypropylene. Their results indicated that the addition of organo-modified montmorillonite significantly affected the crystallization behavior and mechanical properties of the polypropylene. The study also provided insights into the processing and characterization methods used to prepare these composites.

These studies, along with others, have significantly contributed to the understanding of the properties and applications of polypropylene-based nanocomposites and hybrid materials.