Micropropagation of *Tolumnia* Orchids through Induction of Protocorm-like Bodies from Leaf Segments

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Abstract. A protocol for plant regeneration via direct induction of protocorm-like bodies (PLBs) from leaf segments of *Tolumnia* Snow Fairy was developed as a basis for mass production. Ten-month-old, in vitro–grown donor plantlets were obtained by inducing shoots from buds on the flower stalk. Leaf segments harvested from plantlets of different heights and from expanding leaves at different positions were compared, as were two BA concentrations with 0.5 mg L⁻¹ NAA. The greatest rate of PLB induction (16.7%) was observed when leaf segments taken from 1- to 2-cm height plants were cultured in Murashige and Skoog (MS) basal medium supplemented with 2 mg L⁻¹ BA and 0.5 mg L⁻¹ NAA after 16 weeks of culture. When using leaf explants, only inner, expanding leaves cultured on MS basal medium supplemented with 4 mg L⁻¹ BA and 0.5 mg L⁻¹ NAA resulted in PLB induction, at an average rate of 25.5 PLBs per explant. After 16 weeks of culture, histological and scanning electron microscopy (SEM) observations revealed that PLBs originated from epidermal cells of leaf explants. PLBs of 1 to ≤2 mm in diameter continued to proliferate after 4 weeks of culture. These secondary PLBs could be produced from either whole PLBs or the upper side of PLBs. Finally, PLBs were regenerated into plantlets. After ~14 months of culture, fully developed plants exhibiting well-developed roots and shoots were acclimatized. These plants grew well, with 1-year survival rates of nearly 73%, for plants originating as explants taken from inner leaves. Some mature plants flowered 1 year after transplantation. This study presents a simple system that can provide a large number of PLBs for mass propagation in a short time that can be converted into plants and also used for the new cultivars of *Tolumnia* orchids.

The group of orchids now called toolumnia was once called Oncidium section Variegata and was commonly referred to as the “equitant oncidiums” (Aldrich, 1994). Plants in this genus are endemic to the Caribbean islands. These plants are miniature, sympodial epiphytic orchids with triangular succulent leaves that form a small fan and usually lack a pseudobulb (Aldrich, 1994; Baker and Baker, 2006). In addition, their hybrids present a surprising array of colors and, despite being subjected to hybridization for a relatively short time, breeders have made important advances in a few years (Aldrich, 1994). Furthermore, hybridization in breeding programs has proven to be a very credible technique for producing a wide range of successful cultivars with an attractive combination as flower colors, shapes, fragrances, and seasonality. Certainly, clonal mass propagation hybrids of commercial value were made possible by in vitro propagation (Pack et al., 2011).

Normally, an apical meristem or shoot tip is used for in vitro clonal propagation of orchids. However, removal of shoot tips can endanger the mother plant. Leaf explants are easy to obtain, are more expendable to the mother plant (Chugh et al., 2009; De and Sil, 2015), and are available in any season, unlike inflorescence explants (Chugh et al., 2009). Presently, commercial growers of most orchid genera prefer propagation through PLB formation because of the large number of PLBs that can be achieved within a relatively short period of time using this method (Antony et al., 2014; Ng et al., 2010). Plant regeneration through PLB induction from leaf explants has been developed for *Acampe* (Nayak et al., 1997), *Aerides* (Murthy, 2005; Murthy and Pyati, 2001), *Coelogyne* (De and Sil, 2015), *Dendrobium* (Chung et al., 2005, 2007; Goswami et al., 2015; Martin and Madassery, 2006), *Oncidium* (Balilashaki et al., 2015; Chen and Chang, 2001, 2003, 2004; Chen and Hong, 2012; Chen et al., 1999; Chung et al., 2010; Hong et al., 2008; Mata-Rosas et al., 2011; Mayer et al., 2010; Mengxi et al., 2011), *Phalaenopsis* (Chen and Chang, 2006; Gow et al., 2009; Kuo et al., 2005; Park et al., 2002a, 2002b), *Spathoglottis* (Teng et al., 1997), *Vanda* (Decruse et al., 2003), and *Vanilla* (Janarthanam and Seshadri, 2008; Tan et al., 2011). In tissue culture, cytokinins are a group of hormones that promote cell division, especially; BA is widely used for tissue culture research because of its low price and effectiveness (George et al., 2008). Until now, there have only been a few reports describing PLB induction in toolumnia orchids (Shen et al., 2018). However, not all of cultivars can respond to the same protocol under the same culture conditions. In this study, we describe a simple and suitable protocol for clonal propagation of toolumnia orchids from leaf segments via PLB induction. After PLB induction, PLBs were used for further proliferation via a PLB bisection method. PLBs were converted into plantlets and then full plants, which were then acclimatized and allowed to flower.

