Static Liquid Culture of *Doritaenopsis* Seedlings

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Abstract. Methods for static liquid culture are described to improve the growth of *Doritaenopsis* (commercially known as *Phalaenopsis*) seedlings in vitro. The results showed that seeds not only germinated, but also grew faster in liquid medium. No hyperhydric seedlings were observed in liquid culture when liquid level was accurately controlled by culture density, medium volume, and sealing materials. Although the germination percent was unaffected by medium phase (liquid or solid), sowing density, medium volume, or sealing material, the growth of seedlings decreased as density increased or medium volume decreased. Seeds of 1.5 mg mixed with 20 mL of liquid medium per 9-cm petri dish sealed with two layers of parafilm prompted optimal results. Shoot growth also was enhanced while 75-day-old seedlings were subcultured in liquid media with or without support. Seedling growth was enhanced by adding 20 mL liquid media to 36 seedlings without support after 45 days of culture. It was expected that by static liquid culture, the period from sowing to ex vitro would be 1.5 months shorter than the traditional solid culture.

Phalaenopsis and Doritaenopsis, hybrids between *Phalaenopsis* and *Doris*, are collectively referred to as *Phalaenopsis*, the major floral crop in Taiwan for exporting. Asymbiotic culture on an agar-solidified medium is the main method for mass production. However, the high price of agar and the amount of time needed for preparing the medium result in increasing costs. Moreover, plantlets do not grow uniformly on solid medium and require added labor for grouping before subculture.

Seeds of epiphytic orchids, including *Phalaenopsis*, can germinate on a specific host tree in nature. However, the bark surface of the host tree plays an important role in determining whether seeds can germinate on it. Soft, spongy, water-filled bark with an uneven surface is more suitable than a smooth surface (Arndt, 1992). In addition, epiphytic orchids may also rely on mycorrhizal fungi (mycotrophy) as an additional surface of water (Yoder et al., 2000). In our laboratory, *Doritaenopsis* seeds germinated on the walls of the vessel where nutrient liquid remained and seeds germinated faster than those on a solid medium.

Given that in vitro liquid culture has been used successfully to cultivate a number of important nonorchids such as rhododendron (Douglas, 1984), conifers (Pâques et al., 1992), roses (Wong and Chu, 1995), and garlic (Kim et al., 2003), we explored the possibility that orchids, namely *Doritaenopsis*, could be cultured in this manner with optimal results. However, hyperhydricity (or vitrification) was a serious problem in long-term liquid culture (Ettienne and Berthouly, 2002; Ziv, 1994). The hyperhydric materials were unable to grow into normal plants, and it was hard to recover to a normal state (Zhou, 1995). In this study, we investigated raising normal *Doritaenopsis* seedlings from seeds in a static liquid medium.

Materials and Methods

Materials

Capsules of *Doritaenopsis* [(Dtps. Tinny Antique × Dtps. Sinica Peeress) × Dtps. Sogo Beach] were harvested 4.5 months after artificial hand-pollination. Capsules were surface-sterilized for 10 min with 2% sodium hypochlorite and then rinsed three times with sterilized deionized water. Seeds were collected and weighed under aseptic condition. Seeds were stained for 24 h with 0.5% triphenyl tetrazolium chloride at 25°C; seeds over 90% viability were used. The stock suspension was prepared by mixing 150 mg seeds and 100 mL autoclaved sowing medium containing Knudson C (Knudson, 1946) inorganic salts, 20 g L⁻¹ sucrose, and 34 g L⁻¹ potato paste. There were 500 ± 50 seeds per milligram.

Initial culture. Sterile, disposable, 9-cm plastic petri dishes (Alpha Plus Scientific Corp., Taoyuan, Taiwan) were used for liquid sowing, and, without stating otherwise, dishes were sealed with two layers of parafilm (American National Can Co., Menasha, WI). For the control, 1 mL of stock suspension was spread over 80 mL of solid medium containing 8 g L⁻¹ agar in a 500-mL flask. The flask was capped with a rubber stopper, which was 3 cm in height and had a 7.5-mm hole that was plugged with 0.1 g dried cotton. The juncture of the stopper and flask was sealed with six layers of parafilm.

Two weeks after sowing, germinated seeds were identified by enlarged embryos that had burst out of the testa. Two hundred seeds in each petri dish/flask were randomly chosen to determine the germination percent.

Effect of sowing density on the growth of seedling

To 1, 2, or 3 mL of stock seed suspension, 9, 8, or 7 mL of sowing medium was added, respectively. Thus, the total amount of medium was 10 mL in each petri dish and the amount of seed was 1.5, 3.0, or 4.5 mg per petri dish, respectively. The petri dish was sealed by two layers of parafilm. The diameters of 50 random seedlings were measured weekly from each dish under a stereo microscope (Nikon SMZ-10, Tokyo) for 5 weeks.

