ASYMBIOTIC SEED GERMINATION AND PLANTLET DEVELOPMENT
OF Dendrobium spectabile (Blume) Miq.

Perkecambahan asimbiotik biji dan perkembangan planlet Dendrobium spectabile (Blume) Miq.

Eka Martha Della Rahayu*, Melza Mulyani
Research Center for Plant Conservation and Botanic Gardens
Indonesian Institute of Sciences
Jl. Ir. H. Juanda No. 13, Bogor, West Java, 16003, Indonesia
*Email: emdrahayu@gmail.com

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Abstract

Dendrobium spectabile (Blume) Miq. is one of Papuan orchids that is of economic value and threatened. At present, the optimal protocol for asymbiotic seed germination and plantlet development of D. spectabile is not yet available. This research aimed to develop an optimal and comprehensive protocol for in vitro culture of D. spectabile to support the conservation and reintroduction of this species. The experiment was conducted using a completely random design, including seed sowing, protocorm subculture (subculture 1), plantlet subculture (subculture 2), and plantlet acclimatization. The highest germination rate at 3 MAS (months after sowing) was found in modified Knudson’s C (92.59%). The first subculture at 4 MAP (months after planting) showed that the highest leaf and root growths were found in modified Vacin and Went (4.12 and 2.13, respectively). The second subculture at 6 BST showed that the highest leaf growth was found in full strength of Murashige and Skoog supplemented with 100 g/l banana homogenate (5.49), while the highest number of roots and average root extension were found in half strength of Vacin and Went supplemented with 100 g/l banana homogenate (7.05 and 0.47 cm, respectively). The plantlets were best acclimatized in media consisting of tree fern fibre and sphagnum moss (2:1).

Keywords: asymbiotic germination, Dendrobium spectabile, in vitro, plantlet development
INTRODUCTION

Orchids is one of the most diverse plant families. According to Dressler (2005), there are 2500 species of orchids in the world, with the genus Dendrobium is the most popular as ornamental plants. Species of Dendrobium are highly prized ornamental assets, primarily as potted plants with showy flowers that tend to have a long vase life (Vendrame et al. 2007). Some Dendrobium species also has medicinal and pharmaceutical values, such as Dendrobium candidum Wall ex Lindl. (as Dendrobium moniliforme (L.) Sw.), Dendrobium nobile Lindl., Dendrobium officinale Kimura & Migo (as Dendrobium catenatum Lindl.) (da Silva & Ng 2017), Dendrobium crumenatum Swartz, Dendrobium pachyphyllum (Kuntze) Bakh. f., Dendrobium planibulbe Lindl., Dendrobium purpureum Roxb, Dendrobium salassense (Blume) Lindl., and Dendrobium subulatum (Blume) Lindl. (Teoh 2019).

One important species of the genus is Dendrobium spectabile (Blume) Miq. This epiphytic orchid can only be found in Indonesian Papua and Papua New Guinea, Bougainville and Solomon Islands (Cribb 1983, Lavarack et al. 2000). It grows in swampy lowland forest and lower montane forest up to about 1000 m above sea level (Lavarack et al. 2000). This species is noted for its spectacular twisted flowers. The flowers are large, yellow, usually with maroon mottled sepalss and petals, maroon lined lip, and with a white callus (Cribb 1983). The inflorescence consists of 20 flowers, each 4–8 cm across, distinctively twisted in all their parts, and blooms for several weeks (Lavarack et al. 2000).

The unique characteristics of D. spectabile’s flower make this orchid has high commercial value. Consequently, this species is threatened by over collecting in the wild, in addition to deforestation. According to Global Forest Watch (2020), 85% of tree cover loss in Papua occurred within natural forests from 2013 to 2018. This species has also been listed in the CITES Appendix II (CITES 2020). Therefore, this species requires conservation efforts to maintain its population in the wild.

