Fabrication and Characterization of Nanostructured Polymer Composites Prepared by Melt Compounding

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Abstract—In this research we have investigated the effect of different percentages of nanoclay and matrix compositions on the characteristics of NR/BR blend prepared by open two-roll mill. The dispersion of the layered silicate in rubber matrix was characterized by X-ray diffraction (XRD). The results showed that the nanocomposites structure consisting of intercalated structure as well as partially exfoliated structure are obtained. Properties such as permeability and complex viscosity as well as storage modulus are tested to assess the effect of addition of the nanoclay and matrix composition. Permeability and compression set of the nanocomposite samples received markedly improvement. These results were confirmed by Gel content measurements since a noticeable decrease in the gel values was observed when the organoclay was added to the rubber compounds. The properties improvement caused by nanoclay are attributed to intercalation of the organophilic clay by NR/BR macromolecules and established interaction between nanoclay and polymer matrix which was evidenced by SEM photomicrographs. These results were accompanied by enhanced rheological properties.

Index Terms—Nano clay, nanocomposite, x-ray diffraction, SEM photomicrographs, rheological properties

I. INTRODUCTION

Polymer blends have been intensively studied because of their theoretical and practical importance. Among the blends widely investigated in rubber industries are natural rubber (NR)/styrene-butadiene rubber (SBR), NR/butadiene rubber (BR), and NR/BR/SBR.

Synthetic rubbers are classified into general purpose rubbers and special purpose rubbers. Polybutadiene is classified as general purpose rubbers intended for the manufacture of tires and general mechanical products. However, one type of rubber may not possess all the physical properties desired in a finished product. Much attention is currently being devoted to the simplest route for combining the outstanding properties of different existing polymers, i.e., the formation of polymer blends [1]. For example, in tread compounds, the high abrasion resistance under certain conditions conferred by the use of BR is desirable, but the poor road holding and rib tearing properties are not, hence blends of BR with NR and styrene–butadiene rubber are employed. Compatibility is the fundamental property, deciding the practical utility of a polymer blend.

If the two elastomers in a blend are incompatible, it will exist in the form of two separate phases and the cured blend will show inferior properties. In NR/BR blends, BR shows improved tread wear and groove cracking resistance without reduction in resilience. The most important advantage of adding BR to NR is the improved abrasion resistance.

The reinforcement of rubbers is expressed by enhancement of the modulus, failure properties (tensile and tear strength), and abrasion resistance of the vulcanizates. The main aim for filler addition is to improve certain properties and cheapen the compound. In the last two decades, research was focused on the development of other reinforcing agents to replace carbon black in rubber compounds. The synthesis of inorganic–organic complex materials has been widely done for the purpose of preparing noble materials with various special properties. Nanocomposites are composites in which an inorganic filler is dispersed within a polymer matrix at a nanoscale level, thus the clay layer could be intercalated or exfoliated into the polymer to form clay/polymer nanocomposites. In other words, the polymer is nanoscopically confined. Nanocomposites exhibit greatly improved mechanical and thermal properties because of the nanometre-scale dispersion of the reinforcements and the high surface-to volume ratio [2]. Clay silicates, such as montmorillonite, hectorite, Bentonite, etc, have been used as reinforcing fillers for polymers because of their potentially high aspect ratios [3], [4], [5]. The crystal structures of clays are usually interesting in their own right. Their crystal lattice consists of two-dimensional layers where a central octahedral sheet of either aluminum or magnesium is fused to two external silica tetrahedra by their tip so that the oxygen ions of the octahedral sheet also belong to the tetrahedral sheets. The layer thickness is around 1 nm and the lateral dimensions of these layers may vary from 300 Å to several microns and even larger, depending on the particular silicate. The layers are stacked by a weak dipole force and exhibit a net negative charge on the lamellar surface; the interlayers between the galleries are normally occupied by cations such as Na+ and Ca++. As the forces that hold the stacks together are relatively weak, the intercalation of small molecules between the layers is easy. The metal cations of the interlayer can be exchanged with organic quaternary alkylammonium salts with long chains, thus producing organophilic clays, also called organoclays, to give surfaces that are less ionic or polar. Organophilic modification improves the compatibilization between the surface of the clay and the hydrophobic polymer matrix and thus the polymers penetrate more easily into the galleries. These entering guest molecules can either simply increase the distances between the still-parallel layers in an intercalation process or randomly disperse the separate sheets entirely in an exfoliation process.

