Sulfur nanoparticles transform montmorillonite into an inorganic surfactant applicable in thermoplastics processing

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1. Introduction

Montmorillonite (MMT) based polymer nanocomposites usually have enhanced mechanical, thermal, barrier and flame-retardant properties in comparison to pristine polymers [1,2]. MMT was extensively used as filler for preparing polymer/MMT nanocomposites due to their unique properties like nanometer scale thickness and high aspect ratio [3,4]. MMT is a 2:1 phyllosilicate mineral with two tetrahedral silica sheets that sandwich a central alumina octahedral sheet [5]. The effective dispersion of MMT filler into the polymer matrix is a key factor in determining the final properties of the polymer/MMT nanocomposites. High-density polyethylene (HDPE) is the most commonly used thermoplastics due to its regular chain structure, high melting temperature, excellent thermal and electrical resistance, good tensile strength, low cost and high abrasion resistance [6-8]. Unfortunately, HDPE is a hydrophobic viscously melted material, which leads to incompatibility while blending with MMT with hydrophilic properties. Therefore, it is very important to promote the adhesion between the HDPE and MMT, which can be done by either modifying HDPE or MMT. However, modifying HDPE can result in obvious increase in cost and complexity of preparation process. Thus, modification of MMT is useful approach to improve the compatibility between MMT and HDPE. As MMTs are hydrophilic in nature, it is compatible with hydrophilic polymers. In order to be compatible with HDPE, the hydrophilic MMT lamellae must be hydrophobized. Commonly, this surface modification is done by ion exchange reactions with cationic surfactants such as alkyl-ammonium cations. These cations not only lower the surface energy of the clay but also provide functional groups that can react with the polymer. However, the marginal thermal stability of the alkyl-ammonium cations leads to the degradation of polymer/MMT nanocomposites [9,10].

In this work, we prepared organophilic MMT by adding Sulfur as an inert and cheap inorganic surface modifier to the MMT lamellae. The surface adsorption of the Sulfur particles on the clay lamellae was carried out with negligible amount (~0.03 wt%) of chitosan macromolecules and the obtained filler material with composition dependent hydrophobicity was blended in HDPE polymer matrix. The as-prepared MMT/HDPE composites were characterized by XRD, TEM, SEM, zeta...
potential, static contact angles, TGA-DSC, tensile strength, particle charge detector and X-ray micro-computed tomography analysis. We found that, the CB additive of the HDPE was sufficiently dispersed by adding small amount of the inorganic MMT/S surfactants to the melted CB-containing HDPE during the blending process.

2. Materials and methods

2.1. Preparation of sulfur hydropbized montmorillonite composite particles

To obtain sulfur nanoparticles (SNPs) by a modified Raffo-method, 0.03 g of Sulfur powder (Sigma Aldrich) was dissolved in 4 ml of 10 M aqueous NaOH solution (solution pH = 14.15) at a temperature of 80 °C. Afterwards, the liquid was stirred and diluted with 1 ml of 10 M NaOH solution, then 4.65 ml of 2.5 M H2SO4 solution was added dropwise until the initial solution became turbid (pH = 13.95 at precipitation) due to the formation of sulfur nanoparticles (c = −0.09 g/100 ml).

The precipitated sulfur was centrifuged (8000 rpm, 5 min), thoroughly washed with deionized water and dried overnight at 60 °C. The SNPs were then redispersed in deionized water (c = 0.1 g/100 ml), then 200 μl of 0.01 g/100 ml high molecular weight chitosan (Chit; Mn = 310–375 kDa; Sigma) solution (dissolved in 0.13% aqueous acetic acid; pH~4.3) was added to the dispersion to turn the streamlining potential to positive. To form MMT/S composite, different amounts of aqueous EXM-838 montmorillonite (Süd-Chemie AG) dispersion (c = 0.03 g of Sulfur powder (Sigma Aldrich) was dissolved in 4 ml of 10 M aqueous NaOH-solution (solution pH = 13.95 at precipitation) due to the formation of sulfur nanoparticles (c = −0.09 g/100 ml).

The apparent static contact angles on MMT/S layers were measured according to the sessile drop technique at 25.0 ± 0.5 °C under atmospheric pressure, applying an EasyDrop drop shape analyser (Krüss GmbH, Hamburg, Germany) equipped with DSA100 software, a Peltier temperature chamber and a steel syringe needle of 0.5 mm diameter, and using deionized water as test liquid.

