Characterization of Polyphenols from Various Cocoa (Theobroma cacao L.) Clones During Fermentation

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Abstract

Cocoa bean is a rich source of polyphenols, which are the largest group secondary metabolite with natural antioxidant property. Polyphenols from cocoa beans was reported to possess health benefits. Fermentation, one step in cocoa processing is needed to improve the quality of cocoa in which the concentration of cocoa bean polyphenols might decrease significantly through oxidation and exudation. Cocoa polyphenols content among different cocoa clones might also vary. The aims of this study were to determine total polyphenols, total flavanoid, epicatechin, and catechin content in several cocoa clones, those were Sulawesi 1, Sulawesi 2, ICCRI 03, and KW 617. Until now, characterization of polyphenols from those clones has not been reported. The effect of five days fermentation to those parameters was also studied. The results of the study showed that fermentation and type of clones significantly affected total of polyphenols, total of flavanoids, epicatechin, and catechin content of the cocoa, there is also an interaction between fermentation and type of clones. Unfermented of Sulawesi 1 had the highest total polyphenols of 96.94±5.83 mg/g, total flavanoids of 90.92±1.89 mg/g, epicatechin of 52.50±0.46 mg/g, and catechin of 1.99±0.02 mg/g content compared to other clones. Among five days fermented cocoa beans, Sulawesi 2 showed the highest total polyphenols and total flavanoids content, while ICCRI 03 had the highest epicatechin and catechin content than other clones. Thus, in can be concluded that although fermentation is required to improve the flavor quality of cocoa, it significantly reduced the content of bioactive compounds. This effect varied among different cocoa clones.

Keywords: cacao, catechin, clone, epicatechin, fermentation, polyphenols

INTRODUCTION

Cocoa (Theobroma cacao L.) plants are widely grown in Indonesia. ICCO (2017) reported that Indonesia's cocoa estimated production in 2017 reached 330,000 tons, so Indonesia is the biggest cocoa producer after Ivory Coast (1,900,000 tons) and Ghana (850,000 tons). The high production is not in line with the quality of Indonesia cocoa beans, especially related to fermentation and filth. In Indonesia, only 10% cocoa bean is further processed with fermentation which impacts to the quality of exported Indonesian cocoa (Suharyanto, 2013).

Fermentation is the key process for cocoa beans to produce good flavor quality. Unfermented cocoa beans can not develop specific aroma cocoa when they are roasted,
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even still strong in astringent and bitter taste. These type of beans are not favoured by cocoa manufactures (Misnawi et al., 2002). Polyphenols are compounds which responsible for bitter and astringent flavour. Bitterness and astringent are usually due to lack of fermentation (End & Dand, 2015).

Basically, fermentation in cocoa beans is aimed to remove mucilage from cocoa beans and to develop a number of flavor precursors in cocoa (Nigam & Singh, 2014). Generally, fermentation process is conducted for five days (Towaha, 2014). Spontaneous fermentation of cocoa beans can be conducted in wooden box containers, in baskets, with or without addition of microbial starter cultures. Cocoa beans are processed into product such as cocoa powder, chocolate and other cocoa derivative products (Oracz et al., 2015). Cocoa has higher total polyphenols, flavonoids, and antioxidant activity than black tea, green tea, and red wine (Lee et al., 2003).

Cocoa is rich source of polyphenols, in particular flavanols, also known as flavan-3-ols or catechins, especially with its high content of monomeric (epicatechin and catechin) and oligomeric (procyanidins) flavanols (Rusconi & Conti, 2010; Alanon et al., 2016). These polyphenols, especially flavonoid have beneficial effects for human health. Cocoa polyphenols have role as antioxidant that provide various effect against several pathological disorders, such as cardiovascular disease, inflammatory process, and cancer (Andujar et al., 2012; Oracz et al., 2013). Epicatechin is a major polyphenols in chocolate and chocolate extracts, powerful inhibitor of plasma lipid oxidation (Vinson et al., 2006).

