Non-symbiotic Seed Germination and In vitro Plant Development of Pholidota articulata

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Abstract

Pholidota articulata is an epiphytic orchid mostly used in ornamental cut/pot flower and in traditional medicine. As it has high ornamental and medicinal values, its population from natural habitats is decreasing, therefore, it is listed in the Appendix-II of Convention on International Trade in Endangered Species (CITES). The objective of the present study is to obtain the in vitro plants of P. articulata from seed culture to conserve its germplasm. The in vitro seed germination was carried out in different strengths of Murashige and Skoog (MS) and Knudson C (KnC) medium supplemented with various plant hormones. On the half-strength of MS medium, seeds were started to germinate after 4 weeks of primary culture and they were developed into protocorms with first leaf primordium earlier than on the other medium. Therefore, in vitro developed protocorms were sub-cultured on the half-strength of MS medium supplemented with different concentrations of 6-benzylaminopurine (BAP), gibberellic acid (GA₃) and α-naphtalene acetic acid (NAA). They were successfully developed into shoots on the 1.5 mg/l BAP supplemented half-strength of MS medium. Later, they were inoculated on the half-strength of MS medium supplemented with different concentration of α-napthalene acetic acid (NAA), indole-3-acetic acid (IAA) and indole-3-butyric acid (IBA) for the root formation, where IBA supplemented medium was found effective for the development of roots. Thus, this study provides a reliable protocol for non-symbiotic seed germination and plant production, and reveals that seed-derived protocorms are good explants for the in vitro mass propagation for conservation and sustainable utilization in horticulture.

Keywords: Nutrient medium; Pholidota articulata, Propagation, Protocorms, Seed germination

Introduction:

The genus Pholidota, represented by 46 species in the world and by 5 species in Nepal, are distributed from tropics and subtropics to South-West Pacific (Chase et al., 2015; Rajbhandari, 2014). Pholidota articulata Lindl. is a fragrant, medium-sized, epiphytic or lithophyte found in Nepal, India, Tibet, China, Burma, Thailand, Laos, Cambodia, Vietnam, Borneo, Java, Sulawesi, and Sumatra (Rajbhandari, 2014). P. articulata, commonly known as Articulated Pholidota, is grown in semi-deciduous and evergreen montane forests and highland primary cloud forests at an altitude of 300 to 2000m (Pant et al., 2016). It has the ability of the arising of new pseudobulb from the apex of last year's mature pseudobulb and has ovate to linear-lanceolate leaves with prominent veins. It blooms on a slender, drooping, 6" to 11" long, fructifies inflorescence with broad, brownish floral bracts that drop as the many scented flowers with smelling of musk open in the spring and summer (Pant et al., 2016). The whole plant is used in traditional medicine such as bone fractures, age sustaining tonic and restorative, and in the preparation of 'Chyawanprash'. Its root powder is also used to treat cancer, ulcer and skin eruptions (Jalal et al., 2009; Sharma et al., 2017). This orchid is also
equally important for aesthetic value as an ornamental purpose in the form of pot flower as well as a cut flower. As one of the important ingredients of Chyawanprash and having ornamental value as a cut/pot flower, plants of this species are declining from the nature due to over-collection to meet the demand (Subedi et al., 2013). Sexual reproduction of orchids in nature is being very slow as only 2-5% of their seeds can germinate after symbiosis with a special mycorrhizal fungus (Pant et al., 2017). So, plant tissue culture technique is one of the alternative ways to propagate the plants through seeds without fungal association those cannot be reproduced sexually in nature easily (Pant, 2013). In vitro propagation protocol by plant tissue culture has not been developed yet in this orchid. Thus, in this research, we have reported its in vitro non-symbiotic seeds germination and the plant production from seed-derived protocorms. In vitro propagation of this orchid could be an alternative to fulfill its demand on horticulture and traditional medicine as well as for its conservation.

Materials and Methods

Plant material and its sterilization

Healthy green capsules (16 weeks old) were collected from mother plant *Pholidota articulata* (Figure 1 A and B) grown in the botanical garden of Central Department of Botany, Tribhuvan University, Kathmandu (at an altitude of 1300 m asl). Seeds of a capsule were used as initial explant. The capsule was cleaned thoroughly under tap water followed by washing with few drops of Tween-20 (liquid detergent) for 30 minutes and finally rinsed with distilled water. It was surface sterilized with 70% ethanol for 5 minutes followed by an aqueous solution of 1% sodium hypochlorite for 10 minutes and finally rinsed with sterile distilled water for five times.

