Ovule Culture to Obtain Progeny from Hybrid Seedless Bunch Grapes

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Abstract. Ovules of seedless bunch grapes (Vitis spp.) fertilized by controlled pollination increased in size during berry development. More ovules cultured 10 days or 60 to 70 days after pollination became brown compared to those cultured at 20 to 40 days. Cultured ovules developed with and without endosperm. Globular to torpedo stage embryos were recovered. More embryos and plants were recovered from ovules cultured at 40 or 60 days than at 10 or 20 days after pollination. Pollen parent significantly affected both embryo and plant recovery at certain sampling times. BA incorporated into medium significantly increased embryo germination percentage. Electrophoretic analysis of gluosephosphate isomerase in progeny showed that 67% to 88% were hybrids of controlled crosses. Of four vines that fruited thus far, two were seedless. Seedless progeny had smaller seed traces than either parent. Chemical name used: N-(phenylmethyl)-1H-purin-6-amine (BA).

Materials and Methods

Flowers on 4- to 7-year-old vines grown at the Central Florida Research and Education Center, Leesburg, experimental vineyard were emasculated immediately before anthesis and pollinated by placing fresh or stored (–14C) pollen directly onto stigmas. Some of the ‘Thompson Seedless’ and ‘Flame Seedless’ pollen used was kindly provided by D. W. Ramming, USDA/ARS, Fresno, Calif. Pollen was kept in vials and transferred to flowers via a glass rod. Emasculated and pollinated clusters were encased in paper bags to exclude random pollination and then allowed to develop. Berries were harvested at 10, 20, 30, 40, and 60 days after pollination and surface-disinfested for 5 min in 1.3% sodium hypochlorite containing a drop of surfactant and rinsed twice in sterile deionized water. Ovules were dissected so as not to damage ovular tissue and then cultured, 20 ovules per 100 × 15-mm petri dish containing 25 ml of autoclave medium. Culture medium consisted of Nitsch and Nitsch (1962) medium containing 3% (w/v) sucrose and 0.7% (w/v) MS medium containing 3% (w/v) sucrose and 0.7% (w/v) agar, 10 µmol gibberellic acid (GA), 10 µmol indoleacetic acid (IAA), 2% (w/v) sucrose, 0.8% (w/v) TC agar (Hazelton Biologies, Lenexa, Kan.), and 0.2% (w/v) washed, activated charcoal. Charcoal was omitted for the 10-day post-pollination culture date.

Ovules were cultured at 26 ± 2°C under 16-hr cool-white fluorescent illumination (165 µmol·s·m⁻¹) for 12 ± 1 week, after which they were dissected and examined for embryos. Embryos were placed on autoclave Murashige and Skoog (1962) (MS) medium containing 3% (w/v) sucrose and 0.7% (w/v) agar, 10 embryos per 100 x 15-mm petri dish containing 25 ml of medium, to induce germination. Groups of five germinated embryos were placed on 80 ml of the same MS medium in GA7 culture vessels (Magenta Corp., Chicago) and maintained under the aforementioned conditions. After ~8 weeks, resulting plants were placed in potting mix in 25 × 50-cm flats with sealed clear plastic covers and moved to a greenhouse mist chamber. After new growth was evident, flats were removed from the mist chamber to the greenhouse bench, covers were unleashed, then gradually removed over 2 weeks, and individual plants were placed in 1-liter pots. Well-developed potted plants were eventually established in vineyard progeny tests.

The crosses made and their responses are given in Table 1.

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Specific crosses and sample times were used in experiments designed to evaluate treatment effects on the recovery and development of embryos and plants. Treatments that were examined for their effect on embryo and plant recovery included ovule sampling time (10, 20, 40, and 60 days after pollination), four pollenizers crossed with ‘Orlando Seedless’, and reciprocal crosses between ‘Orlando Seedless’ and ‘Missouri Red No. 2’. The effect of cytokinin on germination was evaluated by placing somatic embryos on medium with or without 1 µM BA. Relative vigor of plants obtained from various crosses was compared by rating shoot growth after 2 weeks. Polyembryony was infrequently encountered (Durham et al., 1989); however, for defining effects of test variables on ovules, data analysis was based on recovery of only one embryo per ovule. Each experiment consisted of a factorial arrangement of treatments in a completely random design. Replicates consisted of either single ovules or embryos. Data were analyzed with the Statistical Analysis System using categorical modeling with maximum-likelihood estimates, except for vigor ratings, which employed a general linear model (SAS Institute, Inc., 1985).

