Telomere Length Determination Based on Different Sex and Leaf Development of Snake Fruit (Salacca zalacca GART. VOSS.) Revealed by Polymerase Chain Reaction

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ABSTRACT
Telomere has special roles at the end of eukaryotic chromosomes consisting of repeated DNA sequences. It protects chromosomes and DNA from damage. The plant sex and the leaf development may change in telomere length. Snake fruit (Salacca zalacca GART. VOSS.) is dioecious plants that have female and male organs on separate plants. The research aimed to determine the telomere length of snake fruit from different plant sex and the leaf development. In this research, we observed telomere length in snake fruit (female and male plants) using Polymerase Chain Reaction (PCR) procedure. The results from this research showed that telomere lengths of male and female were equal. Telomere lengths in the leaf development showed that younger leaves of both male and female leaves were longer compared to older and dried leaves.

INTRODUCTION
The aging is a process that occur in all of the cells. Aging and death are closely related to telomeres. In the chromosomes at the end of the eukaryotic cell, there is a segment of DNA called telomeres (Wong and Collins, 2003). Telomere consists of a specific sequence of nucleotides. Most plants have TTTAGGG sequence. Telomeres have an important role to protect DNA from damage and maintain chromosome stability in every cell division (Greider and Blackburn, 1996; Shay et al., 2001).

Changes of telomere length (shortening and lengthening) are linked as markers for cell division and aging processes (senescence) in cells (Flanary and Streit, 2003). Telomere length is affected by the age of the cell. The length of the human telomere shortens proportionally with age (Purwaningsih, 2010). Senescence will result in shortening of telomeres (Bernadotte et al., 2016). In addition, sex differences also have different telomere lengths. Telomere length in men is shorter than women (Mayer et al., 2006). The research of Mu et al. (2015) on the telomere length of the dioecious plant stated that the male willow has a shorter telomere length than the female and the female ash has a longer telomere than the male (Mu et al., 2015).

It is necessary to conduct a research using other dioecious plants to find out whether differences in telomere lengths that occur in leaf development and sex will occur in all dioecious plants. In addition, studies related to telomere lengths in different angiosperms of sex are still not widely practiced (Mu et al., 2015). The dioecious plant that is often found in Indonesia and other tropical countries is snake fruit. This research aimed is to determine telomere length of snake fruit on leaf development and sex difference with Polymerase Chain Reaction (PCR).

MATERIALS AND METHODS
Snake fruit (Pondoh variety) aged about 25 years (three male and female plants) were chosen as experimental materials. Samples of young, older, and dried leaves were collected from Turi, Sleman, Yogyakarta. This research had two stages of observation; the first was to observe the leaf condition of the
sample using chromameter to measure the color intensity. Color intensity analysis with chromameter obtained L* (lightness), a* (redness), b* (yellowness). L* color parameter denoted the darkness of a color from 0 (black or dark) to 100 (white or light), a* color parameter measured from 0 to 80, positive (+) was indicated by red and negative (-) was indicated by green, and b* was from 0 to 70, positive (+) was indicated by yellow and negative (-) was indicated by blue color (Suyatma, 2009).

Detection of telomere length with Polymerase Chain Reaction (PCR) begins with leaf DNA extraction using CTAB method (Doyle and Doyle, 1990). The leaves were weighed and cut by 0.10.5 grams, then crushed with mortar and added with 800 μl of CTAB buffer solution which was previously incubated in a water bath at 65 °C for 30 minutes. The CTAB solution consisted of 2% CTAB, 1.4 M NaCl, 100 mM Tris-HCl pH 8, 20 mM EDTA pH 8, and 1% PVP-40, and 1% mercaptoenol).

The mixture was incubated in a water bath at 65°C for 60 minutes and in every 10 minutes was turned back to keep it homogeneous. After incubation, each sample was added with 500 μl of a mixture of 24 chloroform: 1 isooamic alcohol (CIAA) and vortex for 5 min and then centrifuged for 15 minutes at a rate of 12,000 rpm. The forming supernatant was carefully removed and transferred to a new microtube and recorded in volume. 3M sodium acetate was added with 1/10 of the supernatant volume and mixed well. Cold isopropanol was added with 2/3 of the total volume (supernatant + sodium acetate) and mixed well by flipping it through the tube. The mixture was kept in the freezer at 4°C for 1─24 hours. The mixture was centrifuged at a rate of 12,000 rpm for 10 minutes. The supernatant was removed and the precipitated DNA was washed with 500 μl of ethanol 70% and centrifuged for five minutes at a rate of 12,000 rpm. The supernatant was discarded and the precipitate of the DNA was aerated, then stored in the refrigerator at 4°C.

