Effect of Small-Scale Box-Fermentation on Catechin and Epicatechin Content of Lampung Cocoa Beans Varieties

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Abstract—Cocoa is one of the food commodities that is favored by its high in polyphenols. The polyphenol and flavonoid in cocoa beans can prevent arteriosclerosis, diabetes, and their related risk factors. This research aimed to analyze the total polyphenolic contents (TPC) and flavonoid contents (TFC) of three cocoa bean varieties, namely LAM, TSH, and SUL1 collected from Lampung, Indonesia. The study also aimed to understand the effect of adding starter culture on TPC and TFC contents during cocoa beans fermentation. The TPC and TFC were extracted using ethanol from ground-dried beans and measured three times using a spectrophotometric method based on the Folin-Ciocalteu method at 760 nm and aluminum reagents at 510 nm. LAM variety contained the highest TPC and TFC among other samples at 62.8 mg GAE/g and 21.9 mg QE/g, respectively. The TPC and TFC decreased during both fermentation treatments with starter culture and non-starter culture. The TPC of cocoa bean with starter culture and non-starter culture decreased to 58% and 59% after fermentation. The TFC of cocoa beans with starter culture and with non-starter culture decreased after five days of fermentation, up to 58% and 75%, respectively. Remarkably, the TPC and TFC in fermented cocoa beans with starter culture were higher than those with non-starter culture. Therefore, it is concluded that three days of fermentation with starter culture can produce fermented cocoa beans with the preferred flavor/quality and maintain the TPC and TFC. However, the polyphenols and flavonoids content were reduced during the fermentation process.

Keywords—Theobroma cacao L.; fermentation; starter culture; total polyphenolic content; total flavonoid content.

I. INTRODUCTION

Cocoa (Theobroma cacao L.) is one of the major commodities for the food, beverage, and additives industries [1]. Cocoa is processed through several lengthy steps, including fermentation. First, the fresh cocoa beans were harvested from selected cocoa pods, fermented for three to five days. Later, the fermented beans were dried and processed through several post-harvest and manufacturing steps to yield products or semi-products, e.g., paste, cocoa butter, and cocoa cake or powder.

The cocoa bean is rich in polyphenols and alkaloids consisting of about 14 to 20% of total bean weight [2]. There are three groups of polyphenols: catechins, anthocyanins, and pro-anthocyanidins. In the type of Forastero cocoa beans, fat-soluble polyphenols constitute about 15 to 20% of the bean, whereas 60% of Criollo cocoa beans polyphenols are procyanidins. The primary catechin is epicatechin, with 35% of total polyphenols. The cocoa polyphenol content varies by cocoa cultivar, area of origin, and cocoa production processes such as fermentation [3]. After the fermentation process, the number of fat-soluble polyphenols decreased to 5%, and the number of epicatechins decreased by ca. 90% from its initial content [4]. During fermentation, polyphenol diffuses from cotyledons and undergoes aerobic oxidation by the activity of polyphenol oxidase. Polyphenols react with proteins converting them to an insoluble form. This process increased the brown color [5], [6].

Presently, society is well-aware of achieving better health by consuming natural products. Cocoa has been one of the promising natural products consumed as a functional food due to the high concentration of antioxidant polyphenols, which makes it leaves astringent after taste [6], [7]. Flavonoids such
as catechins in cocoa also function as antioxidants to prevent oxidative stress damage caused by free radicals [8]. Antioxidants donate one electron for free radicals, which have one unpaired electron to reduce their activity. The ability of cocoa flavonoids to neutralize free radicals outperforms vitamin C and vitamin E by a hundredfold. Catechins are also known to have the ability to inhibit the oxidation of low-density lipoprotein (LDL), thus prevent the thickening of the wall of arteries, with the result of prevention of arteriosclerosis [9]. Other benefits of cocoa polyphenols are neuroprotection, cognitive modulation, cardiovascular disease, protective effect against lung cancer and colon cancer [10]. Cocoa polyphenols are also known to have the potential to protect against diabetes and its related risk factors [11].

