Growth and Photosynthetic Capability of Momordica grosvenori Plantlets Grown Photoautotrophically in Response to Light Intensity

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Abstract. Momordica grosvenori plantlets were cultured in vitro for 26 d on sucrose- and hormone-free Murashige and Skoog (MS) medium with four levels of photosynthetic photon flux density (PPFD), namely 25, 50, 100, or 200 μmol m−2 s−1, and a CO2 concentration of 1000 μmol mol−1 in the culture room [i.e., photoautotrophic micropropagation (PA) treatments]. The control treatment was a photomixotrophic culture using MS medium containing sucrose and NAA with a CO2 concentration of 400 μmol mol−1 in the culture room and a PPFD of 25 μmol m−2 s−1. Based on the results, a second experiment was conducted to investigate the effects of α-naphthaleneacetic acid (NAA) and sucrose on callus formation. For this, plantlets were grown in the absence and presence of either NAA or sucrose. Compared with the control, the PA plantlet had a well-developed rooting system, better shoot, greater chlorophyll content, and higher electron transport rate and the ex vitro survival percentage was increased by 31%. Both sucrose and NAA stimulated callus formation on the shoot bases of control plantlets, whereas calluses did not form on the plantlets grown in sucrose- and hormone-free medium. The stronger light intensities increased the fresh and dry weight of plantlets. A PPFD of 100 μmol m−2 s−1 was more suitable for the growth of M. grosvenori plantlets. Therefore, photoautotrophic plantlets grown at high light intensities would be better suited to the intense irradiance found in sunlight.

Momordica grosvenori Swingle or Lo Han Kuo (Chinese) is a perennial vine native to southern China in Guangxi province. The fruit has been used in traditional Chinese medicine to treat colds, coughs, sore throats, and gastrointestinal disorders. The fruit contains a natural sweetener called Mogroside V, which is 300 times sweeter than cane sugar and extremely low in calories (Cheng, 1987). Mogroside V has the potential to be more than just a beverage sweetener. Takasaki et al. (2003) and Yao et al. (2008) reported that Mogroside V suppressed carcinogenesis in mice and reduces oxidative stress in the body.

Momordica grosvenori propagated by layering, grafting, and dividing of tuberous roots often have low multiplication rates and cause a variety of degeneration. These methods also result in low yields of fruits. Variety degeneration has been altered by using tissue-cultured plantlets. These plantlets are conventionally propagated on sugar-containing medium with growth regulators, especially α-naphthaleneacetic acid (NAA) added to the rooting medium. Tissue-cultured plantlets use the sugar in the medium as a carbon source, has several advantages over photomixotrophic micropropagation (PM). The advantages include minimal microbial contamination, increased photosynthesis, growth and rooting in vitro, and survival percentages ex vitro when the in vitro environment is controlled to maximize photosynthesis (Aitken-Christie et al., 1995; Kozai, 1991). High photosynthetic photon flux density (PPFD) and a high CO2 concentration in the culture vessel significantly enhance photosynthesis of plantlets in vitro (Fujiiwara et al., 1988; Kubota and Kozai, 2001). A high CO2 level in the vessel can be achieved by increasing CO2 concentration in the culture room and the number of air exchanges in the vessel. The latter, defined as the hourly vessel ventilation rate divided by the air volume of the vessel (Kozai et al., 1986), can be increased or decreased by changing the gas-permeable filter disks attached to the holes in the lid and walls of the vessel. In addition, photoautotrophic conditions would promote growth and rooting of the plantlets without growth regulators in the medium (Nguyen et al., 1999; Xiao and Kozai, 2004).

Practitioners of photoautotrophic culture have investigated the effects of light intensity on the development of various plant species. These include in vitro plantlets of Solanum tuberosum (Kitaya et al., 1995), Cymbidium (Kozai et al., 1990), and Gardenia jasminoides (Serret et al., 1996). These reports showed that light intensity affected the photosynthetic characteristics of plantlets in vitro. However, there are no reports about the effects of light intensity on growth and photosynthetic capability of M. grosvenori plantlets. Various plant species in nature show a number of acclimatization responses to light intensity (Anderson et al., 1995). For example, light intensity changes the number of chloroplasts and leaf thickness (Bailey et al., 2001) and chlorophyll content (Hirashima et al., 2006; Lindahl et al., 1995). The responses of photoautotrophic plantlets in vitro to high levels of irradiation may be similar to plants in a natural environment.

