In Vitro Shoot Proliferation and Root Induction of Shoot Tip Explants from Mature Male Plants of *Casuarina cunninghamiana* Miq.

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Abstract. This study examined the effects of plant growth regulators, explant types, and their orientations on in vitro shoot proliferation of *Casuarina cunninghamiana* Miq. and also the subsequent rooting ability of shoots. Results showed that shoot proliferation occurred only in shoot tip explants cultured vertically on Murashige and Skoog (MS) medium supplemented with 2 or 4 μM thidiazuron (TDZ). Neither 6-benzylaminopurine alone nor in a combination with 1-naphthylacetic acid (NAA) or gibberellic acid had any effect on shoot proliferation. TDZ at 4 μM resulted in the greatest percentage of axillary bud sprouting (70%) and mean number of sprouts per explant (2.3). Additionally, no shoot proliferation was observed from detipped or single-node explants or from horizontally placed shoot tip explants when cultured on the same TDZ-containing medium. The induced shoots produced adventitious roots on MS medium supplemented with 2.5, 5, or 10 μM indole-3-butyric acid (IBA), not with indole-3-acetic acid and NAA. Although the mean number of roots per explant was not significantly different between 2.5 and 5 μM IBA, the highest rooting percentage (68%) and mean length of roots per explant (0.7 cm) was achieved at 5 μM IBA. The current study provided preliminary information toward commercial in vitro propagation of *Casuarina cunninghamiana* male plants.

*Casuarina cunninghamiana* Miq., a large, woody forest tree of the family Casuarinaceae, has many uses related to its rapid growth, large size, good branching, and survival when planted in impoverished environments less suited for many other trees. It also is an important plant for agroforestry because of its good timber and use as firewood (Midgley et al., 1983). In Florida, *Casuarina* has been present for approximately a century but recently became regulated because of concern about invasiveness (Castle, 2008). However, for the reasons mentioned, there is now interest in using these plants as a windbreak for citrus groves to help manage citrus canker disease, a windblown bacterial infection caused by *Xanthomonas axonopodis pv. citri*. A recent change in its legal status allows *C. cunninghamiana* male plants to be propagated from local sources only for restricted use as windbreaks (Castle et al., 2008).

Conventional propagation methods have been problematic for *C. cunninghamiana*, especially in large-scale commercial operations. Seed germination rates of this species are often extremely low (less than 50%), and seedlings exhibit undesirable variability in growth and form (El-Lakany and Shepherd, 1983; Shen et al., 2009a). In preliminary experiments, we found that seeds from local sources did not germinate whether the seeds were harvested from recently matured or older cones (unpublished data). Elsewhere in the world, where *C. cunninghamiana* is grown commercially for citrus windbreaks, seeds are the preferred propagation method and germination rates are apparently not a problem. Low success rates were also reported when propagation was attempted by rooting cuttings and air layering (Lundquist and Torrey, 1984). We also experienced problems of poor rooting of cuttings and little axillary bud proliferation in propagation trials with male plants of *C. cunninghamiana* (unpublished data). Root suckers are another means of vegetative propagation for some *Casuarina* species (Husain and Ponnuswamy, 1980). However, *C. cunninghamiana* trees in Florida do not produce suckers. The trees are dioecious, so seeds are not an option for propagation of male trees as required in Florida. Seed populations are mixed and the limited literature on the subject suggests that it requires 4–6 weeks to 7 years for the plants to flower and reveal their gender (Castle et al., 2008). In vitro procedures offer a promising means of propagation for species that are difficult to propagate by conventional methods. Research on in vitro propagation of *Casuarina* is limited, but experiments have been conducted on some species, including the related * Allocasuarina verticillata* Lam (Phelep et al., 1991), *C. equisetifolia* (Duhoux et al., 1986; Seth et al., 2007), *C. glauca*, and *C. cunninghamiana* (Aboel-Nil, 1987). In most of studies, seedlings or young immature trees were used as explant sources. There have been no reports on successful in vitro propagation of *C. cunninghamiana* using tissues from mature male trees. We have established a protocol for micropropagation of a *Casuanina* hybrid (*C. equisetifolia* L. × *C. glauca* Sieber ex Spreng) using epicotyl explants excised from seeds germinated in vitro. Shoot proliferation was obtained on Murashige and Skoog (MS) medium (Murashige and Skoog, 1962) supplemented with 6-benzylaminopurine (BA) at 0 to 35.6 μM and root induction on MS medium supplemented with indole-3-butyric acid (IBA) from 4.3 to 17.4 μM (Shen et al., 2009b). However, this in vitro protocol was not successful using shoot tips taken from mature male *C. cunninghamiana* trees in our preliminary experiment. Therefore, our objectives were: 1) to investigate the effects of plant growth regulators (PGRs) (type, concentration, and combination), explant type (entire shoot tip, detipped, and single node), and explant orientation (vertical and horizontal) on shoot proliferation; and 2) to examine the effects of three types of auxin (indole-3-acetic acid (IAA), IBA, and 1-naphthylacetic acid (NAA)) on in vitro rooting of *C. cunninghamiana* shoots.