Diversity in varieties, geographical location, climate condition, and post harvest processing may affect the chemical composition and organoleptic characteristic of cocoa beans, for example cocoa trees grown on West Africa is characterized as bitter in flavour and less aromatic, than the seeds form the plantations in Indonesia (Oracz et al., 2013). Niemenak et al. (2006) reported total polyphenols content in cocoa beans from Cameroon was diverse and ranged from 67.0 to 149.2 mg/g, and cocoa beans from Indonesia was 82.3 mg/g (Towaha, 2014).

During the transformation of fresh cocoa beans to finished products, the polyphenols content can be affected by a variety of biological and processing conditions (Oracz et al., 2013). Thus, the study of polyphenols characterization from various cocoa clones, fermented and unfermented, is necessary to be conducted.

MATERIALS AND METHOD

Cocoa beans of Sulawesi 1, Sulawesi 2, ICCRI 03, and KW 617 clones were obtained from Kaliwining Research Station of Indonesian Coffee and Cocoa Research Institute, Jember with elevation 45 m asl., rainfall 2.130 mm/year, and air temperature 19.6°C–32.5°C. The research used completely randomized design with two factors as four clones and two treatment of fermentation and each combination treatment was repeated two times. Each measurement was also repeated two times.

Cocoa beans used in this study were taken from ripe cocoa pods were cleaned and the beans were taken out from the pods. Preparation of the cocoa bean samples was conducted at Kaliwining Research Station, Indonesian Coffee and Cocoa Research (ICCRl) Jember. Cocoa beans were divided into two groups: five days fermented and unfermented groups. Fermentation of cocoa beans was conducted using batch insert fermentation.
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Cocoa beans were wrapped with shading net, then put either in wooden box (40 cm x 40 cm x 50 cm) at 45–48°C (fermented group) or directly dried in the dry house (unfermented group). Then, cocoa beans were dried for five days and peeled manually to obtain the nibs. Cocoa nibs were ground to form a cocoa liquor.

Cocoa liquor was defatted with Soxhlet method using petroleum ether for seven hours. The defatted residue (cocoa powder) was air dried and stored in refrigerant (Luna et al., 2002). Method of cocoa powder’s total polyphenols and flavonoids extraction refers to Misnawi (2003). A total 250 mg of defatted cocoa powders was sonicated using 40 mL of 80% aqeous acetone (Smart Lab) in a Sonicor (Ultrasonic LC 30 H) for 30 minutes. During sonication, the mixture was kept cold by filling the sonicator vessel with ice water. Then, these crude extract was filtered using vacuum filtration through Whatman no. 1 filter paper to obtain purified extract (residue). The residue was washed with the 80% aqueous acetone and total volume of the extract was made up 50 mL. Method of cocoa powder extraction to obtain epicatechin and catechin followed a method of Shumow & Bodor (2011). Defatted samples were extracted using a mixture of acetone : water : acetic acid (70 : 29.5 : 0.5) followed by sonication at 40°C for 10 minutes and centrifugation for 10 minutes. This steps were repeated twice. Total volume of the extract was made up 25 mL. Then, the extract was filtered with Whatman no. 14 filter paper and PTFE 0.45 µm paper, respectively.

A total 1 mL of purified extract was diluted with 70 mL of distilled water. The extracted polyphenols were then reacted with 5 mL of 2 N Folin – Ciocalteau reagen for two minutes. Then 15 mL of saturated Na₂CO₃ solution was added to stabilize the color formed. The blue color was allowed to develop for two hours and it’s absorbance was measured at 765 nm. Acid gallate (mg/g) standard was used for calculation (Misnawi, 2003)

A total 125 µL of extract was added to 75 µL NaNO₂ 5% solution. The mixture was allowed to stand six minutes, then 150 µL AlCl₃, 10% was added and incubated for five minutes followed by the addition of 750 µL NaOH 1 M. The final volume was adjusted to 2500 µL with distilled water. After 15 minutes of incubation and it’s absorbance was measured at 510 nm. Catechin (mg/g) standard was used for calculation (Rebaya et al., 2014).

Epicatechin and catechin content were determined using the modified method of Shumow & Bodor (2011). 5 µL was injected to the LC-20AD (Shimadzu) detector PDA 278 nm. Separation of epicatechin and catechin was accomplished in C18 reversed phase analytical column using water (0.2% acetic acid) : acetonitril (0.2% acetic acid) (85 : 15) as a mobile phase at flow rate 0.5 mL/min with column temperature 40°C.