Ex situ conservation for this orchid can be done by collecting a few plants from the wild and propagating it in botanic gardens. Dendrobium is conventionally propagated by splitting bulbs and keikis, or vegetative cuttings, but these are very slow and laborious methods that result only a few propagules in a year (Venturieri & Pickscius 2013). Tissue culture, on the other hand, provides an alternative solution for producing a large number of plantlets within a limited period of time.

Asymbiotic seed propagation through tissue culture has major importance for the conservation and propagation of wild species because it can maintain the genetic diversity of the species. Material derived from seed, is representing a broader sample of genetic diversity which is important for the reintroduction of the species to the wild (Akeroyd & Jackson 1995). Orchid propagation by seed results in high genetic variation of progeny (da Silva et al. 2015).

Various efforts using seeds explants of other Dendrobium species had been carried out by many researchers, such as Dendrobium tosaense Makino (as Dendrobium catenatum Lindl.) (Lo et al. 2004), Dendrobium fimbriatum Hook. (Sharma et al. 2005), Dendrobium nobile Lindl. (Vasudevan & van Staden 2010), Dendrobium aphyllum (Roxb.) C.E.C. Fischer (Hossain et al. 2013), Dendrobium nobile hybrids (Udomdee et al. 2014), Dendrobium ‘Iriana Jokowi’ (Rahayu 2016), and Dendrobium lasianthera J.J. Sm. (Utami et al. 2017).

In vitro asymbiotic germination of Dendrobium seeds were influenced by several factors, such as medium composition, culture conditions, and explant age (da Silva et al. 2015). Murashige and Skoog (MS), half strength Murashige and Skoog (1/2 MS), Knudson’s C (KC), and Vacin and Went (VW) were commonly used as basal culture medium for Dendrobium seed germination (Lo et al. 2004, Vasudevan & van Staden 2010, Hossain et al. 2013, Udomdee et al. 2014, da Silva et al. 2015, Utami et al. 2017). Diverse organic compounds (malt, yeast, casein, peptone) and natural supplements (banana homogenate, coconut water, potato homogenate, bean sprout extract, and tomato juice) added to the media could affect (improve or inhibit) Dendrobium germination (Hossain et al. 2013, da Silva et al. 2015, Rahayu 2016, Utami et al. 2017).

Periodicity of light is a crucial factor that influences seed germination. Periodicity of light frequently used for inducing orchid seed germination are 12/12, 14/10, 16/8, (day/night)
The maturity of seeds affects the germination rate of orchid seeds. Mature seeds were usually used in vitro germination of *Dendrobium*. About 3.3–4.7 months are required for *Dendrobium* seed to mature (da Silva et al. 2015). *Dendrobium tosaense* (as *D. catenatum*) took 3–4 months to become mature (Lo et al. 2004), *D. 'Iriana Jokowi'* took 4.5 months (Rahayu 2016), *D. lasianthera* took 3.5 months (Utami et al. 2017), while *D. spectabile* took 8.5 months to become mature (Santoso et al. 2014). Fruit maturity of *D. spectabile* can be identified from seed capsules with yellowish green coloration, supple fruit texture, and yellow seeds (Santoso et al. 2014).

According to Udomdee et al. (2014), seed size increased and seed color changed from white to light green to creamy yellow during development of capsules of *D. nobile* hybrids from 2–5 months.

Kebun Raya Bogor (KRB) has *D. spectabile* as both living plant collection and in vitro plantlets. According to KRB collection database, the species was collected from Papua in 2000, 2004, and 2008. However, only one specimen is currently available in living collection which never flowered. It is very unfortunate since this orchid was also reputed for its rarity in terms flowering and fruiting (Lavarack et al. 2000). The *in vitro* plantlets collections of this species at KRB tend to grow shoots only, rather than develop into normal healthy plantlets with robust roots system to support the acclimatization process.

