Pharmacognostic and Phytochemical Standardization of White Tea Leaf (Camellia sinensis L. Kuntze) Ethanolic Extracts

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ABSTRACT

Background: Tea or also known as Camellia sinensis (Theaceae family) is the most popular plant and beverage in the world because of the sensory properties, prices are relatively cheap, stimulant effects, and their potential health benefits but white tea is not widely known. White tea is made from unfermented tea leaves young shoots protected from sunlight to avoid polyphenols degradation which inhibits of the chlorophyll formation and causing the white color on the leaf buds. Objective: The objective of research and development of herbal medicine is to improve the quality and safety of natural products. Materials and Methods: Macroscopical and microscopical features of the leaf have been analysis using an optical microscope and fragment analysis under scanning electron microscopy (SEM). Phytochemical and physico-chemical analysis were evaluated. The observation of the FTIR spectrum profiles is done by interpreting the typical peak that appears. Results: The leaf has actinocytic stomata, unicellular trichomes, heterogenous mesophyll which is characterized by the presence of calcium oxalate crystals and sclereid cells. Phytochemical analysis indicated resources the presence of tannins, flavonoids, glycosides and saponins. The content of polyphenol from white tea leaves ethanolic extract is 35.73% with the largest concentration of catechins is 18.84% and 17.43% tannins. The derivative content of catechins is EGCG with 73.74%. FTIR analysis showed functional groups of O-H, C-H, N-H, C=O, C=C, and C-O. Conclusion: Pharmacognostic and phytochemicals features established in this study may be used as part of the pharmacopoeial standard which can play an important role in its standardization. Key words: Characteristic, Macroscopic, Microscopic, Physico-chemicals, Phytochemicals, Theaceae.

INTRODUCTION

The development of herbal medicines for health purposes in Indonesia at this time is likely to increase rapidly. The use of herbal medicine is intended to improve health (promotion), restoring health (rehabilitative), disease prevention (preventive) and healing (curative) and is expected to support the development of public health. This condition supported by the potential of Indonesia’s natural resources which is consisting of 30,000 species of medicinal plants. More than 6000 species of medicinal plants have been used as a source of natural medicine that empirically has been used as traditional medicine.1 Tea is the most popular plant and beverage in the world because of the sensory properties, prices are relatively cheap, the stimulant effects and their potential health benefits.2 Tea or also known as Camellia sinensis (Theaceae family) are widely cultivated commercially but white tea is not widely known. Based on their processing, tea can be divided into unfermented tea (white tea and green tea), semi-fermented tea (oolong tea), and fermentation (black tea).3 White tea has a simpler process by withering and drying. White tea leaf is taken from the shoots or very young tea leaves or buds and covered with silvery hair. The young shoots are protected from the sunlight which inhibiting of the chlorophyll formation and causing the white color on the leaf buds. White tea is reported to have a high polyphenol content and showed antiseptic and antioxidant properties.4 In addition, various types of bioactive found in white tea leaves such as polyphenols, caffeine, theogallin, gallic acid, theaflavins, flavonoids glycoside, and catechins especially epigallocatechin (EGC), epigallocatechin gallate (EGGC), epicatechin gallate (EGC), and epicatechin (GEC).4,5 The antioxidant properties of white tea can prevent free radicals and inhibit oxidative stress and inflammation.6 Oxidative stress and inflammation associated with various diseases including obesity disease, dyslipidemia, diabetes, cardiovascular, neurodegenerative and cancer.7,8 The previous studies reported that ethanolic extract of white tea has anti-diabetic in vitro activity by inhibition of alpha-amylase enzyme activity (% inhibition of 99.11 ± 0.01), alpha-glucosidase (IC50 10.54 mg/mL) and dipeptidyl peptidase IV (% inhibition 30.57 ± 0.08).9 It is necessary to develop and further studies of white tea leaves. The objective of research and development of herbal medicine is to improve the quality and safety of natural products. Therefore, it is necessary to standardize raw materials and extracts to maintain the uniformity of quality, safety and efficacy. Raw material quality parameters include moisture content, ash content, acid insoluble ash, water soluble extract content, content of ethanol soluble extract and concentration of substances identity. It also conducted organoleptic, microscopic, macroscopic, chemical content identification, fingerprint profiles, and contamination analysis.9 Standardization of white tea leaf extract is
necessary to provide information on quality standards of herbal medicine development.

