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Structural and thermal degradation behaviour of reclaimed clay nano-reinforced low-density polyethylene nanocomposites

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Abstract

In this study, a novel reclaimed clay nanofiller was used to manufacture low-density polyethylene (LDPE)/Oil based mud filler (OBMF) nanocomposites by a melt compounding process. The mechanical testing samples were manufactured using injection moulding. The effect of reclaimed clay minerals influencing the crystallinity and the dispersion characteristics of this clay in LDPE matrix affecting the structural and thermal properties of the nanocomposites was investigated. It was observed that OBMFs were compatible with LDPE matrix which implies a strong interfacial interaction between the clay layers and polymer and that the influence of clay minerals present in OBMFs formed chemical bonds within the microstructure of the nanocomposites. The char yields of nanocomposites increased with OBMFs content. The $T_{D5\%}$ and $T_{D50\%}$ (onset degradation temperature at 5 wt% loss and 50 wt% loss, respectively) of the LDPE nanocomposite with 10.0 wt% OBMFs was the highest (27 °C higher in $T_{D50\%}$) among the nanocomposites. The tan δ spectra presented a strong influence of the filler contents on the relaxation process of LDPE and its nanocomposites. An enhancement of mechanical properties of composites was identified which showed a gain of 14% Young's modulus and a gain of 18% tensile strength at 10.0 wt% OBMFs loading compared to those properties of neat LDPE. The effect of filler dispersion in LDPE polymer matrix in relation to thermal stability was investigated and heat capacity data is employed to characterise changes in thermal characteristics relating to the nanomorphology of the materials.

Keywords OBMFs · Thermal stability · Thermal degradation · Crystallinity; rigid amorphous fraction; mobile amorphous fraction

Introduction

Incorporation of fillers in polymer matrix is an interesting area of research in recent years [1, 2]. This addition of fillers is not

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limited to saving costs, it improves the mechanical, thermal and flammability properties too [3]. The addition of fillers is function-specific [4, 5] such as adding clay in enhancing physical, thermal, and mechanical properties of polymer matrix [6]. Polymer/clay nanocomposites have drawn an enormous interest recently due to significantly influencing on the physical, thermal, mechanical and flame retardancy properties of nanocomposites [7–10]. Findings from the literature highlight that layered silicate fillers provide the most successful results by acting as nano-filler precursors [11–13]. However, the literature findings are very limited resulting in an information gap to understand the effect of this nanoclay in influencing the physical, thermal, mechanical, and flame retardancy properties of low density polyethylene (LDPE) polymer in details.

LDPE is produced on a large industrial scale and one of the most widely consumed polymer materials due to its excellent and desirable properties such as light weight, low cost, low dielectric constant and losses, high chemical resistance and easy manufacturing process [14, 15]. However, the use of LDPE is still limited due to its inherent chemical nature [16], poor thermal stability and flammability [17, 18]. Improvement of the thermal and flammability properties of synthetic polymeric materials such as LDPE is desirable to replace more traditional inorganic and natural polymeric materials [16, 19]. For this improvement, it is common practice to apply flame retardant additives within the polymer matrix. However, some flame retardant additives are considered harmful for the environment and hence have been phased out [20, 21]. Considering the stringent environmental regulations and the need to improve thermal and flammability properties of the polymers, development of an alternative environmentally friendly flame retardant material is becoming an area of polymer research interest [22].

Utilising industrial by-products as a filler in nanocomposite manufacturing is an area or research interest in recent years. Waste fly ash [23], waste iron ore tailing [24], waste pineapple leaves [25], coir [26], waste rubber powder [27], waste husks ash [28] etc. are predominantly considered as a sustainable source of filler for improving different properties in polymers. In view of the different influential effects of clay minerals on mechanical, thermal and flame retardancy behaviour of nanocomposite materials, the recovery of significant amount of nanoclays in waste drilling fluids warrant investigations. This study was inspired by the use of less explored LDPE polymer matrix incorporated with addition of reclaimed clay from oil based mud waste in the oil and gas industry as a nanofiller.