Materials and Methods

Initiation of in vitro–grown donor plantlets. Tissue culture materials were obtained from the flower stalk nodes of the *Tolumnia* cultivar Snow Fairy (in vivo-grown) (Fig. 1A). After 10-month-old, in vitro–grown donor plantlets of *Tolumnia* Snow Fairy were obtained (Chookoh, 2015; Chookoh et al., 2016) (Fig. 1B–C). Leaves from these in vitro donor plantlets were used as explants for further experiments (Fig. 1D) and to evaluate the effects that plant height and leaf position at the time of leaf harvesting had on PLB induction.

Leaf culture medium and growth conditions. For the culture of leaf segments, MS basal medium (Murashige and Skoog, 1962) modified with half-strength mineral macroelements and microelements and apart from full-strength NaFe-EDTA and FeSO₄, vitamins, and glycine, containing 100 mg L⁻¹ myo-inositol (Sigma-Aldrich Co., St. Louis, MO), 170 mg L⁻¹ NaH₂PO₄ (Hayashi Pure Chemical Ind., Co., Ltd., Osaka City, Japan), 30 g L⁻¹ sucrose (Taiwan Sugar Corporation,
Tainan City, Taiwan), and 8 g L\(^{-1}\) agar (Trade Mark, Tainan City, Taiwan) were prepared. The pH of the medium was adjusted to 5.2 with 0.1 N NaOH or HCl before autoclaving. The medium (10 mL) was dispensed into culture tubes (20 × 15 mm; PYREX Glass Test Tube 9820; Corning, USA). The culture vials containing the medium were autoclaved with 1.2 kg/cm\(^2\) of pressure at 121 °C for 20 min.

Leaf segments were placed horizontally on the surface (one explant per tube) of this MS basal medium. All cultures were maintained in a culture room at 25 ± 2 °C for 8 weeks in the dark. After 8 weeks of culture, they were transferred to dim light (5 m\(^{-2}\)·s\(^{-1}\)·L\(^{-1}\) 1-Naphthaleneacetic acid (NAA) and 2 or 4 mg L\(^{-1}\) 6-Benzylaminopurine (BA)). For each of these treatments, ≈30 leaf segments were cultured. One explant was inoculated in each culture vial, and five test tubes were considered one replication. Six replications were established for each treatment. Explants were subcultured to fresh medium every 4 weeks during the culture period. After 16 weeks of culture, PLB induction rate, necrotic explant rate, and the number of PLBs per explant were recorded. Cultures were examined and photographed with a stereozoom microscope (SZH; Olympus, Tokyo, Japan). Further proliferation of PLBs induced from leaf explants. PLBs still attached to leaf were fixed in 2.5% glutaraldehyde in 0.1 M phosphate buffer (pH 7.0) for 4 h at 4 °C, then dehydrated in an ethanol series, dried in a critical-point dryer (HCP-2, Hitachi, Japan), and finally coated with gold using an ion coater (E1010; Hitachi, Tokyo, Japan) (Chang et al., 2010). A scanning electron microscope (S-3000N; Hitachi) was used for examination and photography of the specimens.

