Somatic Embryogenesis in Three Magnolia Species

S.A. Merkle and A.T. Wiecko
School of Forest Resources University of Georgia, Athens, GA 30602

Abstract. Cultures were initiated from immature seeds of three species of magnolia: sweetbay magnolia (Magnolia virginiana L.), fraser magnolia (M. fraseri Walt.) and yellow cucumber tree (Magnolia acuminata var. cordata Sarg.). Immature seeds were bisected longitudinally and cultured on a solidified conditioning medium containing 2 mg 2,4-D liter, 0.25 mg BA/liter, 40 g sucrose/liter, and 1 g casein hydrolysate/liter. Cultures were maintained in the dark at 22C and transferred to fresh medium at monthly intervals. Within 2 months of culture, somatic embryos or proembryogenic masses proliferated from one end of the endosperm mass. Somatic embryos and proembryogenic masses of each species were cultured on a hormone-free version of the conditioning medium to complete maturation and then transferred to the same hormone-free medium, minus casein hydrolysate, to initiate germination. Germinants were transferred to a hormone-free plantlet development medium for conversion. Plantlets of all three species survived transfer to soil mix and continued to grow. Chemical names used: (2,4-dichlorophenoxy) acetic acid (2,4-D), N-(phenylmethyl)-1H-purin-6-amine (BA).

Materials and Methods

Because sweetbay magnolia trees were readily accessible while fraser magnolia and yellow cucumber tree were not, plant material for explanting into culture was handled differently for each species.

Sweetbay magnolia. Developing fruits (aggregates of follicles) were sampled from 10 ornamental sweetbay magnolia trees growing on the Univ. of Georgia campus at weekly intervals from 2 to 7 weeks postanthesis (2 June -14 July 1989). Fruits only contained a few filled seeds; therefore, collections from trees were bulked for each sampling date. For the first three sampling dates, whole aggregate fruits were disinfested using the following sequence: 70% ethanol for 20 sec, 10% Roccal (National Laboratories, Montvale, N. J.) for 1 min (repeating the ethanol and Roccal steps once), full-strength Clorox (5.25% sodium hypochlorite) for 5 rein, sterile water rinse for 3 rein, 0.01 M HCl rinse for 3 rein, and three additional sterile water rinses. Aggregates were dissected aseptically to remove seeds. For the last four sampling dates, when aggregates were 5 to 7 cm long, seeds were excised from the fruits, then disinfested using the same sequence as was used for the fruits. At least 30 seeds from each tree sample date were explanted for each sample date.

Fraser magnolia. Developing fruits were sampled only once from three fraser magnolias growing naturally near Brasstown Bald in Towns County, Ga. All three trees were within 20 m of each other and were sampled on 7 July, ~5 weeks postanthesis. Fruits were 8 to 10 cm long and mostly green, but beginning to show some pink. Seeds were removed for disinfestation as with sweetbay magnolia seeds. Thirty seeds were cultured from each of the three trees.

Yellow cucumber tree. Developing fruits were sampled twice from a single yellow cucumber tree in Oglethorpe County, Ga. Samples were collected ~3 (7 June) and 9 weeks post-anthesis (17 July). A very low percentage of the seeds were filled because this tree is insect-pollinated and was the only one in the vicinity. For the first sample, whole fruits were disinfested and seeds excised aseptically using the same method as for sweetbay magnolia fruit. For the second sample, seeds were excised and disinfested. Thirty-seven seeds were cultured from the first sampling date, but only nine seeds were cultured from the second one.