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For true nanocomposites, the clay nanolayers must be uniformly dispersed (exfoliated) in the polymer matrix, as opposed to being aggregated as tactoids or simply intercalated. When nanolayers have been exfoliated, the improvement in properties can be seen as an increase in tensile properties, as well as enhanced barrier properties, decreased solvent permeability, and increased thermal stability and flame retardance. [6], [7].

In recent years, Polymer/clay nanocomposites (PCNs), emerged as a new class of material, have attracted negligible interest from academia and industry [2] - [15]. Compared with their relative micro and macro scale counterparts and their pristine polymer matrices, PCNs exhibit significantly improved performances such as tensile strength, modulus, heat resistance, gas barrier, swelling resistance, etc. Nanoclay is a hydrophilic mineral which would not have an appropriate compatibility and adhesion to non-polar polymers. This problem could be solved by modification of clay using alkyl ammonium or alkyl phosphonium cations.
Substitution of hydrated metallic ions by organic cations has an important role in balancing the negative ions and modification of clay from hydrophilic to hydrophobic. Such a modification on the clay causes the surface energy to get defaced and the nano-filler will be compatible with organic polymers. It seems that the interaction between adjacent silicate layers having negative charge has the main role in the possibility of positive organic ions to enter the galleries. As the modified clay is dispersed in the polymeric matrix, ammonium ions and polymer chains attract each other. On the other hand the repulsion between clay layers facilitates the entrance of polymer chains into the galleries of silicate. Very recently, increasing attention has been paid on rubber/clay nanocomposites (RCNs). According to reports, several RCNs based on one or two elastomers has been prepared via solution intercalation, melt intercalation, and rubber-latex compounding [6] - [15]. But to our knowledge, no studies have been published on NR/BR nanocomposites prepared by melt intercalation in the literature.

II. EXPERIMENTAL

A. Materials

NR (SMR 20), Mooney Viscosity ML (1+4) at 100 °C, 55, BR (97% 1, 4(cis)), Mooney Viscosity ML (1+4) at 100 °C, 58, nanoclay were supplied by Malaysia, China and Southern clay Ltd respectively. Compounding ingredients (zinc oxide (3.0), stearic acid (2.5), aminox (2.0), PV1, n-cycloexlythioliphalimide (2.0), sulfur (2.5), TBBS, t-butyl-2-benzothiazolsulfenamide (0.6). The nanoclay used was Cloisite 15A which was a natural montmorillonite modified with a dimethyl dehydrogenated tallow quaternary ammonium having a cation exchange capacity of 125 mequiv/100 g.ing ingredients.

B. Samples Preparation

Nanocomposite samples were prepared by a laboratory open two-roll mill (Polymix 200-L) for 18 min at room temperature at the rotor speed of 80 rpm. It should be mentioned that first nanoclay was pre-dried at temperature of 80°C for 24 hr, then NR and BR were blended and nanoclay and curing agents were added to the compound.

<table>
<thead>
<tr>
<th>Sample</th>
<th>NR (wt%)</th>
<th>BR (wt%)</th>
<th>Nanoclay (wt%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>S1</td>
<td>75</td>
<td>25</td>
<td>0</td>
</tr>
<tr>
<td>S2</td>
<td>75</td>
<td>25</td>
<td>1</td>
</tr>
<tr>
<td>S3</td>
<td>75</td>
<td>25</td>
<td>3</td>
</tr>
<tr>
<td>S4</td>
<td>75</td>
<td>25</td>
<td>5</td>
</tr>
<tr>
<td>S5</td>
<td>75</td>
<td>25</td>
<td>7</td>
</tr>
<tr>
<td>S6</td>
<td>100</td>
<td>0</td>
<td>5</td>
</tr>
<tr>
<td>S7</td>
<td>50</td>
<td>50</td>
<td>5</td>
</tr>
<tr>
<td>S8</td>
<td>25</td>
<td>75</td>
<td>5</td>
</tr>
<tr>
<td>S9</td>
<td>0</td>
<td>100</td>
<td>5</td>
</tr>
</tbody>
</table>

