Abstract  Ex-situ conservation of the ornamental and medicinal orchid, *Coelogyne stricta*, was performed by mass propagation using seed culture. Propagation stages were optimized using full- and half-strength solidified MS medium with different phytohormones. Maximum seed germination (88 ± 0.5% over 6 weeks of culture) was achieved on half-strength MS medium supplemented with 15% coconut water. Maximum shoot numbers were found on full-strength MS medium supplemented with 1 mg/L BAP, 2 mg/L Kinetin, and 10% coconut water, while the longest root was developed on full-strength MS medium with 1.5 mg/L IBA. A 2:1:1 combination of coco-peat, pine bark, and sphagnum moss was found to be a suitable potting mixture resulting in 80% seedling survivability. The cytotoxic activity of extracts of both wild plants and in vitro-developed protocorms was determined using an MTT assay on a cervical cancer cell line. The wild plant extract inhibited the growth of 41.99% of cells, showing that this extract has moderate cytotoxic activity toward cervical cancer cells.

Keywords  *Coelogyne stricta*, Conservation, HeLa cell line, Micropropagation, MTT assay

Introduction

Orchidaceae is being the second-largest family in angiosperms with roughly around 28,484 species (Govaerts et al. 2017) that set up nearly 10% of the total flowering species (Roberts and Dixon 2008; Tsai et al. 2013). Orchidaceae is considered to be highly advanced as proven by their unique floral architecture, condensed pack of pollinia, habitat ecology, and their particular pollination mechanism (Jersáková et al. 2006; Park et al. 2018).

Orchids are famous among cut flowers and potted plants with most beautiful flowers leaving outstanding royalty in the horticulture industry (Murthy et al. 2018). Orchids are of considerable economic increasing indicator particularly in horticulture and floristry for their striking attractiveness and long-lasting blooming age but also they are equally important in the pharmaceutical and perfume industries (Pant 2013). Owing to high demand in the national and international market places, over-collection from its natural habitat and gentle growth rate in nature, their species are delimited only to narrow pocket areas in nature (Pant et al. 2016). Complex life cycle, seed without endosperm, specificity with fungus (Chand et al. 2020; Pant 2013; Shah et al. 2019), habitat-specific and specialized pollinators (Cazzolino and Widmer 2005), illegal and uncontrolled collection and trade, deforestation and defragmentation of habitat are measures for their rarity in nature (McCormick et al. 2004; Shefferson et al. 2007). The IUCN red list of threatened species has 3.3% of the estimated 28,484 orchid species worldwide (Govaerts et al. 2017), but already 56.5% of these were found to be threatened with extinction.

Due to the exposed borders with neighbouring countries, the illegal trade of raw orchid plants has fast-tracked their extinction process in Nepal. Thus, in vitro mass propagation technique is being applied to raise plants in the laboratory to save them from extinction (Pant et al. 2019; Pant et al.
ever since the development of a protocol for non-symbiotic seed germination of orchids by Knudson (1951).

Plants can produce certain bioactive chemicals in response to the influence of their physical and chemical environments. They are used to overcome biotic and abiotic stresses by employing in defence and secondary metabolism (Andrew et al. 2007; Sudha and Ravishankar 2003). This facility of plants respond to physical and/or chemical stimuli can be used for the elicitation of pharmacologically active compounds by exposing an intact plant to stress factor/s (Chand et al. 2020; Kuzel et al. 2009). Therefore, in vitro techniques are very useful in safeguarding sustainable optimized sources of plant-derived natural products that can be used against vicious diseases like cancers. With the recent noteworthy anticancer activity of orchid based novel compounds like moscatilin, denbinobin, erianin, dendrochrysanene, fimbratone, cirrohopetalanthrin (Chen et al. 2007; Heo et al. 2007; Paudel et al. 2020; Peng et al. 2007; Wu et al. 2006; Xia et al. 2005) made orchids become a plausible candidate in the battle against cancer.