The objectives of this study were to: 1) produce M. grosvenori plantlets of good quality and high survival percentages by comparing plantlets grown under photomixotropic and photoautotrophic conditions; and 2) investigate the effects of light intensity on growth, photosynthetic ability, and light acclimation of the photoautotrophic plantlets.

Materials and Methods

Plant material, treatments, and culture conditions. Single shoots, each with two unfolded leaves excised from in vitro cultures of M. grosvenori, were used as explants and cultured for 26 d. The average leaf area, fresh

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and dry weight, and stem length per shoot was 602 ± 47 mm², 98.2 ± 11.3 mg, 8.4 ± 1.4 mg, and 2.1 ± 0.3 cm, respectively. These in vitro cultures of *M. grosvenori* had been cultured on hormone-free Murashige and Skoog (MS) (1962) medium containing 30 g·L⁻¹ sucrose for three subcultures (5 weeks for each subculture) to remove the exogenous hormone accumulation before experiments. Four explants were cultured in each Magenta-type vessel (370 mL; Verde Co., Ltd., Toyohashi, Japan) containing 70 mL MS medium with the pH adjusted to 5.8 before autoclaving. Gelrite™ (Sigma G1910; Sigma, St. Louis, MO) was used as the gelling agent (3.2 g·L⁻¹) because it was more transparent than agar and allowed visualization of root formation.

**Expt. 1.** The first experiment included four PA treatments, L25, L50, L100, and L150, in which L indicates light intensity and subscripts denote PPDF, and one PM treatment as a control at 25 μmol·m⁻²·s⁻¹. Light intensity was measured on the surface of the empty shelves with a light meter (Model LI-250A; LI-COR, Lincoln, NE). The PA explants were cultured on hormone- and sucrose-free MS medium, and CO₂ concentration was maintained at 1000 μmol·mol⁻¹ in the culture room. The CO₂ concentrations were measured and controlled by a CO₂ controller with a nondispersive infrared detector CO₂ sensor (T6004; Telaire Co., Ltd., Shoreview, MN). The number of air exchanges of the culture vessels measured by the method of Kozai et al. (1986) were increased with culture time by increasing the area of gas-permeable filter disks (10 mm in diameter, pore size 0.5 mm; Milli-Seal, Nihon Millipore, Ltd., Tokyo, Japan) attached to the holes in the lid and walls of the vessel. To get a feasible ventilation rate for photosynthesis, the number of air exchanges of the culture vessels were 0.2 h⁻¹ from Days 0 to 3, 1.8 h⁻¹ from Days 4 to 7, 2.7 h⁻¹ from Days 8 to 17, and 3.8 h⁻¹ from Days 18 to 25.

The PM or control explants were cultured on sucrose-containing (30 g·L⁻¹) MS medium with 2.7 μM NAA at a CO₂ concentration of 400 μmol·mol⁻¹ in the culture room and a number of air exchanges of 0.2 h⁻¹. The control treatment was very similar to conventional micropropagation of *M. grosvenori* plantlets.

The air temperature and relative humidity in the culture rooms were maintained at 25 ± 1 °C and 80% ± 5%, respectively. The photoperiod was 12 h per day supplied with cool-white fluorescent lamps. Twelve replications (12 vessels) were used for each treatment. The experiment was conducted twice.

**Expt. 2.** Based on the results of the first experiment, an experiment was conducted to investigate the effects of sucrose and NAA on callus formation. This experiment included two PM and two PA treatments at 25 μmol·m⁻²·s⁻¹ light. The PM plantlets were cultured on sucrose-containing (30 g·L⁻¹) medium with 2.7 μM NAA (PMN treatment) or without NAA (PM treatment). The PA plantlets were cultured on sucrose-free medium with 2.7 μM NAA (PAN treatment) or without NAA (PA treatment). The plant material, culture conditions, and treatment replications were the same as the first experiment for the PM and PA treatments, respectively. The experiment was conducted once. Chlorophyll content, the ratio of variable to maximum chlorophyll fluorescence (Fv/Fm), and electron transport rate (ETR) were not measured in the second experiment.

