Aluminum Pillared Clay (Al–PILC) for Adsorption of Dyes in Red Fruit Oil

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1. Introduction

The red fruit (Pandanus conoideus Lam.) is an endemic plant that grows on Papua island. Red fruit contains high levels of carotenoids [1], β–cryptoxanthin, α-tocopherol, and unsaturated fatty acids [2, 3]. Carotenoids are a group of pigments that are yellow, orange, or orange–red. It is found in plants, skins, shells/exoskeletons of aquatic animals, and other marine products such as mollusks (calm, oyster, scallops), crustaceans (lobsters, crabs, prawns), and fish (salmon, trout, sea bream, red snapper, and tuna). Carotenoids are also found in many bacteria, fungi, algae, and green plants [4].

![Figure 1. Carotene structure [5]](image)

Red fruit oil contains carotene, which is dark orange, so it is not very popular. Therefore, it is necessary to reduce the intensity of the color. Yesterday, in this study, the synthesis and characterization of Al-pillared clay (Al–PILC) from light and heavy clay fractions were carried out. The absorption capacity of red fruit carotene dyes was studied. The research stages included preparation, fractionation, activation, pillaring of clay with aluminum, characterization, and adsorption test for carotene dyes on red fruit. Characterization was carried out using X-ray Diffractometer (XRD) and Gas Sorption Analyzer (GSA). The results showed that basal spacing of natural clay, Al pillared heavy fraction–clay, Al pillared light fraction–clay, activated light fraction were 15.08 Å, 15.27 Å, 16.84 Å, and 16, respectively. 22 Å. The GSA results showed that the surface area and pore volume of the Al pillared light fraction–clay of 0.3 was higher than the heavy fraction. The average pore size of Al pillared light fraction–clay and the heavy fraction was found in the mesoporous range of 30–100 Å, and the adsorption isotherm is type IV. Al–pillared light fraction–clay had higher adsorption ability than a heavy fraction and light fraction before pillaring. When the pillaring agent’s concentration was 0.3 M, Al pillared heavy fraction–has absorption capacity is 58.66%, while Al pillared light fraction–clay is 90.4%.
adsorption temperature, the adsorption increased. The maximum adsorption percentage of 95.108% was obtained at an adsorbent/CPO ratio of 1/3 and a temperature of 60°C.

On the other hand, clay is an abundant natural mineral. It is cheap and can be used as an adsorbent [9]. Clays have been used for β-carotene reduction in palm oil [10] and soybean oil [11]. However, the adsorption capacity of clay is limited. This weakness can be overcome by activation and modification of the clay, such as through clay pillarization. Clay pillarization can increase the thermal stability, porosity, specific surface area, and acidity of the clay surface [12]. Pillared clays have several advantages, including a larger surface area and a larger total pore volume. The superior properties of pillared clay make it potential for use as an adsorbent even as a selective adsorbent [13].

Activated and pillared clays are proven to have better adsorption abilities. Nurhayati [14] reported that natural clay activated using HCl increased the surface area from 298 m²/g to 306 m²/g. Salawudeen et al. [15] conducted a study on removing carotene content in vegetable oils and obtained 33.3% higher yields when using alkaline-activated clays. The development of research on pillared clay shows that pillared clay has been widely used to adsorb dyes. Ti pillared clay increased its adsorptivity properties due to an increase in surface area from 65 m²/g to 216 m²/g, pore volume from 1 x 10⁻³ cm³/g to 8 x 10⁻² cm³/g, and an increase in the number of Lewis acid sites [16]. Al-pillared clay (Al-PILC) from Tunisian bentonite was used to adsorb methylene blue [17]. Al-PILC with Pd metal doping was also used to decolorize azo and triarylmethane dyes [18].

According to Najafi et al. [19], Al pillared clay has a specific area of 00-300 m²/g and basal spacing (d001) of 1.7-2 nm, while Ti pillared clay has a specific area of 100-350 m²/g and basal spacing (d001) of 1.2-1.8 nm.

Based on some of these studies, this study prepared Al pillared clay to adsorb β-carotene in red fruit oil. β-carotene has a relatively large size with the molecular formula C₅₀H₅₀O, which consists of combining eight isoprene units (C₅H₈) [20]. The choice of Al pillared clay is because Al produces pillared clay with higher basal spacing. Clay minerals have a particle size of <4 μm. Small clays have a larger surface area, and characters such as cation exchange capacity, catalytic properties, and plastic properties are higher than larger clays [21]. There has been no study on the pillarization of the clay fraction, even though natural clay contains clay particles of various sizes. In our opinion, it is necessary to study the character of pillared clay from different clay fractions. This work reports on the preparation and characterization of aluminum pillared clays in light and heavy fractions of clays, originating from Boyolali, Central Java, and their use to adsorb dyes in red fruit oil.