*Coelogyne stricta* (D. Don) Schltr. (Fig. 1a and b), native orchid to Nepal, commonly known as ‘The Rigid Coelogyne Pseudobulbs’ is an epiphytic orchid found on trees along steep riverbanks in lower and upper montane forests and also on mossy rocks as lithophytes at elevations of 1400 to 2000 m in Nepal (Raskoti 2009; Rajbhandari 2015). The plant has sparse pseudobulbs on the rhizome, leaves are leathery and inflorescence hysteranthous and up to 10-flowers. Flowers are white, lip with yellow spots, lamellae tinged with red in apical part. It has high aesthetic value, is often used as an ornamental plant in many gardens, nurseries, hotels. *Coelogyne stricta* has been mentioned to be beneficial against headache and fever by using the paste of pseudobulb (Baral and Kurmi 2006; Cragg and Newman 2013; Newman et al. 2003; Pant and Raskoti 2013).

Protocorms cultures have been established for the mass propagation and the production of bioactive compounds of valuable orchid species (Park et al. 2000). In vitro grown protocorms could be a good source of bioactive compounds and possess good biological activity. We have reported for the first time the cytotoxic activity of in vitro grown protocorms and wild plants of *C. stricta* in the present paper. Also, an effective conservation strategy is optimized for this orchid through tissue culture.

**Materials and Methods**

**Plant material for in vitro culture**

The 8 months old capsules of *C. stricta* (Fig. 1c) were collected from Godawari Lalitpur. They are about 2.5 ~ 3.5 cm long and 1.5 ~ 2.0 cm wide, have a mass of dark brown-yellow seeds inside were used for in vitro seed culture materials.

**Sterilization of plant material**

The capsule was washed in tap water with tween-20 detergent to remove the waste adherents. Then, inside the contamination-free chamber of laminar air hood, it was dipped in 0.6% sodium hypochlorite (Merck, India) for 10 min, 95% ethanol (Merck, India) for 1 min, and washed with sterile distilled water.

**Culture medium**

MS medium was selected to optimize the growth of *C. stricta* in vitro. A 30 g/L table sugar was used as a carbohydrate source and 0.01 g/L myoinositol as a vitamin in the MS medium (Hi-Media, India). Full and half strength’s MS medium was additionally supplied with coconut water (CW) and Adenine sulfate (Ad). The medium was solidified with 8 g/L agar (Hi-Media, India). The pH of the medium was adjusted to 5.8 with 0.1 N NaOH or HCl before autoclaving. The MS medium was autoclaved at 120°C and 15 lb for 15 minutes for effective sterilization.

**Culture initiation**

The capsule was cut longitudinally in a sterile petriplate with the help of a sterile No. 10 surgical blade. Seeds were inoculated on the above-mentioned culture medium for the initiation of culture. They were considered to have germinated upon the emergence of the embryo from the testa. Germination percentage of seeds in different culture medium was determined by examining the seeds microscopically after 60 days of culture.
Culture differentiation

After 8 weeks of inoculation, seeds gave enough mass of the green and fully developed protocorms, subcultures of them were performed for further development. The shoot proliferating medium was prepared by using different combination and concentration of plant growth hormones and additives in the full- and half-strength MS medium. Once the shoots reached about 2.5 cm long, they were further subculture on medium varied with auxin type and its concentrations.

Culture conditions

All inoculations were carried out under aseptic conditions in a laminar airflow chamber. The cultures were maintained at 25±2°C under white fluorescent light with a 16-h photoperiod with a light intensity of 3000 lux (fluorescent tubes 40 W, Philips, India). The number and length of seedlings in the culture medium were recorded. Each treatment consisted of at least six cultures replicas for an effective reproducibility check.

Acclimatization of plantlets

Plantlets above 2.5 cm in height with well-developed roots were removed from the medium, washed gently with tap water to remove the medium, and transplanted to appropriate plastic pots containing different substrates. The infected roots were treated with 2% of Bavistin. The pots were covered with plastic bags and its coverage was reduced weekly by 20% to decrease humidity. All the transplanted pots were maintained at greenhouse temperature (22 ~ 25 °C) under natural light for 3 months sprayed with water daily and external vitamins fortnightly.

