PREPARATION AND CHARACTERIZATION OF SODIUM CARBOXYMETHYL CELLULOSE FROM KAPOK (CEIBA PENTANDRA) ALPHA-CELLULOSE

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ABSTRACT

Objective: This study aimed to obtain sodium carboxymethyl cellulose (NaCMC) from α-cellulose of the hulls of kapok (Ceiba pentandra) and determining its identity and characteristics base on compendial requirements and compared to the reference (standard).

Methods: α-Cellulose was isolated from kapok hulls and used to generate NaCMC powder through alkalization and carboxymethylation. Alkalization was performed using 2.5% NaOH (containing sodium tetrahydroxide), whereas carboxymethylation was using sodium monochloroacetate. Identification and characterization were performed through infrared spectrum analysis using Fourier transform infrared, qualitative analysis, organoleptic examination, morphologic examination, and topographical analysis using scanning electron microscopy (SEM), and X-ray diffraction. Tests conducted were pH determination, sulfated ash content, moisture content, loss on drying, particle density, and viscosity.

Results: The obtained NaCMC powder was yellowish-white with a similar infrared spectrum as the NaCMC standard. The powder had a degree of substitution of 0.57 and a pH of 8.5. According to SEM, the obtained powder had a similar morphology as the NaCMC standard, although the synthesized NaCMC had a rougher surface. The obtained NaCMC also had a similar diffractogram as the NaCMC standard, which was characterized by the presence of crystalline and amorphous structures. Besides, the NaCMC powder had a similar moisture content (8.50%), sulfated ash content (36.43%), and loss on drying (9.87%) as the standard, whereas its 1% viscosity value (20.6 cP) was substantially different.

Conclusion: NaCMC generated from α-cellulose of kapok hulls fulfills compendial requirements and has similar characteristics to reference.

Keywords: Kapok, Sodium carboxymethyl cellulose, Alkalization, Carboxymethylation, Characterization.

INTRODUCTION

Cellulose is an important commodity for various industries, such as the pharmaceutical industry. A commonly used cellulose derivative in the pharmaceutical industry is sodium carboxymethyl cellulose (NaCMC), which used in pharmaceutical preparations, including oral and topical preparations. NaCMC is used to increase viscosity and stabilize emulsions. Besides, NaCMC also functions as a binder and disintegrant in tablet preparations [1].

Initially, NaCMC was produced from wood because it contains high cellulose levels (42–47%) [2]. However, the current trend is the production of NaCMC from non-wood lignocellulosic materials due to the limited availability of wood and increasing prices. One of them is kapok (Ceiba pentandra), which is abundantly available in Indonesia.

Kapok has the potential for use as a raw material for producing NaCMC because it contains α-cellulose with a purity level of 94.05% [3]. Cellulose isolated from kapok hulls can be synthesized into NaCMC through alkalization using NaOH and carboxymethylation using sodium monochloroacetate (NaMCA).

This study aimed to identify the optimal reaction conditions for synthesizing NaCMC from kapok α-cellulose to fulfill compendial testing requirements.

MATERIALS AND METHODS

Raw material
The raw material used in this study was kapok hulls powder obtained from Balitro, Bogor, West Java Province, Indonesia.

Chemical material
The chemicals used were 96% ethanol, nitric acid, sodium hydroxide, sodium sulfite, sodium hypochlorite, NaMCA, sodium tetaborate, isopropyl alcohol, KBr powder, glacial acetic acid, methanol, sulfuric acid, orange methyl P, phenolphthalein, α-naphthol, anhydrous sodium carbonate, potassium bisulfate, and standard NaCMC powder as a comparator which were from Merck (Germany). Besides, distilled water was obtained from Brataco (Indonesia).

Instruments
The instruments used in this study included an infrared spectrophotometer (FTIR-8400S Shimadzu, Japan), analytical balance (Sartorius, Germany), a bulk density tester (BDT M-005/04, Indonesia), a furnace (Cole-Parmer, USA), a hot plate and stirrer (IKA Type Hr-7, Germany), a scanning electron microscope (SEM) (QUANTA 650, USA), and X-ray diffraction (XRD) system (Panalytical X’Pert Pro MPD, Europe), a moisture analyzer (ADAM AMB50, USA), a viscometer (Cole-Parmer), a sintered funnel (Pyrex, Germany), filter paper (Whatman, Germany), aluminum foil (Total Wrap, Indonesia), a crucible (PRC, China), an oven (Heraeus, Germany), a desiccator (Duram, Germany), and a burette (Pyrex) and other glassware commonly used in laboratories.