Data analysis were performed by the software SAS for Windows (Version 9.1.3) and SIMCA 13.0.2. If the result of variance was significant difference between tratement, then continued with Duncan Multiple Range Test with α = 5%.

RESULTS AND DISCUSSION

Various components from raw cocoa beans contribute in specific cocoa flavors formation by changes during processing, such as alkaloids (methylxanthines), polyphenols, protein, and carbohydrates (Aprotosoeie et al., 2016). Fermentation of cocoa beans
is one of steps in cocoa post harvest processing. It is generally carried out through a spontaneous fermentation either in heaps, box, baskets, tray, or on platform (Papalexandratou et al., 2011). Fresh cocoa beans contain pigment in white to deep purple colour, depending on the amount of anthocyanin (Camu et al., 2008). During fermentation anthocyanin is converted into anthocyanidin, results in bleaching of the purple color of the bean (Vuyst & Weckx, 2016).

The microorganism is responsible in spontaneous cocoa fermentation and their physiological roles during process. Papalexandratou et al. (2011) reported that in cocoa bean fermentation process, there is a colonization by yeast, lactic acid bacteria, and acetic acid bacteria, which produces ethanol, lactic acid and acetic acid. These compounds then diffuse into the beans, and cause the death of the seed embryo. Next, a complex physical processes and biochemical reactions are initiated in the beans to form the required flavor and color precursors. Other microbial metabolites such as ester and pyrazines, may enter the cotyledons of bean and act as flavour precursors or directly as flavour compound (Camu et al., 2008).

This study showed that fermented and unfermented cocoa beans and their clones exhibited significantly different result of total polyphenols, total flavonoids, catechin, and epicatechin content. During fermentation process, cocoa beans polyphenols are oxidized to form condensed high molecular weight compounds, such as tannin (Nazaruddin et al., 2006; Camu et al., 2008). The oxidation is catalyzed by polyphenoloxidase enzyme. The polyphenols are oxidized into o-quinones which then forming a complex with amino acids, proteins, and undergo polymerization with flavonoids to form tannins that produce brown colour (Afoakwa et al., 2012). The enzymes cause a substrate destruction, then giving brown colour (Wahyudi et al., 2015).

The results of this study showed that fermented and unfermented cocoa beans showed significantly different polyphenols content regardless different clones (Table 1). Decrement of total polyphenols, total flavanoids, epicatechin, and catechin content during fermentation time were affected by the fermentation time. Hurst et al. (2011) reported that fermentation decreased the level of epicatechin. Albertini et al. (2015) also reported that fermentation process for two days decreased polyphenols content significantly, but changes in polyphenols content after six days fermentation were less significant.

Table 2 showed that quantity of these compounds also depended on the varietal characteristics of cocoa. Total polyphenols and total flavonoids of Sulawesi 2 were the highest among all clones, while KW 617 was the lowest. Total polyphenols of Sulawesi 2 was significantly different compared to Sulawesi 1 and KW 617, but not significantly different from ICCRI 03, which total polyphenols of Sulawesi 2 > ICCRI 03 > Sulawesi 1 > KW 617. Total flavonoid of Sulawesi 2 was significantly different from other clones, but Sulawesi 1 was not significantly difference with ICCRI 03, which is total flavonoids Sulawesi 2 > ICCRI 03 > Sulawesi 1 > KW 617. The clone with the highest epicatechin and catechin content was ICCRI 03, while the lowest was KW 617. Epicatechin and catechin content of ICCRI 03 was significantly difference with other clones, which is epicatechin and catechin content ICCRI 03 > Sulawesi 1 > KW 617. In other study, Niemenak et al. (2006) reported that different cocoa clones, such as SNK 10, SNK 413, and ICS 84 had different phenolic compound content, that content could be an interaction of several factors such as genetics, physiological, agronomic, and environment factor.
To explain the effect of fermentation on polyphenols content of different cocoa clones, principal component analysis (PCA) was applied to all samples. The PCA bi-plot is presented in Figure 1. In the bi-plot, fermented cocoa and unfermented cocoa groups are clearly separated from each other, which means that fermentation significantly changed cocoa polyphenols content. Fermented cocoa is grouped on the left part of the plot while the unfermented one was on the right. KW 617 clone was the exception. Unfermented KW 617 is clustered close to fermented KW 617 which means that fermentation of KW 617 cocoa beans had less effect on its polyphenols content. It can be seen in Table 3 that for example total polyphenols of unfermented Sulawesi 1 was reduced more than 50% after fermentation (96.94±5.83 mg/g to 41.68±2.81 mg/g), whereas total polyphenols of unfermented KW 617 only reduced slightly after fermentation (48.80±1.94 mg/g to 46.16±0.87 mg/g).