Materials and Methods

Plant materials. Plant materials were collected from three naturalized mature male trees of *C. cunninghamiana* estimated to be more than 40 years of age located in Fort Pierce, FL (lat. 27.4487, long. 80.3759). The male nature of the source trees was determined by examining flower type. Lower branches of the source trees were pruned in Dec. 2008 to produce new growth. When newly formed branches were ≥3 months old and 2 to 8 cm long, tip cuttings were taken (Fig. 1A), placed in plastic bags with water, and kept in a cooler over ice. Explants were placed on culture medium within 24 h of sample collection.

Sterilization procedure. In preliminary experiments, we experienced high contamination rates when initiating cultures. The following sterilization procedure worked best for reducing contamination rate and increasing explant survival. Shoot tips (2 to 3 cm) long, with no visible axillary buds, were taken from cuttings and placed under running water for 30 min followed by 20-min soaking in 20% v/v regular bleach (1.2% sodium hypochlorite) with the addition of 2 drops of Tween-20 and then rinsed three times, 5 min each, with sterile water. The basal end of shoot tips was trimmed to remove bleach-damaged tissue. Shoot tips ≥1.5 cm long (Fig. 1B) were used as initial explants for in vitro culture.

General culture media and conditions. Culture medium consisted of MS mineral...
salts, 0.4 mg L⁻¹ thiamine, 2.0 mg L⁻¹ glycine, 100.0 mg L⁻¹ myo-inositol, 0.5 mg L⁻¹ pyridoxine, 0.5 mg L⁻¹ nicotinic acid, 30 g L⁻¹ sucrose, supplemented with different types and concentrations of PGRs, with 20 g L⁻¹ activated charcoal (AC) (Fisher Scientific, Fair Lawn, NJ) for shoot proliferation or without any AC for rooting, respectively. The medium was adjusted to pH 5.8 with 0.1 N NaOH before the addition of 8 g L⁻¹ TC agar (PhytoTechnology Laboratories, Shawnee Mission, KS) and autoclaved at 1.2 kg cm⁻² for 20 min. Cultures were kept in GA-7 vessels containing 40 mL of medium. Cultures were maintained at 22 ± 3 °C with a 12/12-h light/dark photoperiod at 40 μmol m⁻² s⁻¹ provided by cool white fluorescent lamps (Lithonia Lighting F40W/SS, Conyers GA).

Shoot proliferation: plant growth regulator effect. Four experiments (E.1 to E.4) were conducted to examine PGR types, concentrations, or combinations for shoot proliferation using entire shoot tip explants placed vertically on the medium: E.1—BA at 0, 1, 2, 4, 8, or 16 μM; E.2—BA at 0, 1, 2, 4, 8, and 16 μM; E.3—thidiazuron (TDZ) at 0, 1, 2, 4, 8, or 16 μM; and E.4—gibberellic acid (GA₃) at 0, 1, 2, 4, 8, or 16 μM. PGR-free medium served as the control. We selected BA because it is a commonly used cytokinin that was effective in inducing shoot proliferation from epicotyl explants excised from in vitro-germinated seedlings of a Casuarina hybrid (Shen et al., 2009b); TDZ is also a cytokinin known to induce axillary bud proliferation in recalcitrant species (Lu, 1993); and GA₃ can aid in breaking bud dormancy (Suzuki and Kitano, 2006).

A random mixture of shoot tips from the three source trees was placed vertically on media with the basal end inserted 2 to 3 mm deep. There were five explants per GA-7 vessel (Magenta Corporation, Chicago, IL) and eight replicates for each treatment. During a 6-week culture period, the explant responses were recorded. If shoot proliferation occurred, the number of explants with sprouted axillary buds and the number of sprouts per explant were recorded. The percentage of axillary bud sprouting was calculated as the number of explants with sprouted axillary buds out of the total number of explants cultured.

Only TDZ (E.3) was effective in inducing shoot proliferation. The following two experiments were conducted in an attempt to increase shoot proliferation rate using TDZ.

Explant orientation. Entire shoot tip explants were placed horizontally on the surface of medium and pressed ≈1 mm into media. Explants were cultured on MS medium supplemented with TDZ (to be comparable with the shoot proliferation experiment) at 0, 1, 2, 4, 8, or 16 μM. There were five explants per GA-7 vessel and eight replicates for each treatment. Explant responses were recorded at the end of 6 weeks culture.

Explant type. We also tested detippled explants in which 1 to 2 mm of the shoot apex was removed and single-node explants formed by separating the shoot explants at each node. These two types of explants were cultured on MS medium supplemented with TDZ at 0, 1, 2, 4, 8, or 16 μM. There were five explants per GA-7 vessel and eight replicates for each treatment. Explant responses were recorded at the end of 6 weeks culture.

