Abstract. Somatic embryos derived from walnut (Juglans regia L.) ovule tissues were evaluated to determine whether they were of zygotic or maternal origin. Molecular markers were used to permit evaluation at an early stage, before whole plant development. Somatic embryos developed from potentially apomictic ‘Sunland’ and ‘Cisco’ ovule tissue isolated from bagged putatively unpollinated flowers. Phosphoglucomutase (PGM) isozyme analysis showed that all of these embryos, except one from each cultivar, carry the same zymotype as the maternal tissue. However, restriction fragment length polymorphism (RFLP) analysis combined with isozyme evaluation demonstrated that the tested embryos originated from zygotic rather than maternal tissues. This study demonstrates the application of molecular marker analyses, particularly RFLPs, to evaluate the origin of somatic embryo origin.

Commercially important walnut cultivars may be improved by adding traits such as disease and pest resistance to their genetic makeup. Incorporation of novel genes by genetic transformation can complement conventional breeding, but plant regeneration from genetically manipulated cells is a prerequisite to this approach. In walnut, repetitively embryogenic somatic embryos have been used as the target tissue for genetic transformation and regeneration of transgenic plants (McGranahan et al., 1988, 1990). However, somatic embryo cultures have only been obtained from immature walnut embryo tissues (Tulecke and McGranahan, 1985) and from endosperm (Tulecke et al., 1988), but not from clonal (nonzygotic) tissues of commercially desirable cultivars. Use of clonal (vegetative) tissues as a source of somatic embryos is advantageous to avoid meiosis and genetic recombination and produce plants that are identical to the parents except for the introduced genes.

The origin of somatic embryos can be determined from histological analysis of the embryos and associated tissues, karyotype, and/or morphological analysis in many crops. Histological methods result in destruction of the embryo. Karyotype analysis is limited to detection of gross chromosomal differences and is not feasible in walnut due to small chromosome size (Woodworth, 1930). Morphological analysis, especially of fruit or flower, is prolonged due to the long generation time of walnut. Analysis using molecular genetic markers is an alternative that can be instrumental for rapidly determining the parentage of plants regenerated in vitro. Protein markers have proven their utility in ascertaining the genotypic parentage of tissue culture-derived plants (Zamir et al., 1981). The detection of restriction fragment length polymorphisms (RFLP) by single copy or cDNA probes has allowed diagnostic identification of strains of fungi (Castle et al., 1987). DNA “fingerprints” have been used to differentiate among rice (Oryza sativa L.) cultivars (Dallas, 1988) and several other plant species (Rogstad et al., 1988). RFLP is potentially useful for determining the parental nature of tissue-culture regenerants, particularly in Juglans regia, where only two polymorphic isozyme markers are available (Arulsekar et al., 1986).

This paper demonstrates the application of molecular markers to ascertain the origin of some walnut somatic embryos that resulted from a study to induce somatic embryos from somatic or apomictic tissues of commercial walnut cultivars. Isozyme and RFLP analyses were used for early determination of the parentage of somatic embryos.

Embryo culture. Su1-Su3 and Ci1-Ci4 somatic embryos were obtained from immature female flowers of ‘Sunland’ and ‘Cisco’, respectively. The flowers were enclosed in polyester nonwoven pollination bags in May 1990 to exclude pollen, before they reached the receptive stage. Three weeks after anthesis, fruits were collected, dissected immediately, or stored in plastic bags at 4°C for up to 30 days before dissection.

Fruits were washed with detergent, rinsed with distilled water, immersed in 70% ethanol for 10 sec, and placed in a stirred 0.1% sodium hypochlorite solution (2% commercial bleach), pH 8 (Sauer and Burroughs, 1986) for 5 min, rinsed three times in sterilized distilled water, blotted on sterile filter paper, and placed on slightly moistened filter paper to prevent desiccation during dissection.

Two ‘Sunland’ ovules were aseptically removed and cultured, one per petri dish, on DKW-C basal medium (McGranahan et al., 1988), supplemented with 1.7 mM glutamine, 4.6 µM zeatin, 0.45 µM thidiazuron (TDZ), 17.0 µM indole-3-acetic acid (IAA), and solidified with 0.5% Seaplaque agarose (FMC, Rockland, Maine), pH 5.7. The dishes were individually wrapped with parafilm and incubated at 25°C in darkness or under a 16-h photoperiod under cool-white fluorescent lamps at 87 µmol·m⁻²·s⁻¹.

