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Banana Cultivars Microshoot Induction and Plantlet Formation using Cytokinin and Auxin

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Abstract

Banana is a horticultural plant with very high potentials, which contains carbohydrates and vitamins that are useful in fulfilling people's food and nutritional needs. Hence, this study aims to produce superior banana seedlings and develop a protocol for their mass production using a plant *in vitro* culture technique. This was a two stage-experiment i.e. microshoot production and plantlet formation. The result showed that Gebyar cultivar produced more shoots than the Kepok Kuning cultivar, with an average of 4.25 microshoots explant⁻¹. However, Kepok Kuning produced more leaves than Gebyar, with an average of 4.64 leaves plantlet⁻¹. Banana shoots cultured on the media containing Indole-3-acetic acid (IAA) at a concentration of 2.5 μ M produced the highest leaves number. Meanwhile, those cultured on the media containing 1-Naphthalenesacetic acid (NAA) at a concentration of 7.5 μ M produced the highest roots number. A Murashige and Skoog (MS) medium supplemented with 6-Benzylaminopurine (BAP) up to 30 μ M and the one supplemented with 7.5 μ M of NAA are suitable for Kepok Kuning and Gebyar cultivars micropropagation with regard to microshoot induction and plantlet formation, respectively.

Keywords: BAP; Gebyar; in vitro culture; Kepok Kuning; NAA; sword sucker

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INTRODUCTION

Banana is one of the horticultural plants with great potential due to its high nutritional value and being an income source for local farmers. This crop is also importantly cultivated by the people living in tropical countries and it is the world's fourth agricultural commodity after rice, wheat and maize (Satuhu and Supriyadi, 2007). According to Singh et al. (2014), almost all modern edible bananas originate from two wild diploid species, namely *Musa acuminata* designated as genotype AA and *Musa balbisiana* as BB. Some are found to have an AA or AB genotype but the vast majority is triploid. Many domesticated bananas are proven to be triploid with a genotype of AAA, AAB or ABB. There are also seedless cultivated AA and AB diploid, AAAA, AAAB, AABB and ABBB tetraploids.

Indonesia's total banana production has been relatively stable in the last five years with an average of 7.202 million tons year⁻¹. This reached 7.280 million tons in 2019, with only 0.22% increase from 2015 (BPS - Statistics Indonesia, 2020). The Directorate of Horticultural Production Development and the Horticulture Research and Development Center prioritize banana as one of the fruits to be developed and studied. Production in the last 10 years has been influenced by plant pests and diseases

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(Prasetyo et al., 2020), especially the outbreak of Panama disease (Fusarium wilt), Bacterium Blood disease caused by *Pseudomonas celebensis* and Moko disease (bacterial wilt) (Mondal et al., 2012; Ploetz, 2015). Furthermore, leaf wilt diseases such as Panama caused by the fungus Fusarium oxysporum Schlecht sp. Cubans and Moko caused by the bacterium Raisotonia solanacearum are the most dangerous to bananas. The two pathogens are soil-borne and easily transmitted, which enhance their spread to almost all banana plantations in Indonesia (Ploetz, 2015). They form chlamydospores in the soil which survive for a very long time (Ghag et al., 2015), therefore making the control of the diseases produced to be extremely difficult.

Most consumable bananas are triploid and sterile, making genetic improvement through conventional crossing difficult. Consequently, the integration of *in vitro* technique for genetic improvement is crucial. The provision of superior banana seedlings is an important strategy to increase production. High-quality seedlings can be developed using *in vitro* culture technique. The factors influencing this process include plant genetic composition and its expression, nutrition, physical growth factors and the utilization of Plant Growth Regulators (PGRs) and vitamins (Amoo and van Staden, 2013; Kadhimi et al., 2014).