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halves were placed cut surface downward in 60-mm plastic petri dishes (three seeds per dish) on yellow-poplar conditioning medium (Merkle and Sommer, 1986) containing 2,4-D at 2 mg-liter\(^{-1}\), BA at 0.25 mg-liter\(^{-1}\) casein hydrolysate (CH) at 1 g-liter\(^{-1}\), and sucrose at 40 g-liter\(^{-1}\). Cultures were maintained at 22°C in darkness and transferred to fresh medium monthly. Following the appearance of proembryogenic masses (PEMs) or embryos, some were transferred for further development to yellow-poplar basal medium that lacked hormones, still in darkness. Mature embryos were transferred to a medium of the same formulation, but lacking CH, to promote germination and grown under fluorescent light. We had previously found that removal of CH from the basal medium promoted germination of mature yellow-poplar somatic embryos, probably by reducing the water potential of the medium (unpublished data). Germinants were moved to test tubes containing 20 ml of yellow-poplar plantlet development medium (Merkle and Sommer, 1986) containing no hormones and 2% sucrose, also under fluorescent light. To initiate suspension cultures, some PEM-like embryos were in- oculated into liquid conditioning medium and grown in darkness at 22°C and with shaking at 90 rpm. Also, we attempted to induce cultures to proliferate in a less-differentiated state (i.e., PEMs instead of globular embryos) by initiating subcultures of each embryogenic line on solidified conditioning medium with the 2,4-D level increased to 4 mg-liter\(^{-1}\).

**Results and Discussion**

**Sweetbay magnolia.** Most sweetbay magnolia explants produced large amounts of nonmorphogenic callus within 4 weeks after being placed in culture. In addition, some cultures produced somatic embryos that originated directly from one end of the explanted seed halves and that we presumed to be the locations of the zygotic embryos. However, we cannot be certain that the zygotic embryos were the source of the somatic embryos, because at the time of explanting, we could not locate embryos in the seeds. Globular-stage somatic embryos were first observed proliferating from 3-weeks post-anthesis seed 5 weeks after explanting. Additional embryogenic cultures arose over the next few weeks, mainly from seeds explanted on the same date (12 embryogenic cultures out of 45 seeds), but also from seeds explanted 2 weeks postanthesis (two embryogenic cultures out of 48 seeds) and 5 weeks post-anthesis (one embryogenic culture out of 36 seeds). Embryogenesis took two forms that appeared to depend on explanting date. Embryos from the two earliest cultures (2 weeks post-anthesis) appeared most similar to PEMs seen in many indirect embryogenic systems, such as carrot *Daucus carota* L. (Halperin, 1966), and proliferated at the highest rate. Somatic embryos from later cultures appeared to be produced by direct embryogenesis, with no intermediate callus or PEMs (Fig. 1A). PEM-like embryos inoculated into liquid medium proliferated rapidly but did not form cell suspensions. Instead, they grew as clusters of globular embryos that would break apart and initiate new clusters. Subcultures to the higher auxin level (2,4-D at 4 mg-liter\(^{-1}\)) appeared to promote production of PEMs along with more rapid proliferation of the cultures.

Somatic embryos maintained on solid conditioning medium, although mostly malformed, often continued development through the torpedo stage and continued to swell but would not germinate. Instead, cotyledons remained fused and secondary embryos formed at the radicle ends. PEMs and globular stage embryos transferred to basal medium appeared to complete development normally, resulting in torpedo-stage embryos with two to three separated cotyledons (Fig. 1B). However, germination of these mature embryos remained inhibited until they were transferred to basal medium without CH and placed in light, whereupon cotyledons became green and expanded and radicles elongated (Fig. 1C). Embryos continued germination (i.e., root elongation) following transfer to plantlet development medium, but apical development was very slow and ceased in about 75% of the germinants. About 25% of the embryos completed conversion to form seedling-like plantlets (Fig. 1D). Plantlets transferred to Peat-Lite (Fafard, Springfield, Mass.) potting mix, placed in a humidifying chamber, and fertilized with a modified Hoagland’s solution weekly continued to produce new leaves at a rate of one every 2 to 3 weeks.

**Fraser magnolia.** Fraser magnolia explants responded very quickly to culture on conditioning medium, with PEMs first appearing only 3 weeks after explanting. Within 5 weeks, 11 of 90 seeds, representing all three sample trees, had produced PEMs. PEMs proliferated slowly and, unlike sweetbay, magnolia embryogenic cultures, tended not to differentiate to globular-stage embryos while on conditioning medium. PEMs inoculated into liquid conditioning medium proliferated more slowly than did those of sweetbay magnolia. Following transfer to solid basal medium, globular embryos differentiated within...