Anatomical observations

Fresh seedlings in liquid (sown as 1.5 mg seed per petri dish) or on solid medium 35 d after sowing were used for anatomical observations by cryosectioning without fixation and staining. Seedlings were put on a stainless-steel plate and mounted with water-soluble Tissue Freezing Medium (Leica Instruments GmbH, Heidelberg, Germany), and then put into Cryocut 1800 (Leica Instruments GmbH), which was precooled to −15°C to solidify the medium. A section of 45 μm was cut and transferred to a microscope slide. A few drops of reverse osmosis water was added to dissolve the medium at ambient temperature, and the tissue section was observed under a light microscope.

Effect of liquid medium volume on seedling growth.

The stock seed suspension of 1 mL was added to 4, 9, 14, 19, or 24 mL of sowing medium. Thus, with a common amount of 1.5 mg seeds per petri dish, the total amount of medium was 5, 10, 15, 20, or 25 mL, respectively. Three months after sowing, the growth of seedlings was investigated. A total of 150 seedlings in each dish/flask was randomly selected and separated into three groups. Seedlings in each group were classified into three growth stages: stage I: heart-shaped to torpedo-shaped embryo; stage II: first leaf 3 mm or less; and stage III: first leaf greater than 3 mm. The fresh weight of seedlings were recorded; also, dry weight was determined after drying in an oven at 60°C for 48 h.

Effect of liquid medium volume and sealing material on seedling growth.

One milliliter stock seed suspension was diluted to 10 or 20 mL of sowing medium. Then the petri dishes were sealed by two layers of parafilm or micropore tape (Micropore surgical tape; 3M Co., St. Paul, MN). The diameters of 50 randomly selected seedlings in each petri dish were recorded weekly for 8 weeks. The average rate of evaporational water loss from the petri dishes sealed with

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the two types of materials was presented as the weight loss per day.

Subculture. After sowing in liquid medium for 75 d, seedlings with a 1- to 2-mm leaf length were selected for subculture. In the subculture medium, the inorganic salts were replaced by one-fourth strength of MS (Murashige and Skoog, 1962) as well as replacing the petri dish with GA-7 vessel (length × width × height = 77 × 77 × 97 mm³) (Magenta Co., Chicago). In parts of the liquid culture, support of microporous membrane raft (Magenta Co.) (LR) or four layers of 5 × 5 cm² nonwoven fabrics (length × width × height = 5 × 5 × 1 cm³; Shin-Li Co., Banchiau, Taiwan) covered with one layer of paper towel (5 × 5 cm²; Kleenex; Kimberly-Clark Co., Taipei, Taiwan) (LN) was placed in the vessel. There was also solid medium in flasks as control. The flasks were capped with rubber stoppers as described previously. The growth of seedlings was investigated 45 and 75 d after subculture. The fresh weight of seedlings was recorded after 75 d of subculture and the dry weight was determined after drying in an oven at 60 °C for 48 h.

Pretransplanting. After 45 d of subculture, seedlings from the control and the best liquid culture during subculture (S and LO) in Table 2) were randomly chosen and transferred to a 650-mL flask containing 120 mL of solidified pretransplanting medium containing 3 g L⁻¹ Hyponex No. 1, 2 g L⁻¹ tryptone, 0.1 g L⁻¹ citric acid, 90 g L⁻¹ banana, and 15 g L⁻¹ sucrose for rooting. Twenty-five seedlings were transplanted to each flask. The flasks were capped with a rubber stopper as described previously. The number of leaf and root and length of the longest leaf and root of each plant were recorded after 120 d of culture.

Culture condition
Temperature in the culture room was maintained at 24 ± 2 °C. Photosynthetic photon flux at 35 ± 5 μmol m⁻² s⁻¹ for a 12-h photoperiod was supplied by a cool white fluorescent lamp (FL40D/38; China Electric Mfg. Co., Taipei, Taiwan).

Statistical analysis
The experiments were conducted in a complete randomized design and there were three replications in each treatment. The average values were analyzed by Costat (CoHort Software, Minneapolis, MN), and the differences were compared by analysis of variance and Duncan’s multiple range test at P ≤ 0.05.

Results
Two weeks after sowing, the germination percent ranged from 89.8% to 94.3%. It was not significantly affected by the types of medium or sowing density. However, seeds sown on solid medium germinated slower than those in liquid medium (Fig. 1). The growth of seedlings in liquid medium significantly decreased as sowing density increased. The growth curve of seedlings on solid culture was parallel to that of liquid culture 2 weeks after sowing at a density of 1.5 mg per vessel, but the seedling diameter of the former was much smaller than the latter (Fig. 1).