The three banana cultivars used in this study were Ambon Nangka (Musa acuminata × Musa balbisiana AAB), Gebyar (Musa acuminata × Musa balbisiana AAB) and Kepok Kuning (Musa acuminata × Musa balbisiana ABB). They are widely cultivated in Indonesia, especially in Banyumas Regency, Central Java. Studies related to the microshoot induction of bananas with AAB and ABB genotypes were previously carried out by Hui et al. (2012), Kindimba and Msogoya (2014) and Yatim (2016). During the induction of axillary shoots growth, 6-Benzylaminopurine (BAP) and kinetin are commonly used due to being very active, longlasting, easily translocated, stable and heat resistant. Meanwhile, cytokinins and auxins addition to the in vitro culture triggers cell division to form new cells, controls apical dominance and inhibits tissue aging (Schaller et al., 2014; Feng et al., 2017). The success of banana shoot multiplication using BAP has been reported by Ali et al. (2011); Vishnevetsky et al. (2011) and Govindaraju et al. (2012). In the plantlet formation phase, auxins such as Indole-3-acetic acid (IAA), Indole-3-butyric acid (IBA) and 1-Naphthalenesacetic acid (NAA) are used. Banana shoots rooting in *in vitro* culture has been successfully carried out by adding auxins according to Muller and Leyser (2011); Müller et al. (2015) and Hossain et al. (2016).

This study aims to produce superior banana seedlings and develop a protocol for their mass production using *in vitro* plant culture technique, as well as to evaluate the influence of cytokinins and auxins types and concentrations on the three banana cultivars plantlet formation. The mass production of disease-free plantlets provides a means whereby plants including bananas (Daniells, 2007), are improved for sustainable cultivation (Viljoen et al., 2004).

MATERIALS AND METHOD

Plant material

The materials used in the microshoot induction phase were the sword suckers, i.e., vigorous shoots arising from the rootstock of banana plants, of three banana cultivars namely Gebyar, Ambon Nangka and Kepok Kuning. They were obtained from a local banana nursery in Purwokerto City, Banyumas Regency, Central Java, Indonesia. The use of sword sucker in banana micropropagation was also reported by Ali et al. (2011) and Govindaraju et al. (2012). During the plantlet formation phase, the Gebyar and Kepok Kuning cultivars microshoots obtained from the induction phase were used.

Explant preparation

Explant isolation was carried out by selecting banana sword suckers with an approximate size of 7 cm \times 6 cm \times 15 cm. The shoots were excised from the sword suckers and cut into $2 \text{ cm} \times 2 \text{ cm}$ \times 2 cm size and subsequently soaked in distilled water. Then, the explants were washed using sterile distilled water supplemented with few drops of Tween 20 to reduce surface tension. They were sterilized in a Laminar Air Flow cabinet (LAF) through immersion in 70% ethanol for 2 minutes, followed by double immersion in 0.2% HgCl₂ for five minutes. Sinha and Deka (2016) reported that HgCl₂ and Tween as effective chemicals for sterilizing Malbhog banana cultivar. Next, the explants were rinsed in sterile distilled water for 30 seconds and this was repeated three times. Each side of the sterilized explants was cut to remove residues and dead tissue which resulted in $1 \text{ cm} \times 1 \text{ cm} \times 1 \text{ cm}$

size as described by Govindaraju et al. (2012). Furthermore, they were inoculated and grown on a Murashige and Skoog (MS) medium supplemented with 15 μ M BAP for axillary shoot induction, then the cultures were incubated at 24 °C with continuous light.

Microshoots induction and plantlet formation

MS basal medium (Sigma-Aldrich-M5519) supplemented with 20 g L⁻¹ sucrose, 2 µM IAA (Sigma-Aldrich-I2886), BAP and kinetin (Sigma-Aldrich-K0753) as treatments, solidified with 2.5 g L⁻¹ phytagel (Sigma-Aldrich-P8169), was used for microshoot induction. The treatments were arranged in a Spit-Split Plot Design, with three replications. The main plot was banana cultivars consisting of Ambon Nangka, Gebyar and Kepok Kuning. The sub-plot was type of cytokinin consisting of BAP (Sigma-Aldrich-B3408) and kinetin (Sigma-Aldrich-K0753). The sub-sub-plot was concentration of cytokinin consisting of 15, 20, 25 and 30 µM. The cytokinin concentrations used are a modification of the results reported by Prayoga and Sugiyono (2010). The explants were sub-cultured into the same medium after six weeks of culturing.