The ‘Sunland’ ovules did not develop past an initial nucellus expansion. After 8 weeks, the nucellus tissue was cut open and examined microscopically to determine whether or not it contained differentiating tissues. These ovules contained white, solid, spherical masses of interior tissues (suspected embryos) after 10 weeks of culture. These masses (Su2 and Su3) were isolated from the ovules and cultured separately on DKW-C basal medium, supplemented with 1.7 mM glutamine, 0.05 µM TDZ, 5.7 µM IAA, 10 µM silver nitrate, Cefotaxime (500 mg·liter⁻¹), and solidified with 0.24% Geltine (Merck, Ruwahy, N.J.), pH 5.7. In addition, a torpedo-shaped somatic embryo (Su1) was found on the underside of the Su2 ovule. The site of origin of this torpedo stage embryo on the
was initiated from a somatic tissue of ma-
bryo was cultured on DKW-C basal medium
bryos were cultured immediately. One em-
These were aseptically opened and their em-
postanthesis were suspected to be apomictic.
fruits, all in one bag, that had not abscised
ternal type.
outside of the ovule strongly suggests that it
was initiated from a somatic tissue of ma-
type.
Four putatively unfertilized 2-cm ‘Cisco’
fruits, all in one bag, that had not abscised
postanthesis were suspected to be apomictic.
and these eventually callused.

Table 1. Band patterns of walnut somatic em-
byros and their maternal parents.

<table>
<thead>
<tr>
<th>Genotype</th>
<th>RFLP locus*</th>
<th>pFP9/ EcoRV*</th>
<th>pFP10/ HindIII</th>
<th>Isozyme* (PGM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cisco</td>
<td>1</td>
<td>1</td>
<td>2</td>
<td></td>
</tr>
<tr>
<td>G1</td>
<td>NT*</td>
<td>1</td>
<td>6</td>
<td></td>
</tr>
<tr>
<td>G2</td>
<td>2</td>
<td>2</td>
<td>2</td>
<td></td>
</tr>
<tr>
<td>G3</td>
<td>3</td>
<td>1</td>
<td>2</td>
<td></td>
</tr>
<tr>
<td>Sunland</td>
<td>3</td>
<td>3</td>
<td>6</td>
<td></td>
</tr>
<tr>
<td>Su1</td>
<td>4</td>
<td>4</td>
<td>6</td>
<td></td>
</tr>
<tr>
<td>Su2</td>
<td>---</td>
<td>---</td>
<td>6</td>
<td></td>
</tr>
<tr>
<td>Su3</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
</tr>
</tbody>
</table>

*RFLP patterns 1–4 refer to band patterns as shown in Fig. 2.
*Cl = ‘Cisco’ somatic embryos, Su = ‘Sunland’ somatic embryos.
*Walnut DNA probe-restriction enzyme combi-
nations.
*Zymotypes as shown in Arulasekar et al., 1986.
*Not tested.
*Missing data.

outside of the ovule strongly suggests that it
bryos developed somatic embryos only when
later subcultured to DKW-C basal medium,
and these eventually callused.

Isozyme procedures. Tissues from the so-
matic embryos were analyzed for phospho-
glucomutase (PGM; EC 2.7.5.1) and esterase
(EST; EC 3.1.1.2) isozyme electrophoresis
patterns to determine whether they origi-
nated from maternal or zygotic tissues. Es-
terase isozymes were not clearly resolved and
were not used for determination of paren-
tage. ‘Sunland’ and ‘Cisco’ leaves were used
to provide standard isozyme patterns for the
maternal genotypes. Extraction, electropho-
resis, and staining protocols are described in
Arulasekar and Parfitt (1986). Reference pat-
terns and inheritance for PGM are presented
in Arulasekar et al. (1986).