Isolation of α-cellulose from kapok hulls
In total, 200 g of kapok shell powder were mixed with 2.67 L of 3.5% nitric acid (containing 26.7 g of sodium nitrate) at 90°C for 2 h. Then, the residue was immersed in 2 L of a solution containing sodium hydroxide and sodium sulfite at a concentration of 2% b/v at 50°C for 1 h and bleached through boiling in a 1.3 L of a 1:1 mixture of water and 3.5% sodium hypochlorite. The obtained residue was heated in 1.3 L
of sodium hydroxide (17.5% v/v) at 80°C for 30 min. The residue was then dried at 60°C and crushed to obtain α-cellulose powder [4].

**Preparation of NaCMC from kapok α-cellulose**

First, 3 g of α-cellulose were weighed and placed in a glass beaker containing 60 mL of isopropl alcohol. Then, 25% NaOH (containing 0.17 g sodium tetrahydrate) was added to a volume of 10 mL, and the mixture was stirred for 1 h at 25°C. Furthermore, the carboxymethylation process was optimized using the amount of NaMCA (3.75, 3.90, and 4.05 g) with constant stirring at 55°C. The reaction time was 3.5 or 4 h for 3.75 g of NaMCA, 2.5 or 3 h for 3.90 g of NaMCA, and 1.5 or 2 h for 4.05 g of NaMCA. The obtained product was filtered and suspended in 60 mL of methanol. The slurry was neutralized with glacial acetic acid. The final product was washed with 80% ethanol and methanol, followed by filtration and drying in an oven at 60°C to obtain NaCMC powder, which was stored in a desiccator [4].

**Organoleptic examination**

The product examinations were shape, color, taste, and odor in comparison with the NaCMC standard commonly used as a pharmaceutical excipient.

**Identification using Fourier transform infrared (FTIR)**

After mixing 99 mg of KBr with 1 mg of each sample, scanning was over the wavelength range of 4000–400 cm$^{-1}$. The IR spectrum of each sample was compared with that of the standard.

**Qualitative analysis**

One gram of sample was added to 50 mL of water. A sample solution of 1 mL was mixed with 1 mL of distilled water, and five drops of α-naphthol. Then, 2 mL of concentrated sulfuric acid was added to the solution, which turned purple. The results were compared to the NaCMC standard following the same procedure [5].

**Degree of substitution (DS)**

One gram of NaCMC was put into 20 mL of 95% ethanol and stirred for 5 min. Then, 5 mL of 2 M nitric acid was added, and the mixture was brought to a boil followed by continuous stirring at 80°C for 10 min. The liquid phase that formed was removed, and the solid phase was washed 5 times, with 10 mL of 80% ethanol at 60°C. The precipitate was washed with 10 mL of methanol. Finally, the precipitate was dried, and the mixture was heated to a boil for 15 min. After the sample dissolved, two drops of a phenolphthalein indicator were added, causing the solution to turn pink, and the solution was hydrated with 0.3 M HCl until the pink color disappeared [6]. The DS of NaCMC was calculated as follows:

$$ DS = \frac{162 \times %CM}{5900 - (58 \times %CM)} $$

$$ %CM = \frac{\text{Blank volume} - \text{HCl volume} \times \text{HCl molarity} \times 0.059 \times 100}{\text{Sample mass (g)}} $$

**pH test**

One gram of NaCMC was dispersed into 100 mL of distilled water, and the pH was measured using a pH meter.

**SEM**

SEM was performed to assess the surface morphology of each sample. Imaging was performed after selecting a certain part of the object (sample) and the desired magnification (×150 and ×600) to ensure that a clear photo was obtained. This analysis was conducted for both synthesized samples and the NaCMC standard.

**XRD analysis**

XRD analysis was performed to assess the crystalline and amorphous forms of NaCMC. First, 2 g of each sample were smoothed and placed on the glass with the help of an adhesive, followed by characterization.

**Moisture content**

Water content can be measured using the moisture content (Adam) at 105°C. First, 1 g of each sample was weighed on top of an aluminum plate that was previously anchored. The tool then measured the sample water content.

**Sulfated ash content**

Ash content was measured using a furnace. A crucible was heated in a furnace at 600±50°C for 30 min, cooled in a desiccator, and weighed. As much as, 1 g of each sample was weighted in the crucible. The sample was then wetted using sulfuric acid (1 mL) and heated until it was charred. After cooling, the residue was wetted with a small amount of sulfuric acid and heated it until the white smoke disappeared. Then, the crucible was heated at 600±50°C until the residue burned completely. The crucible was placed in a desiccator until it was cold and weighed, and then the percentage of ash was calculated [7].

**Loss on drying**

Samples of NaCMC (1 g) were weighed in a crucible and then dried in an oven at 105°C until they reach a constant weight.

**Density test**

Samples of NaCMC (15 g) were weighed and placed in a 50-mL measuring cup with a polygonal bottom. The volume was recorded as the bulk volume. Next, the measuring cup was placed on the tool, which was activated. When the appliance stopped, the final volume was recorded as the tapped volume.