Unfermented Sulawesi 1 had a higher total polyphenols, total flavonoids, catechin and epicatechin content than other fermented or unfermented clones, followed by ICCRI 03, Sulawesi 2, and KW 617. Sulawesi 1 was reported to be more resistance against VSD disease than other clones (Hadi et al., 2014). Prawoto et al. (2013) reported that polyphenols content of resistant clones was higher, it might be because a secondary metabolite was inducible resistance mechanism. Resistant clones contained 22% more components than the susceptible.

Polyphenols content of all clones decreased after fermentation process at different decrement level (Table 3). This might be explained by the difference of polyphenols oxidase (PPO) activity level. PPO is catalyzing reactions for several phenols to produce o-quinones from o-hydraxylation of monophenols (monophenolase activity) and the oxidation of o-diphenols into o-quinones (diphenolase activity) with oxygen as the primary oxidant (Yoruk & Marshall, 2003). Among others, Sulawesi 1 showed the most prominent decrease in all parameters measured in this study (total polyphenols, total flavonoid, epicatechin and catechin content).

PPO enzyme activity is mostly determined by external factors, such as pH. Polyphenol oxidase activity is sensitive against pH changes (Misnawi, 2008). The pH optimum of PPO varies widely with plant source but is generally in the range of 4.0–8.0, such as cherry has pH optimum with estimated 7.4 (Misnawi, 2008). PPO activity is higher at low pH; it is inhibited at high pH. This might be one of the reasons why Sulawesi 1 showed high polyphenols content.

Table 1. Total polyphenols, total flavonoids, epicatechin, and catechin content in fermented and unfermented cocoa powder

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Total polyphenols (mg/g)</th>
<th>Total flavonoids (mg/g)</th>
<th>Epicatechin (mg/g)</th>
<th>Catechin (mg/g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Unfermented</td>
<td>75.635 *</td>
<td>70.570 *</td>
<td>37.653 *</td>
<td>1.501 *</td>
</tr>
<tr>
<td>Fermented</td>
<td>55.939 *</td>
<td>41.012 *</td>
<td>18.280 *</td>
<td>0.871 *</td>
</tr>
</tbody>
</table>

Note: Numbers within the same column with same letter are not significantly different at 5% level according to Duncan test.

Table 2. Total polyphenols, total flavonoids, epicatechin, and catechin from different clones of cocoa powder

<table>
<thead>
<tr>
<th>Clones</th>
<th>Total polyphenols (mg/g)</th>
<th>Total flavonoids (mg/g)</th>
<th>Epicatechin (mg/g)</th>
<th>Catechin (mg/g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sulawesi 1</td>
<td>69.311 *</td>
<td>59.319 *</td>
<td>31.485 *</td>
<td>1.332 *</td>
</tr>
<tr>
<td>Sulawesi 2</td>
<td>73.764 *</td>
<td>64.225 *</td>
<td>27.276 *</td>
<td>1.252 *</td>
</tr>
<tr>
<td>ICCRI 03</td>
<td>72.595 *</td>
<td>59.467 *</td>
<td>39.021 *</td>
<td>1.462 *</td>
</tr>
<tr>
<td>KW 617</td>
<td>47.480 *</td>
<td>40.153 *</td>
<td>14.084 *</td>
<td>0.696 *</td>
</tr>
</tbody>
</table>

Note: Numbers within the same column with same letter are not significantly different at 5% level according to Duncan test.
maximum at about 4.5 and the ratio of monophenolase to diphenolase activity varies depending on plant sources (Yoruk & Marshall, 2003). In cocoa Macedo et al. (2016) reported that pH optimal for PPO in pulp extract was 6.5–6.6 and for seed extract the optimal pH was 5.8–6. Clone and fermentation are not the only factors influenced cocoa polyphenols content, other factors such as drying, roasting, and dutch processing, can affect a flavanol content (Hurst et al., 2011).