DNA was quantified using GeneQuant to determine the concentration and purity of the obtained DNA. Dilutions were performed to obtain the required DNA concentration in the amplification protocol (PCR). Optimization of the primer annealing temperature was carried out at a temperature of 50 °C to 69 °C. DNA amplification was done by polymerase chain reaction (PCR) to multiply DNA sequences and it used telomere primer of Arabidopsis thaliana (CCCTAAACCC-TAAACCCCTAAACCCCTAACC). Table 1 presents the stages of the DNA amplification reaction. The amplification result was electrophoresed in 100 volts for 55 min using 1.5% agarose in electrophoresis tank containing pH 8 TBE buffer which was heated in microwave until it was completely dissolved, then added with 4μ fluorosafe DNA staining. Furthermore, agarose was removed from the electrophoresis tank and visualized using UV light and photographed with a digital camera.

Telomere length was measured by the length of bands of amplification. Data analysis for leaf color intensity and telomere length was analyzed by variance analysis with 95% confidence level. If there is a significant difference then it would proceed to Tukey Test using SAS (Statistical Analysis System for Windows 9.1.3) software.

RESULTS AND DISCUSSION

Young leaves were collected from the light green youngest leaves that had been opened perfectly (first frond). Older dark green leaves were collected from the fourth frond. Dried brownish yellow leaves were collected from the seventh frond. The appearance of various leaf development in male and female snake fruit is presented in Figure 1.

The color intensity results of female and male leaves are presented in Table 2. The L*, a*, and b* values of male snake fruit leaves were 44.34; -0.32;
and 15.54 respectively. The L*, a*, and b* values of female snake fruit leaves were 45.30; 0.54; and 16.32 respectively. L*, a*, and b* values between male and female leaves were not significantly different.

The various leaf development results are presented in Table 3. The highest L* value leaves were the dried leaves (52.92). The young leaves had higher L* value (42.68) than the older leaves (38.86), so the dried leaf had lighter color than the older and young leaves. L* indicated color saturation, since the dried leaves experienced a reduction in the color density causing the color to become lighter. High pigment content affected brightness (Wahyuni and Widjanarko, 2015). The older leaves had high chlorophyll (green pigment), so it had a lower lightness than the young and dried leaves. Further test results showed that L* value of snake fruit leaves were significantly different in young, older, and dried leaves.

The young leaves had negative a* color parameter which were -5.49 and -2.78, while dried leaves had positive a* (8.60). The value of a* in young and older leaves were negative indicated by the color tended to be green. The a* values of snake fruit leaves were different significantly in young leaves with dried leaves and the older leaves with dried leaves.

The young leaves had higher value of b* (20.25) than older leaves (15.15) and dried leaves (12.39), so the young leaves tended to be yellow compared to older and dried ones. The b* values of snake fruit leaves were different significantly in young leaves with dried leaves.

Leaf is one part of the plant that can be aging because it has a limited life. The process of photosynthesis is influenced by the age of the leaves which then affects the leaf color changes, because there are pigments associated with leaf color in photosynthesis (Susanto, 2008). The leaf development increases leaf photosynthesis ability at the beginning, then decreases along with the senescence process caused by chlorophyll recast and decreasing of chloroplast function. The process of deciduous leaves will be followed by changes in leaf color. Young leaves change color into dark green leaves. The older the green, the higher the chlorophyll content, and vice versa. Furthermore, the older leaves will turn color again into yellowish and then into brownish red. Afterwards, the leaves will fall. As the leaves progress, the ability of the leaf photosynthesis will also increase and then begin to decline slowly (Sestak, 1981). These color changes can occur because the chlorophyll content found in older leaves decrease. The lower chlorophyll content will cause the photosynthesis rate to decrease. The development of leaves is characterized by a change of light green to dark green along with the synthesis of chlorophyll b formed from chlorophyll a. Older dark green leaves have chlorophyll content of 72% larger than the light green leaves (young leaves) (Wolf and Price, 1960 in Pandey and Sinha, 1979). The senescence process produces reactive oxygen species (ROS) in high quantities, as well as the increase in DNA damage due to oxidation (Chen et al., 1995; Song et al., 2005). ROS is known to cause damage to lipids, proteins, and DNA that will accelerate the aging process. According to Furumoto et al., (1998);

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### Table 2. Color intensity of snake fruit leaf (female and male)

| Snake Fruit | Color intensity |
|-------------|-----------------|
| Male        | L* 44.34a | a* -0.32 a | b* 15.54 a |
| Female      | L* 45.30 a | a* 0.54 a  | b* 16.32 a |

Remarks: The means in one column and row followed by the same letter were not significantly different according to Tukey (α 5%)

### Table 3. Color Intensity of snake fruit leaf on various leaf development

| Leaf Development | Color intensity |
|------------------|-----------------|
| Young            | L* 42.68 b | a* -5.49 b | b* 20.25a |
| Older            | L* 38.86 c | a* -2.78 b | b* 15.15ab |
| Dried            | L* 52.92 a | a* 8.60 a  | b* 12.39b |

Remarks: The means in one column and row followed by the same letter were not significantly different according to Tukey (α 5%)
Lorenz et al., (2001); Saretzki and von Zglinicki, (2002); von Zglinicki, (2002) this oxidative compound can also speed up telomere shortening.

