INTRODUCTION

Researchers have tried to understand the mechanism that leads to Rafflesia anthesis for many years. Unlike any other endoholoparasitic plant (i.e. holoparasitic plant with its tissue embedded into the host plant), Rafflesia has fragmented endophytes scattered inside the host body, Tetrastigma (Vitaceae) (Nikolov et al. 2014; Mursidawati et al. 2019). These scattered parasitic tissues hypothetically come from one germinating point and spread into the whole host plant tissue (Wicaksono et al. 2017), most probably as clones (Wicaksono et al. 2020). The parasitic endophytic cells proliferate by the push of the forces from the surrounding dividing cells of its host and these actively dividing cells reside within the vascular cambium tissue (Mursidawati et al. 2019). Alternatively, it has been suggested that the surrounding root new cortex tissue in the Tetrastigma...
might also contribute to the endophyte “growth vessel”. This suggestion comes as the new cortex or the pericycle tissue replaced the original cortex tissue when the secondary root is developed as observed in another Vitaceae, Vitis sp. (Gambetta et al. 2013). However, between the scattered endophytes, no study so far has unraveled the physiological and molecular signaling between the endophyte clones, which might trigger the anthesis stage of the bud.

There are several plant growth regulators (PGR) that contribute to flower development, like auxin and cytokinin. Auxin as observed in Arabidopsis thaliana (L.) Heynh. (Alabadi et al. 2009) and in Camellia azalea C.F. Wei (Fan et al. 2015) initiates primordial flower growth. Cytokinin triggers the growth of flower organs, e.g. flower bud and ovule formation, and regulates flower and seed size (van der Krieken 1989; Bartrina et al. 2011). The effect of these regulators on the growth of Rafflesia cantleyi Solms-Laubach was observed in a transcriptomic study of genes and transcription factors related to auxin, cytokinin, gibberellin, abscisic acid, and jasmonic acid (Amini et al. 2019). As Rafflesia endophyte growth is cryptic, observations on the effects of PGR on flower growth can only be done via two possible ways, i.e. culture of the Rafflesia tissue and in vivo manipulation.

While several studies have been conducted on culturing Rafflesia tissue (Sukamto 2001; Mursidawati & Handini 2009; Sukamto & Mujiono 2010; Wicaksono & Teixeira da Silva 2015; Molina et al. 2017), only one has successfully induced the cultured bud tissue of R. arnoldii R.Br into callus tissues (Sukamto & Mujiono 2010). The callus tissues proliferated into the whitish strands under the picloram (synthetic auxin) treatments, but no response was observed on the zeatin (cytokinin) (Sukamto & Mujiono 2010). The strands might be consistent with Nikolov et al. (2014) uniseriate strands of Rafflesia in the early stage of endophyte growth inside the host tissue, which later may form cellular clusters before differentiating into a flower bud (Mursidawati et al. 2019; Wicaksono et al. 2020). From the results in Sukamto & Mujiono (2010) so far, this in vitro study method could offer an effective way to visually observe the vegetative growth of the endophytic tissue because no other study has successfully managed to differentiate the Rafflesia callus into a flower bud. On the later stage of flower development, the generative stage, an in vivo study performed in this short study might reveal the effect of the PGR on the growing flower bud of Rafflesia.

This study was an attempt to replicate one method from Mariani et al. (2011) in Aglaonema on R. patma Blume, although the procedure was performed in a non-analogous organ (Aglaonema stem vs. Rafflesia flower bud in Mariani et al. (2011) vs. this study, respectively). The study on Aglaonema involved an injection of benzyladenopurine (BAP) (synthetic cytokinin) into the axillary node of the stem to induce the axillary bud growth. For the present study, indoleacetic acid (IAA) (auxin) and kinetin (cytokinin) were used. Hypothetically, based on Sukamto & Mujiono (2010), the auxin could induce bud growth better than cytokinin. However, it was unclear if any other effects could be rendered since the whole flower bud organ was involved instead of only an undifferentiated callus. Nevertheless, this study is the first preliminary attempt of in vivo PGR induction on Rafflesia.

MATERIALS AND METHODS

Plant materials

The Rafflesia patma plant grown on its host plant Tetrastigma leucostaphylum (Dennst.) Alston ex Mabb was a result of grafting between the grown host vine rootstock in Bogor Botanical Garden and host vine scion from its natural habitat in Pangandaran Nature Reserve, West Java, Indonesia, as described in Mursidawati et al. (2015). This same plant was also used in previous studies (Wicaksono et al. 2017; Mursidawati et al. 2019; Mursidawati et al. 2020; Wicaksono & Mursidawati 2020). The experiment started on July 28, 2020 on six selected flower buds in cupule stage (Susatya 2020), approximately 5–6 months old on T. leucostaphylum roots. The number of buds was limited due to their availability at the time of the experiment at Bogor Botanical Garden.