Previous *in vitro* culture experiments for *D. spectabile* had been carried out by Nuraini et al. (2011), Santos et al. (2014), and Deswiniyanti (2015). Nuraini et al. (2011) used globular protocorms aged seven months after sowing (7 MAS) and cultured in Hyponex (1 g/l) combined with yeast extract (1 and 1.25 g/l). This method resulted in high number of leaves (6), roots (2.2), length of roots (0.6 cm) and plantlet height (1.6 cm) at four months after planting (4 MAP). They showed that the plantlets required subculture and could not be acclimatized since the plantlets were small without a robust root system.

Santoso et al. (2014) used MUS medium (which was a modified form of VW basic medium) for *in vitro* germination of 8.5 months after pollinating seeds of *D. spectabile*. MUS medium was composed of 200 mg of tricalcium phosphate (Ca₃(PO₄)₂), 525 mg of potassium nitrate (KNO₃), 250 mg of monopotassium phosphate (KH₂PO₄), 500 mg of ammonium sulphate ((NH₄)₂SO₄), 28 mg of ferric tartrate (Fe₃(C₆H₇O₄)₂), 250 mg of magnesium sulphate (MgSO₄.7H₂O), 7.5 mg of manganese sulphate (MnSO₄.4H₂O), 10 mg of imino-inositol, 0.5 mg of thiamin HCl, 0.5 mg of pyridoxine, 0.5 mg of niacin, 20 g of glucose, 10 g of fructose, 20 ml of coconut water, and 7 g of agar. Germination process of *D. spectabile* seeds took up to 5 MAS. Germinated seeds were then subcultured into MUS media for three times, each took three months. Santoso et al. (2014) did not provide data on the growth of *D. spectabile*, but showed that it took 14 months from seed germination to seedling development until it was ready to be acclimatized.

Deswiniyanti (2015) sowed five months after pollinating seeds of *D. spectabile* on VW medium. One month after sowing (1 MAS) germinated seeds were then subcultured to VW medium modification added with 0.5 g/l charcoal and 150 ml/l coconut water. Leaf number and seedling height of *D. spectabile* at 4 MAP were 3.45 and 2.65 cm, respectively. No acclimatization was conducted.

No optimal protocols in asymbiotic seed germination and plantlet development of *D. spectabile* is currently available. The objective of this research was to develop an optimal and comprehensive protocol of *in vitro* culture of *D. spectabile* to support its conservation and reintroduction programs.

### MATERIALS AND METHODS

#### Seed germination test of *Dendrobium spectabile*

Capsule of *D. spectabile* was donated by an orchid enthusiast, where the parent plant originated from Asmat District, Papua, Indonesia. Even though the capsule age was not known, it was considered as a mature capsule based on the characteristics proposed by Santoso et al. (2014) and Udomdee et al. (2014). The *D. spectabile* capsule was dipped in 95% ethyl alcohol and flamed, repeated three times. The sterilized capsule was then dissected longitudinally to extract its seeds. The seeds were sown on different media and kept in the culture room at 25 ± 2°C and 80% relative humidity under white fluorescent tubes.
with a 12/12-h (day/night) photoperiod. The seed germination experiment used a completely randomized design with one factor, namely four germination media commonly used at KRB Tissue Culture Laboratory: modified Hyponex (modified HS), modified Vacin and Went (modified VW), modified Knudson’s C (modified KC), and Knudson’s C (KCA) (Puspitaningtyas & Handini 2014, Rahayu 2016).

Modified HS consisted of Hyponex fertilizer 0.5 g/l, potato homogenate 40 g/l, peptone 2 g/l, and active charcoal 1 g/l. Modified VW consisted of macronutrients and micronutrients Vacin and Went with addition of active charcoal 1 g/l, bean sprouts extract 100 g/l, tomato homogenate 100 g/l, coconut water 150 ml/l, and NAA 10 mg/l. Modified KC consisted of macronutrients and micronutrients Knudson’s C with addition of coconut water 150 ml/l, bean sprouts extract 150 g/l, and active charcoal 1 g/l. KCA only contained macronutrients and micronutrients from Knudson’s C. Each treatment (medium) consisted of four replications, with each replication consisted of 100–200 sown seeds (Puspitaningtyas & Handini 2014).