**Acclimation ex vitro and survival percentage.** After 26 d of in vitro culture, plantlet growth was determined in half of the plantlets from each treatment of both experiments. The remaining plantlets were taken out from the vessels and washed to remove any medium adhering to the root system. The washed plantlets were transplanted into a tray with peat (Growing Mix I; Fafard Co. Ltd., Inkerman, Canada) and placed in a greenhouse with a simple shading screen for acclimatization. During acclimatization, relative humidity in the first week was kept at 80% to 90% with an automatic sprayer and then decreased gradually to 70% to 70% in the second week and 70% to 60% for the next days. The average air temperature was 22 °C day and 17 °C night. Twenty days after transplanting, the dead plantlets were counted in each treatment, and the survival percentages were determined.

**Measurements, calculation, and statistical analysis.** Rooting percentage was recorded every day until Day 16. Number of roots, stem and root length, fresh and dry weight of root and shoot, and leaf area were measured on Day 26. Leaf area was determined using image and quantify analysis software (LIA32 for Windows 95; K. Yamamoto, Nagoya, Japan) with an image scanner. Dry weight of the plantlets was determined after drying at 80 °C for 72 h.

Chlorophyll content was measured after cutting pieces of the second and third fully expanded leaves (counted down from apex), soaking in 5 mL N, N-dimethylformamide solution, and shaking once every 10 min for 2 h at 4 °C (Moran and Porath, 1980). Absorbance was measured with an ultraviolet-visible spectrophotometer (Helios Gamma; Thermo Spectronic Co., Ltd., Cambridge, U.K.) at wavelengths of 664 nm, 647 nm, and 603 nm. The chlorophyll content, including chlorophyll a and chlorophyll b, was determined on a fresh weight basis (μg·g⁻¹) and calculated using the formula of Moran (1982).

The Fv/Fm and ETR were measured and calculated with a Maxi-Imaging-PAM chlorophyll fluorescence measuring system (Walz, Effeltrich, Germany). After 24 d of culture, the second and third fully expanded leaves (counted down from apex) were used for the measurement. Six replications or six leaves were measured in each treatment. For chlorophyll fluorescence measurement, the plantlets were placed in a dark room to adapt for 15 min. Next, randomly chosen intact leaves from five treatments were placed in the measuring head (10 × 13 cm) to measure the Fv/Fm. The leaves without dark adaptation were used to evaluate ETRs. Each ETR value was obtained using actinic irradiance for 10 s. All relevant fluorescence parameters, including actinic irradiance and leaf temperature, were recorded and ETRs were calculated in the Maxi-Imaging-PAM system.

Statistical significance was determined using Sigma Stat (Systat Software Inc., Chicago, IL). Differences between means were assessed with the Holm-Sidak test and a significance level of P ≤ 0.01.

**Results**

**Expt. 1.**

*Root growth.* The pattern of the root system in control and PA treatments differed significantly. Large callus mass formed on the shoot bases of control plantlets, whereas callus was not found on the shoot bases of PA plantlets (Fig. 1). In the control treatment, callus formation was observed on the shoot base on Day 3, callus volume then enlarged, and the roots emerged on Day 8. In contrast, roots in all PA treatments were formed directly from plantlets without an intermediate callus phase. The control plantlets had the highest number of main adventitious roots (Table 1) but were accompanied by large masses of callus and produced sparse lateral branching. The roots of control plantlets were firmly encased in callus and therefore difficult to separate. Accordingly, we determined the fresh and dry weights of roots that were outside of callus. On the other hand, the main adventitious root of the PA plantlet gave rise to a denser growth of fine lateral roots. This was especially true where the L100 treatment for the length and dry weights of the roots were 6.5 and 1.6 times greater than those in the control treatment, respectively.

Light intensity affected root formation. Rooting percentage reached 100% for all plantlets by Day 16 (Fig. 2). However, rooting occurred first in the L25 treatment on Day 7 and last in the L200 treatment on Day 10. Rooting percentage reached 100% in the L25 treatment on Day 11, which was 5 d earlier than the L100 treatment. Root growth was greatest in the L100 treatment, in which the fresh weight of roots was 3.4 times greater than that in the L200 treatment, and the dry weight was 2.5 times greater than that in the L25 treatment (Table 1). Stronger light intensities also increased the ratio of root dry weight to root fresh weight (DW/FW) in PA-treated plantlets.