Effects of plant height, leaf position, and physical wounding on their subsequent growth and proliferation. PLBs induced from leaf explants were sorted into two sizes, small diameter (1 to 2 cm) and large diameter (>2 to 3 cm), to determine whether the size of the PLB affected later PLB proliferation. Other single PLBs were left whole or were bisected transversely into an upper and lower side before reculture to determine whether physical wounding of a PLB affected PLB proliferation.

Histology of PLBs induced from leaf explants. PLBs still attached to leaf were fixed in 2% paraformaldehyde and 2.5% glutaraldehyde in 0.1 M phosphate buffer (pH 6.8, at 4 °C overnight), dehydrated in an ethanol series, and embedded in Technovit 7100 resin (Kulzer & Co., Wehrheim, Germany) as described by Yeung and Chan (2015). Serial sections (3-μm thick) were cut using a rotary microtome (RM2245; Leica, Nussloch, Germany). Sections were stained with periodic acid-Schiff reagent for total insoluble carbohydrates and counterstained with amido black 10B for proteins (Yeung, 1984). These sections were observed, and pictures were captured digitally using a charge-coupled device camera attached to the light microscope (Axiolab; Carl Zeiss Microscopy, GmbH, Jena, Germany).

SEM. For SEM, PLBs still attached to the leaf explant were fixed in 2.5% glutaraldehyde in 0.1 M phosphate buffer (pH 7.0) for 4 h at 4 °C, then dehydrated in an ethanol series, dried in a critical-point dryer (HCP-2, Hitachi, Japan), and finally coated with gold using an ion coater (E1010; Hitachi, Tokyo, Japan) (Chang et al., 2010). A scanning electron microscope (S-3000N; Hitachi) was used for examination and photography of the specimens.

Further proliferation of PLBs induced from leaf cultures. The PLBs that developed on leaf explants after 20 weeks were used as explants for PLB proliferation experiments (Fig. 1F). A stereo microscope was used for dividing the PLBs. PLB explants were placed on the surface (one explant per tube; Fig. 1G) of MS basal medium with one-eighth of NH\(_4\)NO\(_3\) and KNO\(_3\), half strength of the other mineral macroelements and microelements, and full strength of NaFe-EDTA, FeSO\(_4\), vitamins, and glycine. Most of contents were similar to the PLB induction medium, whereas proliferation medium was added containing 20 g L\(^{-1}\) sucrose, 150 mL L\(^{-1}\) coconut water, 1 g L\(^{-1}\) peptone, 6 g L\(^{-1}\) potato powder, and 1 g L\(^{-1}\) activated charcoal. The pH of the medium was adjusted as described previously. All cultures were maintained in a culture room at 25 ± 2 °C for 6 weeks in dim light (5 μmol m\(^{-2}\)·s\(^{-1}\)) provided by cool white fluorescent tubes under a 12-h photoperiod.

Acclimatization and transplantation. After 4 months of in vitro culture, encompassing
Results and Discussion

PLBs outwardly look like somatic embryos in form and development. Lee et al. (2013) investigated the early stages of PLB formation and confirmed that PLBs are truly somatic embryos. In our experiments, at 16 weeks of culture, the number of explants forming PLBs and the mean number of PLBs per explant were determined. This was done because the PLBs were too small to count, even with a stereomicroscope, at 8 weeks.

Effect of plant height and BA concentration on PLB induction. The induction of PLBs from leaf explants from in vitro–grown donor plantlets was influenced by both plant height and the BA concentration in the induction medium (Table 1). The rate of PLB formation on an explant was generally low. However, the greatest rate of explants forming PLBs was observed on leaves taken from plantlets 1 to 2 cm in height that were cultured on MS basal medium supplemented with 0.5 mg L\(^{-1}\) NAA and 2 or 4 mg L\(^{-1}\) BA. Only inner expanding leaves (first to third) cultured on MS basal medium supplemented with 4 mg L\(^{-1}\) BA and 0.5 mg L\(^{-1}\) NAA resulted in PLB induction, at an average of 25.5 PLBs per explant. Likewise, young leaves from Coelogyne flaccida showed better regeneration relative to older explants, owing to their less-rigid cell walls, and retained competence for regeneration and proliferation (De and Sil, 2015).