The DSC curves of the MMT/S composites were recorded applying Mettler Toledo DSC822e device. In the tests, the solid powder samples were heated from 25 °C to 150 °C and cooled down to 25 °C again at a rate of 5 °C·min⁻¹ in sealed aluminium sample holders with a hole at the top. Dry compressed N2 was used as a carrier gas at a flow rate of 50 mL·min⁻¹. The obtained thermoanalytical data was processed using STAR® software.

The thermogravimetric analyses were performed on a TA Instruments TGA Q500 thermogravimetric analyzer equipped with a standard furnace. 1–5 mg of the samples were analyzed in platinum pans at a heating rate of 10 °C·min⁻¹ to 600 °C in an atmosphere of air flowing at 60 mL·min⁻¹.

The tensile tests were performed using a Galbadini Quasar 100 tensiometer according to the ISO 527 procedure, using dumbbell specimens with a gauge length of 108 mm, a width of 10 mm and a thickness of 4 mm. The stretching tests were performed at room temperature at 10 mm·min⁻¹. All tests were carried out on a minimum of 5 samples and the reported results are average values.

The size distribution of carbon black particles in the HDPE matrix was obtained via X-ray micro-computed tomography analysis, which was conducted using the X-ray Microtomography equipment (Skyscan 2211). The sample dimension of ~8.0 × 2.0 × 1.0 mm³ was scanned using 11Mp cooled CCD camera by applying the source voltage of 60 kV and the source current 800 μA with an exposure time of 650 ms. The voxel size of this dataset was 1 μm. NRecon reconstruction software was used to reconstruct the projected images with the pixel size of 4032 × 2688 and Ctan and CTvol software were used to represent the 3D models. Important to note that the smallest particles what we see with CT under these parameters is 216 μm³. Although aggregation of clay minerals is inevitable, we believe that they do not exceed 100 μm³ thus these particles are invisible and CB particles are visible during CT measurement. Only particles larger than 6 * 6 * 6 voxels were considered in the calculation.

For the optical characterization of polymer samples, diffuse reflectance UV–VIS spectra were recorded applying a CHEMKRON UV–VIS (USB2000–UV–VIS, Ocean Optics Inc) spectrophotometer, equipped with an integrating sphere.

3. Results and discussion

3.1. Structural characterisation of composite nanoparticles

The hydrophobization of the different polymer filler materials with initial hydrophilic properties is a possible route to provide improved mechanical properties to polymer based composites through increasing compatibility between the hydrophobic polymer matrix and the initially hydrophilic filler materials [11]. In this study the EXM-838 MMT clay sheets as well-known polymer filler material was hydrophobized with cheap and easy-available inorganic elemental S. To achieve this goal, S
NPs with a maximum diameter of 40 nm were synthesized applying a simple precipitation method. Despite there are many reported bottom-up routes to prepare S NPs, including e.g. the Weimarn [12], and Raffo-methods [13] or the sulfur vapour deposition on cold water [14], only the Raffo-method offers particle yields high enough (~600 g/l) to be industrially profitable besides requiring more environmentally friendly aqueous media [15]. Our version of the applied Raffo-method is
based on the following chemical reactions:

\[
4 \text{S} + 6 \text{NaOH} = \text{Na}_2\text{S}_2\text{O}_3 + 2 \text{Na}_2\text{S} + 3\text{H}_2\text{O} \quad (1)
\]

\[
\text{Na}_2\text{S}_2\text{O}_3 + \text{H}_2\text{SO}_4 = \text{H}_2\text{S}_2\text{O}_3 + \text{Na}_2\text{SO}_4 \quad (2)
\]

\[
x \text{H}_2\text{S}_2\text{O}_3 = y \text{H}_2\text{O} + z \text{SO}_2 + \text{H}_2\text{SnO}_6 \quad (3)
\]

\[
\text{H}_2\text{SnO}_6 = \text{H}_2\text{Sn}_{n-1}\text{O}_6 + \text{Sn}_n \quad (n > 5) \quad (4)
\]

The core step of all Raffo-methods is the colloidal S formation, happening upon the acidification of \(\text{Na}_2\text{S}_2\text{O}_3\) solutions with sulfuric acid (4. reaction step). We obtained our \(\text{S}_2\text{O}_3^{2-}\)-containing solution as a result of dissolving elemental S powder in 10 M \(\text{NaOH}\)-solution at 80 °C, then S-nanoparticles with a maximum mean diameter of 40 nm formed upon the following addition of 2.5 M \(\text{H}_2\text{SO}_4\) solution. According to Equations (1)-(4), the dissolution of sulfur in such media is a disproportionation reaction, while the reverse precipitation reaction is a synproportionation. According to the literature [15], the \(x, y, z\) stochiometric coefficients can vary, depending on the reaction conditions, and the \(m\)-value is not well defined by them because polythionates with different chain-lengths can form in the reaction. However, in our case as we aim for the synthesis of sulfur nanoparticles the exact constitution of polythionates and sulfur homomolecules is less relevant and therefore were not studied in details.