**Shoot growth.** PA treatments improved shoot growth over the control treatment (Table 2; Fig. 1). Leaf area, stem length, and fresh and dry weight of shoots were greatest in the L100 treatment, which were 2.7, 1.2, 3.0, and 3.7 times greater, respectively, than those of control plantlets, but there was no significant difference in dry weight between the L200 and L100 treatments. Leaf area as well as fresh and dry weights of shoots were, respectively, 2.0, 1.8, and 1.8 times higher in PA-treated plantlets compared with the control plantlets grown under the same low light intensity (25 μmol·m⁻²·s⁻¹).
Among all PA treatments, growth was improved as the light intensity increased from L25 to L100, but the L200 treatment had the lowest leaf area and stem length (but not lowest dry weight). Stronger light intensities also increased shoot DW/FW and leaf thickness and produced longer and denser epidermal hairs in all PA-treated plantlets. The branch, thickset stem, and small, thick, and yellow leaves of the plantlets were observed in the L200 treatment (Fig. 1).

**Chlorophyll content and chlorophyll a/b ratio.** The chlorophyll content of plantlets was significantly influenced by the different culture methods (PA or PM) (Fig. 3). Among all the treatments, the plantlet chlorophyll content was greatest in the L50 treatment, being 2.1 times higher than that of the control. PA-treated plantlets (L15) treatment had 1.6 times more chlorophyll content than did the control even at the same low light intensity (25 μmol·m⁻²·s⁻¹).

Light intensity also had a significant influence on chlorophyll content. The chlorophyll content increased to a maximum as the light intensified to 50 μmol·m⁻²·s⁻¹ and then decreased to a minimum at 200 μmol·m⁻²·s⁻¹, creating a parabola (Fig. 3). In contrast, the ratio of chlorophyll a to chlorophyll b increased consistently as the light intensified.

**Ratio of variable to maximum chlorophyll fluorescence ratio and electron transport rate.** The Fv/Fm ratio, reflecting the maximal photochemical efficiency of photosystem II (PSII), is usually steady with fluctuations from 0.75 to 0.85 for nonstressed plants (Bolhar-Nordenkampf et al., 1989). In the present study, the Fv/Fm (determined using the second or third leaf from the apex) ranged from 0.78 to 0.82 except for the L200, which had a ratio of 0.57 in the second leaf (data not shown). This suggests that photoinhibition had probably occurred in the second leaf taken from the L200 treatment, and consequently, it had the lowest maximum ETR (Fig. 4A).

ETRs allow for the rapid and noninvasive assessment of the light response of PSII and are used to study the photosynthetic performance and photoacclimation of photosynthetic organisms (Serôdio et al., 2006). The maximum ETR was related to the maximum photosynthetic capability, which was obtained when the rate of photosynthesis was limited by the activity of the electron transport chain or Calvin cycle enzymes (Behrenfeld et al., 2004; Ralph et al., 2005). Except for the second leaf from the L200 treatment, all the ETRs in the PA plantlets were higher than those of the control plantlets, and greater light intensity increased the maximum ETRs of the PA plantlets (Fig. 4).

**Survival ex vitro.** In the present study, survival percentage was 65% lowest in the control treatment and 100% highest in the L100 and L200 treatments (Table 2). The control plantlets, especially the smaller ones, wilted immediately when placed in the greenhouse. Approximately 85% of the wilted plantlets recovered, but some died in the middle of acclimatization as a result of root decay. In contrast, the PA plantlets wilted very little during the first day of acclimatization, recovered the next morning, and had over 95% survival on Day 20.

**Expt. 2**

**Effects of sucrose and α-naphthaleneacetic acid on callus formation.** Calluses only formed on the shoot base of PM plantlet. The difference between the PM and PA treatments was that the PM plantlets used sucrose as a carbon source with 2.7 μM NAA in the medium, but the PA plantlets used CO₂ as a carbon source without NAA in the medium.