Experimental

Preparation of Material

White tea leaves (Camellia sinensis L. Kuntze) obtained from the Tea Plantation and Quinine Research Center in Gamboeng, West Java, Indonesia. White tea leaves were sorted, collected, and then dried under sunlight. Furthermore, the tea leaves are withered with a dryer. The white tea leaves powder are made by grinding dried white tea leaves by using a grinder.

Preparation of White Tea Extract

White tea leaves extract prepared by reflux method using ethanol 70% as a solvent over 3 hours in 60°C. After 3 hours the extract was filtered and the supernatant was removed and transferred to a volumetric flask then be repeated reflux for 2 (two) times. The ethanol extract then evaporated using a vacuum rotary evaporator.

Identification of Raw Materials

Identification of raw materials conducted by observation of organoleptic, macroscopic and microscopic identification. Macroscopic identification of white tea leaves was observed based on the length and width of leaves, plant height, color and description of the leaves and stems. Leaf color is identified before and after the drying process by using a Microscope IX70. Microscopic identification is done by using cut crosswise and lengthwise of leaf and dried powder white tea leaves. Identification of raw materials powder is also done using field emission scanning electron microscopy (FE-SEM). Previously, powder coating with gold for 1 minute then identified with the FE-SEM.

Extract Characterization

Characterization of the extract was conducted on the determination of moisture content, ash content, acid insoluble ash content, water-soluble extract content, and ethanol-soluble extract content. The extract characterization method conducted to the WHO guidelines.9

Phytochemical Analysis

Phytochemical analysis conducted by screening chemical substances and determination of major compounds. Identification of the presence of alkaloids, flavonoids, tannins, saponins, glycosides, terpenoids and anthraquinone was carried out according to the procedures in WHO guidelines and Harborne.10

Determination of polyphenol content was measured by spectrophotometry method using gallic acid as a standard, and the results expressed as a percent of gallic acid equivalents. The amount of 10 g sample is weighed and put into a flask, dissolved in 50 ml of distilled water and sonication for 10 minutes, then add with distilled water until 100 ml. one milliliter aliquot transfers to a flask. Added 5 ml of a reagent of Folin Ciocalteau 10% and let stand for 3-8 minutes, then added 4 ml of 7.5% sodium carbonate and homogenized. The mixture incubated for 2 hours protected from light. Furthermore, the absorbance of the solution is measured by spectrophotometer at 750 nm.

Determination of total tannin content was measured by spectrophotometry method. The amount of 1 g sample is weighed and put into a flask of 100 ml and diluted with distilled water. One milliliter aliquot put into a flask, then diluted with 75 ml of distilled water. Added 5 ml of reagent Folin-Denis, 10 ml of saturated sodium carbonate solution, and distilled water till 100 ml and then homogenized. This mixture incubated for 30 minutes and absorbance was measured at 760 nm.

Determination of Catechin and EGCG

Determination of total catechin was measured by using spectrophotometry at 210 nm. Catechin as a standard, the results expressed as a percent of catechins equivalents. While EGCG assay was conducted using reverse phase HPLC using the mobile phase isocratic elution system that is a mixture of orthophosphate 0.1%, water, acetonitrile, methanol (14:7:3:1 v/v) at pH = 4.00 with a flow rate of 1.2 mL/min and detected at 280 nm.