As the TEM images and the corresponding histogram in Fig. 1 shows, the size distribution of the obtained spherical sulfur nanoparticles is relatively polydisperse. The mean diameters range between 1 and 40 nm. This size is comparable to the size of the lamellar EXM-838 MMT lamellae with ~2 nm thickness and lateral size of about few hundred nanometers.

The fixation of the S NPs on the surface of the MMT particles was executed exploiting electrostatic interactions. Despite elemental S does not contain ionic functionalities, Coehn’s rule implies that the surface of low permittivity S particles is slightly negatively charged in aqueous medium (point of zero charge at pH = 2). As MMT lamellae also have negative surface charge excess even in weak acidic media (point of zero charge at pH = 4–5.9 [16,17] we used positively (in pH = 4.3 solution) charged chitosan biopolymer macromolecules (Chit; \(\text{Mw} = 310–375\) kDa) as cationic binding agent between the negatively charged inorganic components.

As the characteristic particle charge titration curve of Fig. 2a shows, the initial streaming potential (~1010 mV) of the 0.1 wt% S NP dispersion was continuously increased upon the addition of 0.01 wt% chitosan solution and at about 30 μl added volume it was reached the 0 mV potential value (charge neutralization point). However, it also can be seen that the initially negatively charged S NPs can be overcharged by the further addition of the cationic polyelectrolyte. In the second step 0.1 wt% MMT aqueous dispersion was added dropwise to this overcharged S based dispersion resulting decreasing streaming potentials and the charge neutralization was occurred after the addition of 220 μl.
MMT solution (Fig. 2b). It is important to note that considering the concentrations and added volumes of the components, the composite contains less than 0.03 wt% Chit polymer, which means that we could completely ignore the presence of Chit during sample characterization.

3.2. Structural and interfacial properties

In the previous chapter it was presented that the S NPs overcharged with negligible amount (<0.03 wt%) of chitosan macromolecules could be successfully immobilized on the surface of MMT lamellae via electrostatic interactions. In the continuation, we systematically changed the ratio of the components to get samples with varied behaviors.

To study the crystalline structure of the resulted MMT/S composite powders, X-ray diffractograms were recorded. Fig. 3 shows the diffractograms of initial S, MMT and their 1:1 composite. As the sulfur-content increases, the characteristic peaks of MMT at $2\theta = 7.54^\circ$, $19.78^\circ$, $22.20^\circ$, $26.60^\circ$, $28.32^\circ$, $35.00^\circ$ and $36.08^\circ$ [18,19] decrease in intensity and disappear, while other peaks appear, attributed to the orthorhombic $\alpha$-sulfur (S) (with higher intensities: 23.10$^\circ$ [222], 25.88$^\circ$ [026], 26.76$^\circ$ [311], 28.96$^\circ$ [313]) [20]. Moreover, some peaks are characteristics to monoclinic $\beta$-sulfur (S-$\beta$) (27.74$^\circ$, 31.08$^\circ$, 37.84$^\circ$) [21]. The corresponding Miller-indices of the identifiable reflections are indicated in Fig. 3 however, in the case of MMT, only the [001] crystal plane was identified according to JCPDS card 29–1499 in the absence of other related literature. These observations indicate the crystalline nature of the S NPs, since no broad bands of amorphous phase were detected. As it can be seen in the inserted pictures of Fig. 3, S NPs are barely dispersible in water, but the addition of hydrophilic MMT resulted in better water dispersibility.

As the original reflections of MMT are barely identifiable in the XRD patterns of the composites, we propose complete or at least partial delamination for MMT. The interlayer spacing of unfunctionalized swollen MMT in aqueous medium ranges from 1 to 2 nm [22,23], and as Fig. 1 shows, a little less than 10% of the detectable particles have 2 nm or smaller mean diameters, therefore the intercalation of smaller polyelectrolyte-covered S NPs is theoretically possible.

Zeta potential measurements (Fig. 4.) showed that the initially lower negative zeta potential values (~16 mV) of MMT lamellae exceed 0 mV as the S content of the composites increases. The decrease in zeta potential also indicates the increasing S-coverage of clay lamellae.