In oil and gas industry, driling through rock generally requires the use of some type of fluid to clear cuttings from the borehole formed by the drill. In some applications, the drilling fluid can be as simple as compressed air. However, when drilling is conducted to tap fossil fuel resources, the drilling fluid has to be an actual fluid [29]. The drilling fluid (also called drilling mud) is an essential component of the drilling process. In most deep drilling operation, oil based fluids (OBMFs) are used instead of water based fluids (WBFs) depending on the drilling conditions for efficient and cost-effective operations [30]. It is quite common practice to use both WBFs and OBMFs in drilling the same well, with WBFs for shallow portion of the well and OBMFs for deep drilling [31]. The well drilling process produces waste drilling fluid composed of drill cuttings and spent mud waste. When the drilling process is finished, the drilling fluid waste is composed e.g. of drilling fluid, cuttings, water, oil and many additives. The sources of drilling wastes largely depend on the cuttings based on the geological condition of the borehole, the depth of the well, and the fluid used in the drilling operation [32].

This study focuses on the reclamation and usage of layered silicates from oil based mud (OBM) waste, in particular their utilisation in engineering nanocomposite materials, specifically in low-density polyethene (LDPE) matrix. Detailed investigations were performed to identify the dispersion characteristics of OBMFs in LDPE polymer matrix based on the thermal analysis results, which also lead to highlighting the rigid amorphous fraction (RAF), mobile amorphous fraction (MAF) and percentage of crystallinity along with the mechanical property study.

Experimental

Materials

Lupolen 1800 S (trade name of LDPE) polymer was supplied by Northern Polymers and Plastics Ltd., UK. It has a melting point of 106 °C and a V-2 rating in accordance with UL 94 (vertical burning test) at 1.6 mm thickness. The spent oil based mud (OBM) was donated by a local oil and gas company. Montmorillonite, K10 powder was purchased from Sigma-Aldrich, UK and used as reference material in sample analysis.

OBMFs manufacturing process

Filler was recovered from OBM waste in a series of thermal treatments. The first treatment involved heating the OBM at 50 °C for 12 h to eliminate as much of the petroleum fraction as possible. The temperature was then increased to 80 °C for a further 12 h. Recovery of the solid content was enhanced by decanting oil-water mixture from the top of the sample. In the second stage of recovery, the furnace temperature was increased to 700 °C and held for a further 12 h to vaporise the remaining heavy oil fractions. Only the solid content of the OBM waste was recovered. The recovered mud was milled to a powder using ball mill (Ultra Turrax, IKA).

LDPE/OBMFs nanocomposite manufacturing process

LDPE/OBMFs nanocomposites were manufactured by using a commercially available LDPE and recovered OBM clay. The filler concentrations used in this study were are 2.5 wt%, 5.0 wt%, 7.5 wt% and 10.0 wt%. LDPE Polymer in pellet form and OBMFs were dried at 90 °C overnight prior to melt compounding. Melt compounding of different wt% of concentrations of OBMFs with LDPE was carried out using TwinTech Extrusion Ltd. 10 mm twin screw extruder at 60 rpm over five heating zones: 1st zone (120 °C), 2nd zone (200 °C), 3rd zone (210 °C), 4th zone (200 °C) and die/5th zone (200 °C). The compounded granules were prepared by using a pelletiser which were then injection moulded into a bar mould (dual cavity) for different



Fig. 1 Comparison of ATR-FTIR common scale spectra of LDPE and its nanocomposite

analyses using a temperature of 230 °C with a moulding pressure of 10 bar. The samples were then left to cool to room temperature before submitting to different analysis and characterisation processes.

Sample characterisation

To observe the morphology of OBMFs and their dispersion characteristics in LDPE polymer matricesx, a cryogenic sampling technique was has been followed for sample preparation. The fractured samples were gold coated using sputter deposition for 2 min prior to the

 Table 1
 ATR-FTIR peak assignments of LDPE and its nanocomposites

Wave number (cm^{-1})	Assignments		
2914	CH ₂ stretching		
2850	CH ₂ stretching		
1465	bending of CH ₂		
1377	inter layer carbonate ions		
1082	stretching of Si-O		
718	rocking of CH ₂		
640, 630, 608	oxide bands of metals like Si, Al, Mg, etc.		
525-580	δ (M-O-M) clay		

experiment. The fractured sample sections were observed using a Zeiss EVO LS10 Scanning Electron Microscope (SEM) with a magnification of 250x, 4.5 mm working distance and 25.00 kV accelerating potential. To determine the approximate elemental composition of the samples, energy dispersive x-ray analysis (EDXA) (Oxford Instruments INCA Energy) was carried out. The morphology of cryo-fractured samples were analysed using SEM and EDXA. Attenuated Total Reflectance-Fourier Transform Infra-Red spectroscopy was performed using a Thermo Scientific Nicolet iS10 FTIR Spectrometer setting for 32 scans between 4000 and 400 cm⁻¹. The influence of water vapour and carbon dioxide in air was minimised by subtracting peaks from blank measurement.