Seed germination and protocorm developmental stages were examined using a stereomicroscope with an Olympus digital camera attached. Seed morphology was observed before and after germination. Seeds germination and protocorm development stages followed Vasudevan & van Staden (2010). Germinated seeds, protocorms with rhizoids, and protocorms with pointed shoot apex (stage 4) of *D. spectabile* were counted based on germinated seeds out of the 100–200 sown seed. The data were then analyzed with ANOVA and Duncan Test.

Subculture 1: Induction of leaves and roots from *Dendrobium spectabile* protocorms

The 3 MAS protocorms with rhizoids and pointed shoot apex (stage 4) of *D. spectabile* were subcultured to induce leaf and root growth. Four media were used for this purpose, namely modified Growmore (T1A) consisted of Growmore fertilizer 0.5 g/l, peptone 2g/l, banana homogenate 20 g/l, and active charcoal 1 g/l; VW supplemented with 20 g/l banana homogenate and active charcoal 1g/l (T1V); modified KC; and modified VW. The pH of the medium was adjusted to 5.6. Experiments for leaf and root induction used a completely randomized design with one factor, namely the four media: T1A, T1V, modified KC, and modified VW. Each treatment consisted of three replications, and each replication consisted of five culture vessels; each culture vessel consisted of 10 protocorms. The number of leaves, roots, shoots, and length of roots were counted at four months after planting (4 MAP) (Utami et al. 2017). Data were analyzed with ANOVA. If the results were significantly different, then a Duncan Test was done in order to see the best media.

Subculture 2: Induction of further growth of leaves and roots from *Dendrobium spectabile* plantlets

Four months after planting (4 MAP) plantlets of *D. spectabile* resulted from subculture 1 were subcultured in four different media to induce further growth of leaves and roots in order to support the acclimatization process. The media used for this purpose were half strength of Murashige and Skoog supplemented with 100 g/l banana homogenate (1/2P), half strength of Murashige and Skoog supplemented with 100 g/l banana homogenate and 10 mg/l NAA (1/2P10), full strength of Murashige and Skoog supplemented with 100 g/l banana homogenate (MP), and half strength of Vacin and Went supplemented with 100 g/l banana homogenate (1/2VT).

Experiments for further growth of leaves and roots used a completely randomized design with one factor, namely the four media: 1/2P, 1/2P10, MP, and 1/2VT. Each medium consisted of three replications, and each replication consisted of five culture vessels, each culture vessel contained five plantlets. The number of leaves, roots, shoots, and the length of root at the beginning of culture (0 MAP) were measured. The increase in the number of leaves, roots, shoots, and the length of roots were measured at 6 MAP in order to optimize the growth of the plantlets. Data were then analyzed with ANOVA. If the results were significantly different, then a Duncan Test was done in order to see the best media.
Acclimatization of plantlets

The plantlets from each treatment with well-developed leaves, roots, and shoots were washed thoroughly under tap water for 2–3 minutes to remove traces of agar-gelled medium. These plantlets were then planted in plastic pots (8 cm diameter) containing four different potting media: charcoal, a mixture of charcoal and tree fern fibre (2:1), tree fern block, and a mixture of tree fern fibre and sphagnum moss (2:1). The acclimatized seedlings were kept in the greenhouse. They were kept inside a plastic UV to maintain high humidity (above 90%) and watered every day.

Each treatment consisted of three replications, and each replication consisted of 10 plastic pots; one seedling in each pot. The growth of new leaves, roots, and shoots were counted after six weeks of planting (6 WAP). Data were then analyzed with ANOVA and Duncan Test.