Injection and observation

The injections followed the Aglaonema protocol by Mariani et al. (2011). Six flower buds were used for the present study, with two buds per treatment/control. Plastic labels were attached to the nearby root to identify the bud and the type of treatment or control it received. Every bud within the same treatment/control were on the bud and the type of treatment or control were on separate T. leucostaphylum roots to prevent any possibilities of cross-contamination effects between treatments or between treatment and control. All injections used three different short-needle sterile syringes (insulin syringe) per treatment with a maximum volume of 100 ml. The first treatment was 3 mg/l IAA, the second was 1 mg/l kinetin, and the third was sterile distilled water as the control. According to Mariani et al. (2011), the injected cytokinin was 30 mg/l and the tissue culture auxin and cytokinin was 3 mg/l for both. However, we used the low concentrations used in the tissue culture to prevent any shock to the tissue, as high cytokinin level may lead to hypersensitivity response similar to the condition during pathogenic attack by killing the cells at the infected site (Novák et al. 2013). In similar manner, high auxin concentration possesses herbicidal effect.
(Grossmann 2007). To make the stock, 3 mg of IAA powder was first dissolved with 1 ml of NaOH (0.2 M) to dissolve it before being added to the 100 ml of sterile distilled water. Similarly, 3 mg of kinetin powder was first dissolved with 1 ml of NaOH (0.2 M) before being added to the 100 ml of sterile distilled water. In the end, 30 mg/l of each IAA and kinetin were acquired as the stocks. To obtain final 3 mg/l concentration, each stock was diluted 10 times using sterile distilled water.

The syringe needle was administered at the base (proximal) closest to Tetrastigma of flower bud cupule to avoid damaging the growing flower bud, especially since the R. patma bud meristematic tissue growth is more active at the distal region while the proximal region already has matured parenchyma cells (Mursidawati & Wicaksono 2020). The needle only pierced the bud approximately halfway the length of the whole needle (i.e. the needle is 1 cm long, only 0.5 cm is pierced) to minimize the damage to the bud and subsequent stress responses. Upon injection, the R. patma flower bud was very robust, so the injection was performed to be as slow as possible to maximize the possible intake of PGR (and water) by the buds, despite any drawback leaks caused by the resisting hard tissue of the flower bud. The injected buds were measured at the start of the experiment and then every week using Vernier caliper until stagnant growth was reached or death occurred. The growth data were averaged and divided per unit of time (week) to obtain the difference in growth rate (cm/week) per treatment/control.

RESULTS AND DISCUSSIONS

General growth pattern

From the initial week (week 0) of the experiment, the buds appeared to grow before they entered stagnation on the third week (Table 1). The stagnation conditions across kinetin and IAA treatments might indicate that the PGR effects only last for two weeks. Our results were similar to Mariani et al. (2011) which showed stagnation of axillary growth within two weeks. Compared to the control, bud growth was slow, observed towards the first week, and from the third week towards the fourth week.

Table 1. Flower bud measurements of R. patma per variable.

| Week | Kinetin | IAA | Control |
|------|---------|-----|---------|
|      | K1      | K2  | K3      | K4      | K5      | K6      |
| 0    | 2.07    | 2.50| 2.77    | 3.75    | 2.15    | 2.15    |
| 1    | 2.08    | 2.90| 2.78    | 3.80    | 2.19    | 2.19    |
| 2    | 2.18    | 3.20| 3.10    | 3.84*   | 2.19    | 2.19    |
| 3    | 2.18    | 3.28| 3.30    | 3.90*   | 2.19    | 2.19    |
| 4    | 2.20    | 3.30| 3.34    | 3.90*   | 2.21    | 2.22    |
| 5    | 2.20    | 3.30| 3.35    | 3.94*   | 2.21    | 2.22    |

Note: *) The bud reached the cupule-bract stage, as defined by Susatya (2020).

Growth rate comparison

The kinetin-injected buds showed the highest growth rate from the first-week to the second-week interval, despite a slight slow-down in the second-week interval, and drastic slow-down afterward (Fig. 1A). On the other hand, IAA-injected buds showed a burst of growth from the first-week to the second-week interval before slowing down to stagnation towards the fourth-week interval (Fig. 1B). Of the two IAA-treated buds, one showed rapid growth (Table 1, K3) and the other a transition from cupule stage to the cupule-bract stage (Table 1. K4; Fig. 2), despite both having the same growth age of approximately 5–6 months old (Mursidawati & Wicaksono 2020). Comparatively, the control buds showed only slight growth in the first and fourth weeks with growth stagnation between the second and third weeks.
Auxin and cytokinin functionality on the bud growth

In plant growth, auxin works by altering the cell wall rigidity, allowing water transport into the cell and resulting in the elongation of the cell (Majda & Robert 2018). Auxin plays a significant role in apical growth (upwards and downwards) and dominance (Taiz et al. 2015). In comparison, cytokinin regulates the cell cycle especially the interphase-mitosis transition in plant tissue (Schaller et al. 2014). In the present study, it appeared that the effect of cytokinin (kinetin) was almost immediate as shown by high initial growth in the first-week interval (week 0 towards 1; Fig. 1A), while auxin (IAA) delayed growth until the second-week interval (week 1 towards 2; Fig. 1B).