Preparation of extract

The pseudobulbs of \textit{C. stricta} (Fig. 2a) were collected from Godawari Lalitpur for the wild extract (CsW) preparation. For in vitro extract (CsI) preparation, in vitro developed protocorms (Fig. 2b) were used. The collected pseudobulbs and in vitro protocorms were washed, air-dried and powdered. A sonication extraction was done with methanol in the ratio of 1:10 of the weight of material and volume of methanol (w/v) three times until the powder broth becomes discoulour. The extracts were concentrated in rotavapor (Eyela, Japan) to obtain dry extract and the extract was stored at 4°C for further use.

Cytotoxicity assay

Cytotoxic activity of extracts was evaluated by using the MTT (3- [4, 5-dimethylthiazole-2-yl]-2, 5-diphenyl-tetrazolium bromide) colourimetric assay with slight modification. For this purpose, about 8000 HeLa cells (cervical cancer) were cultured in T-flasks containing Minimum Essential Medium Eagle (EMEM) medium (Caisson Lab, USA) supplemented with 10% of fetal bovine serum (FBS) (Caisson Lab, USA), 1% of penicillin/streptomycin (Caisson Lab, USA) and 1% L-Glutamine (Caisson Lab, USA). The culture was kept in 5% CO₂ incubator at 37°C (Mosmann 1983). Following the attachment and cell confluence, the cells were treated with different concentrations (50, 100, 200, and 400 µg/ml) of the plant extracts for 48 hours in the 96 well cell culture plate. Following 48-hour incubation supernatant was removed and 50 µL of 5mg/ml MTT (prepared in EMEM medium) was added to each well. Following 4 hours of incubation, a purple formazan product was produced. About 100 µL DMSO (2.5%) (Merck, India) was added to dissolve formazan crystals. The absorbance was measured in a microplate reader at 630 nm. The percentage of the cytotoxic activity was calculated by using the following formula:

\[
\% \text{ cytotoxic activity} = \frac{(A_0 - A_t)}{(A_0)} \times 100
\]

Where, \(A_0\) is the absorbance of cells except plant extract, and \(A_t\) is the absorbance of extract-treated cells.

Statistical analysis

The average percentage of germinated seeds was taken after the eighth week of seed culture. Then, growth parameters were noted and analyzed every two weeks after the required subcultures. Data for the shoot and root development were presented as the mean of their respective numbers and lengths with ± standard error. The data of the percentage of cell growth inhibition was recorded as
the mean of triplicate. All the data were analyzed in Microsoft Excel 2019.

Results and Discussion

In vitro seed germination

The seeds of *C. stricta* (Fig. 3a and b) were inoculated on the different combinations of MS medium (FMS with 10% CW and 1mg/L adenine sulfate, half MS, half MS with 10% CW, and half MS with 15% CW). The inoculated seeds were started to swell up after 4 ~ 8 week of inoculation. After few weeks of incubation, green swelled seeds were changed into large globular protocorms. HMS medium with 15% coconut water was found suitable where the maximum seeds were germinated (Fig. 3c). Similarly, both combination formed with 10% coconut water in MS and HMS was also found quite responsive toward seed germination. In these conditions, more than 70% of seeds were germinated (Fig. 4). From all these data, we found that coconut water (CW) was found effective additives for the seed germination. This was supported by the findings of Reddy et al. (1992), who studied the seed germination and seedling growth in four different species of orchids (*Cymbidium aloifolium, Dendrobium crepidatum, Epidendrum radicans, and Spathoglottis plicata*) and found the seed germination after 5 weeks. It was also supported by Hoshi et al. (1994), who worked on the seed germination of four species of *Cypripedium* and Pradhan and Pant (2009) in *Dendrobium densiflorum*.