To identify the degradation and decomposition behaviour of neat LDPE and LDPE/OBMFs nanocomposites, Thermo-Gravimetric Analysis (TGA) was carried out using a TA TGA Q500 instrument. This analysis represented the weight variation of a measured sample on ramp mode from room temperature (31.1 °C) to 1000 °C at a rate of 10 °C per minute. To ascertain the glass transition, melting and crystallisation phases of the materials, Differential Scanning Calorimetry (DSC) was carried out using a TA Q100 instrument by adopting the heat-cool-heat method. The principle of this method is to determine the heat exchange difference between



Fig. 2 EDXA spectra of (a) LDPE; (b) LDPE+2.5 wt% OBMFs; (c) LDPE+5.0 wt% OBMFs; (d) LDPE+7.5 wt% OBMFs and (e) LDPE+10.0 wt% OBMFs

the sample and the reference pan (an empty aluminium pan). The temperature was set from -20 °C to 250 °C at a rate of 10 °C per minute.

XRD analysis was carried out on a Siemens D5000 diffractometer with Cu K α radiation ($\lambda = 0.15418$ nm) and 0.05° 2 θ step size. The scan angle covered the range from 3° to 60° 2 θ for layer materials and nanocomposites at a scan speed of 2° min⁻¹. The dispersion morphology of the nanocomposites was measured by Transmission electron microscopy (TEM), obtained on a JEOL JEM2010, 200 kV. An 80 nm thickness sample was prepared by using a Leica Ultracut-UCT microtome without any staining.

Dynamic shear measurements were obtained with a rheometer (AR 1000, TA Instruments) using a parallel plate geometry (8 mm diameter and 3.1 mm gap). Samples were tested to identify the viscoelastic properties as a function of temperature. The temperature range used in this study was varied from 0 to 90 °C, with a heating rate of 3 °C/min the samples were scanned at a fixed frequency of 1 Hz, with a strain of 0.2%.

Testing

Tensile and flexural (3-point bending) tests were carried out using Instron 3382 universal testing machine connected with Bluehill 3 software. The load was measured using a 100kN load cell. Tensile experiment was conducted using the dog-bone shaped samples which were prepared as per ASTM method D-638 and flexural experiment was conducted according to The European Standard EN ISO 14125. All tests were conducted at constant strain rate of 2 mm/min and the data presented here is the average of 5 samples.



Fig. 2 (continued)

Results and discussion

Elemental analysis of nanocomposite materials using ATR-FTIR study and EDXA

The ATR-FTIR spectrum analysis of neat LDPE and its nanocomposite has been carried out and the resulting spectra are presented in Fig. 1.

A summary of ATR-FTIR peak assignments is presented in Table 1 based on information published in the literature [16, 33–36].

The chemical structure of LDPE/OBMFs nanocomposites has been identified by using ATR-FTIR which shows the IR transmittance peaks at 2914 cm⁻¹ and 2850 cm⁻¹ corresponding to CH_2 stretching, the peaks at 1465 cm⁻¹ and 718 cm⁻¹ representing to the bending and rocking of CH_2 , respectively. The peaks at 1377 cm⁻¹ highlight the presence of interlayer carbonate ions. The peak at 1082 cm⁻¹ represents the stretching of Si-O, and the peaks at 640 cm^{-1} , 630 cm^{-1} and 608 cm^{-1} correspond to the oxide bonds of metals such as Si, Al, Mg etc. The peak at 580–525 cm^{-1} represents the crystalline forms of rare earth oxides which is observed in LDPE nanocomposite samples with 5.0, 7.5 and 10.0 wt% OBMFs content. It can articulate that the peaks at lower bands between 640 and 540 cm⁻¹ correspond to the chemical structure formed by different minerals present in the OBMFs nanoclay.

To identify the elemental composition and explain the addition of OBMFs in the LDPE polymer matrix, Energy Dispersive X-ray (EDXA) analysis was carried out as presented in Fig. 2a–e.