2. Methodology

2.1. Materials and Tools

Materials: Natural clay from Boyolali, Central Java, HCl (Merck, 37%), distilled water, AlCl₃·6H₂O (Merck), NaOH (Merck), red fruit oil from Papua, alcohol (Merck, 99%). All reagents were in analytical grade. Equipment: glassware, 170 mesh sieve, oven (Binder), furnace (Vulcan 3-130), desiccator, analytical balance (Ohaus), magnetic stirrer, Whatman filter paper, shaker (Maxq 2000), X-ray Diffractometer (Shimadzu), Gas Sorption Analyzer (Quantachrom Instruments version 11.0), UV-Vis spectrophotometer (Perkin Elmer).

2.2. Clay preparation and fractionation

Natural clay is washed with water, oven-dried at 105°C for 3 hours. Clay fractionation is carried out by making a clay suspension in water until the light fraction is at the top and the heavy fraction is at the bottom. The two fractions were separated by decantation, then each dried at 105°C. Each clay fraction, the heavy fraction (HF), and the light fraction (LF) that had dried were crushed and sieved in 170 mesh size.

2.3. Clay Activation

Every 100 grams of HF and LF clays were reacted with 200 mL of 2 M HCl. The mixture was stirred for 24 hours, then allowed to stand for 24 hours, then filtered. The solid was washed with distilled water to remove Cl- and oven-dried at 105°C for 3 hours. The resulting solid was characterized by XRD (X-ray Diffraction).

2.4. Clay pillarization with aluminum cations

The pillar solution was made by mixing the NaOH solution into the AlCl₃·6H₂O solution with a mole ratio of 1:1 to obtain a concentration of 0.3 M aluminum pillar solution. The mixture was continuously stirred for 24 hours using a magnetic stirrer. This pillar solution was mixed respectively into the clay suspension of an activated heavy fraction (HF) and a light fraction (LF) and stirred for 24 hours, then left to stand at room temperature for 24 hours. The results of intercalation/pillarization were filtered, then washed with water and dried in an oven at 105°C for 3 hours. Then the dry solid was calcined at a temperature of 450°C for 2 hours with a temperature ramp rate of 1°C/minute. From this stage, heavy fraction and light fraction Al pillared clay (Al-PILC) were obtained. Each aluminum pillar clay was characterized by XRD (X-ray Diffraction) to determine basal spacing and GSA (Gas Sorption Analyzer) to determine pore and surface characteristics.

2.5. Performance test of Al-PILC on β-carotene absorption of red fruit oil

Red fruit oil was diluted with alcohol. The ratio of Al-PILC adsorbent: red fruit oil adsorbate was 5% (w/v). The adsorption was carried out with all Al-PILC adsorbents against diluted red fruit oil with a contact time of 10 minutes. The adsorption results were filtered to obtain the filtrate. Then the filtrate was analyzed using a UV-Vis spectrophotometer at a maximum wavelength of 460 nm. With the same procedure, red fruit oil adsorption was also carried out using the initial clay adsorbent before fractionation, activated HF, and LF clays. From this step, information on which adsorbent has the best adsorption obtained.
3. Results and Discussion

3.1. Characters of Aluminum pillared clay (Al–PILC)

In this study, the natural clay fraction from Boyolali was separated into a heavy fraction (HF) which precipitated, and a light fraction (LF) floated. After being activated, the two clay fractions were polarized using an Al pillared. Characterization of both clay fractions before and after pillared using X-ray diffraction meter (XRD). It aims to determine the mineral type through the typical peaks in XRD and compared with the mineral database in the JCPDS program (Joint Committee on Powder Diffraction Standards) and see the $2\theta$ angle peak shift and associated basal spacing.

![Figure 2. XRD diffraction pattern of (a) activated heavy fraction clay, (b) aluminum pillared clay of heavy fraction](image)

Figure 2 shows that the activated heavy fraction (HF) has the highest intensity at $2\theta = 26.372^\circ$ and Al–PILC heavy fraction at $2\theta = 26.45^\circ$, almost corresponding to illite minerals ($I = 100\%$, $2\theta = 26.75^\circ$). Apart from illite, this clay also contains montmorillonite and chlorite and a small amount of kaolinite.