Development of protocorms into the shoots

The shoot proliferation from the protocorms was tested in various media (MS, MS with 10% CW, MS with 10% CW and 0.5 mg/L Adenine sulfate, MS with 15% CW, MS with 1 mg/L BAP with 2 mg/L Kinetin and 10% CW and HMS with 10% CW). Among all these tested medium, maximum shoot number was found in MS with 1 mg/L BAP and 2 mg/L Kinetin plus 10% CW medium (Fig. 5 and 6). For complete plantlet formation from 10 weeks old protocorms, it took 24 to 30 weeks. The present result was supported by Pant et al. (2011) in *Phiaus tancarvilleae* which took 24 weeks to develop into complete plantlets and Paudel and Pant (2012) in *Esmeralda clarkei* which took 25 weeks. Similarly, Parmar and Pant (2016) found that the MS medium with plant hormones (1 g/L NAA and 1 g/L BAP) were rather similar to this finding.

Development of roots on the shoots

Once the shoots reached about 2.0 ~ 2.5 cm long, they were cultured on full MS (FMS), full MS with 1 mg/L NAA (F1N) and full MS with 1.5 mg/L IBA (F1.5I) and full MS with 1 mg/L NAA and 1 mg/L adenine sulfate (F1N.1Ad). Among all these tested medium, the maximum root was
found in MS with 1.5 mg/L IBA (Figs. 7 and 8). Whereas only the MS medium showed the lowest rate of root proliferation and development. Parmar and Pant (2015) found that MS medium with NAA was more suitable for root proliferation. Similarly, Basker and Bai (2006) found that the MS medium with NAA showed significant results in root initiation and development. However, in the present study, full strength’s MS medium with 1.5 mg/L IBA was found more effective for root development.

Acclimatization of plantlets

Transplantation of the delicately raised in vitro plants had not yet been ill coped with the weather and necessary nutrients essential to develop resistance against physical, chemical, and biological factors. A good substrate has the optimum properties like water holding capacity, porosity, and drainage for the survivability of in vitro grown plantlets. In the present study, in vitro raised plantlets above 2.5 cm with well-developed roots were selected for acclimatization. They were transferred to the earthen pot containing different acclimatization substrates. The combination of coco-peat, pine bark, and sphagnum moss in ratio 2:1:1 was found to be a suitable potting mixture for hardening. Eighty percentage of plantlets were successfully survived under this condition. Hence, this result suggests that the mixture of coco-peat, pine bark, and moss will be favourable for the acclimatization of epiphytic orchid, *C. stricta* (Fig. 9).

**MTT assay**

The cytotoxic activity of wild plant (CsW) and in vitro protocorms (CsI) of *C. stricta* was assessed by MTT. The percentage of HeLa cells growth inhibition for wild plant’s extract (CsW) was 41.99% and in vitro plant’s extract (CsI) was 20.78% at 400 µg/ml (Fig. 10).
Previous studies made it clear that orchids are a potent source of anticancer agents. Several studies about the pharmacological properties of plant metabolites support the findings of this study. The majority of plant-based secondary metabolites like flavonoids, triterpenoids, and steroids (Gupta et al. 2004; Uddin et al. 2009; Wong et al. 2006) possess diverse pharmacological properties, including cytotoxic and cancer chemopreventive effects. In particular, they exert multiple biological effects due to their antioxidant and free radical-scavenging abilities (Gupta et al. 2004).

Many orchid species have shown cytotoxic activity against different human cancer cell lines, some examples like *Dendrobium nobile* has shown cytotoxic activity toward lung carcinoma, ovary adenocarcinoma, and promyelocytic leukemia cell lines (You et al. 1995), *D. chrysanthum* has shown cytotoxic activity against HL-60 cells (Li et al. 2001), *D. amoenum, D. longicorne, D. moniliforme* and *D. crepidatum* against cervical and brain cancer cell lines (Paudel and Pant 2017; Paudel et al. 2017, 2018 & 2019), *Bulbophyllum kwangtungense* against cervical and leukemia cell lines (Wu et al. 2006), and *B. odoratissimum* was found to have cytotoxicity against leukemia, human hepatoma, human lung adenocarcinoma and human stomach cancer cell lines (Chen et al. 2007). *Dendrobium transparens* and *Vanda cristata* have also shown the cytotoxic activity toward cervical cancer and brain tumour cell lines (Joshi et al. 2020). The previous findings of anticancer activity of the extracts support the cytotoxic activity of *C. stricta* toward HeLa cell line.

