Somatic Embryogenesis from Roots of Camellia japonica Plantlets Cultured in Vitro

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Abstract. Somatic embryos were induced on the roots of Camellia japonica L. plantlets regenerated from an in vitro clone of juvenile origin. The embryos appeared to differentiate from epidermic cells and to be connected with the root via a few parenchymatous cells. Somatic embryogenesis occurred on basal medium and with or without various combinations of zeatin, BA, and IBA. Secondary embryos were induced on cotyledons and/or hypocotyl regions of somatic embryos. Two morphological types of somatic embryos were developed, seed-like and bud-like types, and their formation was influenced by the presence of BA in the medium. Embryogenic capacity has been maintained for more than 24 months by subculturing secondary embryos at 7- to 8-week intervals. The best gibberellin/auxin combination for inducing the germination of isolated somatic embryos was GA, at 5 mg·liter−1 G.A., and IAA at 1 mg·liter−1. Plantlets were successfully established in planting medium and have continued to grow in a greenhouse.

Chemical names used: N-[(phenylmethyl)-1H-purine-6-amine (BA); (1α, 2β, 4α, 4bβ, 10β)-2,4a,7-trihydroxy-1-methyl-8-methylene-1-ene-1,10-dicarboxylic acid 1H-acetamide (GA); 1 H-indole-3-acetic acid (IAA); 1 H-indole-3-butyric acid (zeatin).

Shoot cultures based on axillary branching remains the preeminence in vitro method of camellia clonal multiplication from seedling-derived (Samartín et al., 1986) and from mature explants (San-José and Vieitez, 1990; Vieitez et al., 1989a, 1989b). Somatic embryogenesis might be an alternative mass propagation system. Somatic embryogenesis has been achieved from zygotic embryos of some Camellia spp. (Kate, 1989), but its extension to a wider range of older explants would be desirable. We found no reports of the regeneration of camellias by somatic embryogenesis from plantlets cultured in vitro.

Regeneration of plants from root tissue is not common. However, somatic embryos have been obtained on callus derived from intact roots of Prunus incisa × P. serrula and horse-chestnut (Aesculus hippocastanum L.) (Druart, 1981). Also, embryoid structures were induced on callus nodules developed on the roots of micropropagated cherry rootstock Colt (Prunus avium × P. pseudocerasus) plantlets (Jones et al., 1984). McCown (1986) developed highly competent embryogenic and organogenic systems based on the formation of nodules on the roots of Populus shoots rooted in vitro.

We report the induction of somatic embryos from roots of Camellia japonica L. plantlets raised in vitro, the maintenance of embryogenic capacity through secondary embryogenesis, and germination of somatic embryos into plants.

Materials and Methods

Micropropagated plantlets of three clones derived from in vitro shoot cultures of C. japonica were used as source material. Clones 1 and 2 were derived from 3- to 4-month-old seedlings and clone 3 from adult tissues obtained from a 50-year-old specimen of C. japonica ‘Alba Plena’. The establishment of the shoot cultures from axillary buds has been described elsewhere (Samartín et al., 1984; Vieitez et al., 1989a, 1989b). Shoot multiplication cultures had been maintained for 2 years by subculturing at 7- to 8-week intervals on basal medium (see below) supplemented with BA at 1 mg·liter−1 and IBA at 0.01 mg·liter−1.

For generating whole plantlets, rooting was induced by dipping the basal ends of 2- to 3-cm-long shoots in IBA at 1 g·liter−1 in 16% ethanol for 15 min, after which the shoots were placed in 300-ml jars containing 60 ml of basal medium, but with half-strength macronutrients and no growth regulators. Cultures were incubated in the dark for 10 to 12 days and, thereafter, with 30 µmol·s−1·m−2 of light produced by cool-white fluorescent lamps for 16 h·day−1. The light/dark cycle was 26/20C.

Basal culture medium consisted of mineral salts of Murashige and Skoog’s medium (1962) for clones 1 and 2 and Lloyd and McCown’s medium (1980) for clone 3 (mg·liter−1) 1 zeatin, 0.1 nicotinic acid, 0.1 pyridoxine-HCl; (g·liter−1) 30 sucrose and 6 Difco agar. The pH was adjusted to between 5.5 and 5.6 before autoclaving.

After 1 month in rooting medium, rooted shoots were transferred to basal medium or basal medium supplemented with (mg·liter−1) 1 zeatin or 1 or 2 BA, alone or in combination with 0.1 or 1 IBA. Sixteen plantlets of each clone were tested in each medium combination (160 plantlets per clone). Cultures were incubated with 30 µmol·s−1·m−2 of light provided by cool-white fluorescent lamps for 16 h·day−1, with a day/night of 26/20C. Percentage of plantlets exhibiting somatic embryos was recorded after 8 weeks. The experiment was conducted twice.