![Figure 3. XRD diffraction pattern of (a) activated light fraction clay, (b) aluminum pillared clay of light fraction](image)

Figure 3 shows that the light fraction diffractogram pattern is not much different from the heavy fraction, which both show peaks in the illite mineral area, namely at $2\theta = 26.26^\circ$ and Al–PILC of the light fraction at $2\theta = 26.29^\circ$. Figure 2 and Figure 3 show that the clay’s composition in this study is montmorillonite, illite, chlorite, and kaolinite. The presence of Al$_2$O$_3$ was not explicitly indicated because there was no new peak after the clay was pillared with Al. This is possible because the small amount of added alumina oxide in Al–PILC does not affect the XRD diffractogram pattern. Also, the clay itself already contains alumina tetrahedra in its skeleton. There was no change in the clay diffractogram pattern before and after pillarization, indicating that the clay structure did not change much [22]. When viewed from the basal spacing, both the pillared heavy and light fraction experienced an increase in basal spacing when compared to activated clays, as presented in Table 1.

| Samples                              | $2\theta$ (°) | I (%) | d (Å) |
|--------------------------------------|---------------|-------|-------|
| Activated heavy fraction clay         | 5.96          | 29.98 | 14.81 |
| aluminum pillared clay of heavy fraction | 5.78          | 25.77 | 15.27 |
| Activated light fraction clay         | 5.44          | 24.64 | 16.22 |
| aluminum pillared clay of light fraction | 5.24          | 40.07 | 16.84 |

Natural clays have interlayer areas. There are various kinds of cations such as K$, Na$, Ca$, Mg$, and Fe$^2+$, which function as a load balancer for the clay layer is negative, which causes the size between the clay layers to be non-uniform. The light fraction contains montmorillonite, illite, kaolinite, and chlorite, each with different properties. According to Raharjo and Sarmili [23], the fine fraction with a size of <0.002 μm has larger plastic properties. Thus it is easier to experience swelling and is lighter. The light fraction has an illite component where K$^+$ ions dominate this type as a counterpart to the clay charge. In this case, the size of the radius of the K$^+$ ion is larger than that of the other ions, which results in a larger basal spacing.

![Figure 4. XRD diffraction pattern of (a) activated heavy fraction clay, (b) aluminum pillared clay of heavy fraction](image)

The increase in basal spacing in the heavy fraction (HF) and the light fraction (LF) pillared was caused by a change in the balancing cation from the small cation, which was then substituted by the aluminum pillar polycation. The polycation of the Keggin Al$_3$ ion has a large size. When the Al$_3$ polycation enters the area between the clay layers, the distance between the layers increases. The calcination process at a temperature of 450°C converts the Al polycation into a more stable metal oxide. In this study, the light fraction of Al–PILC had a peak at $2\theta = 5$, 26° or basal spacing 16.84 Å. This result is similar to Kostriat’s study [22], which was Al–pillared bentonite calcined at 500°C.

![Figure 5. XRD diffraction pattern of (a) activated light fraction clay, (b) aluminum pillared clay of light fraction](image)

Figures 2 and 3 prove that the basal spacing values of Al–PILC for the heavy and light fraction and the activated light fraction were higher than the activated heavy fraction. Based on Bragg’s law, $n \lambda = 2d \sin \theta$, if there is a shift at $2\theta$ to the right, the basal spacing value gets smaller, and if there is a shift to the left, the basal spacing value will be higher.

Data on the surface area, pore volume, and pore diameter of pillared clay are presented in Table 2. Table 2 shows that the heavy fraction of the Al–PILC clay has a
smaller surface area and pore volume. The pore volume of the heavy fraction of Al–PILC was 0.0516 cc/g, while the light fraction was 0.995 cc/g. Meanwhile, the mean pore diameter of the heavy fraction of Al–PILC was greater than that of the light fraction. This shows that the light fraction of Al–PILC has smaller pores, but a lot, or can be said to be more porous than the heavy fraction.

Table 2. Pore and surface characters of aluminum pillared clays of heavy fraction and light fraction

| Parameter                     | aluminum pillared clay of heavy fraction | aluminum pillared clay of light fraction |
|------------------------------|----------------------------------------|----------------------------------------|
| Surface area (m²/g)          | 11.243                                 | 55191                                  |
| Average pore width (Å)       | 183.690                                | 136.067                                |
| Pore volume (cc/g)           | 0.05163                                | 0.9915                                 |

Figure 4. Adsorption isotherms of (a) aluminum pillared clay of heavy fraction and (b) light fraction

Figure 4 shows the adsorption isotherm curve of a light fraction of Al–PILC. A heavy fraction of Al–PILC is of type IV in the IUPAC classification indicating mesoporous material. Type IV isotherms are characterized by a hysterical loop caused by capillary condensation. Figure 4 shows that at \( P/P_0 = 0.01 \), the gas adsorbed very little so that only a little monolayer. When the pressure is increased \( P/P_0 < 0.2 \), the adsorbed gas begins to saturate the monolayer. When the pressure is increased to 0.6, the gas adsorption in the multilayer begins to occur. At \( P/P_0 = 1 \), gas molecules fill the mesoporous, after which the pressure begins to decrease for gas desorption. When the pressure is lowered, there is an isothermal graph showing the hysteresis loop. The desorption isotherm shows that the amount of desorbed gas is not the same as the amount initially adsorbed due to capillary condensation. The volume of nitrogen gas adsorbed by Al–PILC LF was much greater than that of Al–PILC HF. This result is in line with carotene dyes’ adsorption power in red fruit, which will be explained later.