In vitro rooting. The effects of IBA, IAA, and NAA on in vitro rooting were investigated. Shoots produced on MS medium containing TDZ (optimal for shoot proliferation) were used for rooting experiments. Shoots were removed from GA-7 vessels and the basal ends were trimmed 2 to 3 mm to remove any callus or browning tissues and then cultured on root induction media composed of MS medium supplemented with IBA, IAA, or NAA at 0, 2.5, 5, 10, 15, or 20 μM. PGR-free medium served as the control. There were five shoots per GA-7 vessel and five replicates for each treatment. At the end of 6 weeks culture, the number of shoots forming roots, root number, and the length of the longest root of each shoot were recorded. The rooting percentage was calculated as the number of shoots forming roots out of the total number of shoots cultured.

Experimental design and statistical analysis. All experiments were established in a completely randomized design. Data were subjected to analysis of variance using SAS (SAS Institute, Inc., 1999). Mean separation was achieved by the least significant difference test at the 95% level.

Results

Shoot proliferation: plant growth regulator effects. PGR effects on shoot proliferation were only observed on MS medium supplemented with TDZ (E.5). BA, BA in combination with NAA, and GA₃ at the concentrations tested did not induce any shoot proliferation. Inclusion of NAA at 0.05 μM in media resulted in profuse callus formation at the base of shoot tips. Shoot proliferation occurred only when the medium was supplemented with 2 and 4 μM TDZ.

Discussion

Shoot proliferation. PGR effects on shoot proliferation of Casuarina cunninghamiana were similar to those reported by others for other species (Liu and Li, 2001; Perez-Tornero et al., 1999). BA has been the most popular and widely used cytokinin for stimulating shoot multiplication in a broad range of species (Gaspar et al., 1996). Seth et al. (2007) reported that BA from 0 to 11.1 μM induced 38.5% to 73.86% axillary bud sprouting from mature tree shoot tip explants of Casuarina equisetifolia Forst. However, in our study, BA at concentrations ranging from 0 to 16 μM failed to result in any shoot production from Casuarina cunninghamiana. TDZ at 2 and 4 μM was essential to promote shoot proliferation in the present...
Table 1. Effect of thidiazuron (TDZ) on shoot proliferation of vertically placed entire shoot tip explants (n = 40) taken from mature male trees of Casuarina cunninghamiana Miq. cultured on Murashige and Skoog medium.1

| TDZ (µM) | Sprouted axillary buds (%)2 | Mean no. of sprouts per explant | Shoot length (cm) |
|----------|-----------------------------|---------------------------------|------------------|
| 0        | 0 c                         | 0 c                             | 0                |
| 1        | 0 c                         | 0 c                             | 0                |
| 2        | 53 b                        | 1.7 b                           | 2.0 a            |
| 4        | 70 a                        | 2.3 a                           | 1.6 a            |
| 8        | 0 c                         | 0 c                             | 0                |
| 16       | 0 c                         | 0 c                             | 0                |

1Data from shoot proliferation trial E.3.
2Means followed by the same letter in each column are not significantly different at the P = 0.05 level.

Fig. 2. Shoot proliferation and root formation of Casuarina cunninghamiana Miq. (A) Swelling of internodes after 2 weeks culture on Murashige and Skoog (MS) medium supplemented with 4 µM thidiazuron. Bars = 0.5 cm. (B) Sprouted lateral buds after 4 weeks. Bars = 0.5 cm. (C) Lateral buds at the distant position from apical meristem developed into a branch with many branchlets. Bars = 0.5 cm. (D) Single shoot formation from axillary buds at basal nodes. Bars = 0.5 cm. (E) Apical meristem developed into a branch with multibranchlets. Bars = 0.5 cm. (F) Root formation from a shoot cultured on MS medium supplemented with 5 µM indole-3-butyric acid for 6 weeks. Bars = 1 cm.

Table 2. Effect of indole-3-butyric acid (IBA) on in vitro root induction from shoots derived from entire shoot tip explants taken from mature male trees of Casuarina cunninghamiana Miq. cultured on Murashige and Skoog medium.

| IBA (µM) | Rooting (%)3 | Mean no. of roots per explant | Mean length of roots per explant (cm) |
|----------|---------------|-------------------------------|-------------------------------------|
| 0        | 0 c           | 0 b                           | 0 c                                 |
| 2.5      | 52 b          | 0.6 a                         | 0.4 b                               |
| 5        | 68 a          | 0.7 a                         | 0.7 a                               |
| 10       | 12 c          | 0.1 b                         | 0.1 c                               |
| 15       | 0 c           | 0 b                           | 0 c                                 |
| 20       | 0 c           | 0 b                           | 0 c                                 |

3Means followed by the same letter in each column are not significantly different at P = 0.05 level. Data represent means of 40 explants, including eight replicates and five samples per replicate.
species. Attempting to increase shoot proliferation rate with different explant types (detipped shoot tips and single node) and different explant orientations (horizontally) failed. Acclimatization of in vitro-regenerated plants to the greenhouse as well as to the field is the final step for commercial propagation. This propagation protocol must be substantially improved before such effort can be explored to examine the feasibility for large-scale propagation of this recalcitrant species.

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