The cocoa bean is one of the pre-eminent agricultural export products of Indonesia in addition to CPO. Indonesian cocoa sector has experienced tremendous growth, managed by a spurt of grower farmer contribution in the past 25 years. In general, there are two types of cocoa beans in Indonesia, first is kakao mulia/fine flavor cocoa, and the second is kakao lindak/bulk cocoa. Kakao mulia/fine flavor cocoa mostly from Criollo varieties and some Trinitario varieties. Several clones of fine flavor cocoa in Indonesia, namely DR 1, DR 2, and DR 38. This clone has fine flavor but less productivity and less disease resistance. Kakao lindak/bulk cocoa mostly from Forastero varieties and some Trinitario varieties. There are several clones of bulk cocoa in Indonesia, namely ICS 60, Sulawesi 1, Sulawesi 2, Sca 6, Sca 12, TSH 858. This clone has astrigent flavor, higher productivity, and more resistant to disease [12], [13]. Another cocoa clone was also develop or introduce by local farmer namely MCC01 and MCC02 (Masamba Cocoa Clone) from Luwu, South Sulawesi; BCL (Bayek Clone Langkat) from Langkat, Nort Sumatra; BL 50 (Balubus 50 Kota) from Lima Puluh Kota, West Sumatra and other [13]. Most commonly cocoa variety growth in Lampung is TSH 858, Sulawesi 1, Sulawesi 2, MCC 01, MCC 02, and some local cocoa varieties.

The cocoa export accounts for Indonesia’s largest foreign exchange gross from the agricultural section besides palm oil, rubber, and coconut. Nevertheless, the preponderance of Indonesia’s cocoa exports mainly in the form of raw beans instead of processed cocoa, signifying that Indonesia foregoes on added value capital. Therefore, it is important to process beans domestically; for instance, applying a fermentation process could increase the brown color [5], [6].

Meanwhile, fermentation and other manufacturing processes are known to influence the phytochemical contents of cocoa products and the cocoa bean’s origin [14]. Therefore, it is mainly important to examine the influence of bean’s origin and post-harvest processes on the phytochemical content of cocoa beans. Therefore, the main objective of this study was to evaluate the effect of cocoa beans fermentation on the phenolic content. The information will be of significant use to the development of cocoa products for health purposes from local Indonesian cocoa cultivars.

II. MATERIALS AND METHODS

A. Material

Cocoa beans samples TSH 858 (TSH), Sulawesi 1 (SUL1) were obtained from Pesawaran (Lampung), and local variety Lampung (LAM) were obtained from Pringsewu Regency, (Lampung), Indonesia. The chemicals used were ethanol, Folin-Ciocalteu reagent, sodium carbonate, gallic acid, aluminum chloride, sodium chloride, sodium nitrite, quercetin, catechin, and epicatechin. Some equipment such as rotary evaporator (Büchi, Flawil, Switzerland), centrifuge (Sigma Laborzentrifugen, Osterode am Harz, Germany), spectrophotometer, and high-performance liquid chromatography (HPLC) (Shimadzu, Kyoto, Japan), were used in this study. All chemicals were of analytical grade.

B. Fermentation and sampling

The research flowchart is shown in Fig.1. The beans were extracted from selected cocoa pods, which were then dried in a solar dryer without fermentation. To study the effect of adding starter culture and non-starter culture to the fermentation process on total polyphenolic content (TPC) and total flavonoid content (TFC), two boxes of cocoa beans mass were applying for each treatment. Each box contained 40 kg of mix selected fresh cocoa beans. The fermentation was conducted for five days. The cocoa beans mass was mixed using a plastic shovel every two days to give aeration to the cocoa bean mass. Sampling was taken every 24 hours, and each sample was dried using vacuum drying ALPHA 1-4 LDplus (Martin Christ, Osterode Harz, Germany) and subsequently ground to a powder with a mortar and pestle. Subsequently, cacao powder was evaluated for TPC and TFC [15].

C. Total Polyphenol and Flavonoid Extraction

All samples were weighted appropriately for 5 g and subsequently grounded to powder using mortars. Afterward, all samples were extracted using 15 mL of 80% (v/v) ethanol (Merek, Darmstadt, Germany) three times and pooled to 45
mL. The solutions were homogenized and centrifuged at 10,000 g for 15 minutes at 4°C. The pellet was removed, leaving the supernatant for collection in new tubes. The solutions were evaporated using BUCHI rotary evaporator (Büchi, Flawil, Switzerland) at 40°C and 200 rpm to dryness. The dried sample was dissolved with 5 mL distilled water (DW) [16].