Secondary embryogenic capacity was evaluated for the two clones that produced somatic embryos (clones 1 and 2). Primary somatic embryos were isolated and subcultured on the medium on which they had been obtained to increase the number of embryogenic cultures by repetitive embryogenesis. The competence of secondary embryogenesis was evaluated by transferring cotyledonary stage somatic embryos to basal medium alone or basal medium supplemented with BA at 1 or 2 mg·liter−1 and/or IBA at 0.1 or 1 mg·liter−1. For each treatment/clone combination, 40 somatic embryos were used (four replicate jars with 10 embryos each). Incubation conditions were the same as described previously. After 8 weeks, the percentage of embryos that had produced secondary embryos and the number of secondary embryos per responsive culture were recorded. Percent-age data were transformed using arcsin of the square root and subjected to two-way analysis of variance (ANOVA). Significant differences between means was determined using Duncan’s multiple range test.

Germination experiments were carried out with 5- to 9-mm-
Induction of somatic embryogenesis

Upon transfer to embryogenic medium, regenerated camellia plantlets each possessed four to six roots 10 to 12 mm long. Eight weeks later, average root length ranged from 14 mm for BA at 2 mg·liter⁻¹ to 26 mm for the basal medium.

No treatment induced somatic embryogenesis on plantlets of mature origin (clone 3). Somatic embryogenesis occurred with all treatments, including basal medium without growth regulators, for clone 2 with success rates ranging from 15% to 35% without any apparent relationship to treatment. The overall embryogenesis rate was 26% among the 160 plantlets of the first experiment and among the 154 of the second experiment (six plantlets were lost due to contamination). Similar results were obtained for clone 1, with an overall embryogenesis rate of 10% for 155 plantlets.

Yellow-white somatic embryos differentiated directly on plantlet roots either singly or in groups of up to seven to 3 weeks after...
transfer to embryogenic medium (Fig. 1A). Most embryos developed in the proximal region of a single root. Subsequent development of embryos was not synchronized, although most had reached the cotyledonary stage by 8 weeks on embryogenic induction medium. Somatic embryos appeared to differentiate from epidermal cells and to be connected to the root via a few parenchymatic cells, making them easily removable. No vascular connections were observed between root and embryo (Fig. 1B, C). Some rounded, hypertrophic epidermal cells that were \( \approx 20 \) times larger than normal epidermal cells and possessed visible nuclei were also observed.

**Secondary embryogenesis.** Secondary somatic embryos developed on the hypocotyl and/or cotyledonary regions of previously differentiated embryos after 3 to 4 weeks in culture (Fig. 2A-B). There were two morphological types of embryos: “seed-like” embryos had large, white, opaque cotyledons, whereas “bud-like” embryos had green cotyledons resembling true leaves (Fig. 3A-D). Anatomical examination revealed the presence of shoot and root meristems in both types of embryos (Fig. 2C). Anomalies such as polycotyledony, cotyledon hypertrophy, or fascination (fusion of embryos at the hypocotyl/root level) were observed to different extents in both kinds of embryos (Fig. 3E).

Secondary embryos 8 to 9 mm long were subcultured every 7 to 8 weeks. Embryogenic capacity declined when individual embryos <3 to 4 mm long were subculture, but successful subculture of groups of such immature embryos was possible.

For both embryo types of clone 2, secondary embryogenesis rate and number of secondary embryos per responsive embryo varied significantly \((P < 0.005)\) with treatment factor, independently of the embryo type (Table 1). The results for clone 1 were similar. Best results were achieved in medium without growth regulators or medium supplemented with IBA at 0.1 mg·liter\(^{-1}\). Lowest rates of embryogenesis were recorded in medium containing BA at 2 mg·liter\(^{-1}\), although the most abundant, had high fascination rates. Five weeks on germination medium produced not only whole regenerated plantlets (Fig. 4) but also incompletely regenerated plantlets lacking either roots or shoots.

In preliminary experiments we observed poor germination (9%) of bud-like embryos on basal medium. Germination was generally improved by treatments with GA\(_3\) either alone or in combination with IAA (Table 3). Media supplemented with BA at 1 mg·liter\(^{-1}\) + IBA at 0.1 mg·liter\(^{-1}\), although more abundant, or with GA at 1 mg·liter\(^{-1}\) + IAA at 0.1 or 2 mg·liter\(^{-1}\) resulted in a significantly higher \((P = 0.05)\) germination rate than the media with GA alone or with GA at 1 mg·liter\(^{-1}\) + IAA at 0.1 or 2 mg·liter\(^{-1}\). In general, length of shoots produced increased with increasing GA\(_3\) concentrations (significant in whole plantlets, \(P = 0.05\)). Root length was not significantly affected by growth regulator treatments.

Germination. Germination experiments used bud-like embryos because seed-like embryos had exhibited less germination capacity in preliminary experiments. Embryos developed on BA at 1 mg·liter\(^{-1}\) + IBA at 0.1 mg·liter\(^{-1}\) were used because those developed on BA at 2 mg·liter\(^{-1}\) although more abundant, had high fascination rates. Five weeks on germination medium produced not only whole regenerated plantlets (Fig. 4) but also incompletely regenerated plantlets lacking either roots or shoots.