Chromatography Analysis

Chromatography analysis was done using a Thin Layer Chromatography (TLC) system. Ten microliters of white tea ethanolic extract and five microliters catechins standard were applied on chromatographic precoated silica gel plates (Merck, TLC grade) as the stationary phase. The chromatograms were developed in twin trough glass chamber containing toluene, acetone, and formic acid (5:4:1 v/v) as the mobile phase. The plates were removed after the solvent front has moved from the original position to the finish line and subsequently allowed to dry. After drying, the plate was sprayed with iron (III) chloride (FeCl₃) 1%. The plate then visualized under visible light (white) and short UV (254 nm) light. The amount that each component of a mixture travels can be quantified using retention factors (Rf). Values were calculated for each spot using the formula:

\[ R_f = \frac{\text{Distance from the starting point to the center of the spot}}{\text{Distance from the starting point to the solvent front}} \]

Contamination Analysis

Contaminant analysis was determined by measuring heavy metal content using AAS method, and pathogenic microbial contamination such as E. coli, Salmonella sp., and Pseudomonas aeruginosa.

Examination of Fourier Transform Infrared Spectroscopy (FTIR)

FTIR spectrum components conducted by squirting liquid extract on the surface of a very thin salt plate, for example, potassium bromide, then a second plate is placed on the first plate to spread a thin layer of liquid between the plate. Furthermore, the spectra obtained with the aid of spectrophotometer and software that is attached to the tool.

RESULTS

The characterization of organoleptic observation including shape, smell, color and taste of Camellia sinensis L. Kuntze leaves. White tea has a distinctive aroma and astringent taste as same as the other tea. The macroscopic examination of the leaf consists of botanicals characteristic observation. However, the microscopic examination was carried out with cross-section observation of leaf or specific part fragment of leaf powder. The purpose of this observation is to examine specific parts contained in the organs of plants used.11,12

Macroscopic characteristic of Camellia sinensis (L.) Kuntze showed that the leaves are glossy dark green, alternate, short-petiole, coriaceous, lanceolate or elongated-ovate, and roughly serrate. The leaf blade is 6-18 cm long, 2-6 cm wide, short-stemmed, alternate layout, stiff leaves, elongated ellipse-shaped with finely toothed edges, and pinnately netted venation.11 Mature leaves are bright green colored, smooth, and leathery (Figure 1B). While young leaves appear silver because of the covering of short downy hair on the underside (Figure 1C). White tea leaves are as same as the Camellia sinensis (L.) Kuntze leaves but still rolled up, oval, like a needle with a length of 1.5-3 cm, diameter 1-2 mm, and the surface is covered with hairs which are very delicate (Figure 2A). Dried white tea leaves color are silvery-white (Figure 2B).

The observation of cross sections of Camellia sinensis (L.) Kuntze leaves under a microscope, found the cuticle, epidermis, mesophyll, crystals of Calcium oxalate, sclereid cell, and trichomes (Figure 3). Upper epider-
mal cells are large, distinctly visible with undulating walls, and simple hairs ( unicellular trichomes non-glandular) pointed, undivided, and also very long but not found stomata. The lower epidermis larger than those of the upper surface. The mesophyll is heterogeneous and asymmetrical, it is characterized by the presence of sclereid cells and crystals of Calcium oxalate (Figure 3E and Figure 3F). The type of sclereid cell is astro sclereid with branched, pointed, irregular (often star-shaped). Parenchymal cells similar to those of most other leaves, and not very distinctive (Figure 3D).

While in the observation of a longitudinal cross section of the tea leaves showing trichomes (hairs on the lower epidermis cells), stomata, and sclereid cell (Figure 4). The type of stomata is actinocytic, the variant type of diacytic stoma with two reniform cells (guard cells), oval or nearly round, regularly surrounded by 4 or more tangential elongated cells (Figure 4B). Microscopic observation of the powder found any fragments of leaves and a layer quiet lot of fine hairs, (Figure 5A) and it is showed under field emission scanning microscopy (FE-SEM) analysis (Figure 5B). Young buds leaves are cover with a lot of hairs and it giving the distinctive of tea aroma.

Extracts of white tea leaves were obtained using ethanol 70% with reflux extraction method at 60°C. The yield of the extract is 57.08%. The crude extract obtained was red-brownish with a distinctive aroma of tea. The extract examination was conducted by physicochemical and phytochemical examination. TLC profile is one of the examination methods to analyze the presence of marker compound from the extract. The TLC profiles of white tea extract and catechin standard were obtained under UV 254 light after post-derivatization with FeCl₃ 1%. Figure 6 showed the catechin spot in the white tea extract specifically. Distinct TLC spot on the silica gel plate representing isolated compound with specific Rf values (Rf = 0.56).