Figures 2a–e show the EDX spectra and also the elemental composition of neat LDPE, LDPE with 2.5 wt% OBMFs, LDPE with 5.0 wt% OBMFs, LDPE with 7.5 wt% OBMFs and LDPE with 10.0 wt% OBMFs nanocomposite. Elemental analysis of this filler has been reported in a Naxos 2018 conference proceedings which showed the presence of O, Na, Al, Si, S, Cl, K, Ca, Fe and Ba. The EDXA study here also confirmed the presence of different elements exist in clay minerals distributed in the LDPE matrix, as highlighted in Fig. 2. The lower limit of detection of EDXA is 0.01 wt%.

Morphology study of OBMFs and LDPE/OBMFs nanocomposite materials

To identify the surface topography of reclaimed clay particles from oil based mud waste and LDPE/OBMFs nanocomposite, morphological studies were carried out by using SEM. The SEM micrographs were used to analyse the changes in morphological structure of OBMFs in LDPE matrix.

Figure 3a shows the presence of tightly stacked clay platelets with size ranges up to 1000 nm. Figure 3b shows the cavity and uneven surfaces due to the cryogenic fracture of the sample. However, Fig. 3c shows that there are loosely dispersed OBMFs in the LDPE matrix which has higher Fig. 3 SEM images of (a) OBMFs; (b) neat LDPE; (c) LDPE with 2.5 wt% OBMFs; (d) LDPE with 5.0 wt% OBMFs; (e) LDPE with 7.5 wt% OBMFs and (f) LDPE with 10.0 wt% OBMFs



concentration of OBMFs, noticeable in the LDPE matrix in Fig. 3d. Figure 3e shows a mix of agglomeration of OBMFs and local scattered filler in LDPE matrix indicating intercalation of clay platelets. However, Fig. 3f shows poor dispersion of OBMFs with a remarkable agglomeration of OBMFs in the LDPE matrix.

Micro-structure by XRD and TEM

The XRD analyses are illustrated in Fig. 4 addressing the diffractograms at (a) wide angle X-ray diffraction (WAXD); (b) small angle X-ray diffraction; (c) comparing XRD patterns between MMT (as a reference) and OBMFs and (d) illustrating a few extra peaks for the LDPE polymer matrix potentially caused by adding these phases through OBMFs addition to the polymer matrix.

In Fig. 4, XRD patterns of LDPE and its nanocomposites are reported. A clear shift of the diffraction peaks of the planes (001) of OBMFs towards lower angles for the LDPE/OBMFs nanocomposites is noted. However, it can be seen that the basal spacing of OBMFs increases with different nanocomposites in different ratios. The d₀₀₁ spacings were calculated using Bragg's law $n\lambda = 2dsin\theta$, where λ is the wavelength of X-ray radiation used in the experiment, d corresponds to the distance between diffraction lattice planes and θ is the half of the diffraction angle. Analysing different peaks from these XRD data, it can be highlighted that the basal spacing increases with the increase of filler contents in nanocomposites. The diffraction peak of OBMFs was observed at $2\theta = 11.40^{\circ}$ which corresponds to a d-spacing of 7.75 Å. The d-spacings of LDPE with 2.5 wt% and 5.0 wt% OBMFs nanocomposites were identified at 9.6° and 5.6° which corresponds to the value



Fig. 4 Different XRD patterns of LDPE and its nanocomposites at (a) WAXD; (b) SAXD; (c) WAXD patterns of MMT and OBMFs and (d) potential peak initiation in LDPE polymer matrix due to addition with OBMFs

of 9.02 Å and 15.77 Å, respectively. Moreover the *d*-spacings of LDPE with 7.5 wt% and 10.0 wt% OBMFs nanocomposites were determined as 9.06 Å and 16.06 Å.

Analysing the *d*-spacing data of the corresponding peaks of reclaimed clay and LDPE/OBMFs nanocomposites, it can be highlighted here that the trend of *d*-spacings among OBMFs and LDPE/OBMFs nanocomposites are not consistent. However, the peak positions of LDPE with 5.0 wt% and 10.0 wt% OBMFs nanocomposites are noticeable. In both of these materials, the peaks of the planes (001) are shifted towards lower angles, which corresponds to the highest *d*-spacing value of these materials. This finding also agreed with the findings of the SEM (in Fig. 3) study. Based on this SEM study, it is important to mention here that LDPE with

10.0 wt% OBMFs nanocomposites indicates both local exfoliation and also shows agglomeration of clay minerals in the morphology study. Considering the data obtained in this XRD analysis, it can be inferred that OBMFs is exfoliated in LDPE with 5.0 wt% and apparently OBMFs might locally exfoliated in the LDPE polymer matrix. Transmission electron microscopy analysis is believed to clarify the nanomorphology in more detail in the next section.

