Initiation and Maturation of Somatic Embryos of Squash (Cucurbito pepo)

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Abstract. Plant regeneration from tissue cultures of summer squash (Cucurbita pepo L. ev. YC60) has been observed. Embryogenic callus tissues were initiated when cotyledons of mature seeds were excised and cultured on Murashige and Skoog (MS) medium supplemented with either 22.7 µM 2,4-D or a combination of 4.7 µM 2,4,5-T, 4 µM BA, and 0.5 µM kinetin. Clusters of somatic embryos were found in callus tissue. Maturation of these somatic embryos was effected by transfer of embryogenic callus tissues to MS supplemented with 0.5 µM NAA and 0.25 µM kinetin. Regenerated mature plants were morphologically normal and set fruits containing seeds that germinated normally. Chemical names used: 6-benzylaminopurine (BA); 2,4-dichlorophenoxyacetic acid (2,4-D); α-napthaleneacetic acid (NAA); 2,4,5-trichlorophenoxyacetic acid (2,4,5-T).

There have been very few studies concerning tissue culture of Cucurbita pepo. Schroeder (1968) reported the production of embryogenic tissue from pericarp of zucchini squash. Jelaska (1972, 1973, 1974, 1980) reported somatic embryogenesis in hypocotyl- and cotyledon-derived callus of pumpkins and demonstrated that embryos could develop into normal plants. Pink and Walkey (1984) reported a micropropagation method for pumpkin through apical meristem culture. Chee (1991) reported somatic embryogenesis with plant regeneration from shoot apex-derived callus of summer squash. This report describes a method for induction and maturation of somatic embryos of 'YC60' summer squash from cotyledons of mature seed.

Mature seeds of 'YC 60' summer squash (Asgrow Seed Co., Kalamazoo, Mich.) were surface sterilized for 10 min with a solution of 5.25% sodium hypochlorite and rinsed three times with sterile water. After removal of the seedcoats, the seeds were again treated with 8.75% sodium hypochlorite for 25 min and rinsed three times with sterile water. The seeds were then treated with 70% alcohol for 1 min and rinsed three times with sterile water.

Sterilized embryos were cut transversely into two unequal sections. One section consisted of the embryonic axis and one-third of the cotyledons; the other, the remainder of the cotyledons. The explants, including the embryonic axis, were cultured vertically with the cut surface facing upward; cotyledon explants were cultured with a subaxial surface in contact with the culture medium. All cultures were kept in the dark for 9 weeks at 26°C. Unless otherwise stated Murashige and Skoog (1962) (MS) medium supplemented with 3% sucrose and solidified with 0.8% Phytagar (Gibco, Grand Island, N.Y.) was used. The pH of all media was adjusted to 5.8 before autoclaving at 121°C for 20 min.

In the first experiment, the effect of 2,4-D (2.3, 4.5, 9.1, 13.6, 22.7, 45.4, 113.5, and 227 µM) was determined in combination with five concentrations of kinetin (0, 2.3, 4.6, 9.2, and 13.8 µM). Between 100 to 170 explants of each type were used per treatment. In the second experiment, the effect of β-indoleacetic acid (IAA) (0, 8, and 16 µM) was determined in combination with kinetin (0, 6.9, 13.8, 18.9, and 27.9 µM). For each treatment, 120 explants of each type were used. Finally, a combination of 4.7 µM 2,4,5-T, 4 µM BA, and 0.5 µM kinetin (TBK medium) was used. For this treatment, 531 explants of each type were used.

After 9 weeks on the induction media, including one subculture at 4.5 weeks, callus containing putative somatic embryos were transferred to maturation medium composed of MS salts and vitamins with 0.5 µM NAA and 0.25 µM kinetin. The cultures were incubated at 28°C under diffuse cool-white fluorescent lamps (80 µmol·m⁻²·s⁻¹) with a 16-photoperiod. When the apical region grew, the tissues were transferred to GA7 boxes (Magenta, Chicago) containing 50 ml of half-strength hormone-free MS medium. After an extensive root system developed, the plantlets were transplanted to 0.9-liter plastic pots containing planting medium and covered with clear plastic storage bags for 1 week for acclimation. Subsequently, the regenerated plants were potted in 7.6-liter pots containing a soil mixture (Metro Mix 300R; A.H. Hummert Seed Co., St. Louis) and grown in a greenhouse.

Representative embryogenic callus tissues on individual somatic embryos were prepared for microscopic examination by fixing in 10% neutral buffered formalin for 45 min, dehydrated in an ethyl alcohol-tertiary alcohol series, and embedded in Paraplast (Monoject Scientific, St. Louis) at 56°C. Embedded tissues were sectioned at 5 µm and stained for 12 h with 1% (v/v) safranin O (Sigma Chemical Co., St. Louis) dissolved in 50% alcohol, followed by a 20-sec exposure to 1% (v/v) fast green (Sigma Chemical Co.) dissolved in 95% alcohol.

In the first experiment, the only effective treatment for induction of embryogenic callus was 22.7 µM 2,4-D without kinetin. Only 5% of the explants produced embryos when the cotyledon was present, but 28% did when the cotyledon and embryonic axis were present.

After 6 weeks of culture on MS containing 22.7 µM 2,4-D, both types of explants formed embryonic-type callus that was translucent, smooth, and gelatinous in appearance. Most of this callus developed at the embryonic axis area and at the site of explant contact with the medium. After an 8- to 9-week incubation, ~10 to 20 putative somatic embryos had formed on the surface of the gelatinous callus (Fig. 1). Similar callus morphology...
A longitudinal cross-section of a bipolar somatic embryo before transfer to maturation medium is shown in Fig. 2. The development of the somatic embryos on maturation medium initially appeared abnormal. The embryos were elongated with distorted morphology (Fig. 3); however, -75% of these embryos developed into normal plantlets (Fig. 4).

Root formation was obtained when the mature embryos were grown on half-strength MS basal medium without growth regulators. The success of transfer of the plantlets from agar to planting medium was 100%. Two-hundred plants regenerated in this manner displayed no gross phenotypic abnormalities; they flowered and set viable seeds.

In the second experiment (IAA combined with kinetin), callus tissues grew very slowly on all test media. Some tissues became markedly nodular, rhizogenic, and brownish, but embryogenic, gelatinous-type tissues were absent. Jelaska (1974) reported that embryogenic callus tissues were induced on pumpkin hypocotyl explants cultured on media supplemented with 5.6 to 11.2 µM IAA. However, with summer squash IAA does not appear to have any induction effect on the explants.

With TBK medium, somatic embryogenesis was observed only on the explants containing the embryonic axis; 8% of the explants produced embryogenic tissue. Embryoids initially appeared abnormal; however, nearly 50% of them grew into plantlets. Regenerated plants were morphologically normal, flowered, and set seeds.

To my knowledge, this is the first report of the induction of somatic embryos, and their maturation to plants, of summer squash from mature seeds. Mature seeds provide a convenient and readily available explant source. Initiation of embryogenic cultures is simple and rapid. Putative embryogenic tissue that developed on the surface of the explant was easily identified by a characteristic translucence. Plantlets were obtained within 4 months of culture initiation. Further studies are needed to determine if this protocol is genotype-specific.

**Literature Cited**