C. Characterization

Curing process were carried out on the Diffenbecher hot press at 160°C and 150 bar. To measure the change of gallery distance of organo-MMT before and after intercalation, X-ray diffraction (XRD) was performed at room temperature with a Philips X-ray diffractometer model X’Pert. The X-ray beam was a Cu Ka radiation (λ= 1.540598 Å) using a 50 KV voltage generator and a 40 mA current. The rheological measurements of prepared samples were carried out using RPA oscillatory rheometer at temperature of 80 °C and frequency of 0.01-80 Hz. In order to study the morphological properties, SEM micrographs of cryogenically fractured surface of gold coated samples were taken with a TESCAN electron microscope. The permeation experiment of nitrogen was carried out with gas permeability-measuring apparatus. The pressure on one face of the sheet (about 1 mm in thickness and 8 cm in diameter) was kept at 0.57 MPa and the other face at zero pressure, and nitrogen permeated through the sheet. The rate of transmission of nitrogen at 40 °C was obtained by gas chromatography and the nitrogen permeability was calculated from it. Gel content values were calculated according to the weight loss of the sample washed with acetone after a 24 hrs dipping in boiling toluene. Compression set testing measures the ability of the rubber to return to its original thickness after prolonged compressive stresses at a given temperature and deflection. As a rubber material is compressed over time, it loses its ability to return to its original thickness. This loss of resiliency may reduce the capability of a rubber seal to perform over a long period of time. The resulting permanent set that a seal may take over time may cause a leak. Compression Set results for a material are expressed as percentage. The lower the percentage figure, the better the material resists permanent deformation under a given deflection and temperature range. The specimens are compressed by 50% at 100°C for 24 hr.

III. RESULTS AND DISCUSSION

A. X-ray Diffraction

XRD is a powerful technique to follow the intercalation of polymer chains into the silicate layers of clay and dispersion
of organoclay in the polymer matrix. XRD patterns of the NR75/BR25 samples containing 1, 3, 5 and 7 wt% is typically shown in Fig 1. As we know, the organoclay (Cloisite 15A) presents a peak at 20=2.9, which as deduced by the Bragg equation corresponds to an interlayer distance of 31.5 Å. Shift of the organoclay diffraction peak to lower 2θ values indicated that elastomer chains intercalated between consequent silicate layers. This clearly indicates that interlamellar spacing of the clay are enlarged after melt compounding [2].

![XRD results of NR/BR (75/25) containing a) 7 b) 5 (image)](image1)

Fig. 1. XRD results of NR/BR (75/25) containing a) 7 b) 5 c) 3 d)1 Wt% nanoclay

Moreover, XRD results of different compositions of NR/BR containing 5 wt% is shown in Fig 2. As can been seen, as the BR content of the samples increases, the organoclay diffraction peak shifts toward lower angles and consequently, the peak intensity declines. It can be attributed to the higher Mooney viscosity of BR resulting in a higher shear stress undergone by the matrix. This might help the deagglomeration of the clay stocks within the matrix [2].

![XRD results of NR/BR blends a) 100/0 b) 75/25 c) 50/50 d) 25/75 e) 0/100 containing 5 Wt% nanoclay (image)](image2)

Fig. 2. XRD results of NR/BR blends a) 100/0 b) 75/25 c) 50/50 d) 25/75 e) 0/100 containing 5 Wt% nanoclay

B. SEM Results

Further insights on the morphology of these materials are found in Fig. 6, where SEM images of cryogenically fractured and nanocomposite samples based on NR/BR (75/25) with and without clay are reported. As seen the fractured surface of pristine NR/BR (Fig 3-a) is smooth. The harsh surface of NR/BR filled compound by 7wt% nanoclay (Fig 3-b) exemplifies an effective interaction established between polymer matrix / filler [2], [15]. The SEM photomicrographs shown in Figure 6 confirm that smaller particle size of the filler provides a larger surface area for a better filler dispersion and interfacial bond between filler and rubber matrix.