**Viscosity test (1%)**

As much as 1 g of NaCMC was weighed, dissolved in 100 mL of distilled water, and stirred until homogeneous, followed by testing for viscosity using a viscometer.

**RESULTS AND DISCUSSION**

**Preparation of α-cellulose from kapok hulls**

The isolation of α-cellulose from kapok hulls begins with delignification using 3.5% nitric acid containing sodium nitrite. In this process, nitric acid will degrade the lignin bond structure, permitting lignin to separate from cellulose [8]. Sodium nitrite is added to accelerate the degradation process. Nitric acid cannot eliminate lignin, and thus, the second round of delignification was performed using a mixture of 2% sodium sulfite/2% sodium hydroxide followed by bleaching using a mixture of 3.5% sodium hypochlorite and water. Sodium hypochlorite can remove lignin residue in the pulp. Furthermore, α-cellulose, β-cellulose, and γ-cellulose were separated using 17.5% sodium hydroxide [9]. Cellulose was heated at 100°C to obtain α-cellulose from kapok fibers (Fig. 1).

![Fig. 1: α-Cellulose isolated from kapok hulls](image-url)
Optimization and preparation of NaCMC from α-cellulose
NaCMC is synthesized in two stages, namely, alkalization and carboxymethylation. Alkalization was performed by stirring α-cellulose in isopropyl alcohol and then adding 25% NaOH containing sodium tetaborate. Isopropyl alcohol functions as a reaction medium that affects the quality of NaCMC. A smaller polarity of the reaction medium can increase the reaction rate of NaCMC formation [2]. The use of isopropyl alcohol as a reaction medium also results in relatively fewer byproducts (sodium glycolate) [6]. NaOH converts cellulose into alkali cellulose. Sodium tetaborate can improve the quality of NaCMC by indirectly increasing the DS of the synthesized products [10].

The product synthesized through alkalization was alkali cellulose, which was reacted with NaMCA to form NaCMC through carboxymethylation. NaMCA functions as an etherification reagent that induces the substitution of sodium carboxymethyl groups at C2, C3, and C6 [11]. The carboxymethylation stage was optimized by varying the amount of NaMCA and the duration of the reaction, in line with prior research illustrating that varying the conditions of carboxymethylation altered the DS results [4].

After completing both stages, the formed product was separated into its liquid and solid phases. The solid phase was immersed in absolute methanol and neutralized with glacial acetic acid. The mixture was filtered, and then the residue was washed 3 times, with 80% to remove the remnants of the byproducts formed. Finally, the residue was again washed with absolute methanol, filtered, and dried. The DS of the synthesized NaCMC was examined as a quality parameter.

Based on the experiments conducted, the largest DS value of 0.59 was obtained using 3.75 g of NaMCA and a reaction time of 3.5 h. A small DS value was obtained because the carboxymethyl group was not substituted well on α-cellulose from kapok. Therefore, it was necessary to optimize other parameters (e.g., temperature, reaction medium, and NaOH levels) to optimize NaCMC synthesis. Although the DS value did not meet the requirements, the optimal conditions regarding NaMCA treatment (3.5 g of NaMCA, a reaction time of 3.5 h) were selected to synthesize NaCMC from α-cellulose. Furthermore, identification and characterization of NaCMC from kapok cellulose were performed in comparison to the NaCMC standard.

Organoleptic examination
According to the organoleptic examination, the NaCMC standard and NaCMC from kapok α-cellulose have no smell and taste. However, there was a slight difference in color between the two moieties. The NaCMC standard had a white color, whereas NaCMC synthesized from kapok α-cellulose had a yellowish-white color. A difference in color occurred because of the presence of residual lignin that was not completely dissolved in the bleaching process. Besides, the two powders exhibited slight differences in texture. NaCMC synthesized from α-kapok cellulose had a finer powder texture than the NaCMC standard, which had a texture similar to that of cotton fibers.

Identification using FTIR
Infrared spectrometry was performed to determine the suitability of α-cellulose and NaCMC functional groups. The spectrum of kapok α-cellulose was compared with Avicel PH-101 (Fig. 2), whereas NaCMC synthesized from α-cellulose was compared with the NaCMC standard commonly used in pharmaceutical preparations (Fig. 3).

The functional groups in the structure of α-cellulose include OH groups at 3650–3200 cm⁻¹, stretched CH groups at 3000–2850 cm⁻¹, and COC bonds at 1200–980 cm⁻¹ [11,12]. The spectra of Avicel PH-101 and α-cellulose from kapok were similar, and thus, it can be stated that the compound produced in the isolation process was α-cellulose and that it could be used for the next stage, namely, NaCMC synthesis [12].