**Conclusion**

Plant cell and tissue culture offer an alternative source for the rapid propagation of medicinal plants for conservation and else. Protocorm cultures have established for the mass propagation of this orchid species. Half-strength’s MS medium with 15% coconut water, full-strength’s MS medium with 1 mg/L BAP, 2 mg/L kinetin and 10% coconut water, and full-strength’s MS medium with 1.5 mg/L IBA were recorded as suitable for the different stages of propagation from seeds. The in vitro-developed plantlets were successfully acclimatized on the 2:1:1 ratio of coco-peat, pine bark and sphagnum moss. The in vitro-developed protocorms are enriched in bioactive metabolite is likely to be highly useful for commercial production. However, wild pseudobulbs’ extract has shown more growth inhibition of cervical cancer cells as compared to in vitro-developed protocorms. The less cytotoxic activity of protocorms is may be due to the immaturity of protocorms where more bioactive compounds cannot be formed.

**Acknowledgements**

This study was supported by the KOICA/KU-Integrated Rural Development Project (Grant No. 01). The grant was received by Bijaya Pant (Principal Investigator).

**References**

Andrew RL, Peakall R, Wallis IR, Foley WJ (2007) Spatial distribution of defense chemicals and markers and the maintenance of chemical variation. Ecology 88:716-728

Baral SR, Kurmi PP (2006) A Compendium of Medicinal Plants of Nepal. Publisher Rachana Baral, Kathmandu, pp 26

Barker S, Bai VN (2006) Micropropagation of Coelogyne stricta (D. Don) Schltr. via pseudobulb segment cultures. Trop and Subtrop Agroecosys 6:31-35

Chand K, Shah S, Sharma J, Paudel MR, Pant B (2020) Isolation, characterization, and plant growth-promoting activities of endophytic fungi from a wild orchid Vanda cristata. Plant Sig Behav 15:e1744294

Chen Y, Xu J, Yut H, Qin CW, Zhangt Y, Liu Y, Wang J (2007) *Bulbophyllum O doratissimum 3,7-Dihydroxy-2,4,6-trimethoxyphenanthrene*. J Korean Chem Soc 51:352-355

Cozzolino S, Widmer A (2005) Orchid diversity: an evolutionary consequence of deception? Trends Ecol Evol 20:487-494

Cragg GM, Newman DJ (2013) Natural products: a continuing source of novel drug leads. Biochim Biophy Acta 1830:3670-3695

Govaerts R, Bernet P, Kratochvil K, Gerlach G, Carr G, Alrich P, Pridegeon AM, Pfahl J, Campacci MA, Holland BD, Tigges H, Shaw J, Cribb P, George A, Kreuz K, Wood J (2017). World Checklist of Orchidaceae. The Royal Botanic Gardens, Kew. http://wcsp.science.kew.org. Retrieved on 31 October 2017

Gupta M, Mazumder UK, Kumar RS, Sivakumar T, Vamsi MLM (2004) Antitumor activity and antioxidant status of Caesalpinia bonducella against Ehrlich ascites carcinoma in Swiss albino mice. J Pharm Sci 94:177-184

Heo JC, Woo SU, Son M, Park JY, Choi WS, Chang KT, Kim SU, Yoon EK, Kim YH, Shin HM, Lee SH (2007) Antitumor activity of Gastrodia elata Blume is closely associated with a GTP-Ras dependent pathway. Oncol Rep 8:849-853

Hoshi Y (1994) In vitro germination of four Asiatic taxa of Cypripedium and notes on the nodal micropropagation of American Cypripedium montanum. Lindleyana 9:93-97

Jersáková J, Johnson SD, Kindlmann P (2006) Mechanisms and evolution of deceptive pollination in orchids. Biol Rev 81:219-235