![SEM photomicrographs of NR75/BR/25 containing a) no nanoclay b) 5 Wt% nanoclay (image)](image3)

Fig. 3. SEM photomicrographs of NR75/BR/25 containing a) no nanoclay b) 5 Wt% nanoclay

C. Permeability

The gas permeability of gum NR/BR vulcanizate and NR/BR/clay nanocomposites is presented in Table 2. As shown in Table 2, the nitrogen permeability reduces with an increased amount of clay. Compared with the gum NR/BR vulcanizate, the nitrogen permeability of NR/BR/clay nanocomposites with 1, 3, 5, and 7 Wt% clay reduces by 10.675, 9.651, 8.340 and 7.752 respectively. It can be concluded that the clay layers with the large aspect ratio and the planar orientation lead to the great increase of the diffusion distance by creating a tortuous path for the diffusing gas.
TABLE II: NITROGEN PERMEABILITY OF NR/BR/CLAY NANOCOMPOSITES 

<table>
<thead>
<tr>
<th>Material (NR/BR/Nanoclay)</th>
<th>75/25/0</th>
<th>75/25/1</th>
<th>75/25/3</th>
<th>75/25/5</th>
<th>75/25/7</th>
</tr>
</thead>
<tbody>
<tr>
<td>Permeability</td>
<td>12.1</td>
<td>10.675</td>
<td>9.651</td>
<td>8.340</td>
<td>7.752</td>
</tr>
</tbody>
</table>

D. Gel Content

Table 3 illustrates the gel content variation of NR/BR (75/25) nanocomposite samples containing 0, 3, 5, 7 and 10 wt% Nanoclay. As expected, addition of nanoclay particles results in an increase in the crosslink density of the samples and consequently a decrease in the gel content values. Similar trends were reported and observed by other researchers [2].

TABLE III: GEL CONTENT VALUES FOR NR/BR SAMPLES CONTAINING DIFFERENT PERCENTAGES OF NANOCLAY

<table>
<thead>
<tr>
<th>Material (NR/BR/Nanoclay)</th>
<th>75/25/0</th>
<th>75/25/1</th>
<th>75/25/3</th>
<th>75/25/5</th>
<th>75/25/7</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gel Content(%)</td>
<td>91</td>
<td>87.50</td>
<td>84.651</td>
<td>81.40</td>
<td>80.02</td>
</tr>
</tbody>
</table>

E. Rheological Properties

Rheological properties of nanocomposite materials is a great of interest and has been investigated by several researchers [2], [6], [8], [9]. The complex viscosity (η*) and elastic modulus (G') of prepared samples are shown in Fig 4. As seen, complex viscosity of prepared samples in sensibly higher than that of pure NR/BR. It can be attributed to the established interaction between polymer/filler. The falling trend of viscosity with frequency represents the pseudoplastic nature and shear-thinning behaviour of samples. At low and medium range of frequencies, the complex viscosity showed an increase indicating a yield stress. The existence of yield stress, has been indicated as a common characteristic of highly filled polymer melts, associated with interaction between the filler particles [2], [14]. Elastic modulus also increases with clay loading which would be ascribed to the good interactions established between the matrix components and organoclay functional groups [2], [14].

F. Compression Set

Compression set testing measures the ability of the rubber to return to its original thickness after prolonged compressive stresses at a given temperature and deflection. As a rubber material is compressed over time, it loses its ability to return to its original thickness. This loss of resiliency may reduce the capability of a rubber seal to perform over a long period of time. The resulting permanent set that a seal may take over time may cause a leak. Compression Set results for a material are expressed as percentage. The lower the percentage figure, the better the material resists permanent deformation under a given deflection and temperature range. The results of the compression set for the samples of NR/BR/Nanoclay are shown in Fig. 5. Comparison of the samples results indicates that unfilled NR/BR reinforced by 7 Wt% nanoclay has the lowest percentage of compression, which means that it resists permanent deformation better than the other samples.
then providing no elastic contribution to the recovery stage. In the case of NR/BR samples the nanoclay results in restriction in polymer chain movements and lower compression set. Alipour et al. [15] have reported that increased formation of effective network chains or crosslinking in the deformed state decreases compression set. It should be noted that the smaller the size of particles the lower the compression set could become. Therefore addition of nanoclay in the current study with resultant reduced compression set indicates such effects reported by previous researchers.

