Silylated-montmorillonite as co-adsorbent of chitosan composites for methylene blue dye removal in aqueous solution

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Abstract

Industrialization plays an important role in the economy of developing countries, including increasing community welfare. However, the presence of poorly industrial waste disposal system has brought a negative impact on the environment. Therefore, it is deemed necessary to overcome this problem with low-cost technology, namely adsorption. In this research, silylated-montmorillonite (sMMt) has been successfully prepared as supporting material for the adsorption of methylene blue by chitosan. The sMMt was characterized by FTIR, XRD, and elemental mapping techniques. The chitosan/sMMt composites were prepared through dissolution-precipitation method later on producing the rod-like morphology as observed by SEM. The adsorption process was carried out in a batch method by studying the pH and the adsorption contact time. The adsorption kinetic mechanism of the chitosan/sMMt nanocomposite followed pseudo-second order rather than Lagergren model indicating chemisorption predominant. The addition of silylated-montmorillonite into chitosan then enhanced the methylene blue dye removal performance, evidenced by improving the Qe values by 10\% compared to chitosan.

Keywords: Adsorption, chitosan, methylene blue, montmorillonite, silylation

1. Introduction

Plastic, textile, dye, leather, cosmetics, pharmaceutical, food, and paper manufacturing industries are the largest contributor in wastewater containing dyes resulted from the coloring process [1]. Channeling of wastewater with a high content of dyes causes serious environmental pollutions, such as inhibiting the photosynthesis process and hampering the growth of aquatic biota [2]. Methylene blue (MB), one of water soluble cationic dyes [3], is the most common dye consumed in dyeing cotton, wood, and silk. In respiratory, MB causes difficulties in breathing. Moreover, it also can cause a permanent damage to the eyes, a burning effect, mental disorders, or excessive sweating, when it directly contacts to the human body [4,5].

Wastewater containing dyes is very difficult to treat due to its molecular size and structure, which makes it resistant to aerobic digestion, stable against light, heat, and oxidation agents [6]. Therefore, an inexpensive, simple and effective treatment should be applied to treat waste water containing dyes from the batik industry to reduce or eliminate any harmful substances. Several well-known methods in the removal of the dye are coagulation-flocculation [7], physical methods [8], physicochemical treatment [9], photocatalytic degradation [10,11], ion exchange [12], and others. However, most of them require high operation condition and financial cost. Ngah et al. [13] reviewed some techniques for wastewater containing dyes treatments and elaborated the advantages of adsorption method, such as low cost, easy to operate and effective in dye removal. Moreover, the adsorption process can be performed in chemistry, physics or a combination of both chemistry and physics.

Lately, many researchers have studied the possibility of using low-cost material obtained from nature to remove various types of dyes. Lazim et al. [14] employed orange peel and spent tea leaves as low-cost biosorbent for methylene blue and remazol brilliant blue dyes removal \textit{via} batch adsorption method. Other biosorbents, such as pineapple leaf and lime peel powder [15], mangosteen peel-based activated carbon [16] and chitosan [17,18], were also reported as promising biosorbent for dye removal.

Chitosan is derived from chitin through decetylation process. This second most abundant biopolymer is easy to be found in nature and mostly deposited in shells of shrimp, crab, and other shelled animals. Chitosan has good adsorption ability due to its active functional groups such as amines (\textit{\textendash}NH\textsubscript{2}), hydroxyl (\textit{\textendash}OH) and cluster glucosamine [19]. This material is relatively abundant in Indonesia, but still lacks in utilization. The main reason of using chitosan as adsorbent is due to the presence of amines group that could be interacted with dyes molecules, especially anionic dyes, through electrostatic interaction [20–22]. In some cases, such as the
adsorption of cationic dyes, the chitosan polymer could not optimally adsorb those dyes due to repulsion effect caused by their similar surface charges. Therefore, in some researches, chitosan was composited with another material that has negative surface charges such as did Mokhtar and co-workers [23]. Therefore, in this study, to obtain a good performance of chitosan as adsorbent for cationic dyes (methylen blue), modification with negative surface charges materials (such as clay minerals) was performed.