RESULTS AND DISCUSSION

Seed germination test of Dendrobium spectabile

Embryo of D. spectabile is located in the center of the seed. Each seed of D. spectabile consisted of a testa and an embryo without endosperms (Figure 1A). Observation at 3 MAS showed that germination rate ranged between 88.03–92.59 %. The germination rates of D. spectabile in the modified KC, HS, KCA and modified VW were 92.58%, 89.24%, 88.32%, and 88.03%, respectively (Table 1). This showed that the four tested media provided the macronutrients and micronutrients needed for the seeds to germinate.

The high germination rate was also probably due to the capsule used in this study contained mature seeds with a high percentage of viable seeds that could germinate under in vitro asymbiotic conditions (da Silva et al., 2015). A three months old capsule of D. tosaense as D. catenatum) showed germination rates of 90% and 92.59% respectively (Figure 1A). Observation at 3 MAS showed that the seeds germinated well on all four tested media (Table 1). This showed that the four tested media (modified KC, modified VW, HS) is useful for in vitro asymbiotic germination of D. spectabile. These works showed that the best media for orchid seed germination were specific for each species.

Good germination and development trend in modified KC and modified VW was probably due to the addition of various organic material in the medium, such as coconut water, bean sprout extract, and tomato homogenate. According to Thorpe et al. (2008), coconut water contained amino acids, organic acids, nucleic acids, purines, sugars, sugar alcohols, vitamins, growth substances,
Asymbiotic seed germination and plantlet development of Dendrobium spectabile (Blume) Miq.

Figure 1. The development stages of seeds and protocorms of Dendrobium spectabile: A) Seeds of D. spectabile showing testa (t) and embryo (e); A–D showing germination stages: 1) imbibed seed, swollen, still covered by testa, 2) enlarge embryo, seed coat ruptured (germination), 3) protocorm with rhizoids, 4) protocorms with pointed shoot apex and rhizoids (appearance of shoot apex), 5) emergence of first leaf. Magnification of 40x. Scale bar = 0.5 mm

Table 1. Effects of media on seed germination and protocorm development of Dendrobium spectabile at 3 MAS (months after sowing)

<table>
<thead>
<tr>
<th>Media</th>
<th>Germination rate (%)</th>
<th>Mean Germination</th>
<th>Mean Protocorms</th>
<th>Mean Protocorms</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Un-germinated</td>
<td>Germinated</td>
<td>with rhizoids</td>
</tr>
<tr>
<td></td>
<td></td>
<td>seeds</td>
<td>seeds</td>
<td></td>
</tr>
<tr>
<td>HS</td>
<td>89.24</td>
<td>20.25</td>
<td>168.00</td>
<td>35.25</td>
</tr>
<tr>
<td>Modified VW</td>
<td>88.03</td>
<td>24.50</td>
<td>180.25</td>
<td>73.00</td>
</tr>
<tr>
<td>Modified KC</td>
<td>92.59</td>
<td>15.50</td>
<td>193.75</td>
<td>106.75</td>
</tr>
<tr>
<td>KCA</td>
<td>88.32</td>
<td>23.25</td>
<td>175.75</td>
<td>27.50</td>
</tr>
</tbody>
</table>

Values followed by different letters within a column are significantly different at P < 0.05 according to DMRT