Figure 1: *Pholidota articulata* (A), and its capsules (B).

Nutrient medium for culture

Murashige and Skoog (MS) and Knudson C (KnC) medium were used as the basal nutrient medium. Full, half and quarter strength of them were used individually. Different concentration of 6-benzylaminopurine (BAP), gibberellic acid (GA), \( \alpha \)-naphtalene acetic acid (NAA), indole-3-acetic acid (IAA) and indole-3-butyric acid (IBA) have been added in both the nutrient medium for seeds germination, shoots elongation and multiplication, and roots formation. In the entire medium, 3% (w/v) sucrose was added as carbon source and 0.8% (w/v) agar was added as a gelling agent. The pH of all the medium was adjusted to 5.8 with 0.1N NaOH or HCl before autoclaving. All the medium were autoclaved at 121 °C and 15 psi for 20 minutes.

Inoculation and germination of seeds

The surface-sterilized capsule was cut longitudinally apart on sterilized petridish and seeds were spread thinly over the surface of solidified nutrient medium. Full-, half- and quarter-strength of MS and KnC medium, and full-strength of MS and KnC medium supplemented with 0.5 mg/l BAP and 0.5 mg/l NAA was used for non-symbiotic seeds germination.

Elongation and multiplication of shoots

Protocorms with first initiated leaf developed from the germinating seeds were used for shoots multiplication. Half-strength of MS medium with different concentration of \( \alpha \) (0.5, 1.0, 1.5 and 2.0 mg/l) with or without 0.5 mg/L NAA, and different concentration of BAP (0.5, 1.0, 1.5 and 2.0 mg/l) were used for the elongation and multiplication of shoots from the seed derived-protocorms.

Roots formation on shoots

In vitro developed shoots were transferred on the half-strength of MS medium supplemented with different concentrations (0.5 and 1.0 mg/l) each of NAA, IAA and IBA for the development of roots on the shoots.

Culture condition

All the cultures were kept in the culture room at 25±2 °C. The culture room has facilitated with 16 hours of light period by fluorescent white light.
**Statistical data analysis**

All the data were presented as a mean ± standard error of the mean in each treatment. Univariate analysis was done using SPSS.

**Results**

**Germination of seeds on different nutrient medium**

Seeds were inoculated on the full-, half-, and quarter-strength of MS and KnC medium, and full-strength of MS and KnC medium supplemented with 0.5 mg/l BAP and 0.5 mg/l NAA. Their stages of development into protocorms and shoots have shown in Figure 2. Sign of seeds germination is by turning them into light yellowish-green on 4th week, they became swollen and elongated spherules in 6th week on all the strengths of MS medium and in 5th week on all strengths of KnC medium. Protocorms formation was observed in 10th week of seeds culture on full- and half-strength of MS medium (Figure 3 A and B). They were proliferated into shoots on half-strength of MS medium in 22nd week. Similarly, protocorms were observed in 8th week of seeds culture on full- and half-strength of KnC medium. They were globular, light yellowish-green and hairy. They were developed into shoots on half- and quarter-strength of KnC medium in 19th week of primary culture (Figure 3 C and D). The green, globular hairy protocorms changed into pale yellow over time on these medium. No shoot proliferation or further growth of protocorms was observed in all the concentrations and combinations of the KnC medium.

![Figure 2: Germination of seeds on the different strength of MS and KnC medium with plant hormones.](image1)

![Figure 3: Development of protocorms from seeds on the full- and half-strength of MS medium (A and B) and on the half- and quarter-strength of KnC-medium (C and D).](image2)
Multiplication of shoots on the different nutrient medium

Protocorms were transferred on the half-strength of MS medium supplemented with GA$_3$ (0.5, 1.0, 1.5 and 2.0 mg/l) with or without 0.5 mg/L NAA, and BAP (0.5, 1.0, 1.5 and 2.0 mg/l) for their elongation and multiplication. The variation in the development of shoots from the inoculated explants has shown in Figure 4. Among the medium, 0.5 mg/l GA$_3$ supplemented half-strength of MS-medium was favoured for the development of more than 5 shoots per culture after 5 weeks (Figure 5A). Similarly, more than 4 shoots per culture were developed on the GA$_3$ (0.5 mg/l) plus NAA (0.5 mg/l) supplemented half-strength of MS medium after 4 weeks (Figure 5B). However, BAP (0.5 and 1.5 mg/l) supplemented half-strength of MS medium was favoured for development of more than 6 shoots in the 5th week (Figure 5C).