The growth and fresh weight of seedlings increased as the volume of liquid medium increased from 5 to 25 mL, and no treatment excised during the 3 months of culture. However, there was no difference in dry matter when medium volume exceeded 15 mL. The most uniform growth of seedlings was in the 20-mL treatment as 95% seedlings developed into stage II. Also, in this treatment, seedling fresh and dry weight was three and two times, respectively, of that in solid culture (Table 1).

Anatomical observation by cryosection showed that two kinds of cells—the condensed

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**Table 1. Effects of medium phase and support on the growth of seedlings after 45 d of subculture.**

| Medium phase | S (solid) | LO (liquid) |
|--------------|----------|------------|
| Support      | O (without) | R (raft)   |
| LO           | 90.2%    | 90.1%      |
| SLO          | 89.8%    | 90.1%      |
| LO           | 90.2%    | 90.1%      |
| SLO          | 89.8%    | 90.1%      |

**Table 2. Effects of medium phase and support on the growth of seedlings after 45 d of subculture.**

| Medium phase | S (solid) | LO (liquid) |
|--------------|----------|------------|
| Support      | O (without) | R (raft)   |
| LO           | 90.2%    | 90.1%      |
| SLO          | 89.8%    | 90.1%      |
| LO           | 90.2%    | 90.1%      |
| SLO          | 89.8%    | 90.1%      |

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small cells at the anterior and the swollen cells at the posterior—were observed in seedlings from either liquid or solid culture 35 d after sowing (Fig. 2). Differentiation of the first leaf, second leaf, and rhizoid was observed in both seedlings. Root initiation was observed in the seedling from liquid culture only.

When dishes were sealed with Micropore tape, the evaporational water loss rate was 420 mg·d⁻¹. The medium dried up and seedlings stopped growing after 3 or 6 weeks when 10 or 20 mL medium was supplied, respectively (Fig. 3). When sealed with parafilm, water loss was reduced to 14.8 mg·d⁻¹, and seedlings were of similar sizes after 6 weeks regardless of 10 or 20 mL medium was used. However, the growth slowed down after 6 weeks when only 10 mL medium was supplied. The growth rates were similar. The largest seedlings after 6 weeks were those in 20 mL of medium and sealed with parafilm.

Leaf number and length were enhanced by liquid culture whether or not a support was used (Table 2). Furthermore, most (76% to 85%) seedlings cultured in liquid medium without support (LO₁₀ or LO₂₀) developed two leaves, but 28% or fewer seedlings developed the second leaf in liquid culture with support (LR or LN). In addition, the leaf length in LO₁₀ and LO₂₀ was longer than in LR and LN. Seedlings that grew on non-woven cloth (LN) had the most roots (Table 2).

After 75 d, only seedlings in liquid culture without support (LO₁₀ and LO₂₀) developed the third leaf. Seedlings in LO₂₀ treatment had the most leaves and largest plant fresh and dry weight. Plants in liquid culture with supports (LR and LN) had similar root length. Fresh weight and dry weight of LO₂₀ were 3.3 or 7.8 times that of solid culture (S), respectively. Although seedlings in LO₂₀ developed the third leaf, their leaves were much shorter than those in LO₁₀ and their roots were the shortest and turned brown after 75 d of subculture (Table 3).

After subculturing, the seedlings cultured in LO₁₀ or on solid medium for 45 d to the solidified pretransplanting medium, the difference of growth between the two seedling sources increased as time progressed (Fig. 4). There were no differences detected in root number and the longest root length between seedlings from S and liquid culture without support (LO₁₀). However, the latter had more and longer leaves (Table 4).

**Discussion**

It had been reported that seeds of both epiphytic and terrestrial orchids had higher water content and germinated faster in symbiotic culture than that cultured asymbiotically on agar-solidified medium, because mycorrhizal fungi may facilitate the absorption of free water to enhance seed germination (Yoder et al., 2000). This was supported by our results that *Doritaenopsis* seeds in liquid medium germinated faster than that on solid medium within 2 weeks (Fig. 1). However, germination percents were not different between the liquid and solid media.

The epidermal cells of *Phalaenopsis* protocorm are capable of absorbing nutrients (Lee, 1990). Although the volume of the liquid medium was much less than the solid culture in this study, seedlings in liquid culture had greater fresh and dry weight (Fig. 1; Table 1), indicating that the growth of seedlings may be enhanced by increasing the surface of seeds in contact with free liquid. Orchid embryos form protocorms on germination, which accumulate carbohydrates and serve as a storage organ for seedling growth (Edwin, 1993). In our study, although the growth of seedlings stopped as the liquid medium dried, there was no significant change in seedling size (Fig. 3) and no dehydrated seedlings were observed during the drying period. Perhaps the nutrients and water stored in protocorms made it possible for seedlings to survive during the drying period for as long as 5 weeks.