Physicochemical examination of the extract included the content of moisture, ash, acid insoluble ash, water-soluble extract, ethanol-soluble extract (Table 1). Phytochemical analysis of the extract showed the presence of alkaloids, flavonoids, tannins, glycosides, saponins. The result for terpene/steroid and anthraquinone was negative. The content of polyphenol from white tea leaves ethanolic extract is 35.73% with the largest concentration of catechins is 18.84% and 17.43% tannins. The derivative content of catechins is EGCG with 7.37% (Table 2).

Description: + = detected; - = undetected

Figure 1: A. Field view of Camellia sinensis (L.) Kuntze, plant; B. fresh tea leaves; and C. shoots.

Figure 2: A. Fresh white tea leaves; and B. Dried white tea leaves.

Figure 3: The Observation of leaves Camellia sinensis (L.) Kuntze using IX70 Microscope Leaf: Cross section: A. cuticle, B. upper epidermis, C. mesophyll, D. parenchyma, E. crystal of Calcium oxalate, F. sclereid cell, G. lower epidermis cell, H. trichome.

Figure 4: The observation of longitudinal cross section leaves using Microscope IX70 Leaf: Longitudinal cross section: A. trichome, B. stomata, C. sclereid cell.

Figure 5: Microscopic observation of the white tea leaves powder White tea leaf: Powder: A. Hairs fragment observation of white tea leaves powder using microscope IX70 and B. Hairs fragment observation under FE-SEM.

Figure 6: TLC chromatogram of white tea ethanolic extracts Extract: Spot: 1. Catechins standard, 2. white tea ethanolic extract
Table 1: Characterization of ethanolic extract of white tea leaves (Camellia sinensis (L.) Kuntze)

| No. | Physico-chemical Parameter               | Result (%) |
|-----|------------------------------------------|------------|
| 1.  | Moisture content                         | 16.03      |
| 2.  | Ash content                              | 6.04       |
| 3.  | Acid insoluble ash content               | 0.12       |
| 4.  | Water-soluble extract content            | 17.18      |
| 5.  | Alcohol soluble extract content          | 33.31      |

Table 2: Phytochemical Analysis of White Tea Leaves Ethanolic Extract

| No. | Phytochemical Parameter | Result |
|-----|-------------------------|--------|
| 1.  | Alkaloids               | +      |
| 2.  | Flavonoids              | +      |
| 3.  | Tannins                 | +      |
| 4.  | Glycosides              | +      |
| 5.  | Saponin                 | +      |
| 6.  | Terpene/steroid         | -      |
| 7.  | Anthraquinone           | -      |
| 8.  | Total polyphenol        | 35.73% |
| 9.  | Total tannin            | 17.43% |
| 10. | Total Catechin          | 18.94% |
| 11. | EGCG content            | 7.37%  |

Table 3: Contaminant Result of White Tea Ethanolic Extract

| No. | Parameter Test | Result | Unit     |
|-----|----------------|--------|----------|
| 1.  | Pb             | ND*    | ppm      |
| 2.  | As             | ND*    | ppm      |
| 3.  | Cd             | 0.042**| ppm      |
| 4.  | E. coli        | Negative| Colonies/unit |
| 5.  | Salmonella sp  | Negative| Colonies/unit |
| 6.  | Pseudomonas aeruginosa | Negative| Colonies/unit |

Description: * ND = not detected; ** Requirements level of Cd ≤ 0.3 ppm

Table 4: FTIR Region Spectra of White Tea Ethanolic Extract

| Bond | Compound | Frequency Area (cm⁻¹) | Intensity |
|------|----------|-----------------------|-----------|
| O-H  | Phenol, Alcohol | 3500-3650 | Medium, broad |
| C-H  | Aromatic | 690-900 | Strong |
| N-H  | Amine, Amide | 3300-3500 | Medium |
| C=O  | Carbonyl | 1820-1660 | Strong |
| C=C  | Alkene | 1650 | Weak |
| C-O  | Ester | 1300-1000 | Strong |