Another abundant material in Indonesian clay is clay. In the present report, clay can be also employed as adsorbent cationic ions or organic molecules like dye [24], due to its high surface area and cationic exchange capacity as well as porous structure [25,26]. Sarma et al. [27] used montmorillonite clay as an adsorbent in methylene blue dye removal and discovered a good interaction between dye and clay via surface ionic interaction, which then resulted in high adsorption capacity. Gurses et al. [26] reported the dye adsorption mechanism of montmorillonite clay following the pseudo-second order in which chemisorption was predominant. To enhance the adsorption selectivity and capacity, some researchers like Nejib et al. [28] modified the clay by adding an additive. The modified clay mineral has a higher adsorption capacity compared to the unmodified one. The modification of clay minerals with organosilane compounds improved the adsorption ability either to anionic or to cationic dyes. Moreira et al. [29] modified the surface of palygorskite and sepiolite with 3-aminopropyltriethoxysilane (APTES), and found a significant improvement on adsorption capacity of modified-clay mineral towards both metanil yellow (anionic dye) and methylene blue (cationic dye). Another investigation also found that organosilane modified montmorillonite has a rapid adsorption performance towards methylene blue. The modification of clay minerals with organosilane compound is aimed not only to improve their adsorption capacity, but also enhance their compatibility to react with polymer matrix like chitosan to create a synergistic effect during adsorption. Herewith, we modified the surface of montmorillonite with 3-glycidiloxoypropyltrimethoxysilane (GPTMS) to provide an oxirane ring functional group on their surfaces. This leads a chemical reaction with the amine groups of chitosan via opening ring reaction [30–32]. Consequently, the clay distributed on the chitosan polymer matrix and could improve their adsorption performance towards the methylene blue dye.

In this research, the surface modification of montmorillonite clay with organosilane compound aimed to add an active site and to improve their adsorption ability. To the best of our knowledge, the surface modification by silylation method on montmorillonite and further compositing with chitosan is not studied yet. Moreover, the application of silylated-montmorillonite supported chitosan as adsorbent is also not mentioned in another research. Therefore, we studied the effect of silylated-montmorillonite addition on chitosan toward methylene blue dye adsorption in aqueous solution.

2. Materials and Methods

2.1. Materials

Chitosan was obtained from PT. Biotech Surindo, Indonesia. Montmorillonite clay was purchased from Sigma Aldrich (Montmorillonite K 10, surface area 220–270 m² g⁻¹). Other chemicals, such as 3-glycidiloxoypropyl trimethoxysilane (GPTMS, Aldrich), toluene (Merck), ethanol (Merck), hydrochloric acid (Merck), sodium hydroxide (Merck), and Methylene blue (MB) dye were used in this study without any purification or modification except mentioned.

2.2. Silylation of Montmorillonite

The silylation method was adopted from Yuan et al. [33] with a slight modification. About 4 mL of GPTMS was dissolved in 50 mL of dry toluene. Then, 1.2 g of montmorillonite clay (MMt) was added and ultrasonically treated for 30 minutes. The mixture was refluxed for 24 h at 135 °C. The product was filtered and washed several times using fresh toluene to remove any unreacted GPTMS. Then, the final product was dried for 12 h at 135 °C and denoted as sMMt. The sMMt product was characterized using FT-IR (Fourier Transform Infra Red) spectroscopy IR prestige 21 SHIMADZHU. The morphology of MMt and sMMt was observed by Scanning Electron Microscopy (SEM, Phenom FEI) and the surface elemental analysis of them was detected by SEM-EDX OXFORD Vega3 LM TESCAN.