Damaged meristem had been indicated as a typical characteristic of hyperhydric status and the promeristem zone comprising central mother cells had been reported absent in hyperhydric buds (Pâques and Boxus, 1987). It was reported that mature seeds of *Phalaenopsis* are composed of two kinds of cells: the small cells at the upper region and the big cells at the bottom (Edwin, 1993). The former differentiate into meristem and the latter serve as a storage organ, which is exhausted 30 d after sowing (Lee, 1990). Although the volume of the liquid medium dried, there was no significant change in seedling size (Fig. 3) and no dehydrated seedlings were observed during the drying period. Perhaps the nutrients and water stored in protocorms made it possible for seedlings to survive during the drying period for as long as 5 weeks.

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![Fig. 2. Longitudinal sections of seedlings 35 d after sowing. The first leaf (FL), second leaf (SL) and rhizoids (Rh) differentiated from seedlings in liquid medium (A) or solid medium (B). Root (R) was evident on seedling from liquid culture. Bar = 0.1 mm.](image2)

![Fig. 3. Effect of medium volume and sealing material on the growth of seedling. The bars represent SEs, n = 50. ‘P’ or ‘M’ represents parafilm or Micropore ‘M’ tape, respectively.](image3)
no hyperhydric seedlings occurred in liquid culture.

The desiccation or maturation drying of most seeds is the terminal phase of seed development leading to a metabolic quiescence state. Subsequent rehydration of mature dry seeds leads to germination, and hence the loss and gain of water play a role in the transition from a developmental program to a germination program (Dasgupta and Bewley, 1982). By desiccation, the zygotic embryo axis of *Ricinus communis* switched into the germination mode from the development mode by destroying certain mRNA and synthesizing new mRNA (Kermde and Bewley, 1985). Dramatic desiccation of vitreous protocorm-like bodies (PLBs) of *Durwinana* and *Brassomatella* enhanced the number of regenerated plantlets. It was proposed that desiccation of PLBs must have induced a change in direction from enlargement and micropropagation of PLBs to develop leaves and roots (Kishi and Kakagi, 1997). Thus, the control of the liquid level for the desiccation of seedling turning from protocorm enlargement to meristem differentiation was important in liquid sowing. In our previous study, when *Phalaenopsis* seeds were sown on a support of nonwoven fabrics or a microporous membrane raft with 40 mL liquid medium in a GA-7 vessel, 20% or 30% of seedlings were hyperhydrated, respectively (Tsai and Chu, 2004). In this study, 20 mL of medium was supplied and the petri dish was sealed with parafilm, medium desiccation resulted in seedlings protruding above the liquid level 7 weeks after sowing. Subsequently, 98.3% of seedlings developed normal leaves 3 months after sowing (Table 1).

Modification of medium had been used to control hyperhydricity such as increasing the concentration of gelling agent (Debergh, 1983; Marga et al., 1997; Pasqualetto et al., 2001). Thus, the control of the liquid level with medium volume and sealing materials shown in this study.

In conclusion, static sowing of 1.5 mg *Doritaenopsis* seeds in 20 mL of liquid medium per 9-cm petri dish sealed with parafilm for 75 d and subculturing 36 seedlings in 20 mL liquid medium without support are suggested for the fast and uniform growth of seedlings. It was expected that by liquid culture, the period from seed sowing to the ex vitro could be 1.5 months shorter than the traditional solid culture.

### Table 3. Effects of medium phase and supports on the growth of seedlings after 75 d of subculture.

| Subculture | leaf | root |
|------------|------|------|
| S<sup>100</sup> | 5.3 b | 3.7 a |
| LO<sub>36</sub> | 5.9 b | 4.2 a |
| LO<sub>20</sub> | 5.9 b | 4.2 a |

### Table 4. Effects of culture method during subculture on the growth of seedlings after 120 d of pretransplanting.

| Subculture | leaf | root |
|------------|------|------|
| S<sup>100</sup> | 3.8 b | 4.5 a |
| LO<sub>20</sub> | 3.7 a | 4.5 a |

That no hyperhydric seedlings occurred in liquid culture.

3 months (Table 1). After subculturing 75-d-old seedlings in liquid medium for 45 d, shoot growth was enhanced regardless of whether supports were used (Table 2), confirming that no hyperhydric seedlings occurred in liquid culture.

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