![Graph showing the development of shoots from protocorms on half-strength MS medium with different hormones.](image)

**Figure 4:** Development of shoots from the protocorms on half-strength of MS medium with different hormones.

![Images of shoots developed on MS medium with different hormones.](images)

**Figure 5:** Development of shoots from protocorms on the half-strength MS medium with 0.5 mg/l GA$_3$ (A), with 0.5 mg/l GA$_3$ plus 0.5 mg/l NAA (B), and with 1.5 mg/l BAP (C).
Root formation on the different nutrient medium

For the root formation, in vitro developed shoots were transferred on the NAA, IAA and IBA supplemented half-strength of MS medium. The higher number of roots were developed on the shoots in the IBA (0.5-1.0 mg/l) and 0.5 mg/lNAA supplemented half-strength of MS medium (Figure 6). The high concentration of NAA (1.0 mg/l) was found to be ineffective for the formation of roots.

![Figure 6: Formation of roots on the shoots on NAA, IAA and IBA supplemented half-strength of MS medium.](image)

Discussion

Both MS and KnC medium showed varied response on non-symbiotic seed germination of P. articulata. The orchid seeds responded successively in half strength of MS and KnC medium but further development of shoots from germinating seeds was suppressed in different strengths of KnC medium. This is because of MS medium is highly enriched with macro and microelements with different vitamins whereas KnC medium contained a comparatively low amount of macro and microelements without vitamins (Pradhan & Pant, 2009; Vejsadová, 2006). It was found that even though the time taken for germination and initiation of shoot formation was faster in KnC medium, it did not favour further development of shoots, as the shoots became achlorophyllous. The development of orchids requires a balanced supply of both organic and inorganic nutrients (Johnson et al., 2007). Though KnC medium was found to be best for the culture of many orchids, it did not favour Pholidota articulata. MS medium showed good response compared to KnC medium in seed germination and shoots formation which was found similar to the previous study in Esmeralda clarkei (Paudel et al., 2012). This effect may be due to the presence of a high concentration of vitamins, amino acids and other regulatory compounds present in MS medium compared to KnC medium (Paudel & Pant, 2013; Pradhan et al., 2013). Similar types of result have been reported to the previous findings of seed germination of orchids; Habenaria macroceratitis (Stewart & Kane, 2006), Coelogyne mossiae (Sebastianraj et al., 2006), Coelogyne stricta (Parmar & Pant, 2016), Cymbidium aloifolium (Pradhan et al., 2014; Pradhan et al., 2013).

GA3 treatment was found to be much effective in shoot multiplication (Giridhar & Ravishankar, 2004; Maharjan et al., 2020). Half-strength MS medium supplemented with 0.2 mg/l BAP proved better for multiplication of protocorm and healthy shoot proliferation of Dendrobium (Nuraini et al., 2010), Phaius tancarvilleae (Pant & Shrestha, 2011), multiple shoot production of Vanilla planifolia (Palama et al., 2010), multiple shoot productions in Dendrobium aqeuum (Parthibhan et al., 2015), Esmeralda clarkei (Paudel & Pant, 2012a,
2012b), inducing multiple shoots in MS with additives of BA (Pant et al., 2019), as well as multiple shoot production in the combination of BAP and NAA (da Silva et al., 2015; da Silva & Acharya, 2014). Rooting of a shoot in 0.5 mg/l NAA containing medium was the most appropriate as is supported by the previous works (Panwar et al., 2012; Pradhan et al., 2013; Zeng et al., 2012). IAA supplemented medium was found best for the initiation of the root in Dendrobium and Vanilla (Aktar et al., 2007; Geetha & Shetty, 2000), similar to the present result.

Conclusion

Half-strength of MS medium was found to be the most appropriate for the non-symbiotic seeds germination and protocorms formation of Pholidota articulata. Half-strength of MS medium supplemented with 1.5 mg/l BAP was found to be the most effective condition for the multiplication of shoots from protocorms. Root development was found best on half-strength of MS medium supplemented with 1.0 mg/l IBA. Hence, this protocol might be useful for non-symbiotic germination, mass propagation and conservation of Pholidota articulata.

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Declaration of conflict of interest and ethical approval:

Authors declared that there is no any type of competing interest regarding the current manuscript. R. D. Prasad did the experiment and wrote the draft, M. R. Paudel design the experiment, wrote and finalized the manuscript, S. Pradhan and B. Pant conceptualized, designed and supervised the experiment. All authors have read, reviewed and approved the manuscript before submitting to the journal.

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