Figure 2. Effects of media on seed germination and protocorm development of Dendrobium spectabile at 1, 4, 8, and 12 WAS (weeks after sowing): A) germinated seeds, B) protocorms with rhizoids, C) protocorms with pointed shoot apex and rhizoids. HS = Hyponex, mVW = modified Vacin and Went, mKC = modified Knudson’s C, KCA = Knudson’s
minerals, and plant hormones. Winarto & da Silva (2015) stated that coconut water was used to induce cell division, promote morphogenesis and accelerate the multiplication of protocorm like bodies (PLBs) in orchid's in vitro culture. Application of 150 ml/l coconut water in 1/2 MS medium increased the growth and proliferation capacity of Dendrobium ‘Gradita 31’. According to Amilah & Astuti (2006), bean sprout extract contained several essential vitamins and minerals, namely vitamin C, Thiamin, Riboflavin, Niasin, Vitamin B, β-karotein, Vitamin A, Vitamin E; and minerals calcium (Ca), ferrum (Fe), magnesium (Mg), phosphor (P), sodium (Na), zinc (Zn), copper (Cu), and mangan (Mn). Tomatoes homogenate contained sugar and antioxidants including vitamin C that stimulated seed germination and the growth of protocorm (Semiarti et al. 2010). Tomato extract in VW medium promoted protocorm's growth of Dendrobium bigibbum Lindl. (as Dendrobium phalaenopsis Fitzg.) (Setiari et al. 2016). Lycopene in tomato extract acted as an antioxidant that was required as a trigger for embryo development initiation. Tomato homogenate also contained K elements that facilitated the formation of water sac within the cell walls, and made it easier to absorb water (George & Sherington 1984), thus it made embryo more susceptible to swelling. When embryo swelled, testa opened and released the embryo within, thereby accelerating germination (Setiari et al. 2016).

The development stages of D. spectabile’s seeds and protocorms in all four tested germination media (Figure 1) were generally in line with the results by Vasudevan & van Staden (2010), Hossain et al. (2013), Udomdee et al. (2014), and Rahayu (2016) of different species of Dendrobium. All of these works showed that the developmental stages started with imbibed seed, swollen, still covered by testa (stage 1). Stage 2 was indicated by enlarge embryo and ruptured seed coat (germination). At stage 3, protocorms developed rhizoids. Subsequently, at stage 4, protocorms were armed with rhizoids and pointed shoot apex (appearance of shoot apex). Then at stage 5, first leaf was emerged and further development of the plantlet was shown. Stage 1 and stage 2 on germination of D. spectabile were started at 1 WAS, stage 3 at 4 WAS, stage 4 at 9 WAS, and stage 5 after 12 WAS (Figure 2).

Subculture 1: Induction of leaves and roots from Dendrobium spectabile protocorms

Protocorms of D. spectabile that were subcultured in induction media developed plantlets at 4 MAP. The highest mean number of leaves (4.12) and roots (2.13) and length of roots (1.21 cm) were observed on plantlets planted in VW, whereas the highest mean number of shoots (0.09) was observed in T1A (Table 2). It was probably due to the addition of organic material to the medium, such as bean sprouts extract, tomato homogenate, and coconut water. Addition of organic supplements in plant tissue culture had been investigated by a number of workers. The addition of bean sprouts extract in culture media applied to ‘unti sayang’ banana (Musa x paradisiaca L.) showed that the bean sprouts extract provided significant influences on plantlet height, leaf number, root length, and root number (Jufri et al. 2014). The fastest rate of embryo development and the formation of shoot apical meristem prior to the emergence of leaf primordia of Phalaenopsis amabilis Blume was achieved by New Phalaenopsis medium supplemented with both coconut water and tomato homogenate (Semiarti et al. 2010). The addition of tomato homogenate in culture media was also effective for the germination of Geodorum densiflorum (Lam.) Schltr. (Mutukhrisnan et al. 2013). The tomato homogenate contained carotene, vitamin C, K, lycopene and other antioxidants which were not detected in coconut water that affect the growth and development of cells (Semiarti et al. 2010, Mutukhrisnan et al. 2013).

Table 2. Growth of plantlets of Dendrobium spectabile on the roots and leaves induction media at 4 MAP (months after planting)

<table>
<thead>
<tr>
<th>Media</th>
<th>Leaves</th>
<th>Roots</th>
<th>Length of roots (cm)</th>
<th>Shoots</th>
</tr>
</thead>
<tbody>
<tr>
<td>T1A</td>
<td>4.07a</td>
<td>1.56b</td>
<td>1.08ab</td>
<td>0.09b</td>
</tr>
<tr>
<td>T1V</td>
<td>3.98a</td>
<td>1.16c</td>
<td>1.21a</td>
<td>0.04ab</td>
</tr>
<tr>
<td>Modified KC</td>
<td>3.64b</td>
<td>1.28c</td>
<td>0.94b</td>
<td>0.03b</td>
</tr>
<tr>
<td>VW</td>
<td>4.12a</td>
<td>2.13a</td>
<td>1.21a</td>
<td>0.02b</td>
</tr>
</tbody>
</table>