RFLP procedures. DNA was isolated from
1 to 3 g of somatic embryo or callus tissue
using the minipreparation method of Delia-
porta et al. (1983). Leaves of ‘Sunland’ or
‘Cisco’ were used for isolation of maternal
dNA standards. After a 1-h RNAse diges-
tion with 10 µg RNAse/ml (Sigma, St. Louis)
at 37°C, extraction with 24:1 chloroform/
isoamyl alcohol, and ethanol precipitation,
formed from 24 to 48 h. Membranes were
formed from 24 to 48 h. Membranes were
washed twice for 15 min with 2 × SSC (0.3
NaCl, 0.03 m Na citrate) at 25°C, 0.1% sodium
dodecyl sulfate (SDS), followed by two
15-min 1 × SSC, 0.5% SDS washes at
40°C, and a 0.5 × SSC, 0.5% SDS wash for
30 min at 65°C. The membranes were ex-
posed to X-ray film and intensifying screens
for 3 days at -70°C. Probes pFP9 and pFP10
produced reliable hybridization and were
sufficient to differentiate the somatic em-
byros from parental material with the hy-
bridization patterns observed.

Analysis of embryo origin. PGM isozyme
analysis suggested that ‘Sunland’ somatic
embryos from the bagged unfertilized ovules
as well as the torpedo stage embryo (Su1–
Su3) originated from maternal tissue (Fig. 1A;
Table 1). PGM analysis of somatic embryos
from fertilized unbagged ‘Sunland’ ovules
indicated that only one embryo was defi-
nitely of nonmaternal genotype (Fig. 1B).
PGM isozyme analysis was not sufficient to
indicate the parentage of the embryos. Ad-
ditional testing, RFLP analysis, was there-
fore required to determine the exact origin
of the somatic embryos. RFLP analysis of
bagged ovules revealed that the torpedo em-
bro (Su1) was of zygotic origin due to ob-
served differences between Su1 and maternal
pFP9/EcoRV fragments (Fig. 2A; Table 1).

Missing data entries, such as those for Su2
and Su3, occurred because insufficient DNA
was isolated from the embryos. Although
adequate amounts of DNA were obtained by
using the DNA isolation procedure reported
above, some embryos had little or no tissue
mass left to sample after failed DNA isolation
attempts using other techniques. Con-
sequently, the origin of embryos Su2 and
Su3 could not be determined. Embryos from
bagged fruits probably resulted from either
pollination before bagging, pollen that re-
mained viable inside the bag until flowers
became receptive, or poorly sealed bags.

PGM isozyme comparisons of ‘Cisco’ so-
matic embryos from the suspected apomictic
ovules with maternal tissue disclosed that G1
originated from open-pollinated zygotic tis-
sue rather than maternal tissue (Fig. 1A; Ta-
ble 1). RFLP analysis of DNA from the other
three clones with pFP9/EcoRV and pFP10/
Hind111 probe-enzyme combinations re-
vealed that they also originated from zygotic
(nonmaternal) tissue (Fig. 2A and B; Table
1). Value of molecular markers for analysis
of embryo origin. The use of isozyme and
RFLP analyses proved valuable in detecting
the tissue from which the somatic embryos
originated. RFLP analysis, using two walnut
DNA probes × restriction enzyme combi-
nations, and PGM isozyme analysis dem-
onstrated that somatic embryos originated
from zygotic rather than maternal tissue. To
our knowledge, this is the first reported use
of RFLP analysis to assess possible apomixis
in walnuts. DNA fingerprinting with mini-
satellite probes (Dallas, 1988) may also prove
useful for early screening as well as confir-
mation of the origin of the somatic embryos.
However, no significant polymorphisms were
identified in our laboratory among walnut
cultivars during preliminary studies with the
MI3 minisatellite probe (Rogstad et al.,
1988).