TEM analyses were carried out to validate the results obtained from thermal and morphology analyses. It can be highlighted in Fig. 5a that the clay plates are barely connected with each other, representing the exfoliation nature of the clay platelets which is also noticeable in Fig. 5b. Figure 5b clearly shows the tendency of dispersion characteristic from



Fig. 5 TEM micrographs of (a) LDPE with 2.5 wt% OBMFs; b LDPE with 5.0 wt% OBMFs; c LDPE with 7.5 wt% OBMFs; d LDPE with 10.0 wt% OBMFs nanocomposite and (e) schematic diagram of nanoparticles orientation and tensile stress direction

exfoliation towards intercalation. It can be referred as the exfoliation-intercalation interphase.

In Fig. 5c, the loose but structured network of clay platelets is visible which indicates the intercalation nature of these platelets. However, the dark shades and spots in Fig. 5c highlight the agglomeration of platelets, although a few light spots indicate the local exfoliation of clay platelets.

Thermal properties by DSC and TGA

Non-isothermal DSC studies were conducted to elucidate the influence of novel filler OBMFs on the thermal degradation behaviour of nanocomposite materials. Different phases of degradation stages are shown in Fig. 6.

Analysing the DSC thermograms in Fig. 6a, it can be highlighted that there is not any significant change in the glass transition temperature of these nanocomposite materials compared to that of neat LDPE polymer. There is a minor change in glass transition temperature (less than 1 °C) in LDPE with 2.5 wt% OBMFs nanocomposite. However, the melting point remains almost unchanged, only a fraction of a degree Celsius change is noticeable in Fig. 6b. Figure 6c shows a consistent higher recrystallization temperature compare to that of LDPE. These three main heat flow curves in thermograms are discussed in more detail, addressing the potential different thermal properties possessed by these materials.

There is a distinctive peak present in neat LDPE and its nanocomposites in Fig. 6b which represents the melting behaviour of alpha phase crystals associated with the materials. There is not any associated heat curve shoulder noticeable corresponding to the absence of other crystalline phases or the other phases might be very weak compared to the presence of alpha phase in the materials. Considering the thermograms in melting peaks representing the heat capacity value of LDPE and its nanocomposites, the % of crystallinity can be identified using the following equation:

%of crystallinity =
$$[\Delta Hm - \Delta Hc]/\Delta Hm^* * 100\%$$
 (1)

Where ΔHm is the heat of melting, ΔHc the heat of cold crystallisation which is 0 as not present in this experiment ($\Delta Hc = 0$ in this case), and ΔHm° is a reference value if the polymer were 100% crystalline. All the units are in J/g and the value of ΔHm° is 293 J/g [37]. The % of crystallinity value of LDPE and its nanocomposites are presented in Table 3.

It is clearly evident that there is a decreasing trend of % of crystallinity as the percentage of filler content increases. The decreasing trend in the % of crystallinity degree of the nanocomposite materials can be explained by the presence of filler contents in polymer matrix, which can hinder the motion of the polymer chain segments and inhibit crystal growth.

The specific heat capacity value of LDPE and its nanocomposites have been identified by analysing the thermograms in Fig. 6c. The specific heat capacity value can be determined by the following equations:

$$C_p = \left(\delta Q / \delta T\right) \tag{2}$$



Fig. 6 DSC thermograms of LDPE and its nanocomposites at (a) Tg; (b) T_m and (c) Tc

$$C_p = (\delta Q/\delta t) \times (\delta t/\delta T) \tag{3}$$

Where Cp is the heat capacity in Joules per Kelvin (JK^{-1}) , Q is heat energy in Joule and T is the temperature denoted as °C or K. $\delta Q/\delta t$ represents the heat flow and $\delta t/\delta T$ corresponds to reciprocal heating rate. By using these two equations, the analysed specific heat capacities of neat LDPE and its nanocomposites are identified, as presented in Table 2.

The heat capacity data presented in Table 2 shows the negative effects of OBMFs in polymer matrix. However, LDPE with 2.5 and 7.5 wt% OBMFs show higher heat capacity property in nanocomposites, LDPE with 10.0 wt% shows the lowest heat capacity indicating a superior thermal conductivity property of this material among different nanocomposites. The heat capacity value decreases about 21% in LDPE with 10.0 wt% OBMFs nanocomposite.