D. Total Polyphenolic Content (TPC) Measurement

According to the method described elsewhere, the total polyphenol contents (TPC) measurements were conducted using Folin-Ciocalteu reagent [17]. Briefly, five grams of the dried extracts were suspended using 5 mL of distilled water and diluted to a ratio 1:20 with DW in the test tubes. A 500 µL of Folin-Ciocalteu reagent (Merck, Darmstadt, Germany) was added to each tube and incubated for 3 minutes at room temperature. After that, 2 mL of 20% sodium carbonate solution (Merck, Darmstadt, Germany) were added to each tube to form the color complex. A blank of DW was treated similarly with the samples. The absorbance was measured at a wavelength of 760 nm using spectrophotometer UV-1800 (Shimadzu, Kyoto, Japan). A series concentration (0-270 µg/mL) of gallic acid (Merck, Darmstadt, Germany) was prepared to construct the standard curve. The TPC was calculated with a gradient equation from the gallic acid standard curve [16]. All of the experiments were conducted in triplicate, and the values are given as the mean of standard deviation.

E. Total Flavonoid Content (TFC) Measurement

The total flavonoid content (TFC) was measured using aluminium complexation reaction [18]. Five grams of dried extracts were suspended using 5 mL of distilled water (DW) and diluted to ratio 1:20 with DW in the test tubes. Appropriately, 150 µL of 10% NaNO₃ reagent (Merck, Darmstadt, Germany) were added to each tube and incubated for 5 minutes at room temperature. After that, 150 µL of 10% AlCl₃ (Merck, Darmstadt, Germany) reagent was added to each tube and incubated for 5 minutes at room temperature. Subsequently, 1 M of NaOH solution (Merck, Darmstadt, Germany) was added to each tube. A blank of DW was treated similarly. The sample absorbance was measured at 510 nm using spectrophotometer UV. A series concentration of quercetin (Merck, Darmstadt, Germany) was prepared to construct the standard curve. The quercetin was diluted with DW at a concentration of 1000 ppm as much as 2 mL. As many as 50, 100, 150, 200, 250, 300, and 350 µL of that solution were added with 2750, 2700, 2650, 2550, 2500, and 2450 µL of DW. The TPC was calculated with a gradient equation from the quercetin standard curve [19]. All of the experiments were conducted in triplicate, and the values are given as the mean of standard deviation.

F. Measurement of Polyphenols by High-Performance Liquid Chromatography (HPLC)

1) Cocoa beans preparations: The concentrations of phenolic and flavonoid compounds were confirmed by HPLC analysis according to the method described elsewhere [14], [20], [21]. Briefly, 2 g of freeze-dried cocoa beans were powdered into a particle size of ca. 1 µm³ in a laboratory mill using 10 mL n-hexane. The sample was then rinsed twice with 50 mL petroleum ether. Subsequently, the defatted powder was dried in a vacuum oven at room temperature [14], [20], [21].

2) Samples Preparations for RP-HPLC Analysis: HPLC analysis was carried out by dissolving a hundred milligram defatted cocoa powder in 3 mL methanol and stirred for 20–30 s with an agitator. Two mL of methanol was adding to the rinsed agitator. The mixture was cooled for 15 min at 0°C and centrifuged for 10 min at 2,683 x g. Subsequently, the obtained supernatant was evaporated by a rotary evaporator. The residue of the evaporated sample was dissolved in 3 mL methanol and filtered through a 0.22 µL and stored at -20 °C until further analysis [14], [21].