For plantlet formation, microshoots were transferred to the root induction medium, namely an MS basal medium supplemented with 20 g L⁻¹ sucrose and solidified with 0.2% phytagel. The treatments were arranged in a Split-Split-Plot Design, with three replications. The main plot was banana cultivars which consisted of

Gebyar and Kepok Kuning, while the subplot was an auxin type consisting of IAA, IBA (Sigma-Aldrich-I5386) and NAA (Sigma-Aldrich-N0640). The sub-sub-plot was auxin concentration consisting of 2.5 μ M, 5.0 μ M, 7.5 μ M and 10 μ M. NAA and IBA had been reported as the best treatments for banana culture in rasthali cultivars (Govindaraju et al., 2012). The variables observed in this study were the formation of microshoots and plantlets. The parameters measured include the number of shoots, the number of leaves and the number of roots.

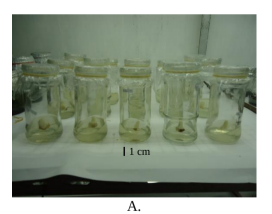
Data analysis

The data obtained were analyzed using an Analysis of Variance (ANOVA), followed by the Honestly Significant Difference (HSD) test with a confidence level of 95%.

RESULTS AND DISCUSSION

Microshoots induction

The *in vitro* culture of several banana cultivars has been successfully carried out, where explants were planted in the treatment medium (Figure 1). New shoots developed after six weeks of incubation (Figure 2) and few microshoots appeared after several sub-cultures (Figure 3). In microshoot induction, Gebyar and Kepok Kuning produced more microshoots than Ambon Nangka. Hence, Ambon Nangka did not proceed to the plantlet formation stage due to its low microshoot production.



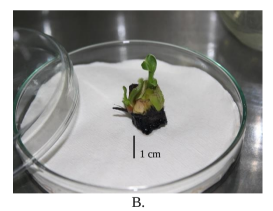


Figure 1. The effect of different banana cultivars and cytokinin types and concentrations on microshoot formation. (A) The appearance of banana explant on treatment medium, (B) Gebyar microshoot development on MS medium supplemented with BAP

The small number of microshoots produced was probably due to the PGR concentration used (Bordoloi, 2016), the explants physiological conditions such as apical dominance (Bhende and Kurien, 2015; Kebrom, 2017) and the culture period (Rahman et al., 2013). The cytokinin

concentration added to the media did not stimulate new shoots formation. According to Ahmed et al. (2014), the time required for explant shoot formation on a combined media (BAP and IAA) was slower than on the media containing only BAP or in combination with kinetin. This result is in agreement with the report by Buah et al. (2010) and Ashraf et al. (2014). BAP's better effect on shoot formation compared to kinetin may be attributed to its high stability within in vitro cultures, because it is not easily broken down and therefore persists in the medium. Possibly, the BAP amount conjugated in the medium was smaller than that of other growth regulators, leading to the presence of more BAP in their free or ionized forms, which were made readily available to plant tissues from the medium (Buah et al., 2010).

Furthermore, the explants in this study were taken at the dry season peak, hence it is suggested that they had high abscisic acid (ABA) content as a response to water stress. ABA is widely known as a stress hormone, and its level increases in the presence of stress, including water stress (Hu et al., 2016). The high concentration of endogenous ABA inhibits microshoots formation and growth (Feng et al., 2012). These conditions are thought to have influenced explants growth and microshoots development. To solve this problem, an extremely high cytokinin concentration have to be used to break the dormancy caused by ABA, which in turn stimulates microshoots development and growth (Goggin et al., 2015; Qiu et al., 2019). Nuraini et al. (2016) and Prasetyo et al. (2020) stated high cytokinin utilization breaks dormancy and initiates shoot growth.



A.



Figure 2. The appearance of banana microshoot formation. (A) Axillary bud development in banana sword sucker explants. (B) Development of banana sword sucker with split open treatment in sub-culture media

The sword sucker explants used were grown intact and not split open, hence, the apical shoots might produce high endogenous auxin and inhibited lateral shoot growth. This inhibition of shoot growth by auxin was also reported by Kebrom (2017). Moreover, apical dominance breakage requires high exogenous cytokinin concentrations (Nuraini et al., 2016; Prasetyo et al., 2020). It is suspected that the exogenous cytokinin level added to the *in vitro* culture did not break the apical dominance as reported by Müller et al. (2015) and Ngomuo et al. (2014).