2.3. Synthesis of Chitosan/sMMt composite

About 2 g of chitosan was dissolved in 90 mL of 1.5% acetic acid and then refluxed at 80 °C. Then, the sMMt (0 and 20% wt) was added to the chitosan solution and stirred for 1 hour. A liquid form of chitosan or Cs/sMMt composite was slowly dripped into a beaker containing 100 mL of NaOH 1 M. The precipitates were washed with distilled water and dried at a room temperature for 4 days. The products were then crushed by ball-milling (1000 rpm, 60 minutes), collected and characterized using FT-IR (IR prestige 21 SHIMADZHU) for functional group analysis and SEM (Phenom FEI) for morphological study.

2.4. Adsorption Study

The adsorption was performed in room temperature. A total of 20 mg of chitosan, sMMt or chitosan/sMMt composite was added into 10 mL of conditioned 30 mg L⁻¹ methylene blue solution at pH 3, 5, 7, 9 and 11. The adsorption process was conducted for 20 minutes. The filtrates were immediately collected and measured by using UV-Vis PerkinElmer Lambda 25 spectrophotometry. From these data, the dye removal was calculated by following equation 1 [14]:

\[
\text{Dye removal (\%) = \left(\frac{C_{i} - C_{f}}{C_{i}}\right) \times 100}\% \tag{1}
\]

Where \( C_{i} \) (mg L⁻¹) and \( C_{f} \) (mg L⁻¹) are initial and equilibrium dye concentration, respectively.

2.5. Adsorption Kinetic

A 30 mg L⁻¹ MB dye solution was prepared at optimum pH condition (pH = 11). Then, about 20 mg of chitosan, sMMt or Chitosan/sMMt composite was added to 10 mL of the MB
dye solution. The adsorption contact time was varied, i.e., 5, 10, 15, 20, 30, 40, 50, 60, 70, 80 and 90 minutes. In this research, the adsorption was performed in room temperature. Afterward, the filtrates of each adsorption time were collected and the remaining concentration was measured by using UV-Vis spectrophotometry.

3. Results and Discussion

3.1. Structural Analysis of Silylated-Montmorillonite

The surface modification of MMT through silylation method aims to improve the selectivity of clay in dye adsorption as a consequence of an additional functional group (oxirane ring). To find out the success of surface functionalization of MMT, the functional group analysis of origin MMT and sMMT by FTIR was examined. Fig. 1 shows the FTIR spectra of MMT and sMMT. Generally, MMT has a typical peak of hydroxyl (−OH), Si−O−Al deformation, and the Si−O−Si deformation at 3434 cm⁻¹, 1060 cm⁻¹, 930 cm⁻¹, 525 cm⁻¹, and 469 cm⁻¹, respectively. Some new peaks were observed in FTIR spectra of silylated-MMT. Those peaks appeared at 2939-2880 cm⁻¹, and 693 cm⁻¹, which correspond to C−H stretching vibration and oxirene ring vibration of organosilane compound. According to Herrera et al. [34], the appearance of the typical peak of C−H aliphatic on FTIR spectra of clay after modification indicates a bond formation between the silane molecules and the clay through SiC−O−SiC linkage. Additionally, a typical of siloxanes peak was also noticed in FTIR spectra of sMMT at around 1190 cm⁻¹. Based on this characterization, it can be concluded that the MMt has been successfully silylated by GPTMS.

Fig. 2 reveals the X-ray diffraction of the MMT and sMMT. The MMt had a typical peak at 8.7, 19.6, 27.5, 34.7, 45.4, 55.4, and 61.8 which showed a good agreement with JCPDS no. 03-0015 having a chemical formula (Na, Ca)₀.₅ (Al, Mg)₂ Si₄O₁₀ (OH)₂ ⋅xH₂O indicating K10 structure [35,36]. The quartz phase (JCPDS 74-1811) was also observed in the MMT XRD pattern as impurities. Generally, the XRD peaks of both MMT and sMMT were similar. However, the intensity of (001) typical peak at 20 8.74° decreased as observed in XRD pattern of sMMT. This indicated that the organosilane compounds mainly attached to this (001) planes. The organosilane formed a siloxane networks on the surface or interlayer of MMT. Consequently, The presence of this poly(siloxane) networks caused the XRD peaks of sMMT seemed broader than MMT, especially in 2θ 5−13°, indicating a decrease in its crystallinity compared to MMT.