Under equal nutritional conditions seem to indicate the importance of the source of explants (Murthy and Pyati, 2001). In contrast, Oncidium Gower Ramsey in vitro–grown donor plantlets 5 to 7 cm in height yielded both the greatest rate of somatic embryo proliferation and more somatic embryos per explant than did plants 2 to 4 cm in height (Chen et al., 1999). In this study, the low number of explants forming PLBs may be because leaves from different position on plantlet were taken as explants. However, this prompted an investigation into the effects of leaf position on PLB induction.

Effect of leaf position and BA concentration on PLB induction. The effects of leaf position within the donor plantlet at two different BA concentrations on PLB induction were tested using in vitro–grown donor plantlets of 1 to 2 cm in height (Table 2). No PLBs formed when outer leaves were grown on MS basal medium supplemented with 0.5 mg L\(^{-1}\) NAA and 2 or 4 mg L\(^{-1}\) BA. Only inner expanding leaves showed better regeneration relative to older explants, owing to their less-rigid cell walls, and retained competence for regeneration and proliferation (De and Sil, 2015).

In contrast, in Acempe praemorsa (Roxb.) Blatter and McCann, both young expanding leaves (first and second) and older expanding leaves (fifth and sixth) failed to regenerate shoot buds, whereas the third and fourth leaves could (Nayak et al., 1997). Meanwhile, Seeni and Latha (1992) have reported that all leaves from in vitro–grown shoots of Renanthera inschoottiana Rolfe produced shoot buds in culture. This rate of success and low rate of PLB formation was similar to results in Acampe (Nayak et al., 1997), Aerides (Murthy, 2001; Murthy and Pyati, 2001), Dendrobium (Chung et al., 2005), Oncidium (Chen et al., 1999; Mayer et al., 2010), and Phalaenopsis (Baililashaki et al., 2015; Chen and Chang, 2006; Kuo et al., 2005; Park et al., 2002a). Nayak et al. (1997) showed that explants exuded copious amounts of phenolic compounds within 3 to 5 d of culture and that transfer to fresh medium at 7-d intervals could decrease the amount of exudates to overcome the inhibition of PLB formation.

Cytokinins play important roles in the control of plant development, especially in plant cell division (George et al., 2008). Trigiano and Gray (2000) reported that BA has remarkably stronger cytokinin activity than the naturally cytokinin such zeatin. Moreover, BA has been shown to efficiently induce

Table 1. Effects of plant height on PLB induction of Tolumnia Snow Fairy after 16 weeks of culture.\(^a\)

<table>
<thead>
<tr>
<th>BA (mg L(^{-1}))</th>
<th>Ht</th>
<th>PLB induction (%)(^b)</th>
<th>Necrosis (%)(^c)</th>
<th>PLBs/explant</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>2–3 cm</td>
<td>16.7 ± 9.5 a</td>
<td>33.3 ± 15.2 a</td>
<td>17.5 ab</td>
</tr>
<tr>
<td>2</td>
<td>3–4 cm</td>
<td>10.0 ± 4.5 ab</td>
<td>43.3 ± 6.1 a</td>
<td>14.3 bc</td>
</tr>
<tr>
<td>4</td>
<td>1–2 cm</td>
<td>13.3 ± 4.2 a</td>
<td>23.3 ± 9.5 a</td>
<td>24.0 ab</td>
</tr>
<tr>
<td>4</td>
<td>2–3 cm</td>
<td>3.3 ± 3.3 ab</td>
<td>43.3 ± 9.5 a</td>
<td>41.0 ab</td>
</tr>
<tr>
<td>3–4 cm</td>
<td>0.0 ± 0.0 b</td>
<td>36.7 ± 8.0 a</td>
<td>0.0 c</td>
<td></td>
</tr>
</tbody>
</table>

\(a\)Six replicates each containing five test tubes (one explant per test tube) were performed for each treatment and cultured on MS basal medium supplemented with different of BA concentrations with 0.5 mg L\(^{-1}\) NAA.