Phenolic and flavonoid concentrations were analyzed using high-performance chromatography (HPLC) (LC-20AB, SPD-M20A photodiode array detector (PDA), Shimazu, Kyoto Japan) equipped with an InfinityLab Poroshell 120 EC-C18 chromatography column, 150 mm length, 4.6 mm width, and particle size 2.7 µm at column oven temperature 26°C. The binary gradient method was used in HPLC analysis incorporated 2% acetic acid dissolved in water (A) and a mixture of concentrated acetic acid, water, and acetonitrile (1:9:40 v/v/v) (B). The total runtime of the analysis was 93 min referring to the method described elsewhere [14, 21], as follows: a) initially 0-25 min, 10-30% B; b) 25-50 min, 30-40% B; c) 50-75 min, 40-90% B; d) 75-93 min, 10% B. As many as 20 µL of samples were injected onto the column, and three wavelengths 280, 360 and 520 nm were chosen for analysis in this investigation using HPLC-DAD. For quantitative purposes, a calibration curve was constructed by analysis of known concentrations of different standard compounds.

III. RESULTS AND DISCUSSION

Theobroma cacao L. consists of three main varieties, namely Criollo, Forastero, and Trinitario. The cocoa varieties have been successfully cultivated in Indonesia are Forastero and Trinitario. There are several clones of cocoa from Forastero and Trinitario varieties in Indonesia, and the clone is highly dependent on regions because cocoa is a self-cross-pollinated plant [22]. The three cacao (Fig. 1) pods used in this study, namely LAM, TSH, Sulawesi 1, are known to grow well in Lampung Province, especially in Pesawaran and Pringsewu regency. All three varieties of cocoa are then mixed and used for cocoa beans fermentation experiments.

A. Total polyphenolic content (TPC)

The TPC of cocoa is affected by variety, geographical location, climate, and methods used in the fermentation and drying process. For example, in an adjacent geographical area with an identical cocoa cultivar, the cocoa bean is distinguished by the bean size, chemical composition, and organoleptic characteristics. These characteristics could lead to various flavors and aromas related to the polyphenol contents [3].

As shown in Fig. 2, every variety of cocoa beans have a different TPC concentration. The TPC concentrations for samples in descending order were observed at 62.8; 56.5; and 29.3 mg GAE/g, for LAM, SUL1, and TSH, respectively.
Fig. 1. The varieties of cocoa used in this study, LAM, TSH 858, SUL1.

The LAM sample has the highest TPC followed by SUL1 and TSH. The differences in total polyphenolic content among cocoa clones were also discovered by Oracz et al. [3]. They found differences in TPC between Indonesian, Cameroon hybrid clones, Venezuela, and Papua New Guinea Trinitario beans. The difference in TPC concentration between cocoa clones may be due to genotype varieties and growth conditions of cocoa trees [3]. Oracz et al. suggest that in addition to variety, the polyphenolic contents are also affected by the tree species, growing region, and climatic conditions [3]. It is suggested that high solar exposure induced the formation of anthocyanins and quercetin glycosides as plant self-protection. It is also observed that the lower altitudes will result in higher polyphenol contents [4]. The growth condition of LAM is in Pringsewu Regency with the geographic condition of 104°48′–105°08′E and 05°12′–33′S, whereas TSH and SUL1 are in Pesawaran regency with the geographical condition of 104.92°-105°34′E and 5′12′–5.84′S, respectively. SUL1 are identified have resistance to the disease, red pod color, bean count about 77, bean color violet; TSH resistance to disease, red pod color, bean count 70, bean color violet; local Lampung, are much more resistant to disease, red pod color, bean count 90, bean color violet. The violet color of the bean indicates the presence of anthocyanins in the cotyledons [23].

The fermentation process can affect the total polyphenol contents of the cocoa bean. Thus, the polyphenol contents of unfermented cocoa beans are higher than fermented cocoa beans [24]. The activity of the polyphenol oxidase, as well as permeation during bean fermentation, can cause polyphenol reduction [24, 25]. Polyphenols react with sugar and amino acids during the fermentation process forming the brown color and reduce the astringency [6].

The results obtained were similar to cocoa bean fermentation from Nigeria, where the TPC decreases from the initial to the fifth day of fermentation. The TPC was 20% and reduced to 5% by the fifth day of fermentation [26]. Another
similar pattern was also observed for Nicaraguan cocoa beans, where the TPC was significantly reduced in fermented cocoa beans than in non-fermented cocoa beans [27]. This case is under the escalation of the brown color from the beginning to the end of the fermentation process. Therefore, shorter fermentation is preferable to retain as much as polyphenolic contents of the cocoa bean.