In addition to the SEM observation, it is also important to understand that different interphases exist between LDPE and layered silicate nanocomposites. It is believed that there are two interphases present in LDPE polymer matrix and layered silicates in OBMFs. One of these interphases represents the interphases between the crystal fraction and amorphous fraction of the polymer matrix and the other is between filler and polymer matrix [38]. To understand the thermal degradation behaviour of any nanocomposite material, it is important to identify the different phases present in the polymer matrix predominantly crystalline and amorphous phases exist in the material. Analysing the heat capacity value, Cp in glass transition temperature, mobile amorphous fraction (MAF) can be identified by the following equation:

$$MAF = \Delta C_p / \Delta C_{pamp} \tag{4}$$

Where $\Delta C_p / \Delta C_{p \ amp}$ are the heat capacity increments at the glass transition temperature of LDPE and its nanocomposites and the pure amorphous LDPE polymer, respectively. Using the MAF value, rigid amorphous fraction (RAF) can also be identified by the following equation:

$$RAF = 1 - crystallinity - \Delta C_p / \Delta C_{p \ amp}$$
(5)

Table 2 Structural composition and thermal properties details of LDPE and its nanocomposites

Material	% of crystallinity	RAF = 1-MAF-CF	MAF ($\Delta C_p / \Delta C_{p(am)}$)	Specific heat capacity (<i>Cp</i>) $JK^{-1} kg^{-1}$
LDPE	16.71	0.13	0.70	4828
LDPE+2.5 wt% OBMFs	13.00	0.29	0.58	3990
LDPE+5.0 wt% OBMFs	12.48	0.28	0.59	3891
LDPE+7.5 wt% OBMFs	12.48	0.23	0.64	4160
LDPE+10.0 wt% OBMFs	11.07	0.31	0.58	3794

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 Table 3
 TGA analysis at

 different decomposition stages of
 LDPE and its nanocomposites

 with different clay contents

	T (1.0)	T (0,0)	5 4 / 6 5	
Material	T _{D5%} (° C)	T _{D50%} (° C)	D 1/2 Time	Residue (% wt) at 1000 °C
LDPE	381.27	448.42	45.03	0.22
LDPE+2.5 wt% OBMFs	321.24	413.14	44.34	1.27
LDPE+5.0 wt% OBMFs	336.26	427.18	43.79	4.01
LDPE+7.5 wt% OBMFs	313.73	395.60	43.81	5.40
LDPE+10.0 wt% OBMFs	340.83	449.72	45.68	7.24

Using the eqs. (4) and (5), the MAF and RAF values are identified and presented in Table 2.

It can be highlighted here that although RAF is non-crystalline, it does not participate in the glass transition due to the parts of the molecules are fixed due to the immobilisation of molecules in the polymer chain [38, 39]. The crystalline fraction in polymer chain, inorganic based filler (OBMFs) (assumed mostly in crystalline phase) and RAF incurs immobilisation in the polymer chain which is evident in the data presented in Table 2. It is also manifested based on the data presented in Table 2 that the RAF value is increased more than 100% in nanocomposites compared to that of neat LDPE. The reason can be explained that the incremental filler content acts as a nucleating agent and increases the nucleation ratio.

Thermal degradation of LDPE/OBMFs nanocomposites has been studied under a dry nitrogen environment using a TA instrument TGA Q500.

The degradation scenario of these materials at different heating stages are analysed and the key findings are presented in Table 3.

The onset degradation temperature of these materials are identified at weight % losses of 5% and 50%. In both cases,



Fig. 7 TGA thermograms of LDPE and its nanocomposites at: a onset degradation at 5% weight loss; b onset degradation at 50% weight loss; c D-half (50% weight loss) time and (d) residue at 1000 $^{\circ}$ C



Fig. 8 Variation of (a) storage modulus, b loss modulus and (c) damping intensity $(\tan \delta)$ with temperature of LDPE and its nanocomposites

LDPE with 10 wt% OBMFs shows the highest onset degradation temperature among the nanocomposites with different wt% filler contents which represents LDPE with 10 wt% OBMFs nanocomposite possessing the highest thermal stability. However, the thermal stability property is significantly affected in LDPE with 7.5 wt% OBMFs. Considering the lowest temperature at D-half (50% weight loss), the presence of peak assignments representing amorphous phase in FTIR analysis and heat capacity value from DSC analysis, it can be inferred here that LDPE with 7.5 wt% OBMFs possesses a good heat capacity property. It is also noticeable that the addition of OBMFs into LDPE matrix significantly reduces the thermal stability of the material based on $T_{D5\%}$ and $T_{D50\%}$ data.