\(b\)Percentage data used angular transformation before analysis, and means ± SE within a column followed by different letters are significantly different according to least significant differences multiple range test at \(P \leq 0.05\).

\(c\)Significance was determined by analysis of variance.

Table 2. Effects of leaf position on PLB induction of Tolumnia Snow Fairy after 16 weeks of culture.\(^a\)

<table>
<thead>
<tr>
<th>BA (mg L(^{-1}))</th>
<th>Positions</th>
<th>PLB induction (%)(^b)</th>
<th>Necrosis (%)(^c)</th>
<th>PLBs/explant</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>Inner</td>
<td>0.0 ± 0.0 b</td>
<td>36.7 ± 9.5 a</td>
<td>0.0 b</td>
</tr>
<tr>
<td>2</td>
<td>Outer</td>
<td>0.0 ± 0.0 b</td>
<td>26.7 ± 8.4 ab</td>
<td>0.0 b</td>
</tr>
<tr>
<td>4</td>
<td>Inner</td>
<td>16.7 ± 6.1 a</td>
<td>10.0 ± 4.5 b</td>
<td>25.5 a</td>
</tr>
<tr>
<td>4</td>
<td>Outer</td>
<td>0.0 ± 0.0 b</td>
<td>26.7 ± 6.7 ab</td>
<td>0.0 a</td>
</tr>
</tbody>
</table>

\(a\)Six replicates each containing five test tubes (one explant per test tube) were performed for each treatment and cultured on MS basal medium supplemented with different of BA concentrations with 0.5 mg L\(^{-1}\) NAA. Leaves were taken from in vitro–grown donor plantlets of 1 to 2 cm in height.

\(b\)Percentage data used angular transformation before analysis, and means ± SE within a column followed by different letters are significantly different according to least significant differences multiple range test at \(P \leq 0.05\).

\(c\)Significance was determined by analysis of variance.

PLB = protocorm like-body; NS = not significant; MS = Murashige and Skoog.
PLBs from leaf explants (Murthy and Pyati, 2001; Sheelavanthmath et al., 2005) and to play an important role in plant regeneration from leaf explants in tissue culture of other orchid species (Deb and Pongener, 2013; Janarthanam and Seshadri, 2008; Martin and Madasserry, 2006; Nayak et al., 1997). Our result showed that both 2 and 4 mg·L⁻¹ BA with 0.5 mg·L⁻¹ NAA in the medium promoted the formation of PLBs, especially when leaf explants were cultured on MS basal medium supplemented with 4 mg·L⁻¹ BA and 0.5 mg·L⁻¹ NAA. Several orchid studies on PLB induction and shoot regeneration from leaf explants used a range of BA concentrations, between 1 and 10 mg·L⁻¹ (Chen and Chang, 2001; Chen and Hong, 2012; Chung et al., 2005; Kuo et al., 2005; Martin and Madasserry, 2006; Murthy and Pyati, 2001; Nayak et al., 1997; Park et al., 2002b; Sheelavanthmath et al., 2005; Su et al., 2006). However, high levels of BA also could inhibit PLB induction (Murthy and Pyati, 2001) as well as which prefer to induce adventitious shoots formation (Narayana et al., 2016; Trigiano and Gray, 2000). Furthermore, Hongthongkham and Bunnag (2014) reported that BA alone induced shoot formation rather than PLB, whereas low concentrations of auxin promoted PLB growth (Khoddamzadeh et al., 2011) and reduced plantlet conversion (Gow et al., 2010).