Another characterization supporting the formation of siloxane network of GPTMS on MMT surfaces is morphological analysis by SEM and SEM-EDX. Fig. 3 depicts the morphology of MMT and sMMT. The surface morphology of MMT resembles a plate structure due to its layered-structure (Fig. 3a). Meanwhile, the morphology of
sMMt seems to be interconnected sheets as a consequence of the formation of poly(siloxane) on the MMT surfaces (Fig. 3b). The presence of the siloxane network is able to link MMT plates. Further, the surface elemental analysis of MMT and sMMt recorded by SEM-EDX is used to identify the atomic content on their surfaces. As shown in Fig. 3c, the elemental mapping images of both MMT and sMMt clearly state that the MMT contains Al (purple), O (green) and Si (red) atoms with Al to Si ratio of 4.9:29.7. On the other hand, sMMt has the Al (blue), O (green), Si (red) and C (yellow) atoms on their surface. The presence of C element on the sMMt surfaces indicates that the surface modification of MMT by GPTMS has been successfully achieved as supported by FTIR and XRD analysis. Based on the EDX spectra (Fig. 3d), the amount of C atoms on the sMMt surface was 29.9%. Meanwhile, other elements such as Si, Al and O were 18.7%, 2.2%, and 49.2%, respectively. Si:Al ratio of sMMt was 39.3% higher than Si:Al ratio in the MMT. The increase of the Si atom occurred due to the contribution of the Si element from GPTMS compound. In addition, the presence of siloxane (Si–O–Si) layers on the MMT surface caused a decrease of the Al ratio.

3.2. Synthesis of Chitosan/sMMt composites

Chitosan/sMMt composite was prepared through the dissolution precipitation method. The presence of oxirane ring on the surface of sMMt facilitated the chemically interaction to the amine groups of chitosan. The nucleophilic addition was involved in reaction between sMMt and chitosan. The amine group acted as nucleophilic and attacked the oxirane ring [30–32]. Consequently, the chitosan and sMMt were linked via β-amino alcohol bonds as illustrated in Fig. 4.

Fig. 5 indicates the FTIR spectra of chitosan and chitosan/sMMt. The chitosan had a typical absorbance peak at 1669 cm⁻¹ corresponding to the bending vibration of N–H functional group. A broad absorbance band at 3300 cm⁻¹ indicated the O–H stretching vibration. All of chitosan typical absorbance band was clearly observed in chitosan/sMMt composites FTIR spectra. However, the N–H bending vibration peak slightly shifted to 1687 cm⁻¹ which was probably due to the chemical reaction between the amines group of chitosan and oxirane group on the sMMt surfaces via opening ring reaction as illustrated in Fig 4. Additionally, peaks at 1062 cm⁻¹ and 468 cm⁻¹ corresponded to Si–O stretching and Si–O–Si deformation vibration, originated from sMMt.
The synthesis of chitosan/sMMt composites was carried out by means of dissolution-precipitation method. The solution of chitosan or chitosan/sMMt composites was precipitated in a basic solution to yield a solid form. The immediate solidification process affected the morphological structure of both chitosan and chitosan/sMMt composites. As revealed in Fig. 6, the rod-like structure of either chitosan or chitosan/sMMt nanocomposites was observed, probably due to the high rate of phase transfer from liquid to solid. The surface of chitosan looked smooth (Fig. 6a). However, it became rough after combining with sMMt (Fig. 6b). An agglomeration of sMMt in some spot was also seen in the SEM image of chitosan/sMMt composite, caused by the attracting force between the same polarities of sMMt.