Joshi PR, Paudel MR, Chand MB, Pradhan S, Pant KK, Joshi GP, Bolhara M, Wagner SH, Pant B, Pant B (2020) Cytotoxic effect of selected wild orchids on two different human cancer cell lines. Heliyon 6:e03991

Knudson L (1951) Nutrient solutions for orchids. Bot Gaz 112:528-532

Kuzel S, Vydra J, Triska J, Vrchołtova N, Hruby M, Cigler P (2009) Elicitation of pharmacologically active substances in an intact
medical plant. J Agric Food Chem 57:7907-7911
Li YM, Wang HY, Liu GQ (2001) Erianin induces apoptosis in human leukemia HL-60 cells. Acta Pharma Sin 22:1018
McCormick MK, Whigham DF, O’Neill J (2004) Mycorrhizal diversity in photosynthetic terrestrial orchids. New Phytot 163:425-438
Mosmann T (1983) Rapid colorimetric assay for cellular growth and survival: application to proliferation and cytotoxicity assays. J Immunol Meth 65:55-63
Murthy HN, Paek KY, Park SY (2018) Micropropagation of orchids by using bioreactor technology. In: Lee YI, Yeung EC (eds) Orchid propagation: from laboratories to greenhouses-methods and protocols. Springer, New York, pp 195-208
Newman DJ, Cragg GM, Snader KM (2003) Natural products as sources of new drugs over the period 1981 – 2002. J Nat Prod 66:1022-1037
Pant B, Paudel MR, Chand MB, Wagner SH (2016) A Treasure Trove of Orchids in central Nepal. Central Department of Botany, Tribhuvan University, Kathmandu, Nepal
Pant B (2013) Medicinal orchids and their uses: Tissue culture a potential alternative for conservation. Afr J Plant Sci 7: 448-467
Pant B, Paudel MR, Chand MB, Pradhan S, Malla BB, Raskoti BB (2012) Medicinal orchids of Nepal. Himalayan map house Ltd., Kathmandu pp 104
Pant B, Shrestha S, Pradhan S (2011) In vitro seed germination and seedling development of Phaius tancarvilleae (L’Her.) Blume. Sci World 9:47-52
Park SY, Huh YS, Paek KY (2018) Common protocol in orchid micropropagation. In: Lee YI, Yeung EC (eds) Orchid propagation: from laboratories to greenhouses-methods and protocols. Springer, New York, pp 179-193
Park SY, Murthy HN, Kee YP (2000) Mass multiplication of plant regeneration in Phalaenopsis. Plant Cell Tiss Org Cult 62:109-124
Pant B, Raskoti BB (2013) Medicinal orchids of Nepal. Himalayan map house Ltd., Kathmandu pp 104
Pant B, Shrestha S, Pradhan S (2011) In vitro seed germination and seedling development of Phaius tancarvilleae (L’Her.) Blume. Sci World 9:47-52
Park SY, Huh YS, Paek KY (2018) Common protocol in orchid micropropagation. In: Lee YI, Yeung EC (eds) Orchid propagation: from laboratories to greenhouses-methods and protocols. Springer, New York, pp 179-193
Park SY, Murthy HN, Kee YP (2000) Mass multiplication of protocorm-like bodies using bioreactor system and subsequent plant regeneration in Phalaenopsis. Plant Cell Tiss Org Cult 62:109-124
Parmar G, Pant B (2015) In vitro seed germination and seedling development of Coelogyne flaccida Lindl. (Orchidaceae). Adv Forestry Sci 2:85-88
Parmar G, Pant B (2016) In vitro seed germination and seedling development of the orchid Coelogyne stricta (D. Don) Schltr. Afr J Biotechnol 15:105-109
Paudel MR, Bhattarai, HD, Pant B (2020) Traditionally Used Medicinal Dendrobium: A Promising Source of Active Anticancer Constituents. In: Merillon JM, Kodja H (eds) Orchid Phytochemistry, Biology and Horticulture. Springer, Cham pp. 16