In the R. patma, flower buds treated with kinetin (cytokinin), it appeared that the cytokinin directly affected cellular division upon injection, which led to the enlargement of the flower buds. Comparatively, in the IAA (auxin)-treated flower buds, the auxin induced the distal meristem of the flower buds, resulting in the bract emergence. For now, it is not understood if the flower bud distal meristem that differentiates into bracts, perigone lobes, and the central disc organ (Mursidawati & Wicaksono 2020) is analogous in function to the plant apical meristem that differentiates into a shoot or root. Auxin has been known to be produced in young developing leaves in the plant apex and transported unidirectionally towards the root tip via the phloem, but later auxin can be synthesized in different plant tissues (Chandler 2009), while most of the cytokinin is synthesized in roots, cambium and actively dividing cells, and transported by xylem (Chen et al. 1985; Campbell et al. 2008). Hence, it is possible that some primordial tissue in Rafflesia plays a role in auxin production. In Rafflesia, auxin and cytokinin actively play a role during flower development, with cytokinin mainly regulating flower development by activating MADS-box transcriptional factor genes (Amini et al. 2019).

Limitations and future challenges

The problem of performing experiments with Rafflesia is the limited availability of plant material in the wild or the botanical garden. Of all available flower buds within a specific area, we can only select a few of them in order to keep the population sustainable. The only solution is to repeat the study in a controlled area (e.g. botanical garden or specific area in a national park with permission), where flower buds may be more readily available within the population to sample. Alternatively, a study could be repeated in separated time frames but in the same period, e.g. one in July 2020, later in July 2021, etc. Careful selection for the sample is also highly encouraged, especially due to Rafflesia population fragility.

The other problem to be considered is the PGR injection to the flower buds. As it causes an open wound to the flower bud, which may render the Rafflesia bud vulnerable due to exposure to microorganisms or herbivores. Environmental factors, like temperature and soil moisture, might also play roles in flower growth (Major 1980). In the wild, sometimes Rafflesia buds never reached the anthesis period and died during the flower bud development period, thus indicating that the flower bud may be sensitive to slight disturbances (Mursidawati...
2014). If a wound in the bud is exposed by injection, it may further lead to death.

Lastly, we have also considered the PGR effectiveness periods. According to Mariani et al. (2011), single injection of cytokinin resulted in a plant response within two weeks. Other studies have revealed that in the cortex and stele of root tissue, auxin lasts for 0.12 and 21 hours, respectively. In a condition where auxin was provided using agar and it is transported in the order of hundreds to thousands of pictograms per hour (Nonhebel et al. 1985; Kramer & Ackelsberg 2015). As for cytokinin, transportation mode is unknown. It is unclear, however, under in vivo conditions with injection treatments as in Mariani et al. (2011) and the present study, the speed and size would be for the metabolic rate and transport of auxin and cytokinin by the Rafflesia exposed tissues. Giving multiple doses to Rafflesia flower bud could open more wounds that damage the bud, hence a better way to administer the PGR doses needs to be considered. Additionally, other than the requirement to test on a larger sample number, this study should probably be tested on other Rafflesia species to see if they provide similar results to R. patma. Also, other PGRs like gibberellic acid (GA) and jasmonic acid (JA) and the combinations between them should be tested to see if different results in flower bud growth might occur. From that, a histological analysis could be done to determine if the given PGRs or their combinations might alter tissue growth compared to the control.

CONCLUSION

Auxin and cytokinin might affect the R. patma flower bud growth. Injecting both IAA and kinetin, which are auxin and cytokinin, respectively, caused flower bud enlargement. However, only IAA treatment showed flower bud transition from the cupule stage to the cupule-bract stage. This preliminary overview might reveal the significance of auxin in flower bud development in Rafflesia compared to cytokinin. Nevertheless, more samples are needed to confirm this claim, with PGR administration needing to be made more effective to reduce stress or damage in the treated flower buds. Additionally, other PGRs and their combinations will be required in future studies, combined with histological analysis of the treated flower bud tissues.

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