However, the degradation trend is distinctly noticeable as it does not show any significant changes of weight loss % until 300 °C. Degradation mainly occurred between 300 and 500 °C and the poor thermal stability of nanocomposites is clearly visible except for LDPE with 10 wt% OBMFs nanocomposite. In Fig. 7c, LDPE with 10 wt% OBMFs shows the longest time needed to reach the D-half point which also indicates superior thermal stability property among the samples. It is also manifested in this figure that at this D-half point LDPE with 10 wt% OBMFs shows the highest temperature which gives an indication of improving of thermal conductivity property of this material. It is also verified with the lowest heat capacity value obtained from DSC study. It is articulated here that the incremental filler contents may have a strong influence on the thermal stability of LDPE/OBMFs nanocomposites. Filler contents up to 10 wt% reduce the thermal stability of the materials whereas LDPE with 10 wt% possesses the best thermal stability and thermal conductivity properties among the nanocomposites with different wt% filler content.

Dynamic mechanical properties

The storage modulus (E') represents the load bearing capacity of a material. The effect of OBMFs in influencing this storage modulus is graphically enumerated in Fig. 8a. It is noticeable from Fig. 8a that there was a significant decrease in the storage modulus of LDPE/OBMFs nanocomposites compared to the storage modulus of neat LDPE matrix. This is perhaps due to the decrease in the stiffness of LDPE matrix with the reinforcing effect of the fillers prone to decrease in stress transfer at the interfacial surface. As noticeable in Fig. 8a, the storage modulus decreased drastically with the increase in temperature and the degree of storage modulus drop was significant in the temperature regions between 20 and 70 °C.

The loss modulus curves of neat LDPE and its nanocomposites are presented in Fig. 8b which shows a relaxation peak at 50 °C in neat LDPE decreased to lower temperatures for nanocomposites such as the peaks were at 25 °C, 10 °C, 5 °C and 15 °C for LDPE with 2.5, 5.0, 7.5 and 10.0 wt% OBMFs respectively. There is a significant changes in loss modulus noticeable in the temperature regions between 40 to 55 °C which clearly shows the loss modulus is higher for LDPE with 2.5 and 5.0 wt% OBMFs compared to that of neat LDPE. However, the loss modulus decreased in LDPE with 7.5 and 10.0 wt% OBMFs in those temperature regions. This trend is also clearly shown between 70 to 90 °C temperature regions. This trend completely agreed with the tensile results mentioned in the next section. The equipment which we used for rheology analysis (TA AR1000) is associated with an open hot plate which is not environmentally controlled and we assume the temperature reading in the software may be not the same as the actual temperature in the sample due to the heat absorption by the samples and heat dissipation from the hot plate to open air. It is noticeable that loss modulus reduced drastically from 40 °C which we assume is nearly in the same room temperature the tensile tests were performed at 25 °C.

The ratio of loss modulus to storage modulus is measured which is graphically presented in Fig. 8c to identify the damping properties of the materials. The damping peak in the nanocomposites showed an increased magnitude of tan δ in comparison to neat LDPE. This graph represents the balance between the elastic phase and viscous phase in the polymeric structure. It is noticeable the maximum peak of neat LDPE at 50 °C which is assumed Tg of neat LDPE shifted significantly to lower temperatures for LDPE/OBMFs nanocomposites (Fig. 9).

Tensile properties of LDPE/OBMFs nanocomposites

The mechanical properties of LDPE/OBMFs nanocomposites are presented in Table 4.

It can be seen that the addition of 10.0 wt% OBMFs can improve tensile strength and elongation at break by 17.5% and 13.7% respectively. However the tensile strength and elongation at break of other nanocomposites are remained almost same as tensile strength and elongation at break of neat LDPE. There is an increasing trend

of improving tensile modulus among nanocomposites, but only LDPE with 7.5 wt% and 10.0 wt% nanocomposites show higher tensile modulus than that of neat LDPE.