Based on its structure, NaCMC has similar functional groups as α-cellulose. However, NaCMC possessed a carbonyl group (C=O) at 1800–1670 cm⁻¹, which indicated the presence of a carboxymethyl substitution. A peak indicated the presence of CH2 at 1425–1421 cm⁻¹ [11,12]. Based on these data, the NaCMC standard and NaCMC synthesized from kapok α-cellulose have similarities.

Qualitative analysis
Qualitative analysis was performed to determine the presence of glucose in NaCMC. The NaCMC standard was used as a positive control, and samples containing no NaCMC were used as negative controls.

The negative control did not have a purple color, unlike the NaCMC standard, and NaCMC synthesized from kapok hulls. This difference in color can be attributed to the addition of o-nitrophenol and sulfuric acid [5]. NaCMC is dehydrated when reacted with sulfuric acid, thus forming a furfural structure that will then condense with o-nitrophenol to produce a purple compound (Fig. 4) [13].

DS
The NaCMC standard had a DS value of 0.77, which met the requirements (0.7–1.2). Conversely, the synthesized NaCMC had a DS value of 0.57. This rather low value of NaCMC from kapok α-cellulose was unique and can be caused by different raw material from NaCMC standard.

pH test
The pH of the NaCMC standard was 6.98, versus 8.50 for the synthesized NaCMC, both of which fell into the desired range (6.5–8.5). Although the requirements were met, some results failed to meet the standard. This negative result, possibly due to some NaCMC samples, contained high levels of NaOH because of a lack of neutralization process.

SEM
SEM analysis revealed similarities between the synthesized NaCMC and the NaCMC standard. Morphologically, the NaCMC standard had a stem-like structure arranged separately with a smooth surface. Meanwhile, NaCMC synthesized from kapok α-cellulose was in the form of rods that formed clot-like masses and had a slightly rough surface (Fig. 5). The morphological differences between these two types of NaCMC can be attributed to the different sources of raw materials (α-cellulose) used.

XRD analysis
The NaCMC standard (Fig. 6) had a typical diffractogram pattern featuring a sharp peak at a 2θ of 20.2825, which denoted the nature of the crystal. Besides, the diffractogram also exhibited a wide peak at 2θ of 38.0465, which denoted its amorphous properties. For the synthesized NaCMC (Fig. 7), a similar diffractogram pattern with a sharp peak at a 2θ of 20.4161 was recorded, illustrating the crystalline structure. The diffractogram additionally displayed a wide peak, although it was not clear; at a 2θ of 39.2665, which reflected its amorphous properties. Thus, crystalline and amorphous forms were identified for both the NaCMC standard and NaCMC produced from kapok α-cellulose.

Moisture content
The NaCMC standard had a water content of 5.75%, compared with 8.50% for the synthesized NaCMC. Both values met the moisture content requirement of <10%.

The sulfated ash content of the NaCMC standard was 23.87%, versus 14.21% for the synthesized NaCMC, both of which fell into the desired range (14–23%). Although the requirements were met, some results failed to meet the standard. This negative result, possibly due to some NaCMC samples, contained high levels of NaOH because of a lack of neutralization process.

Density test
According to the test results, the NaCMC standard had a bulk density of 0.37 g/cm³ and a tapped density of 0.56 g/cm³, whereas NaCMC
Fig. 2: Fourier transform infrared spectra of Avicel PH-101 (blue) and (b) α-cellulose isolated from kapok hulls (pink)

Fig. 3: Fourier transform infrared spectra of sodium carboxymethyl cellulose (NaCMC) standard (blue) and NaCMC synthesized from kapok α-cellulose (pink)

Fig. 4: (a) Negative control, (b) positive control, and (c) sample
Fig. 6: Diffractogram of the sodium carboxymethyl cellulose standard

Fig. 7: Diffractogram of sodium carboxymethyl cellulose synthesized from kapok α-cellulose

synthesized from kapok α-cellulose had a bulk density 0.47 g/cm³ and a tapped density 0.55 g/cm³. These results indicate that neither the standard nor the product met the requirements for bulk or tapped density.

The smaller density of the NaCMC standard can be attributed to its lighter mass, giving it a relatively larger volume but smaller density than the synthesized NaCMC. Furthermore, the shape of synthesized NaCMC particles was more rounded than that of the NaCMC standard, which decreased the space between particles and the resulting volume, leading to greater density [15].

Viscosity test (1%)
The viscosity of the NaCMC standard was 3528.5 cP, versus 20.6 cP for the synthesized NaCMC. Although both values met the stated viscosity requirement, they were substantially different. This difference can be explained by the different sources used to generate the materials.

CONCLUSION
As the conclusion, based on the characteristics data obtained, the synthesized NaCMC had a similar morphology to the standard and also showed identity and characteristics similar to the reference. The substantial difference is only its viscosity that possibly due to the different raw material used.

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CONFLICTS OF INTEREST
None.

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