B. Total Flavonoid Content (TFC)
Cocoa flavonoids are utilized because of their health benefits, such as antioxidative activity and preventing arteriosclerosis [8]. The TFC was measured to understand the potential benefit of cocoa flavonoids. The result was depicted in Fig. 4. LAM variety has the highest level of TFC with (21.8 mg QE/g), followed by SUL1 and TSH (9.7 and 4.8 mg QE/g, respectively). This result followed the same trend since the variety of cocoa beans also affected TFC and TPC.

Flavonoids such as catechins are parts of polyphenols and might be declined during a fermentation process. Catechins are oxidized into quinones by polyphenol oxidase activity. Quinones react with amino acids, peptides, and proteins to form condensed tannins, such as brown pigments [28]. As shown in Fig. 4 there was a significant decrease in the TFC during fermentation. Fermentation with non-starter culture, the total flavonoid contents decreased to 75.2% by the end of fermentation, with a daily range of decrease from 9 to 38%. The TFC on the initial day to the first, second, third and fourth day of fermentation decreased by 9.6%, 24.3%, 53.7%, and 70.6%, respectively. Whereas fermentation with a starter culture TFC decreased by 57.6%, with a daily range of decrease from 11 to 30%. The TFC on initial fermentation to the first, second, third, and fourth fermentation days decreased by 11.4%, 24.8%, 34.8%, and 53.4%, respectively. In conjunction with TPC, the TFC at the first three days of fermentation with starter culture was higher than with non-starter culture.

C. Polyphenol and Flavonoid Compound of Cocoa Beans
Details of polyphenol and flavonoid compounds are affected by the fermentation process, as shown in Table 1. The concentration of polyphenol and flavonoid decreased caused by the fermentation process, particularly on the initial of fermentation to the end of the fermentation process. More than 50 % of polyphenol lost is caused by fermentation. (+) - catechin loosed about 63% after five days of fermentation. Parallel with loosed of (-) – epicatechin about 68% after five days fermentation.

| Polyphenol and flavonoid compound | Before fermentation (mg kg⁻¹, ffdm) | After fermentation (mg kg⁻¹, ffdm) |
|----------------------------------|--------------------------------------|-----------------------------------|
| (+) - catechin                   | 1450                                 | 529                               |
| (-) - epicatechin                | 25800                                | 8200                              |

The epicatechin of the cocoa bean from Nigeria was found also decreased, which might arise due to oxidation, condensation, and diffusion out of cotyledon [26]. The oxidation process causes a reduction in concentration since epicatechin acts as the principal substrate of polyphenol oxidase. Other losses of catechins are due to the diffusion into fermentation sweating and polymerization of simple catechins with anthocyanidins to form complex tannins [29]. These indicate that fermentation can cause the degradation of cocoa polyphenols by physical or chemical means.

Catechin and epicatechin are commonly known to benefit health, as reported elsewhere [30-34]. The phenolic cocoa extract can protect the pancreatic beta cells against oxidative stress [30]. In vitro, an in vivo test using cocoa extract showed increased insulin secretion in BRIN-BD11 cell lines and reduced plasma glucose levels in diabetic rats [31]. Another study reported by Ryan concerning the enzymatic study of the fermented and unfermented cocoa extract against α-glucosidase and α-amylase; both fermented and unfermented cocoa extract showed effectively inhibited α-glucosidase and α-amylase [33]. Further study is needed to determine the beneficial effect of cocoa polyphenol as an antioxidant, diabetic agent, and other beneficial effects.

IV. CONCLUSION
This study has successfully examined the influence of bean’s origin (varieties) and two types of fermentation processes (with starter culture and non-starter culture) on the total polyphenol contents (TPC) and the total flavonoid contents (TFC) of cocoa beans. LAM variety has the highest TPC and TFC. Fermentation of cocoa beans with starter culture produced the highest TPC and TFC, especially after the third day of fermentation. (+) – catechin and (-) – epicatechin was reduced by the fermentation process. To maintain the polyphenol content, the third day of fermentation is recommended to reduce the decrease of polyphenol content during the fermentation process.

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1034