3.3. Effect of pH on Methylene Blue Adsorption

The removal of methylene blue is influenced by several factors, one of which is a pH condition. Fig. 7 presents the dye removal percentage of the adsorbents (chitosan, sMMt, and chitosan/sMMt composites) at a different pH condition. The chitosan worked both in acid (pH = 3) and basic (pH = 9) condition with dye removal of 56.22% and 71.66%, respectively. However, the dye removal of MB by sMMt was predominant at basic condition (pH = 11) about 93.99%. The adsorption of dyes was influenced by the charges of both dye molecules and the surface of the adsorbent. The chitosan had amines (tended to be positively charge) and hydroxyl (negatively charge). Meanwhile, the clay minerals are rich of hydroxyl groups, which are mainly negative charges [27]. Therefore, the chitosan is capable of absorbing MB strongly at both acid and basic condition. Meanwhile, the sMMt dominantly adsorbs the MB at basic condition. Compared to the sMMt, adsorption capacity of chitosan is much lower. Therefore, an addition of sMMt into chitosan polymer is expected to improve the adsorption ability of chitosan.

As desired, the chitosan/sMMt composite had better adsorption ability compared to chitosan as evidenced by an improvement of dye removal to be 77.50%. Besides it was rich of the hydroxyl group, the sMMt also contained the oxirane group, which contributed to increase the basicity of sMMt. Therefore, the chitosan/sMMt composite had good interaction with acid properties of MB dyes at higher pH condition [37]. According to Feddal et al. [38], at higher pH, the –OH groups on clay in solution becomes negative charge due to deprotonation, which facilitates the attachment of methylene blue onto the adsorbent surface.

3.4. Adsorption kinetics

The adsorption contact time is an important parameter to study in adsorption process. This determines the adsorption kinetic mechanism. Fig. 8 clarifies the effect of contact time towards the adsorption capacity of chitosan, sMMt, and
chitosan/sMMt composites. All adsorbents have rapid MB adsorption rate indicated by a high adsorption capacity at the initial stage of adsorption (5 minutes). Chitosan has the adsorption capacity of 9.13 mg g\(^{-1}\) at the initial time, while sMMt and chitosan/sMMt have 13.46 and 10.40 mg g\(^{-1}\), respectively. The adsorption of MB onto chitosan begins to reach equilibrium state after 60 minutes of contact time. However, after adding sMMt, the equilibrium state is achieved at 70 minutes. As revealed, the adsorption capacity at equilibrium state of chitosan/sMMt was higher than chitosan. This condition was also found by Daraei et al. [39], which enhanced the chitosan adsorption performance by adding cloisite nano clay. The data also demonstrated that the adsorption of MB onto chitosan/sMMt is relatively faster at the initial time, but becomes slower afterward. Heraldy et al. [40] described this phenomenon as a two-stage adsorption mechanism, in which at the first stage, the adsorption of dye molecules was instantaneous, so the fast stage occurred. At the second phase, an equilibrium was reached due to relatively small of the total reactive dye molecules or decrease of the adsorbent capacity.

To evaluate the performance of adsorbent, studies of adsorption kinetics was performed. In this research, the Lagergren and pseudo-second order were employed as a kinetic model to determine the adsorption mechanism and its adsorption rate by fitting both of these models. The Lagergren and pseudo-second order equation were expressed as equation 2 and 3 [41]:

\[
\log(q_e - q_t) = \log q_e - (k_1 2.303)t \quad (2)
\]

\[
t/q_e = (1/k_2Qe^2) + (t/Qe) \quad (3)
\]

where \(q_e\) and \(q_t\) is the amount of dye adsorbed at equilibrium and at time (mg g\(^{-1}\)), \(t\) is the contact time (min) and \(Q_e\) is adsorption capacity at equilibrium.