In preliminary experiments we observed poor germination (9%) of bud-like embryos on basal medium. Germination was generally improved by treatments with GA, either alone or in combination with IAA (Table 3). Media supplemented with BA at 1 mg·liter\(^{-1}\) + IBA at 0.1 mg·liter\(^{-1}\) resulted in a significantly higher \((P = 0.05)\) germination rate than the media with GA alone or with GA at 1 mg·liter\(^{-1}\) + IAA at 0.1 or 2 mg·liter\(^{-1}\). In general, length of shoots produced increased with increasing GA concentrations (significant in whole plantlets, \(P = 0.05\)). Root length was not significantly affected by growth regulator treatments.

During the first 2 weeks after transfer of somatic embryos to germination medium, cotyledons grew and thickened, and a single tap root began to emerge; the shoots developed later. Media with only GA, at 1 mg·liter\(^{-1}\) tended to favor the thickening and development into cotyledonal structures of the first leaves developed on the shoot, whereas those cultured with GA,
at 5 mg-liter\(^{-1}\) induced thinner shoots with smaller leaves. Secondary embryogenesis sometimes occurred on the hypocotyl and proximal cotyledonal region during germination; it was more frequent when low GA\(_3\) concentrations were used. Secondary embryogenesis rates were lower and root growth was better when embryos were subjected to an initial 15-day dark period on germination media; with this treatment, 83% of somatic embryos developed primary roots.

Plantlets were successfully transferred into potting medium and plants were moved to the greenhouse 6 weeks after transfer to soil. They appeared morphologically similar to clonal material.

**Discussion**

We have developed an on-going somatic embryogenic system for *C. japonica* from plantlets regenerated from in vitro clones of juvenile origin. The somatic embryos developed directly on the roots of the plantlets, and there was no evidence of differentiation from callus.

Wann (1988) considered the pattern of indirect embryogenesis only if a true callus phenotype was described. In our experiments, primary somatic embryos appeared to differentiate directly from root epidermal cells, although a few parenchymatic cells linked the root to the embryo. A limited cell proliferation process observed before the direct differentiation of somatic embryos on cacao cotyledons might constitute a situation intermediate between direct and indirect embryogenesis (Duham et al., 1989).

Growth regulator treatment had comparatively little effect on induction of somatic embryogenesis, which even occurred in basal medium. However, it is important to emphasize that rooting of camellia shoots has been induced by a pulse treatment in a very concentrated IBA solution. Direct somatic embryogenesis also takes place in hormone-free media from the hypocotyls and cotyledons of mature and immature zygotic embryos of *C. reticulata* and *C. japonica* (Plata and Vieitez, 1990; Vieitez and Barciela, 1990). Both phenomena appear to reflect the inherent capacity of these species to produce pre-embryogenic determined cells (Sharp et al., 1980) when physiologically juvenile. Conversely, the plantlets obtained from the ‘Alba Plena’ clone of adult origin failed to respond to any of the embryogenic treatments.

The long-term maintenance of secondary embryogenic competence by successive subcultures allows the creation of an embryogenic regeneration system that may not possess an embryogenic callus proliferation stage. A similar system using repetitive embryogenesis has been reported in other woody species such as walnut (*Juglans regia* L.) (Tulecke and McGranahan, 1985), eucalyptus (*Eucalyptus citriodora* Hook) (Muralidharan et al., 1989), and red currant ‘F. Hosszúfúrtú’ (*Ribes rubrum* L.) (Zatyko et al., 1975). This system may prove...
more efficient than shoot micropropagation methods (Samartin et al., 1984; Vieitez et al., 1989a, 1989b).

We observed two morphological types of somatic embryos similar to somatic embryos obtained from zygotic embryos of C. japonica (Vieitez and Barciela, 1990). According to Ammirato (1985), the bud-like type was differentiated with relatively high concentrations of cytokinins.

The somatic embryos used in the germination experiments had cotyledons smaller than those of the zygotic embryos, but their hypocotyl-root and shoot axes were well defined. The improvement of germination rates by GA3, has been reported in other woody species such as satsuma (Citrus unshiu Marc.) (Nito and Iwamasa, 1990) and horse-chestnut (Radojevic, 1988). Germination of the latter was enhanced, as in our experiments, from the inclusion of IAA in the medium. Kato (1986) used GA3 at 1 mg·liter\(^{-1}\) to induce germination of somatic embryos of C. japonica obtained from cotyledon cultures, but she did not state the germination rates achieved. The rates obtained in our experiments using GA3 at 5 mg·liter\(^{-1}\) + IAA at 1 mg·liter\(^{-1}\) were quite good compared with those reported for other woody species (Warn, 1988). However, a certain immaturity of germinating embryos was indicated by the tendency to form secondary cotyledons instead of true leaves, which has also been observed in cacao (Theobroma cacao L.) (Duhem et al., 1989) and red oak (Quercus rubra L.) (Gingas and Lineberger, 1989).

The occurrence of secondary embryogenesis during germination is often associated with lack of maturity and failure to develop as a normal plantlet. This phenomenon presumably reflects the escape of certain cells from integrated group control (Williams and Maheswaran, 1986). Further control of embryogenic maturation along the lines employed by DeWald et al. (1989) for mango (Mangifera indica L.) seems to be necessary.

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