A prerequisite for the use of biochemical
genetic markers is the presence of markers
that are polymorphic in the walnut parents or in the general pool of walnut pollinators. For those markers that are heterozygous at a single locus, the probability of identifying a nonmaternal genotype is 0.5 (represented by homozygote genotypes) if selfing or crossing to males with the same alleles occurs. Nonmaternal genotypes will also be detected from the presence of nonmaternal alleles, resulting in detection of a nonmaternal genotype frequency from 0.5 to 1.0. The presence of a maternal genotype at the locus does not imply that the embryo is of apomictic origin, since heterozygotes can also occur from fertilization events. Therefore, when discussing probability of apomixis we can only establish a maximum value. The actual frequency of apomixis may be much less. However, given analysis at several loci with an appropriate selection of alleles in potential male parents, it is often possible to eliminate most or all of the embryos as possible apomicts. This was true for the present study. The maximum probability of apomixis is: $P = (0.5)^k$, where $P$ is the probability of having the maternal genotype and $k$ is number of heterozygous loci tested. A 95% level of confidence that an offspring is identical to its maternal genotype at the locus does not imply that the embryo is of apomictic origin, since heterozygotes can also occur from fertilization events. Therefore, when discussing probability of apomixis we can only establish a maximum value. The actual frequency of apomixis may be much less. However, given analysis at several loci with an appropriate selection of alleles in potential male parents, it is often possible to eliminate most or all of the embryos as possible apomicts. This was true for the present study. The maximum probability of apomixis is: $P = (0.5)^k$, where $P$ is the probability of having the maternal genotype and $k$ is number of heterozygous loci tested. A 95% level of confidence that an offspring is identical to its maternal parent could be established by testing five heterozygous markers ($P = 0.031$). For walnuts, $\approx 20\%$ of our probe/enzyme combinations show heterozygosity. Thus, up to 25 markers could be required to show that apomixis had occurred with $>95\%$ confidence. However, as seen in the present study, relatively few markers may eliminate the possibility of apomixis within a given population of embryos. For situations where the possibility of selfing can be dismissed and only nonmaternal alleles are present in the pollinator population, only one marker is needed to test for apomixis. A second marker could be used to confirm the results.

This study illustrates the direct use of molecular genetic markers at an applied level to facilitate plant improvement programs. In addition to evaluation of somatic embryos, these markers may be used to evaluate the products of protoplast fusion experiments or progeny from crosses at the seed or seedling stage. They provide unambiguous results in a very short time.

**Literature Cited**


**Somatic Embryogenesis in Carnation**

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**Additional index words.** Dianthus caryophyllus, histology

**Abstract.** Somatic embryogenesis was induced from internodal callus of ‘Scania’, ‘Improved White Sim’, and ‘Sandra’ carnation (Dianthus caryophyllus L.). The optimum protocol for the induction of somatic embryogenesis included initiation of callus in liquid basal Murashige and Skoog medium supplemented with 3.0 $\mu$M 2,4-D followed by transfer to liquid basal medium lacking 2,4-D for embryo development. Somatic embryos originated from single cells and early embryonic development proceeded conventionally (i.e., via globular, heart-shaped, and torpedo stages), but clearly developed apical or root meristems were not always formed. A few embryos developed into seedlings and were acclimatized to ex vitro conditions. Chemical name used: 2,4-dichlorophenoxyacetic acid (2,4-D).

Somatic embryogenesis induced from mature somatic tissue is a desirable means of rapid vegetative propagation (Ammirato, 1983; Janick et al., 1989). This study was undertaken to determine the feasibility of inducing somatic embryos from sporophytic tissue of carnation, a species, to our knowledge, for which somatic embryogenesis has not been reported previously.

Stock plants of ‘Scania’, ‘Improved White Sim’, and ‘Sandra’ carnation were maintained in a greenhouse at 16°C (days) and 10°C (nights) with supplemental light from incandescent bulbs to interrupt winter nights.

In a preliminary study, young internodal tissues from ‘Improved White Sim’ were disinfested for 30 set in 80% (v/v) ethanol and surface sterilized for 30 min in a 0.5% (v/v) solution of NaOCl [10% laundry bleach (v/v)] containing a few drops of Tween 20.

Tissues were then rinsed four to five times with sterile distilled water, cut into explants 8 mm in length, and cultured on callus induction medium (CIM) consisting of Murashige and Skoog (1962) salts, MS vitamins, 87.6 $\mu$M sucrose, and 1.0 g casein hydrolysate/liter. Semisolid media contained 8 g agar/liter (Sigma, St. Louis). Various plant growth regulators (as listed below) were added, pH adjusted to 5.8, and the media were autoclaved for 20 min at 120 kPa. The

**Fig. 1.** Somatic embryos of ‘Improved White Sim’ carnation: (A) globular stage; (B) heart-shaped stage; (C) germinating embryo with poorly differentiated cotyledons.