Values followed by different letters within a column are significantly different at P < 0.05 according to DMRT.
Fully-ripe tomato fruit contained basic nutrients and essential vitamins, as well as trace elements (Semiarti et al. 2010). Coconut water promoted the growth of leaves (3.5) and roots (5.0) of *D. tosaense*’s (as *D. catenatum*) seedlings (Lo et al. 2004). According Utami et al. (2017), VW added with coconut water was able to produce plantlets of *D. lasianthera* with the highest number of roots and leaves.

**Subculture 2: Induction of further growth of leaves and roots of *Dendrobium spectabile* plantlets**

The subculture 2 at 6 MAP resulted in plantlets that were ready for acclimatization (Figure 3). The highest increase in leaf number was observed in MP (5.49), whereas that of new roots and root length was found in 1/2VT (7.05 roots and 0.47 cm, respectively). There were no significant differences in the number of new shoots grown in all four media. The number of new shoots ranged from 2.83 to 3.08 (Table 3).

The highest number of leaves of *D. spectabile* plantlets on subculture 2 was found in MP (5.49). However, the increase of roots in MP was much lower than in 1/2VT (3.28 and 7.05, respectively). MP medium consisted of banana homogenate. Udomdee et al. (2014) showed that the seeds of *D. nobile* hybrids germinated in medium without additional sucrose but with an addition of banana homogenate. Banana homogenate contributed to the basal level of carbon source. According to Sharrock & Lusty (2000), ripe banana contained high sugar content which was 23.43/100 g raw edible portion. During ripening process, the sugar content in banana was in the approximate ratio of glucose : fructose : sucrose = 20 : 15 : 65.

**Figure 3.** Plantlets of *Dendrobium spectabile* at 6 MAP (months after planting) ready for acclimatization, subcultured in: A) 1/2P, B) 1/2P10, C) MP, and D) 1/2VT media

**Table 3.** Growth of plantlets of *D. spectabile* on media for further induction of roots and leaves at 6 MAP (months after planting)

<table>
<thead>
<tr>
<th>Media</th>
<th>Average number of</th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>New leaves</td>
<td>New roots</td>
<td>Length of roots expansion</td>
<td>New shoot</td>
</tr>
<tr>
<td>1/2P</td>
<td>4.73b</td>
<td>5.56b</td>
<td>2.57b</td>
<td>2.95</td>
</tr>
<tr>
<td>1/2P10</td>
<td>4.46b</td>
<td>6.09b</td>
<td>2.53b</td>
<td>3.08</td>
</tr>
<tr>
<td>MP</td>
<td>5.49a</td>
<td>3.28c</td>
<td>1.14c</td>
<td>2.83</td>
</tr>
<tr>
<td>1/2VT</td>
<td>3.18c</td>
<td>7.05a</td>
<td>3.23a</td>
<td>3.05</td>
</tr>
</tbody>
</table>

Values followed by different letters within a column are significantly different at \( P < 0.05 \) according to DMRT.
Acclimatization of plantlets

The survival rate of *D. spectabile* seedlings at 6 WAP in all acclimatization media were 46.67–100% (Table 4). The highest survival rate (100%) was achieved by seedlings planted in a mixture of tree fern fibre and sphagnum moss (2:1) (Table 4, Figure 4), followed by those in a mixture of charcoal and tree fern fibre (2:1) (90%) (Table 4). The lowest survival rate (46.67%) was found in seedlings planted in charcoal. The highest number of new leaves, roots, and shoots were found in the seedlings grown in a mixture of tree fern fibre and sphagnum moss. The average of new leaves ranged from 0.37–2.5 (Table 4).

