MODIFIED EXTRACTION AND PURITY TEST OF ARENGA PINNATA GUM

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ABSTRACT

Objective: The purpose of this study is modified extraction method to obtain gum from Arenga pinnata Merr. and their purity test.

Methods: Gum has been extracted from palm seed using distilled water by centrifugation and warming. Characterization of gum was done by scanning electron microscopy-energy dispersive X-ray and spectrophotometer infrared. Gum refluxed with HCl 2 N for 14 h would be the monomer. Galactose and mannose were used as standard. Purity test was done by the thin-layer chromatography (TLC) and Luff Schoorl method.

Results: The study results obtained that the modification methods (4.8%) and centrifugation (4.6%). Identification of functional groups with the spectrophotometer infrared provided the same spectrum forms between the two methods. Purity test by TLC with the best eluent butanol: ethanol: water 2:2:2. Luff Schoorl's method gives results of 87.46%, whereas the raw comparison galactose-mannose standard 96.69%.

Conclusions: The modification method gives more results with the same purity rate as the centrifugation method.

Keywords: Galactomannan, Palm seed, Arenga pinnata, Modification, Extraction, Luff Schoorl.

INTRODUCTION

Arenga pinnata is an Arecaceae family plant that grows in the tropics. A. pinnata is an enclosed seed plant because its fruit seed is wrapped with fruit flesh. The round-shaped palm fruit, about 4 cm in diameter, has three spaces and has three seeds, arranged in a chain-like string. Each bunch has 10 stalks or more, and each stalk has approximately 50 grains of green to yellowish brown. The young palm fruit is hard and firmly attached to the strands of the fruit, while the fruit is ripe fruit flesh is rather soft. The young fruit palm flesh contains very sticky mucus when it comes to the skin because it contains oxalic acid. Palm seeds are endosperm half-ripe palm fruit seeds after going through a process of soaking with lime water solution for several days so that it becomes soft, supple, and a bit clear white. A. pinnata gum is galactomannan compound [1].

Galactomannan is a polysaccharide not starch extracted from plant seeds by centrifuge method [2] and autoclaved [3] with water solvent at neutral pH. Galactomannan is a carbohydrate reserve and regulates the amount of water in the seeds during germination [4].

The advantage of galactomannan when compared with other types of polysaccharides is its ability to form a thick solution in very low concentrations, and only slightly affected by pH, ionic, and heating strength. The viscosity of galactomannan is very constant in the range of pH 1 – 10.5 which may be caused by its neutral molecular character. However, with high temperatures, and very acidic or alkaline conditions, galactomannan can be degraded [5].

Galactomannan has been widely applied in the food industry, pharmaceutical field, and cosmetics. In the food industry, galactomannan is commonly used as a clot in the ice cream industry to make ice not melt quickly in the cheese-making industry and salad spice. In the pharmaceutical industry, galactomannan was applied in the manufacture of tablets, hydrogels, films, and controlled release drug matrices [6].

The total carbohydrate analysis was done using the method of Luff Schoorl. International Commissions for Uniform Methods of Sugar Analysis considers Luff Schoorl's method as one of the methods used to standardize reducing sugar analysis. The entire compound carbohydrates are broken down into simple sugars (monosaccharides) with the help of the HCl acid and heat. Monosaccharides that is formed is then analyzed by Luff Schoorl's method. The method analysis principal is the reduction of Cu²⁺ to Cu⁺ by becoming a monosaccharide. Free monosaccharides will reduce alkaline solution of metal salts or oxides form becomes free. The unreduced Cu²⁺ is then quantified by iodometric titration [7].

METHODS

The study was conducted of the Organic Chemistry Laboratory, Faculty of Mathematics and Natural Science, Universitas Sumatera Utara. The plant material used in this research is the palm seed which is determined by the Herbarium Medanense of Biology department. The materials used are ethanol, cupric sulfate, citric acid, sodium carbonate, hydrochloric acid, sodium hydroxide, sulfuric acid, potassium iodide, and sodium thiosulfate (Merck). The tools used are analytical balance (Sartorius), centrifuge (Hitachi), heating mantle (Seoh), and water bath. Palm seed gum was characterized by infrared spectrophotometer (Shimadzu) and scanning electron microscope-energy dispersive X-ray (EDX) spectroscopy (Bruker).