Detection of Telomere Length

The telomere length was measured by the length of the bands amplified by male (Figure 2) and female (Figure 3). The telomere length of female leaves was 1,594.4 bp and that of male leaves was 1,444.4 bp. The telomere length between male and female leaves were not significantly different. Telomere length between female and male snake fruit leaves showed no significant difference. The results of research on dioecious plants, either male or female in Ginkgo biloba also showed similar results (Song et al., 2010). Riha et al. (1998) also showed the same length of telomere between the male and female Melandrium albums. While the female ash had a shorter telomere than males, the female willow had longer telomeres than males (Mu et al., 2014).

According to Song et al. (2010), further research on telomere length between males and females is still needed.

The telomere length of young, older, and dried leaves decreased (Figure 4). Telomere length of snake fruit in young leaves was 2,483.3 bp, in older leaves was 1,500 bp, and in dried leaves was 575 bp. Further test results showed that telomere length of snake fruit leaves were significantly different in young, older, and dried leaves. The older the leaves, the more shortened the telomere would be. This is

| Snake Fruit | Telomere Length of Leaf (bp) |
|-------------|-----------------------------|
|             | Young                      | Older                      | Dried                     |
|             | 1  2  3 Mean               | 1  2  3 Mean               | 1  2  3 Mean              |
| Young       | 2,500 2,500 2,600 2,533.33 | 2,200 700 1,700 600 450 550 |
| Older       | 2,400 2,400 2,500 2,433.33 | 950 850 1,300 600 600 600 600 |

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Figure 2. DNA amplification of female leaves (SB): young (M), older (T), and dried (K)
Remark: L = ladder, 1–3 = sample number

Figure 3. DNA amplification of male leaves (SJ): young (M), older (T), and dried (K)
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| Young       | 2,500 2,500 2,600 2,533.33 | 2,200 700 1,700 600 450 550 |
| Older       | 2,400 2,400 2,500 2,433.33 | 950 850 1,300 600 600 600 600 |
consistent with the research results on *Ginkgo biloba* that had shortened telomeres in older leaves (Song et al., 2010), as well as willow and ash plants in older leaves (Mu et al., 2014).

The older leaves of female snake fruit in sample 1 was 2,200 bp, in sample 2 was 2,200 bp, and in sample 3 was 700 bp. The older leaves of male snake fruit in sample 1 was 2,100 bp, in sample 2 was 950 bp, and in sample 3 was 850 bp. The older leaves of female and male snake fruit showed varying telomere lengths. It was presumed that inter-individuals had physiological differences related to aging. Based on research by Liang et al. (2014) in *Panax ginseng*, it was indicated that there was a complex relationship between telomere length and plant growth and it might be affected by environmental factors and telomerase activity.

Variation in telomere lengths might occur due to the relationship between telomere length and telomerase enzyme activity. The enzyme had been observed in several plant species (Fajkus et al., 1996, Kilian et al. 1998, and Mu et al., 2015). In maize and barley, telomerase activity was suppressed during tissue differentiation (Kilian et al., 1998), so telomeres were shortened due to end-replication problems (Kilian et al., 1995; Kilian et al., 1998). The highest telomerase activity in plants was found in meristems and reproductive organs, whereas in endosperms, leaves, and stems there was little or no telomerase activity (Liang et al., 2014).

The shortening of telomere may result from the early stages of apoptosis (Rescalvo-Morales et al., 2016) and the occurrence of DNA damage (Ramirez et al., 2003). Environmental stress factors and physiological disorders associated with aging can increase DNA damage, especially telomeres (Watson dan Riha, 2011). Stressful plants can produce oxidative compounds, where these oxidative compounds can also speed up telomere shortening (Furumoto et al., 1998; Lorenz et al., 2001; Saretzki dan von Zglinicki, 2002; von Zglinicki, 2002). Telomeres can be a marker of apoptosis caused by an increase in oxidative compounds (Shammas et al., 2004; Granato et al., 2009). This is evidenced in the study by Na et al.

| Snake Fruit | Telomere Length (bp) |
|-------------|----------------------|
| Male        | 1,444.4 a            |
| Female      | 1,594.4 a            |

Remarks: The means in one column and row followed by the same letter were not significantly different according to Tukey (α 5%).

| Snake Fruit | Telomere Length (bp) |
|-------------|----------------------|
| Young       | 2,483.3 a            |
| Older       | 1,500 b              |
| Dried       | 575 c                |

Remarks: The means in one column and row followed by the same letter were not significantly different according to Tukey (α 5%).
Telomere length of snake fruit leaves were significantly different in young, older, and dried leaves. The older the leaves grow, the shorter the telomere becomes. Telomere length of snake fruit in both female and male are equal.

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