The above presented results indicated that the presence of the sulfur nanoparticles changed the wetting and structural properties of MMT composite filler material which was also evidenced by contact angle measurements (Fig. 5). As the sulfur content increases, the spray-coated composite films (with $1 \pm 0.5$ mg/cm$^2$ specific layer mass) changed their wetting character from hydrophilic with water contact angle (WCA of 50.1$^\circ$ at 0% S NPs) to hydrophobic (WCA = 130.0$^\circ$ at 100% S NPs), with a sharp contact angle-increase at 80% S NP content.

This composition dependent wetting characteristics was also evidenced by DSC measurements. The DSC-curve of initial MMT in Fig. 6 shows a broad endothermic peak ($T_{end}=101.2^\circ$ C, referring to the evaporation of the adsorbed water content), while this peak is not present in the DSC-curves of the 80% MMT/S composite and S, indicating the increased hydrophobicity. Besides, as S content increases, the characteristic peaks of S appear in the curves: the $\alpha \rightarrow \beta$ transition gives an endothermic peak at $\sim 110^\circ$ C, while the endothermic melting is displayed at the 115–120 $^\circ$C range [24]. Upon cooling, the exothermic peaks of sulfur solidification and/or recrystallization are also presented [24].

As evidenced by SEM-images and energy-dispersive X-ray spectroscopy (EDX) measurements, the surface of the MMT/S composite with 80 wt% S content is microstructured (Fig. 7.) and have a relatively...
homogenous and even distribution of both components.

The elemental composition was also determined based on EDX-spectral data: as Fig. 7b shows, the measured 75.45 ± 5.95% sulfur-content relatively well matches the nominal 80 wt%. These observations indicate even and nearly lossless electrostatic deposition of S NPs with 1–40 nm primary particle size on the surface of barrier MMT lamellae platforms with lateral size of about few hundred nanometers.

To conclude the results, we managed to hydrophobize the MMT lamellae by fixing water insoluble and inorganic S NPs on their surface utilizing electrostatic interactions. The hydrophobicity of the composites sharply increased at 80% S content, while the dispersibility in water and zeta potential absolute values dropped at this composition.

Considering the above presented results at 80 wt% S content the MMT/S composite has relatively high surface charge (ca. −11.6 ± 1.8 mV zeta potential) which is due to the accessible hydrophilic MMT surfaces and it also shows moderate hydrophobicity (WCA = 90.3 ± 7.2°) because of the surface immobilized S NPs.

With the resulting amphiphilic character, the prepared composite particles could potentially be applied as Pickering-type emulsifiers or surfactants, as they may provide a more effective stabilization alternative than molecular surfactants due to their resistance to coalescence and Ostwald-ripening [25]. According to our idea this surfactant-like amphiphilic character of the inorganic composite could be useable for further plastic applications, because it could enhance the interfacial interactions between the different polymer filler materials and the continuous polymer matrix.

To study this assumption, the inorganic composite with 80 wt% S content was added to a commercially available PE with 0.5 wt% CB content and the dispersion degree of the incorporated CB was

| Samples  | Melting point [°C] | Crystallization temp. [°C] |
|----------|--------------------|-----------------------------|
| C-PE     | 133.7              | 112.9                       |
| C-PE-P   | 129.3              | 115.9                       |
| C-PE-C   | 129.0              | 116.2                       |

Fig. 11. TGA curves of (A) C-PE (B) C-PE-P (C) C-PE-C.

Fig. 12. DSC curves of (A) C-PE (B) C-PE-P (C) C-PE-C.

Fig. 13. Tensile strength of the samples.
characterized. As it is reported, many polymers are unsuitable for outdoor use on their own because of rapid degradation through the action of visible and especially UV light [26]. However, CB absorbs both visible and UV radiation converting it to heat and thus makes an excellent stabilizer. As the UV-VIS diffuse reflectance spectra of Fig. 8 show, the 0.5 wt% CB-content in our samples has a considerable contribution to the UV-absorbance, as well. In many cases stabilities spanning decades can be obtained, making the polymers suitable for outdoor applications. UV stabilization by CB in polymers depends on its ability to absorb and to backscatter light. While the absorption increases with smaller particles and aggregates, the backscattering decreases and thus the UV stabilization requires small and evenly dispersed CB particles in the continuous polymer matrix. Therefore, the effect of MMT/S surfactant-like additive on the dispersion degree of CB in HDPE polymer matrix was investigated.