Gum extraction

As many as 500 g sample is cleaned and mashed with blender for ± 3–5 min by the addition of water 1:1 and warmed 5 min in water bath, and then stored in the refrigerator for 24 h. Filtered sediment formed, then washed with ethanol p.a. Sediment was dried in the desiccators [1,2].

Modification method

As many as 500 g is stored in the refrigerator for 24 h, then the sample is cleaned and mashed with blender for ± 3–5 min by the addition of water 1:1 and warmed 5 min in water bath, and then stored in the refrigerator for 24 h. Supernatant obtained is stored in the refrigerator for 24 h, then added ethanol 96% in comparison with the volume of 1:2, then stored in the refrigerator for 24 h. Filtered sediment formed, then washed with ethanol p.a. Sediment was dried in the desiccator [3].
Thin-layer chromatography
About 50 g of palm seed gum was hydrolyzed by addition of 50 mL 1 N HCl using a heating mantle for 14 h, then the solution neutralized with the addition of sodium carbonate, the filtrate evaporated into viscous fluid. Then, a small amount of each aliquot is bottled on Whatman No.1 paper with micropipette. Then, eluted with butanol: ethanol: water (BEW) 4:1:1, 1:4:3, 1:3:4, 1:4:4, and 2:2:2, butanol: acetic acid: water 4:1:5, 20% and 50% acetic acid and 10% and 20% hydrobodic acid. Then, it was dried and detected by spray reagent (1.2 g p-anisidine and 1.6 g phthalate acid in 100 mL ethanol) will give a brown color [8].

**Luff Schoorl’s analysis**
Luff Schoorl’s solution is made by 25 g cupric sulfate dissolved in 100 mL water, 50 g of citric acid dissolved in 50 mL water, and 388 g sodium carbonate dissolved in 400 mL water free carbon dioxide. A solution of citric acid sodium carbonate in solution is poured while stirring carefully next added cupric sulfate and added water to 1000 mL.

About 100 mg palm seed gum hydrolyzed with 100 mL 1N HCl for 14 h using a heating mantle, then neutralized with NaOH 30%, then put in flask 100 mL and sufficient volume with distilled water. Then, plucked 10 mL and added with 10 mL of Luff Schoorl’s solution quantitatively, and 6 mL of distilled water, added some boiling stones and then heated to boiling for 10 min in a water bath, then cooled immediately with ice. After a cold, 10 mL of sulfuric acid 25% is added and 6 mL of potassium iodide 30%, it is immediately titrated with a standard solution of 0.1 N sodium thiosulfate until a weak yellow color is formed. Then, 3 mL of 0.5% starch indicator until the solution was blue, and the titration was resumed until blue color disappeared, recorded the titration volume used. The same experiment was performed on blanks by substituting the sample solution with water. Next is calculated by the formula below [7].

\[ V_{Na_2S_2O_3} = \left( \frac{\text{ml blanko-ml sample}}{0.1} \right) \times N_{Na_2S_2O_3} \]

After getting the value of \( V_{Na_2S_2O_3} \) then equalized the value on Luff schoorl’s table to get the value of \( Z \).

Total sugar content (%) = \( \frac{Z \times DF \times 0.95}{\text{Sample weight (mg)}} \times 100\% \)

\( Z \) is seen in Luff Schoorl’s table to see the sugar content and DF is dilution factor.

**RESULTS**
The study results obtained that the modification methods (4.8%) and centrifugation (4.6%). Fig. 1 shows the functional group identification with the infrared spectrophotometer giving a very similar spectrum between the centrifuge and the modification method, it proves that both are galactomannan compounds. Fig. 2a shows the surface morphology of a hollow Arenga pinnata gum particle using Scanning electron microscopy (SEM), and Fig. 2b shows the elements contained in the gum using Energy Dispersive X-ray Spectroscopy (EDS) are atoms C and O. Fig. 3 show the purity test by TLC with the best eluent butanol: ethanol: water 2:2:2. Table 1 show the Luff Schoorl’s method gives results of 87.46%, whereas the raw comparison galactose-mannose standard 96.69%.