The results from the second experiment showed that both sucrose and NAA stimulated callus formation (Table 3; Fig. 5). The callus texture was more compact in the PMN

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Table 1. Effects of four levels of light intensity on rooting percentage and root growth of *Momordica grosvenori* per plantlet cultured on sucrose-free medium without NAA for 26 d.

| Treatment | Rooting (%) | Number of roots | Root length (cm) | Root wt (mg) |
|-----------|-------------|----------------|-----------------|-------------|
|           |             |                | Fresh           | Dry         | DW/FW (%)  |
| Control   | 100         | 22.8 ± 4.6 a   | 1.2 ± 0.3 c     | 148.6 ± 19.2 b | 5.2 ± 0.6 b | 3.5   |
| L25       | 100         | 2.5 ± 0.3 c    | 6.7 ± 0.6 a     | 81.8 ± 7.9 c | 3.3 ± 0.4 c | 4.0   |
| L50       | 100         | 4.1 ± 0.4 bc   | 7.8 ± 0.5 a     | 138.5 ± 13.1 b | 5.5 ± 0.6 b | 4.1   |
| L100      | 100         | 4.4 ± 0.4 b    | 3.3 ± 0.3 b     | 187.3 ± 12.3 a | 8.3 ± 0.7 a | 4.4   |
| L200      | 100         | 5.0 ± 0.6 b    | 55.0 ± 6.8 c    | 4.3 ± 0.5 bc | 12.3      |

Data are mean ± se, and treatments denoted by the same letter in a column were not significantly different (P ≤ 0.01).

DW/FW = dry weight/fresh weight.
Table 2. Effects of four levels of light intensity on shoot growth of *Momordica grosvenori* per plantlet cultured on sucrose-free medium without NAA for 26 d and survival percentage ex vitro on Day 20.

| Treatment | Leaf area (mm²) | Stem length (cm) | Fresh Shoot wt (mg) | Dry Shoot wt (mg) | DW/FW (%) | Survival (%) |
|-----------|----------------|-----------------|---------------------|------------------|-----------|--------------|
| Control   | 1,412 ± 76 c   | 4.0 ± 0.2 ab    | 281.0 ± 15.1 c      | 26.2 ± 2.5 d     | 9.3       | 65           |
| L₂₂₅      | 2,802 ± 228 b  | 4.0 ± 0.2 ab    | 499.4 ± 27.0 b      | 48.3 ± 2.8 c     | 9.7       | 96           |
| L₅₀       | 3,650 ± 232 a  | 4.5 ± 0.2 a     | 751.0 ± 53.5 a      | 80.1 ± 6.9 b     | 10.6      | 98           |
| L₁₀₀      | 3,740 ± 270 a  | 4.7 ± 0.2 a     | 846.8 ± 49.0 a      | 97.1 ± 5.6 a     | 11.5      | 100          |
| L₂₀₀      | 2,198 ± 121 b  | 3.6 ± 0.2 b     | 612.9 ± 34.7 b      | 100.7 ± 4.1 a    | 16.4      | 100          |

*Data are mean ± SE, and treatments denoted by the same letter in a column were not significantly different (P ≤ 0.01). DW/FW = dry weight/fresh weight.*

Discussion

Rooting and callus formation. The plantlets cultured on sucrose- and NAA-containing medium produced more short roots but had large callus masses at the shoot bases. The results of the second experiment suggest that NAA and, a lesser extent, sucrose caused callus formation and influenced the number of roots. NAA with the presence or absence of sucrose in the medium stimulated callus formation and produced more adventitious roots with less lateral branching. The same was seen for plantlets grown in medium containing sucrose with or without NAA. Plantlets grown on sucrose- and NAA-free medium had no callus and produced more vigorous adventitious roots with well-developed lateral branching. Clearly, sucrose and NAA were unnecessary for root growth of *M. grosvenori* under photoautotrophic conditions. We therefore suggest that photoautotrophically grown plantlets synthesize appropriate endogenous growth-promoting hormones. Xiao and Kozai (2004) argued that endogenous phytohormones such as auxin, which are necessary for rooting and protein synthesis, are produced more by plantlets grown photoautotrophically than by plantlets grown photomixotrophically.

Light intensity influenced growth and development of the roots. The stronger the light intensity, the later the adventitious roots formed in all PA treatments (Fig. 1). Yellow or wilted leaves were observed in higher light-treated plantlets at the beginning of the culture period. The higher light intensities probably increased the temperature inside the vessel, which promoted transpiration in the plantlets and caused water loss from the leaves. Thus, the higher light-treated explants needed a relatively longer period to recover, resulting in delay of root formation. In addition, root formation in plantlets is dependent on the auxin concentration. Light intensity probably affected synthesis and transport of indole-3-acetic acid (IAA) within the plantlets, which consequently influenced root formation. However, the relationship between light intensity and auxin synthesis is not known and additional study is justified.