Figure 7: FTIR Spectra of White Tea Ethanolic Extract

The contaminant analysis was observed by analyzing the heavy metals and pathogenic bacterial content. Parameters on heavy metal analyzing including Plumbum (Pb), Arsenic (As), and Cadmium (Cd) (Table 3). Based on the literature, results obtained that heavy metal contamination has been qualified. Furthermore, neither pathogenic bacteria such as Escherichia coli, Salmonella sp., nor Pseudomonas aeruginosa detected. The FTIR spectra of white tea ethanolic extract were obtained with the infrared radiation region is 4000-400 cm⁻¹ (Figure 7). Observation of the FTIR spectrum profiles is done by interpreting the typical peak that appears. The interpretation of FTIR spectra profile showed that the white tea ethanolic extract composed organic com-
pounds which mostly containing functional groups O–H from phenolic or alcohol with medium and broad spectrum, C–H aromatic, N–H amine or amide, C=O carbonyl, C=C alkene, and C–O ester (Table 4).

**DISCUSSION**

Macroscopic and microscopic characters are one of the important criteria for identification. The analysis of the chemical and physical standards are the confirmatory tests for identification. The leaves size is the main criterion for the classification of tea plants, which is the type of the tea leaves are characterized by intermediate size. The different leaf ages produce different tea qualities since their chemical compositions are different. The results of the stomata distribution showed that between stomata and distribution in the leaves are hypostomatic. Stomata associated to photosynthesis to produce food for plants. Stomata is the entrance of CO₂, which is one of the raw materials in the process of photosynthesis. Photosynthesis also altering the synthesis of secondary metabolites in tea plant including catechin, and also polyphenol was correlated to the weather. It causes the relation between the photosynthesis and secondary metabolites content such as catechins.

The presence of a lot of unicellular trichomes on both sides of the white tea leaf creates a microenvironment of water vapor around the leaf. Calcium oxalate crystals are one of the characteristic of the tea leaves and the presence of the crystalline associated with the photosynthesis rate. The white color of the white tea leaf because the chlorophyll has not been formed. White hair that covers the surface coloring white to the leaves when it has been dried. The result of macroscopic and microscopic characterization of the leaves can determine the anatomical structures of plant metabolite storage. The quality of herbal medicines relies on their bioactive constituents. Therefore, TLC fingerprinting serves as an important and powerful tool for standardization and determination bioactive compounds. White tea leaves extract represented the distinctive TLC spots that similar to the RF values of the catechin. The composition of polyphenol as catechin reported was higher (major compound), it is indicated that astrignency was mainly determined by the content of catechins and the other phenolic compounds. White tea leaves have a simple processing without any fermentation process and higher content of polyphenols especially catechins derivatives. It has been confirmed by the determination of the polyphenol, catechins, and EGCG content.

According to the results of the phytochemical analysis in white tea leaves extract, there is the presence of alkaloid, flavonoid, tannin, and saponin. Various types of bioactive found in white tea same as those found in green tea (polyphenols, caffeine, theogallin, gallic acid, theaflavins, glycoside flavonoids, and catechins). Main catechins found in white tea are EGC, EGCG, ECG and EGC which are higher than green tea.

**CONCLUSION**

The current study revealed that the result of macroscopic and microscopic characterization of the leaves can determine the anatomical structures of plant metabolite storage. The present work can be used to provide data about the pharmacognostic characteristics of the white tea leaves of *Camellia sinensis* L. Kuntze that proves the importance of these results. This study will be helpful in the pharmacognostic and phytochemical identification of white tea leaves extract, also the result of identification will be fingerprinted for the proper identification of crude drugs from white tea leaves and provide pharmaceutical preparation in the future.

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**CONFLICT OF INTEREST**

The authors have no conflict of interest to declare.

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