The improvement in mechanical properties of Polyethylene/clay nanocomposites were reported by using compatibilisers by different researchers [40-42]. However, in our recent studies on LDPE/OBMFs nanocomposites, it is clearly noticeable that there is a little but consistent improvement in tensile property among LDPE/OBMFs nanocomposites. A slight decrease in the tensile strength and modulus can be observed when the OBMFs loading increases from 0 to 5.0 wt%. The significant enhancement in the tensile strength and modulus were obtained with 7.5 wt% and 10.0 wt% OBMFs loading. In addition the decrease trend of elongation at break may be attributed to a reduction in deformability of the rigid interface between nanofillers and matrix [43]. The OBMFs used in this study are nanoparticle in considering size and shape, but observing the micrographs in Fig. 5 it can be highlighted that without adding any grafting compatibilisers, the OBMFs nanoparticles are uniformly dispersed and no agglomeration is noticeable in LDPE with 2.5 wt% and 5.0 wt% fillers loading which are difficult to visualise in SEM images in Fig. 3. However, the orientation of nanoparticles due to the OBMFs compatibility facilitate to act this nanoparticle in a similar manner as fibre which is highlighted in Fig. 5e. After observing carefully the TEM micrographs in Fig. 5a-d, schematic diagrams are presented in Fig. 5e considering the applied force in tensile test and the direction in nanoparticles orientation. It is noticeable that the gradient between nanoparticle orientation and crack propagation plane is decreasing with the increase of filler contents in LDPE matrix which may be attributed to an improvement in materials strength predominantly the fracture strength.



Fig. 9 Tensile stress-strain curves of neat LDPE and its nanocomposites **Table 4** Mechanical properties ofLDPE and its nanocomposites

Sample	Tensile modulus (MPa)	Tensile strength (MPa)	Elongation at break (%)
LDPE	28 ± 1.1	8.0 ± 0.4	66±4
LDPE+2.5 wt% OBMFs	26 ± 1.6	7.5 ± 0.2	64 ± 6
LDPE+5.0 wt% OBMFs	27 ± 1.4	7.2 ± 0.1	62 ± 3
LDPE+7.5 wt% OBMFs	31 ± 1.1	7.9 ± 0.4	61 ± 5
LDPE+10.0 wt% OBMFs	32 ± 1.2	9.4 ± 0.1	57 ± 4

Conclusions

A series of the LDPE/OBMFs nanocomposites with OBMFs content ranging from 2.5-10 wgt% has been prepared by using a melt-blending technique in a twin-screw extruder. The structure of the LDPE/OBMFs nanocomposites was presented using XRD, SEM and TEM. The thermal degradation behaviour of the LDPE/OBMFs nanocomposites was demonstrated by using DSC and TGA. The results of XRD, SEM and TEM indicate that the LDPE/OBMFs nanocomposites are a hybrid material consisting of OBMFs nanoplatelets and LDPE polymer matrix. Moreover, it is highlighted that the OBMFs nanoplatelets exist as aggregates and contribute to different dispersion behaviour with a change of OBMFs content in the LDPE matrix. In this study, OBMFs shows exfoliation characteristics with LDPE with 5.0 wt% OBMFs nanocomposites whereas these show structure intercalation in LDPE with 7.5 wt% OBMFs. In addition, OBMFs create a mix of agglomeration and local exfoliation which was detectable by thermal analysis.

The thermal properties of the LDPE and its nanocomposites were studied by means of DSC and TGA. No changes in melting and recrystallization temperatures due to the addition of OBMFs in the polymer matrix were detected. However, there was a decrease in the percentage of crystallinity noticeable with an increase in filler content. In addition, heat capacity reduced in nanocomposites compared to the neat LDPE matrix which indicate the inclusion of OBMFs in LDPE improves the thermal conductivity of these nanocomposites in general. LDPE with 10.0 wt% OBMFs nanocomposites showed the lowest heat capacity (about 21% reduction) addressing the superior thermal conductivity of this material. A significant increase of thermal stability was also achieved with LDPE with 10.0 wt% OBMFs nanocomposite. Furthermore, the viscoelastic investigation revealed the storage modulus, loss modulus and damping intensity properties of materials which also agreed with the mechanical properties of the material. LDPE with 10.0 wt% of OBMFs content showed both thermal stability and superior tensile properties compared to those properties of neat LDPE and LDPE with 2.5, 5.0 and 7.5 wt% OBMFs content nanocomposites.

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