Figure 5 shows that the pores of all the aluminum pillared clay samples obtained are mostly located in the diameter range of 30–100 Å, which is included in the mesoporous category. The pore size distribution was not uniform, as indicated by the presence of several peaks. This non-uniform pore size is probably caused by the Al₂O₃ pillars formed in the area between the clay layers, which are not uniform and unevenly distributed.

Figure 5. The pore size distribution of (a) aluminum pillared clay of heavy fraction and (b) light fraction

3.2. Adsorption of β-carotene in red fruit oil

In this study, red fruit oil was mixed with ethanol so that it was not too thick. The reduced color concentration in red fruit oil indicates the adsorption power of the adsorbent. The absorption results for clay, heavy fraction (HF), and a light fraction (LF) before and after activation and after pillaring with Al obtained are presented in Figure 6.

Figure 6. Adsorption ability of several absorbents

Figure 6 shows that the adsorbent’s adsorption ability of β-carotene increases when the clay is fractionated and activated, where the activated light fraction has an adsorption ability of 89.15%. The adsorption process factors are the adsorbate concentration and surface area. The greater the adsorbate concentration, the adsorbate, and adsorbent interactions will increase. Furthermore, the larger the adsorbent’s surface area, the greater the adsorption ability to adsorb the adsorbate.

Figure 6 also shows the adsorption ability of Al–PILC. Al–PILC of the light fraction has an adsorption power of 90.4%, where this value is higher when compared to Al–PILC of the heavy fraction, which can only adsorb 58.66%. This adsorption ability is in accordance with the adsorption isotherm in Figure 4, where Al–PILC of a light fraction can adsorb 121.45 cc/g of nitrogen gas, but Al–PILC of the heavy fraction is only 33.22 cc/g. This is also in line with the data presented in Table 2 that the Al–PILC of the light fraction has a larger surface area and pore volume than the heavy fraction so that the light fraction interacts with more β-carotene
dyes. Likewise, if it is related to pore size, the Al-PILC of the light fraction is smaller than the heavy fraction, but its ability to absorb adsorbate is higher.

The adsorption ability of β-carotene in light fraction clay increased with the activation and pillarization treatment, but in clay, the heavy fraction decreased after the Al pillarization process. This decreased adsorption ability may be related to Al pillars’ irregular arrangement in heavy fraction clay, or even agglomeration may occur, thus clogging the pores of heavy fraction clay. The pore size of pillared clay for Al-PILC-HF is 183.69 Å, and this is larger than the light fraction, which is only 136.067 Å. However, the surface area and pore volume are lower. Thus, it can be assumed that the pores in the Al-PILC-HF pillared clay are bigger but shallower because the Al pillar actually covers the inside of the pore. Physically, the color change in red fruit oil before and after adsorption can be seen in Figure 7.

![Figure 7: The color of red fruit oil before and after adsorption](image_url)

The coloration of the red fruit oil occurs after the adsorption of the clay. Possible adsorption that occurs is physical adsorption due to the interaction of the adsorbent with the adsorbate through van der Waals forces. The β-carotene dye, which has a structure with the molecular formula C₉₅H₁₅₂O₂, contains isoprene units. Even though the molecule is large enough, it still allows entry into the pore based on the pore diameter of Al-PILC ranging from 30–100 Å. Another possibility is that β-carotene molecules are adsorbed on the clay’s basal spacing with a size of 14.81 Å. It means that the smaller size of the β-carotene molecules are adsorbed, the color of the red fruit oil will fade.

4. Conclusion

It is concluded that pillarization using aluminium can increase the basal spacing value from 14.81 Å in heavy fraction clay to 15.27 Å in Al-PILC of heavy fraction. Likewise, the basal spacing of 16.22 Å in the light fraction was activated to 16.82 Å in the Al–PILC of the light fraction. The β-carotene dye found in red fruit oil is successfully absorbed by the clay so that the red fruit oil, which has a reddish-orange color, becomes more transparent. The smallest adsorption ability occurred in Al-PILC-HF with an adsorption ability of 58.66%. In comparison, the highest was obtained by Al-PILC-LF with an adsorption ability of 90.14%, where this was following the structural character of the pillared clay.

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