Paudel MR, Chand MB, Pant B, Pant B (2017) Cytotoxic activity of antioxidant-riched Dendrobium longicorne. Pharmacogn J 9:499-503
Paudel MR, Chand MB, Pant B, Pant B (2018) Antioxidant and cytotoxic activities of Dendrobium moniliforme extracts and the detection of related compounds by GC-MS. BMC Comp Alt Med 18:134
Paudel MR, Chand MB, Pant B, Pant B (2019) Assessment of Antioxidant and Cytotoxic Activities of Extracts of Dendrobium crepidatum. Biomolecules 9:478
Paudel MR, Pant B (2012) In vitro micropropagation of rare orchid (Esmeralda clarkei Rehbg.) from shoot tip section. Int J Bio Pharm Allied Sci 1:1587-1597
Paudel MR, Pant B (2017) Cytotoxic activity of crude extracts of Dendrobium amoenum and detection of bioactive compounds by GC-MS. Bot Orient: J Plant Sci 11:38-42
Peng J, Xu Q, Xu Y, Qi Y, Han X, Xu L (2007) A new anticanicer dihydroflavanoid from the root of Spiranthes australis (R. Brown) Lindl. Nat Prod Res 21:641-645
Pradhan S, Pant B (2009) In vitro seed germination in Cymbidium elegans Lindl. and Dendrobium densiflorum Lindl. ex Wall. (Orchidaceae). Bot Orient: J Plant Sci 6:100-102
Rajbhandari KR (2015) A handbook of the orchids of Nepal. Kathmandu, Nepal: Department of Plant Resources, Ministry of Forest and Soil Conservation, Government of Nepal
Raskoti BB (2009) The orchids of Nepal published by Bhakta Bahadur Raskoti and Rita Ale; Quality printers, Kathmandu, Nepal
Reddy PV, Nanjan K, Shanmugavelu KG (1992) In vitro studies in tropical orchids: seed germination and seedling growth. J Orchid Soc India 6:75-78
Roberts DL, Dixon KW (2008) Orchids. Curr Bio 18:325-329
Shah S, Thapa BB, Chand K, Pradhan S, Singh A, Varma A, Sharma J, Pant B (2019) Piformospora indica promotes the growth of the in-vitro-raised Cymbidium aloifolium plantlet and their acclimatization. Plant Sig Behav 14:1596716
Shefferson RP, Taylor DL, Weiß M, Garnica S, McCormick MK, Adams S, Gray HM, McFarland JW, Kull T, Tali K, Yukawa T, Kawahara T, Miyoshi K and Lee YI (2007) The evolutionary history of mycorrhizal specificity among lady’s slipper orchids. Evolution 61:1380-1390
Sudha G, Ravigshankar GA (2003) Elicitation of anthocyanin production in callus cultures of Daucus carota and involvement of calcium channel modulators. Curr Sci 84:775-779
Tsai WC, Fu CH, Hsiao YY, Huang YM, Chen LJ, Wang M, Chen HH (2013) Orchid Base 2.0: comprehensive collection of Orchidaceae floral transcriptomes. Plant Cell Physiol 54:e7
Uddin SJ, Grice ID, Tiralongo E (2009) Cytotoxic effects of Bangladeshi medicinal plant extracts. eCAM. doi:10.1093/ ecam/nep111
Wong CC, Li HB, Cheng KW, Chen F (2006) A systematic survey of antioxidant activity of 30 Chinese medicinal plants using the ferric reducing antioxidant power assay. Food Chem 97:705-711
Wu B, He S, Pan YJ (2006) New dihydrodibenzoxepins from New Phytol 97:705-711
Xia WB, Xue Z, Li S, Wang SJ, Yang YC, He DX, Ran GL, Kong HH, Wu B, He S, Pan YJ (2006) New dihydrodibenzoxepins from New Phytol 97:705-711
You HL, Park JD, Baek NI, Kim S, Ahn BZ (1995) In vitro and in vivo antimural phenanthrenes from the aerial parts of Dendrobium nobile. Planta Med 61:178-180