G. Dynamic Properties

To complete our exploration on the effects of nanoclay in a NR/BR rubber compound we looked also at certain dynamic properties such as the generation of heat under cyclic deformation. We have used the classic test of the Goodrich flexometer and the temperature increase was then divided by the hardness of the sample to have a heat index. The results of this observation for NR/BR (75/25) reinforced by 1, 3, 5 and 7 Wt% nanoclay is shown in Fig 6 and Fig 7. It can be observed that the heat build up decreases by adding the nanoclay. Thus, the nanoclay offers also the beneficial effect of being able to reduce the hysteresis of a rubber compound [15].

![Fig. 6. Value of heat build up for NR 75/BR 25 nanocomposites](image)

![Fig. 7. Values of heat index for NR 25/BR 75 nanocomposite samples](image)

H. Interactions Between Layered Silicates and Rubber

It was emphasized that better results were observed in organoclay partly due to better intercalation. From the results obtained, there were few possible ways that the clay could have interacted with the rubber matrix. Schematic diagrams are drawn in Fig 8 to illustrate the possible formation of composites from the interactions of layered silicates and rubber matrix. In the case of Fig 8-a, the rubber chains are not interacting with the silicates at all and a micro-rubber composite is formed. This is more like to occur when unmodified clay is used. In Fig 8-b, the rubber chains are partially intercalated into the galleries. Such ‘partial’ intercalation could be due to the following reasons:

- The viscosity of the rubber is too high and this restricts the movement of the molecular chains and prevents them from crawling into the silicate galleries.
- The vulcanization reaction is too fast and before the rubber chains can crawl into the galleries the movement is restricted by the formation of network. Thus, the intercalation only occurs at the entrance of the galleries.
- Since the method used for nanocomposite preparation is mechanical blending, the insufficient shear could cause the ‘partial’ intercalation.

Figure 8-c shows excellent interaction between the silicates and rubber as the rubber chains are completely intercalated between the silicate layers. From the interpretation of results, the type of rubber/clay nanocomposites obtained in our experiment is believed to be in between Figs 6-b and 6-c.

![Fig. 8. Schematic diagram of the possible interactions between layered silicates and layers](image)

I. The Reinforcement of Layered Silicates in Rubber

Reinforcement refers to the increase in stiffness and strength imparted to a vulcanize by particulate filler. It is dependent on the shape of hard domains and their interaction with the elastomeric matrix. However, the chemistry of elastomeric materials is very complex and they are capable of undergoing crosslinking to form a three-dimensional network when the right vulcanizing agent is added. The vulcanized or cured rubber is usually reinforced with fillers such as carbon black to enhance its properties and the reinforcement mechanism is known and rather straightforward to understand. However, the reinforcement of layered silicates is far more complicate...
owing to their unique structure and the interaction behaviour with the rubber matrix.

The rubber/clay nanocomposites in Fig 9-a would have superior mechanical properties as a constrained region is formed where the rubber chains have a restricted mobility. The failure is more likely to occur at the layered silicates (failure regions are circled). The properties of the rubber/clay nanocomposite in Fig 9-b depend greatly on the bonding between the layered silicate surface and rubber matrix and thus the failure is likely to occur at the interface of the silicates and the rubber matrix. The amount of layered silicates used and their homogeneity also have an influence on the reinforcement. Agglomeration tends to occur at higher loading of filler and results in the formation of weak spots. Thus, the composites would go through a maximum or critical filler loading before a drastic drop in properties is observed. The clay content used in the experiments was below the ‘critical’ amount as no dramatic drop was observed in the mechanical property results. Instead, the mechanical properties generally increased gradually as the clay content increased. Good dispersion of filler also plays an important role in composites. The mechanical properties which are obtained from an inhomogeneous composite are not the true properties. The reinforcement could be due to the original strength of the polymer matrix or the filler itself. Thus, good dispersion and distribution of the filler in the polymer matrix is critical.

**Fig. 9. Schematic diagram of the possible failure mechanism in nanocomposite samples**

**IV. CONCLUSION**

This work was devoted to investigate the rubber composition and nanoclay content on the properties of NR/BR/Cloisite 15A nanocomposites using different techniques. Experimental results of X-ray diffraction showed expansion of the distance between the silicate layers. Rheological properties, permeability and gel content values as well as compression set of the prepared samples received markedly improvement by clay loading. SEM photomicrographs showed that there is a strong interaction between polymer matrix and nano clay particles in NR/BR filled samples compared to the pristine NR/BR.

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**REFERENCES**