**DISCUSSION**
Infrared gum spectrum of centrifuge seed method is shown in Fig. 1a where the wave number at 3425.58 cm\(^{-1}\) shows the stretching vibration -OH. The wave number at 2924.09 cm\(^{-1}\) shows an aliphatic C-H stretching vibration supported by wave numbers at 1404.18 and 1319.3 cm\(^{-1}\) indicating the presence of an aliphatic C-H bending vibration. Wave numbers at 1635.64 and 1087.85 cm\(^{-1}\) indicate the presence of bending vibrations C-O in C=O bonds.

The infrared spectrum of modified palm seed gum is shown in Fig. 1b where the wave number at 3394.72 cm\(^{-1}\) shows the stretching vibration -OH. The wave number at 2924.09 cm\(^{-1}\) shows stretching vibration of C-H aliphatic supported by wave numbers at 1373.32 and 1311.59 cm\(^{-1}\) indicating the presence of an aliphatic C-H bending vibration. Wave numbers at 1635.49 and 1026.13 cm\(^{-1}\) indicate the presence of bending vibrations C-O in C=O bonds.

Wavenumber at 2800–3000 cm\(^{-1}\) shows stretching vibration C-H and 3100–3500 cm\(^{-1}\) shows vibration stretching O-H. The wide peak at 1134–983 cm\(^{-1}\) is a bending characteristic of C-OH. The peak at 1635 cm\(^{-1}\) indicates a bend between the polysaccharide and water [9].

Fig. 1a and b show the identical infrared spectrum shape, especially in the fingerprint region (2000–500 cm\(^{-1}\)). This infrared spectrum shows galactomannan compounds such as the infrared guar gum spectrum. Guar gum is composed of a mixture of galactose and mannose combined through a glucoside link, which can be described chemically as galactomannan. Galactomannan polysaccharides have a linear general structure, in which the polymer unit (1,4)-β-D-mannopyranose, bonded to (1,6)-α-D-galactopyranose [4].
Fig. 2a shows the particles morphology of palm seed gum by scanning electron microscopy (SEM). Fig. 2b shows the elements contained in the palm seeds gum are 71.47% C atoms and 28.53% O atoms by EDX spectroscopy. Hollow particle surfaces allow these compounds to still be cross-linked with crosslinking agents. SEM-EDX is able to show the particle surface shape and elements found on the particles [1].

Gum has been refluxed with the addition of 1 N HCl for 14 h, then bottled on paper chromatography performed to separate the galactose-mannose contained in the sample. Of all motion phase comparisons, BEW 2:2:2 is the best phase of motion where the stain is formed away from the bottling point. However, with all comparisons of the phase of motion, it has not been able to separate galactose-mannose in sample and standard. Rf obtained after being eluted with BEW 2:2:2 phase of the galactose comparator 0.65 cm and mannose 0.63 cm. The adjacent Rf ensures that the galactose cannot be determined by comparison with the paper chromatography method. However, elution of BEW 2:2:2 ensure that galactomannan of palm seed has been purified [8].

Sugar levels in the sugar palm seed gum successfully established by Luff Schoorl method. In principle by hydrolyzing galactomannan from Arenga pinnata will produce monosaccharide compounds that have the properties of reducing. The Luff Schoorl solution is added in excess so that cupric sulfate (Cu²⁺) is reduced to Cu⁺ by reducing sugars. Excess Cu⁺ can oxidize potassium iodide (I⁻) to iodine (I₂). Iodine is titrated with a standard solution of sodium thiosulfate. Iodine is equivalent to Cu⁺ excess so that the sugar content is the reduction of the titration volume of the blank less the titration of the sample. The total sugar content of sugar palm seed gum was 87.46% while galactose+mannose was 96.69%. The difference in levels is due to the need for optimization of glycosidic bond termination time on palm seed gum [7].

CONCLUSIONS
The modification method gives more results with the same purity rate as the centrifugation method. The shape of the infrared spectrum is similar between the centrifuge method and the heating method. Luff Schoorl’s method can analyze total carbohydrate levels. Optimization of reflux needs to be done to ensure the galactomannan compound has disconnected its galactose and mannose.

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AUTHOR’S CONTRIBUTION
Study conception: Tarigan.
Acquisition of data: Reveny.
Analysis of data: Zebua.
Drafting of manuscript: Zebua.
Critical revision: Kaban.

CONFLICTS OF INTEREST
All authors have none to declare.

REFERENCES