The Lagergren kinetic rate \((k_1)\) was generated from \(t\) versus \(\log(q_e - q_t)\) plots as presented in Fig. 9a, while the pseudo-second order kinetic rate constant \((k_2)\) and adsorption capacity at equilibrium \((Q_e)\) were determined from the slope and intercept of \(t/q_e\) versus \(t\) plot as shown in Fig. 9b. The adsorption kinetic mechanism was resolved based on the correlation coefficient (R-square). Based on the fitting linear of both models, pseudo-second order had higher R-square values than Lagergren model. Therefore, the adsorption kinetic mechanism of all adsorbents followed the pseudo-second order models which enclosed to chemisorption.

The \(t/q_e\) versus \(t\) plots for all adsorbents had a straight line, where the R-square values for each chitosan, sMMt and chitosan/sMMt composite as depicted in Table 1 were 0.999, 0.999 and 0.998, respectively. The higher R-square values indicated the suitability of the adsorption mechanism to those models [42]. Sun et al. [43] described that the adsorption rate was mainly ruled by the dye concentration at the absorbent surface when the adsorption followed the pseudo-second order.

![Fig. 8 Effect of adsorption contact time toward the adsorption capacity of chitosan, sMMt, and chitosan/sMMt](image1)

![Fig. 9 (a) Lagergren and (b) pseudo-second order linear plots of chitosan, sMMt, and chitosan/sMMt](image2)
Table 1. Kinetic parameters resulted from plotting Lagergren and pseudo-second order models

| Models          | Parameters | Chitosan | sMMt | Chitosan/sMMt |
|-----------------|------------|----------|------|---------------|
| Lagergreen      | $k_1$ (min$^{-1}$) | 0.066    | 0.081| 0.127         |
|                 | R-square   | 0.898    | 0.906| 0.678         |
| Pseudo-second   | $k_2$ (g mg$^{-1}$ min$^{-1}$) | 0.031    | 0.047| 0.028         |
| order           | Qe (mg g$^{-1}$) | 12.029   | 15.152| 13.233        |
|                 | R-square   | 0.999    | 0.999| 0.998         |

Table 1 presents the $k_1$, $k_2$ and $Q_e$ values of all adsorbents. The $k_2$ of chitosan was $3.1 \times 10^{-2}$ g mg$^{-1}$ min$^{-1}$. The addition of sMMt into chitosan slightly decreased the $k_2$ value to be $2.8 \times 10^{-2}$ g mg$^{-1}$ min$^{-1}$. However, the $Q_e$ value of chitosan/sMMt increased by 10%. It should be noted that MMt as one of the layered aluminosilicate materials was reported by previous research as a good adsorbent for dyes removal due to providing a high surface area and porous structure [44]. This then led MMt to have a high adsorption capacity toward cationic dyes. Moreover, the presence of negative charges on the MMt surfaces also enhances their ability to absorb and to interact with cationic dyes. In addition, the presence of the oxirane ring of GPTMS organosilane contributes to the adsorption of MB. By these reasons, the chitosan supported by sMMt has high adsorption capacity compared to the chitosan.

4. Conclusion

The silylated-montmorillonite (sMMt) has been successfully synthesized evidenced by FTIR, XRD and elemental analysis characterization. The presence of sMMt on chitosan (chitosan/sMMt composites) had a positive impact on dye removal. The optimum pH condition of the MB adsorption was reached at basic condition (pH 11). The adsorption kinetic mechanism followed the pseudo-second order indicating that the chemisorption played the role. The adsorption kinetic rate $k_2$ and adsorption capacity $Q_e$ of chitosan/sMMt were found to be $2.8 \times 10^{-2}$ g mg$^{-1}$ min$^{-1}$ and 13.23 mg g$^{-1}$, respectively. The presence of sMMt on chitosan matrix was able to improve the $Q_e$ by 10% compared to chitosan.

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