Photosynthesis and chlorophyll content. The slow growth of plantlets has frequently...
Table 3. Effects of sucrose and NAA on growth of *Momordica grosvenori* plantlets on Day 26 and survival percentage ex vitro on Day 20.

| Treatment | Callus | Root | Shoot |
|-----------|--------|------|-------|
|           | Volume (mm³) | DW (mg) | Number | Length (cm) | DW (mg) | DW (mg) | Survival ex vitro (%) |
| PMN** | 1,300 ± 200 c | 40.2 ± 5.2 a | 20.6 ± 5.6 a | 1.2 ± 0.5 c | 5.7 ± 5.1 a | 21.2 ± 3 b | 67 |
| PM | 140 ± 60 c | 5.1 ± 0.3 c | 2.8 ± 1.2 c | 3.3 ± 0.8 b | 0.9 ± 0.9 c | 24.3 ± 2 b | 75 |
| PAN | 360 ± 80 b | 16.3 ± 2.8 b | 8.4 ± 3.2 b | 0.3 ± 0.1 d | 1.4 ± 3.2 c | 41.1 ± 6 a | 87 |
| PA | 0 d | 0 d | 3.6 ± 0.8 c | 6.8 ± 0.9 a | 4.2 ± 0.4 b | 48.7 ± 4 a | 100 |

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NAA ** ** ** ** NS NS

Sucrose ++ ++ ++ ** NS NS

NAA × sucrose ++ ++ ++ ** ++ NS

**PMN and PM were the photomixotrophic treatments using sucrose-containing medium with 2.7 μM α-naphthaleneacetic acid (NAA) and without NAA, respectively. PAN and PA were the photoautotrophic cultures using sucrose-free medium with 2.7 μM NAA and without NAA, respectively.

Data are mean ± se, and treatments denoted by the same letter in a column were not significantly different (P ≤ 0.01).

Analysis of variance was applied to two levels of NAA and sucrose concentrations, respectively.

*dw* = dry weight; NS = nonsignificant.

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been explained by low photosynthetic ability. In the present study, the photosynthetic ability and the growth rate in the control treatment were much lower than those of the PA treatments. The lowest photosynthesis and growth in the control treatment was probably because of a combination of sucrose-containing medium, low light intensity, and infrequent air exchanges in the vessel. Generally, sugar in the culture medium suppresses photosynthesis of plantlets in vitro (Desjardins et al., 1995). Plantlets cultured in sugar-containing medium depend more on the sucrose and less on CO₂ in the air (Kubota 2001; Koniger et al., 2008). In the present study, the chlorophyll content of plants plays an important role in the absorption of light during photosynthesis. In the present study, the presence or absence of sucrose in the culture medium significantly influenced chlorophyll content of the plantlets. When the plantlets were grown under the same light intensity (25 μmol·m⁻²·s⁻¹), the presence of sucrose in the medium reduced the chlorophyll content and lowered the photosynthetic ability in the control treatment (Fig. 3). In contrast, sucrose-free cultures produced more chlorophyll and increased the photosynthetic ability in L₁₂⁵ treatment. This demonstrates that PA could improve photosynthetic ability of plantlets by stimulating chlorophyll synthesis and thus in turn improves adaptation to low light conditions.

The light intensity also significantly affected the chlorophyll content in the PA plantlets. The chlorophyll content increased as the light intensity rose from 25 to 50 μmol·m⁻²·s⁻¹ but decreased at higher intensities, 50 to 200 μmol·m⁻²·s⁻¹. This suggested that the plants could balance light absorption and translation by regulating chlorophyll synthesis (Bailey et al., 2001). The change of chlorophyll content in our study was possibly an acclimation of the plantlets to different light intensity.