**Microscopic observation of leaf-derived PLBs.** PLB formation is a unique characteristic of Orchidaceae. A structure can be called a PLB from the initiation of globular swelling to appearance of shoot primordia without any roots (Batygina et al., 2003). Histological and SEM observations revealed in inner position of leaf on MS basal medium supplemented with 4 mg·L⁻¹ BA and 0.5 mg·L⁻¹ NAA medium that PLBs (primary embryo) originated from epidermal cell division (Fig. 2A), whereas meristematic cells were densely stained and had a smaller size than the leaf cells. In some cases, secondary embryos developed from outer layers of the primary embryos (Fig. 2B). After 16 weeks of culture, multiple PLBs were visible (Fig. 2C). Antony et al. (2014) used histological and SEM observations to report that PLBs can develop from a wounded surface of an explant and can be converted into plantlets, as the PLBs consisted of multiple meristematic centers then can be gradually differentiated into shoot, leaf, and root. Our results support the various other reports that demonstrated that PLBs can form from the epidermal cell layers (Chen and Chang, 2004, 2006; Khoddamzadeh et al., 2011; Kuo et al., 2005). Moreover, most PLBs formed at the base of the explant. Efficient PLB induction on the leaf base has been reported for several other orchid species, such as *Aerides* (Murthy and Pyati, 2001), *Coelogyne* (De and Sil, 2015), *Cymbidium* (Deb and Pongener, 2013), *Dendrobium* (Martin and Madasserry, 2006), *Oncidium* (Chen and Hong, 2012), and *Phalaenopsis* (Gow et al., 2009; Park et al., 2002b). Moreover, our results support those showing that no PLB induction occurred on either side of the leaf explant between the tip and the cut end (Chung et al., 2007). Although some secondary PLBs did form during the PLB induction stage, the number of PLBs that were obtained was still low, prompting the requirement of a PLB proliferation stage to increase PLB numbers.

**Effects of PLB size and physical wounding on PLB proliferation.** Julkiflee et al. (2014) revealed that PGR-free medium could increase PLB proliferation. In addition, Ng and Saleh (2011) suggested that PGR-free medium could obtain genetically stable PLBs. In recent years, secondary PLB development from PLBs has been shown for various orchids, including *Dendrobium* (Julkiflee et al., 2014), *Doritaenopsis* (Amaki and Higuchi, 1989), and *Phalaenopsis* (Huang et al., 2014; Khoddamzadeh et al., 2011). However, the results in this study showed that both large and small PLBs formed secondary PLBs (PLB proliferation) at nearly the same rates over 2, 4, and 6 weeks of culture (Fig. 3). However, there was a difference in generation of secondary PLBs after the first subculture. Small PLBs, 1 to ≤2 mm in diameter, tended to continue generation through 4 weeks after their subculture, whereas PLBs >2 to 3 mm in diameter tended to deteriorate 4 weeks after subculture. In addition, after subculture, shoot germination was observed in PLBs of both sizes. PLB necrosis was also reported by Bustam et al. (2014). Our study indicated that PLBs 1 to ≤2 mm in diameter could proliferate over a longer term, with no deterioration until 6 weeks of culture. Therefore, we selected this size PLB for the physical wounding investigation.

PLBs of 1 to ≤2 mm were physically wounded by cutting them in half, and PLB proliferation was measured after 6 weeks of culture (Table 3). Whole PLBs achieved a 33.3% PLB proliferation rate, in comparison with a 40% PLB proliferation rate for the upper half of bisected PLBs (Fig. 4A–B). The PLB proliferation rates and the average
numbers of secondary PLBs were not significantly different between the upper half and whole PLBs. In addition, secondary PLBs produced an additional 3.1 to 3.5 PLBs. In contrast, no secondary PLBs were observed from the lower half of PLBs. This study indicated that only whole PLBs and the upper sides of PLBs could form secondary PLBs. This may be because the whole PLB and the upper side of the PLB still had many cell clumps around apical meristems that it would protrude and develop after 2 weeks. Others have shown that the lower halves of bisected PLBs are more efficient in the proliferation of secondary PLBs (Amaki and Higuchi, 1989; Huang et al., 2014; Tanaka, 1987).