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*Fig. 1. Somatic embryogenesis in sweetbay magnolia. (A) Somatic embryos arising by direct embryogenesis from end of an immature sweetbay magnolia seed. (B) Torpedo- and cotyledon-stage somatic embryos arising from mass of sweetbay magnolia proembryogenic masses. (C) Germinating somatic embryo. (D) Somatic embryo-derived sweetbay magnolia plantlets. Bar in all photos = 500 µm.*

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placed on top of basal medium, a procedure we have previously used for somatic embryos on conditioning medium (Fig. 2B). As with the other species, PEMs could also be maintained in liquid conditioning medium. However, transfer of PEMs to basal medium failed to induce the PEMs or globular embryos to develop further. Finally, PEMs were transferred to filter paper disks that were placed on top of basal medium, a procedure we have previously applied to obtain mature yellow-poplar somatic embryos (unpublished data). After 5 to 6 weeks, somatic embryos differentiated. As with sweetbay, magnolia, embryos could be germinated in the light on basal medium without CH and ∼10% of the germinants continued to grow upon transfer to plantlet development medium. A few of these plantlets have been transferred to potting mix and have continued to produce new leaves in a humidifying chamber.

The genus Magnolia appears to be amenable to initiation of embryogenic cultures based on the performance of the three species we tested. Both sweetbay and fraser magnolias produced multiple embryogenic cultures, although only three fraser magnolia trees were sampled and developing fruit were collected only on a single date. The inability to obtain more than one embryogenic culture from yellow cucumbertree was probably an effect of very low percentage of filled seed. Although the seeds cultured from this species appeared to be full-sized, indicating they were filled, we found no zygotic embryos. Thus, many of the seeds placed in culture may not have contained viable embryos.

Our results with sweetbay magnolia indicate that an early stage of development may be optimum for explanting the zygotic embryo (and other seed components) to initiate an embryogenic culture. The group of seeds producing the highest percentage of embryogenic cultures was explanted only 3 weeks post-anthesis, when embryos were too small to be found under a dissecting microscope. The globular stage of embryo development was found to be optimum for initiation of embryogenic cultures of the related yellow-poplar species (Merkle et al., 1989).

Although the protocol for initiation of embryogenic magnolia cultures was very similar to that for yellow-poplar (e.g., use of immature zygotic embryos, same media), growth characteristics of the cultures differed. Yellow-poplar cultures invariably formed PEMs on media with 2 mg 2,4-D/liter and produced somatic embryos via indirect embryogenesis when transferred to hormone-free induction medium. PEMs would readily form cell suspensions in liquid 2,4-D-supplemented media. In contrast, the magnolia cultures did not readily form PEMs, even when the 2,4-D level was raised to 4 mg-liter⁻¹. Cultures were mixtures of PEMs and somatic embryos at globular or later stages of development that appeared to proliferate via direct embryogenesis. Magnolia PEMs and embryos transferred to liquid conditioning medium did not form true cell suspensions, but continued to produce new PEMs or embryos in large clusters. These in turn broke apart to initiate new clusters. Finally, in comparison to yellow-poplar embryos given the same conversion treatment, magnolia embryos formed plantlets very slowly or failed to complete conversion. Although root elongation in sweetbay magnolia and yellow cucumbertree proceeded at rates similar to that in yellow-poplar, the magnolia plantlets produced leaves much more slowly than did the yellow-poplar. Height growth rate was similarly reduced. However, it is possible that the relative rates of leaf production and height growth may also hold for seedlings of these species and not be specific to somatic embryo-derived plantlets.

Our results indicate that embryogenic cultures may provide a means of mass propagation of magnolia species. The cultures, which we believe to be derived from immature zygotic embryos in the explanted seeds, have continued to produce somatic embryos and plantlets for >8 months. The potential of this tissue culture system may be especially useful for propagation of some of the rare native species of magnolia, such as yellow cucumbertree.

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