Similarly, Lo *et al.* (2004) found that the combination of tree fern and sphagnum moss was a more suitable substrate for *D. tosaense*’s (as *D. catenatum*) seedlings survival than tree fern alone, with the survival rate was 87% in the former and 43% in the latter. Luo *et al.* (2008) also found that the same mixture was the most suitable substrate for plant survival of *D. densiflorum*. Formation of a functional root system in tissue cultured plantlets appeared to be an essential step for their survival after transplantation. In vitro roots remained functional and continued to grow during ex vitro acclimatization (da Silva *et al.* 2017). Therefore, a strong and robust root system should have been induced in in vitro plantlets before they were transferred outside.

Figure 4. Seedlings of *D. spectabile* acclimatized in a mixture of tree fern fibre and sphagnum moss (2:1) at 6 WAP (weeks after planting)

<table>
<thead>
<tr>
<th>Media</th>
<th>Survival rate (%)</th>
<th>New leaves</th>
<th>New roots</th>
<th>New shoots</th>
</tr>
</thead>
<tbody>
<tr>
<td>Charcoal</td>
<td>46.67&lt;sup&gt;c&lt;/sup&gt;</td>
<td>0.37&lt;sup&gt;b&lt;/sup&gt;</td>
<td>0.00&lt;sup&gt;b&lt;/sup&gt;</td>
<td>0.07&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>Charcoal and tree fern fibre (2:1)</td>
<td>90.00&lt;sup&gt;a&lt;/sup&gt;</td>
<td>1.03&lt;sup&gt;b&lt;/sup&gt;</td>
<td>0.30&lt;sup&gt;ab&lt;/sup&gt;</td>
<td>0.43&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>Tree fern block</td>
<td>70.00&lt;sup&gt;b&lt;/sup&gt;</td>
<td>0.73&lt;sup&gt;b&lt;/sup&gt;</td>
<td>0.27&lt;sup&gt;ab&lt;/sup&gt;</td>
<td>0.47&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>Tree fern fibre and sphagnum moss (2:1)</td>
<td>100.00&lt;sup&gt;a&lt;/sup&gt;</td>
<td>2.50&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0.47&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0.93&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
</tbody>
</table>

Values followed by different letters within a column are significantly different at *P < 0.05* according to DMRT

This research produced plantlets with strong and robust root system, because both potting mixture used in this research, i.e. tree fern fibre:sphagnum moss (2:1) and charcoal:tree fern fibre (2:1), had strong permeability and retained humidity. This was in line with da Silva *et al.* (2017) who reviewed different kinds of media to acclimatize in vitro-raised *Dendrobium* plantlets, namely bricks, charcoal pieces, pine bark, Cycas bark, cocopeat, coconut dirt, sawdust, perlite, vermiculite, peat, and sphagnum moss. They concluded that sphagnum moss performed low pH and absorbed large quantities of water and mineral nutrients, and was best used for the acclimatization of *Dendrobium* plantlets.

**CONCLUSIONS**

An efficient and simple protocol for rapid production of *D. spectabile* using seeds has been established as follows: the high germination rate at 3 MAS was achieved when the seeds were cultured in modified Knudson’s C. The protocorms were then grown on modified VW media for four months. Plantlets were further grown on full strength of Murashige and Skoog supplemented with 100 g/l banana homogenate (MP) or half strength of Vacin
and Went supplemented with 100 g/l banana homogenate (1/2VT) for six months. A total of 13 months was required to produce plantlets that were ready for acclimatization. Seedlings of D. spectabile were then acclimatized on potting media consisting of mixture of tree fern fibre and sphagnum moss (2:1). This protocol can be applied to support ex situ conservation and reintroduction programs of D. spectabile to their natural habitat, or for a large-scale cultivation for the interest of ornamental products as it would ensure high production of plantlets.

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REFERENCES