Plantlet formation

The shoot formation and root induction aimed to form plantlets that have both leaves and roots. The microshoots were separated individually from the original clump and then transferred to the treatment media. At the plantlet formation phase, shoot morphogenesis was influenced by banana cultivar (Table 1) and the interaction between the cultivar and auxin type used (Table 2). Gebyar cultivar produced more microshoots compared to Kepok Kuning and on average, it produced 4.25 axillary buds explant⁻¹. The difference in plants' genotypic responses is observed in their ability to grow and regenerate. Each genotype has different abilities regarding axillary shoot growth, shoot number and rooting emergence (Basri, 2016).

Table 1. The average number of shoots formed			
Types of	The number of		
cultivars	axillary buds		
Gebyar	4.25ª		
Kepok Kuning	1.47 ^b		
NY NY 1 0 11 1 1	.1 1 1		

Note: Numbers followed by the same letters show a significant difference in DMRT (≤0.05)

Gebyar cultivars planted on a media containing NAA produced the greatest shoots number (5.7 shoots explant⁻¹), although this treatment was not significantly different from that of the media containing IBA (Table 2). PGR treatment is used to stimulate cell proliferation and differentiation through organogenesis or embryogenesis (Méndez-Hernández et al., 2019). Not all cells in plant tissue respond to PGR, a cell only responds at a particular stage in the plant growth cycle (Tréhin et al., 1998; Schaller et al., 2014). Besides plant genotypes, explant's physiological conditions such as meristematic ability to grow and cells or tissues growth status determines bud regeneration's success. It is also related to cell metabolism, availability of endogenous PGR and the activity of genes controlling growth and development (Pillay and Tenkouano, 2011; Feng et al., 2012; Remakanthan et al., 2014).

Table 2. The average shoot number formed by
two banana cultivars with different
types of auxin

Type of cultivers	Type of auxin		
Type of cultivars	NAA	IBA	IAA
Gebyar	5.75 ^a	4.58^{a}	2.42 ^b
Kepok Kuning	1.18 ^b	1.58 ^b	1.67 ^b

Note: Numbers followed by the same letters show a significant difference in DMRT (≤0.05)



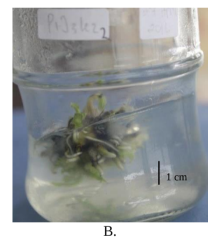


Figure 3. The effect of auxin types and concentration with different banana cultivars on the number of shoots formation. (A) Kepok Kuning cultivar with an added 2.5 μM IAA. (B) Gebyar cultivar with an added 5.0 μM NAA

Leaf formation in the banana plantlet formation phase was controlled by the cultivar used and the interaction between the growth regulator type and PGR concentration used. Kepok Kuning produced more leaves than Gebyar, by producing 4.64 leaves plantlet⁻¹ on average (Table 3). IAA utilization in culture media showed a significantly different effect on the number of leaves formed. Banana shoots grown on the media supplemented with 2.5 μ M IAA produced the greatest number of leaves, namely 5.5 leave plantlet⁻¹. Shoots number is inversely related to the number of leaves formed. The greater the number of shoots produced, the smaller the number of leaves per shoot formed (Figure 4).

 Table 3. The average number of leaves formed by two banana cultivars

Types of cultivars	Number of leaves	
Gebyar	1.75 ^b	
Kepok Kuning	4.64 ^a	
Note: Numbers followed	by the same letters show	

a significant difference in DMRT (≤0.05)

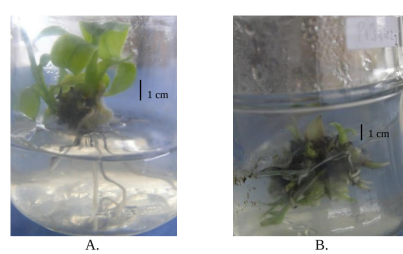


Figure 4. The effect of auxin types and concentration with different banana cultivars on leaf formation. (A) Kepok Kuning cultivar with added 2.5 μ M IAA. (B) Gebyar cultivar with added 5.0 μ M NAA