The 3-dimensional structure of the polymers and the size distribution of the embedded carbon particles were studied by computed tomography (micro-CT) measurement. The representative CT-images in Fig. 9 show the incorporated CB particles in HDPE matrix. The corresponding particle size distributions of the examined specimens are presented in Fig. 10., while the average particle size values obtained are indicated in Table 1. As it appeared in the pictures and from the data, the average size of the particles are really similar in a case of the C-PE and C-PE-P samples, but in the case of the C-PE-C composite with the loading of the as-prepared MMT/S (with 80 wt% of S NP content) value is become significantly smaller. In parallel, the average particle size was significantly decreased indicating higher distribution (small and evenly incorporated particles) of CB in HDPE. The difference between the C-PE-P (simply physical mixture of S and MMT) and C-PE-C (electrostatically adhered S NPs on MMT) containing samples indicates that only this latter one behaves like a surfactant, the physical mixture is not suitable for increasing the dispersion degree of CB. There are documented cases in the literature, which are in good accordance with our results: in an example, the addition of max 6.8% hydrophobized (di-methyl, dihydrogenated tallow-) MMT as a dispersing additive could increase the dispersion degree of multicomponent (matrix: polypropylene; disperse phase: ethylene-octene based elastomer) thermoplastics blends. The authors reported a decrease in the size of the dispersed elastomer particles with increasing MMT-content and fixed elastomer concentration: besides the amphiphilic properties, the addition of functionalized MMT resulted in viscosity build-up, which also contributed to the dispersibility-increase through the retardation of polymer-melt coalescence during mixing [27,28]. While the viscosity build-up is generally considered as a hindering factor during the blending process, in this case it can result in beneficially high dispersion degrees, as well.

In line with this, the particle density also changes, which nearly tripled in the C-PE-C sample compared to the C-PE or C-PE-P samples. (Table 1). These results suggest that the MMT/S nanocomposite (C-PE-C) obviously increase the distribution of the carbon particles in the continuous polymer matrix and decrease their aggregation. This structure with evenly and continuously dispersed CB particles is advantageous in the viewpoint of UV stabilization of the polymer.

TGA was used to study the thermal stability of the HDPE composites (Fig. 11.). It was showed that pure HDPE is thermally stable up to 250 °C. At 250 °C it starts to degrade and fully degraded at 470 °C. Adding MMT/S additives to HDPE has no significant effect (Table 2), only a slight thermal stability was observed as the C-PE-P and C-PE-C samples started to decompose at 240 °C.

DSC of HDPE shows an endothermic process (melting of the polymer) at 134 °C and an exothermic crystallization at 113 °C [29]. The DSC curves for C-PE-P and C-PE-C displayed slightly lower melting and higher crystallization temperature compared to the C-PE composite (Fig. 12.). This indicates that, the addition of MMT and S to the C-PE does not change appreciably the melting and crystallization temperature of the resulting polymer. Considering the related literature, the increase in crystallization temperature can be attributed to the nucleating effect of MMT and S [30].

The tensile properties of C-PE-C, C-PE-P, C-PE-C samples are shown in Fig. 13. The tests revealed that the additives did not significantly alter the tensile strength of HDPE in either case.

In addition to the introduced beneficial properties of the MMT/S polymer additive, it is important to mention, that the composite offers a way cheaper alternative to the commercially available organic surfactants. For instance, the current (2020 January) average bentonite clay and sulfur prices are at approximately 77 and 140 U. S. dollars per ton, respectively. The prices of the synthetic and bio-based organic surfactants depend on their type (anionic, non-ionic, cationic, and amphoteric) and applications (detergents, oil industry, personal care, textile, elastomers & plastics, crop protection, food & beverage, etc.), however, their price is at least one order of magnitude more expensive (>1100 U. S. dollars per ton). Furthermore, as elemental sulfur is an abundant byproduct of the petrochemical industry, and MMT is easily accessible, our composite can reduce the ecological footprint of thermoplastics production, as well.

4. Conclusion

Sulfur modified MMT with different compositions were blended with Carbon Black-containing HDPE composite by different methods. An electrostatically adhered S NPs on MMT composite with a sulfur loading of 80 wt% showed both hydrophobic and hydrophilic nature. This material as a surfactant-like amphiphilic additive was blended into Carbon Black-containing HDPE structure with a low (0.5 wt%) loading concentration. In this case, the basic properties (heat stability, tensile strength) of the composite was not changed significantly, however, the carbon black additive distribution was increased drastically. The average particle size of the CB was 2 times smaller in the C-PE-C composite compared to the basic C-PE matrix. We believe that this method will be a helpful way to disperse additives in highly viscous hydrophobic or hydrophilic materials even at high temperature.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.polymertesting.2020.106419.

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