Six weeks after subculture of the PLBs to the same medium, the shoot germination rate was 33.3% with an average of 3.5 shoots per whole PLB, with corresponding rates of 40% with 3.1 shoots for the upper PLB halves. The PLB upper half formed primordial leaves instead of PLBs (Fig. 4C). Amaki and Higuchi (1989) indicated that the dorsal segment of a PLB had a tendency to generate plantlets, but a basal segment could not. Because a single PLB was divided by scalpel, the lower halves of the PLBs easily died. Bustam et al. (2014) reported that PLBs became brown and that most failed to proliferate further when compact clumps of PLBs were separated by scalpel and forceps. This result revealed that PLBs tended to die easily and that their viability decreased in long-term culture.

Regeneration of PLBs into plantlets. After 16 weeks of culture, some PLBs (still attached to leaf explants) were transferred to proliferation medium for multiplication and were subcultured to fresh medium at 7-d intervals. Shoot buds first appeared as small, green globular swellings during PLB proliferation and eventually developed into shoots (Fig. 5A). After 1 month of proliferation, PLBs were removed from the explant and moved to regeneration medium for further PLB development and plant regeneration (Fig. 5B). Plant regeneration has been induced from PLBs from various orchids (Balilashaki et al., 2015; Chen and Chang, 2004; De and Sil, 2015; Mayer et al., 2010; Teng et al., 1997). Since PLBs are true orchid embryos (Lee et al., 2013), these structures can easily convert into plantlets (Ng and Saleh, 2011).

Table 3. Effect of physical wounding of PLB by bisection on PLB proliferation, shoot germination and necrosis rate of Tolumnia Snow Fairy after 6 weeks of culture.

<table>
<thead>
<tr>
<th>Physical wounding of PLB</th>
<th>PLBs proliferation (%)</th>
<th>Shoot germination (%)</th>
<th>Necrosis (%)</th>
<th>PLBs/explant</th>
<th>Shoots/explant</th>
</tr>
</thead>
<tbody>
<tr>
<td>Whole PLB</td>
<td>33.3 ± 6.7 a</td>
<td>33.3 ± 6.7 a</td>
<td>60.0 ± 11.5 a</td>
<td>3.5 a</td>
<td>0.7 a</td>
</tr>
<tr>
<td>Upper</td>
<td>40.0 ± 11.5 a</td>
<td>40.0 ± 11.5 a</td>
<td>60.0 ± 11.5 a</td>
<td>3.1 a</td>
<td>1.3 a</td>
</tr>
<tr>
<td>Lower</td>
<td>0.0 ± 0.0 b</td>
<td>0.0 ± 0.0 b</td>
<td>86.7 ± 6.7 a</td>
<td>0.0 a</td>
<td>0.0 a</td>
</tr>
</tbody>
</table>

*Three replicates each containing five test tubes (one explant per test tube) were performed for each treatment and cultured on MS basal medium modified for PLB proliferation.

*Percentage data were angular transformed before analysis and means ± SE within a column followed by the different letters are significantly different according least significant differences multiple range test at P ≤ 0.05.

PLB = protocorm like-body; MS = Murashige and Skoog.
Acclimation of in vitro–grown plantlets. After 3 rounds of subculture on plant regeneration medium, fully developed plantlets with well-developed roots and shoots, measuring on average 4 to 5 cm tall, were acclimatized for about 2 weeks on their tissue culture medium in a net house (Fig. 5C). Plants were then transferred to white plastic pots containing sphagnum moss. One year after transplantation, plants were transferred to pine bark and coconut husk chips in plastic commercial pots. These plants grew well (Fig. 5D), and transplants survived at rates of nearly 73%, for explants taken from 1- to 2-year-old plants and nearly 79%, for explants taken from inner leaf positions. Some mature plants flowered after 1 year (Fig. 5E).

In the method described here, Tolumnia orchid explants first formed PLBs during in vitro induction, followed by PLB proliferation and plant regeneration after transfer to PGR-free medium for proliferation and plant regeneration. Under natural conditions, the period from protocorm-like bodies induction to flowering of the Tolumnia Snow Fairy is 2 years. This protocol is easy to carry out, simple, and can provide large numbers of plants for mass propagation in a short time. Finally, all procedures may be conveniently applied for mass multiplication of Tolumnia orchids.

Literature Cited