The chlorophyll content of plants responds to light intensity by changing chlorophyll a/b and adjusting to the ETR. As shown in Figures 3 and 4, an increase in chlorophyll a/b accompanied an increase in the ETR, except when the second leaf from the L₂₀⁰ treatment was sampled (Fig. 4A). An increase in chlorophyll a/b was associated with a decrease in the size of the PSII light-harvesting antenna (Bailey et al., 2001; Leong and Anderson, 1984; Masuda et al., 2002). Moreover, a faster ETR was associated with an increase in the reaction-center content of PSII (Evans, 1987; Leong and Anderson, 1984). Bailey et al. (2001) and Königer et al. (2008) also found changes in the numbers of chloroplasts and leaf thickness associated with changes in chlorophyll content and photosynthetic capacity in *Arabidopsis thaliana* grown under different light intensities.

Photoinhibition may occur when light intensity exceeds what is required for the saturation of photosynthesis. This adverse phenomenon can manifest as a loss of PSII activity (Masuda et al., 2002). This could account for the lower capacity of ETR as shown in the second leaf from the L₂₀⁰ treatment in this study (Fig. 4A). In photoinhibition, the chloroplasts contained much of the same amount of PSII as they did under low light. However, up to 80% of PSII was photochemically inactive because of photodamage (Vasilikiotis and Melis, 1994). During the growth of plantlets, the upper leaves may create a dense canopy that protected the lower ones from photodamage. Thus, photodamage of PSII reaction centers could be repaired and chlorophyll antennae enlarged (Masuda et al., 2002; Neidhardt et al., 1998; Webb and Melis, 1995). Perhaps this is why, in the L₂₀⁰ plantlets, the second and third leaves from the apex had the lowest and highest ETRs, respectively (Fig. 4). The third leaves had the highest ETRs, which contributed to the L₂₀⁰ plantlets having the highest dry weight. The plantlet growth in the L₂₀⁰ treatment, however, was not as good as that in the L₁₀⁰ treatment because of the stunted plantlets with aging stems and small and yellow leaves. The formation of lateral branches in the L₂₀⁰ treatment may have been caused by high light intensity, which could have inhibited the synthesis of IAA in the shoot apex and apical dominance.
Plantlet quality and survival percentage. The intrinsic quality of plants produced in vitro was one of the key factors governing the survival percentage during acclimatization to greenhouse or field conditions (Afreen-Zobayed et al., 1999). The results of our study indicated that micropropagation methods affected plantlet quality. The PA-grown plantlets were of good quality and had a high survival percentage, whereas those grown by PM were of poor quality and few survived transplantation. The effectiveness of a micropropagation system can be measured by the survival percentage of plantlets successfully transferred from culture vessels to a greenhouse or field (Kirdmanee et al., 1995; Xiao and Kozai, 2006). The problems associated with conventional micropropagation of M. grosvenori may be primarily a result of the poor-quality plantlets in vitro.

Poor-quality plantlets influence survival and growth in many ways. First, large callus masses in the root system can reduce the uptake or absorption of water and nutrients into the plantlet and cause root decay, resulting in a low survival percentage. Nguyen et al. (1999) suggested that callus at the shoot base of coffee plantlets grown by the conventional method may affect the survival percentage of plantlets when transferred from in vitro to ex vitro. Second, poor plantlets usually grow slowly during ex vitro acclimatization. An extended early period of acclimatization shortens growth in the middle and later stages. This not only contributes to nonbearing or low yields of fruits, but also to little dry matter accumulation. Third, the improper use of plant growth regulators also causes poor growth and nonbearing or low yields of fruits. Fourth, plantlets grown photomixotrophically usually have low photosynthetic ability, and this causes the low survival and growth rate in the acclimatization stage (Grout, 1978). The finding that the enhanced net photosynthetic rate of eucalyptus plantlets in vitro increased the survival percentage ex vitro supports this conclusion (Kirdmanee et al., 1995).

Conclusion

In comparison with the PM plantlets grown on sucrose- and hormone-containing medium, the PA plantlets grown on sucrose- and hormone-free medium had a more developed rooting system, better shoots, higher chlorophyll content, greater photosynthetic capability, and high survival ex vitro. Therefore, PA is an alternative way to produce high-quality M. grosvenori plantlets. A PPFD of 100 μmol·m⁻²·s⁻¹ was more suitable for growth of M. grosvenori plantlets. The callus-on-plantlet base reduced viability ex vitro. Callus formation was not only caused by NAA, but also sucrose. Plantlets grown on sucrose- and hormone-free medium had no callus on the roots.

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