Figure 1. Principal component analysis on fermented and unfermented cocoa powder on total of polyphenols, total of flavanoids, epicatechin, and catechin indeks (F Sul 1 = fermented Sulawesi 1, F Sul 2 = fermented Sulawesi 2, F ICCRI 03 = fermented ICCRI 03, F KW 617 = fermented KW 617; Sul 1 = unfermented Sulawesi 1, Sul 2 = unfermented Sulawesi 2, ICCRI 03 = unfermented ICCRI 03, KW 617 = unfermented KW 617)
Table 3. Total polyphenols, total flavanoids, epicatechin, and catechin from different clones of cocoa powder

<table>
<thead>
<tr>
<th>Clones</th>
<th>Total polyphenols (mg/g)</th>
<th>Total flavanoids (mg/g)</th>
<th>Epicatechin (mg/g)</th>
<th>Catechin (mg/g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Unfermented</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sulawesi 1</td>
<td>96.937 ± 5.83</td>
<td>90.915 ± 1.89</td>
<td>52.502 ± 0.46</td>
<td>1.991 ± 0.02</td>
</tr>
<tr>
<td>Sulawesi 2</td>
<td>75.388 ± 4.95</td>
<td>66.182 ± 2.64</td>
<td>38.945 ± 0.07</td>
<td>1.676 ± 0.04</td>
</tr>
<tr>
<td>ICCRI 03</td>
<td>81.416 ± 4.09</td>
<td>78.382 ± 1.88</td>
<td>43.120 ± 0.31</td>
<td>1.571 ± 0.03</td>
</tr>
<tr>
<td>KW 617</td>
<td>48.798 ± 1.94</td>
<td>46.799 ± 3.22</td>
<td>16.047 ± 0.04</td>
<td>0.759 ± 0.04</td>
</tr>
<tr>
<td>Fermented</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sulawesi 1</td>
<td>41.684 ± 2.81</td>
<td>27.723 ± 2.08</td>
<td>10.467 ± 0.18</td>
<td>0.675 ± 0.01</td>
</tr>
<tr>
<td>Sulawesi 2</td>
<td>72.140 ± 3.67</td>
<td>62.298 ± 1.89</td>
<td>15.607 ± 0.15</td>
<td>0.829 ± 0.03</td>
</tr>
<tr>
<td>ICCRI 03</td>
<td>63.772 ± 1.08</td>
<td>40.551 ± 2.26</td>
<td>34.923 ± 0.36</td>
<td>1.347 ± 0.02</td>
</tr>
<tr>
<td>KW 617</td>
<td>46.160 ± 0.87</td>
<td>33.507 ± 1.89</td>
<td>12.121 ± 0.07</td>
<td>0.631 ± 0.00</td>
</tr>
</tbody>
</table>

Notes: Means ± standard deviations.

CONCLUSIONS

Total of polyphenols, total of flavanoids, epicatechin, and catechin content were affected by fermentation process and type of clones. Unfermented cocoa beans have higher total polyphenols, total flavanoids, epicatechin, and catechin content than the fermented ones. Cocoa clone of Sulawesi 1 had the highest content of the above mentioned compounds than other clones, but during fermentation the content was dramatically decreased. Among the fermented samples, the fermented beans of Sulawesi 2 clone had the highest total of polyphenols and total of flavanoids, while the fermented beans of ICCRI 03 clone had the highest epicatechin and catechin content. Changes in polyphenols content due to fermentation as described in this study might affect health beneficial effect of the cocoa, such as its antioxidant activity, and it can produce good flavour quality.

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