Auxin plays a very important role in leaf growth, by stimulating the development of prospective leaf meristem tissue (Xiong and Jiao, 2019). The leaves number is closely related to photosynthesis and plant metabolism, as well as nutrient absorption, while their increased growth is due to accelerated cell division and differentiation (Keller et al., 2011; Novak and Whitehouse, 2013). Also, their growth and development processes require PGR, such as auxin and cytokinin, as well as other nutrients contained in the growing media. Auxin and cytokinin work synergistically in stimulating plant growth, hence IAA which is a type of auxin, IAA functions in regulating cell division and stimulating growth to ensure an increase in leaves number (Scarpella et al., 2010; Skupa et al., 2014). Auxin addition affects leaf growth, especially the vascular tissue's length (Aloni, 2010; Zazımalova et al., 2010).

Leaves are essential plant organs, especially for photosynthesis, which facilitate organic materials production and optimum growth. The higher the number of leaves, length and width, the higher the leaf mass per unit area (LMA). LMA is an important leaf morphological trait affecting photosynthesis (Ren et al., 2019). The relationship between leaf number and individual size has metabolic and mechanical consequences that influence energy balances and carbon uptake at the whole plant level. In addition, it is also crucial to light interception and net carbon gain (Sun et al., 2019). Root induction was carried out at the final stage of plant *in vitro* culture. When axillary shoots emerged, they were subsequently subcultured on rooting media. Banana root formation was influenced by the interaction between the type of auxins used and their concentration. Although the addition of the three auxin types, namely IBA, NAA and IAA, did not show a significantly different effect on roots number. Generally, NAA demonstrated quite better results and produced the highest roots number compared to the other two auxins (Table 4). Banana shoots grown on the media containing NAA at a concentration of 7.5 μ M produced the highest roots number (13.3 roots plantlet⁻¹) (Figure 5).

Table 4. Interaction between auxin type and concentration on the plantlet formation

concentration on the plantiet formation					
Type of	Concentration	Number of	Number of		
auxin	of auxin (µM)	leaves	roots		
IAA	2.5	5.50 ^a	4.50^{bc}		
	5.0	2.17^{abc}	5.67 ^{bc}		
	7.5	2.17^{abc}	3.83 ^{bc}		
	10.0	3.83 ^{abc}	3.17 ^{bc}		
IBA	2.5	2.00^{abc}	4.67 ^{bc}		
	5.0	5.17 ^{ab}	7.00^{bc}		
	7.5	3.33 ^{abc}	6.17 ^{bc}		
	10.0	3.17 ^{abc}	4.17 ^{bc}		
NAA	2.5	1.50^{bc}	6.17 ^{bc}		
	5.0	1.17 ^c	2.33°		
	7.5	4.67 ^{abc}	13.33ª		
	10.0	3.83 ^{abc}	9.67 ^{ab}		

Note: Numbers followed by the same letters show a significant difference in DMRT (≤0.05)



Figure 5. The effect of auxin types and concentration with different banana cultivars on roots' formation after a 7.5 μ M NAA treatment

At low concentrations, IAA causes shoots and roots elongation, but at a higher concentration, it inhibits the elongation (Pamungkas, 2015; Lathyfah and Dewi, 2016). According to Govindaraju et al. (2012) and Zhao (2014), NAA in low concentrations produced a higher roots number. Also, root's formation is related to the endogenous auxin and cytokinin contents in plant tissue. Therefore, lower auxin concentration is suggested to be used in inducing rooting on a nicely growing shoot.

CONCLUSIONS

Gebyar cultivar produced more shoots than Kepok Kuning, with an average of 4.25 microshoots explant⁻¹. However, Kepok Kuning produced more leaves than Gebyar, with an average of 4.64 leaves plantlet⁻¹. The media containing IAA with a concentration of 2.5 µM produced the highest leaves number. Banana shoots that were cultured on a media containing NAA at a concentration of 7.5 µM produced the highest roots number. Besides, good acclimatization to produce ready-to-plant seedlings and a study to optimize acclimatization conditions to provide a higher seedling survival rate are needed. The mass production of disease-free plantlets is expected to meet the increasing demand for banana seeds to increase the national production scale.

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