Parentage of plants from ovule culture was analyzed electrophoretically using the polymorphic enzyme glucosephosphate isomerase (GPI) (Durham et al., 1989).

Results and Discussion

Ovule culture

Mean length of ovules from pollinated berries of ‘Orlando Seedless’ increased from 0.75 mm at 10 days after pollination to 2.2, 4.8, and 5.8 mm at 20, 40, and 60 days, respectively. This increase indicates continuous growth and development of ovules during berry maturation and confirms similar observations for seedless V. vinifera where growth was measured by weight increase (Spiegel-Roy et al., 1985). Response of ovules sampled at different times after pollination changed dramatically during culture. Ovules dissected and cultured 10 days after pollination became brown, prompting us to add 0.2% (w/v) activated charcoal to subsequent media. About 75% of ovules cultured 20 days after pollination became green on charcoal-containing
medium; we could not determine whether greening was due to sample time, charcoal, or both. Green ovules were larger than brown ovules at 20 days on charcoal-containing medium (data not shown). Most ovules cultured 40 days after pollination became green (Fig. 1A) and light-green callus frequently grew from the ovular wall (Fig. 1B). However, progressively more ovules cultured at 60 and 70 days became brown, a change that was correlated with berry softening. It was difficult to remove
pulp surrounding ovules from softened berries, and the thin layer of remnant pulp appeared to oxidize and cause ovular browning during culture. Thus, the latest developmental stage at which ovules could be conveniently dissected was immediately before berry softening.

Cultured ovules possessed a papery inner integument surrounding endosperm and/or embryos (Fig. 1C). A multilayered outer integument with both a fleshy outer layer and a slightly sclerified inner layer composed the ovule wall (Fig. 1 D and E). This interpretation is consistent with descriptions of integument development for seeded grapes (Pratt, 1971). Abortive ovules appear to be similar to nonabortive ovules with respect to occurrence of multilayered integuments.

**Embryo development**

Ovules were dissected and examined for embryos after 12 ± 1 week of culture. Ovules developed both with and without endosperm (compare Fig. 1E with 1 C and D). Embryos were white and situated in the micropyle end of the ovule, with cotyledons protruding inward (Fig. 1D). All stages of embryo development, from globular (Fig. 1C) to torpedo (Fig. 1 D and E), were recovered.

Embryos were occasionally found in ovules cultured 10 and 20 days after pollination. After 10 days, a single embryo was obtained from ovules of only one cross, ‘Orlando Seedless’ x ‘Thompson Seedless’ (OS x TS) (Table 2). However, the embryo did not develop into a plant. We could not determine whether or not the absence of charcoal in the 10 day culture medium influenced these results. With culture at 20 days after pollination, 3% of ovules of ‘Orlando Seedless’ x ‘Missouri Red No. 2’ (OS x MR) and 1% of OS x TS crosses contained embryos (Table 2). Plant recovery from these crosses was 0% and 1%, respectively. These results represent the earliest postpollination culture dates from which embryos and plants have been recovered. Previously, 1% “viable embryos” were recovered from ‘Perlette’ ovules cultured at 16 days after pollination, but additional details were not provided (Spiegel-Roy et al., 1985).

In our study, embryo and plant recovery rates generally increased dramatically at 40 and 60 days postpollination culture dates (Table 2) and plants were eventually obtained from most crosses (Table 1).

For culture of ‘Orlando Seedless’ ovules with pollen parents listed in Table 3, there was no clear choice between a standardized 40 and 60 days postpollination sampling time when considering recovery of embryos. Similarly, there was no difference between 42- and 70-day sampling periods when measuring embryo enlargement from cultured open-pollinated ovules of ‘Thompson Seedless’ (Emershad et al., 1989). Currently, we allow berries to develop for up to 60 days, but dissect and culture ovules as early as 40 days if berry softening has commenced. This approach allows the greatest degree of ovular development but also considers ease of dissection.

Comparison of embryo recovery from ‘Orlando Seedless’ with four pollen parents showed ‘Orlando Seedless’ x ‘Lakemont’ (OS x LK) produced fewer embryos than other crosses when cultured 60 days after pollination (Table 3). Both OS x LK and OS x TS produced the most embryos at 40 days (Table 3). Reasons for low embryo recovery from OS x LK at 60 days are unknown. There were no statistical differences in embryo recovery at the 60-day culture date between reciprocal crosses and ‘selfs of ‘Orlando Seedless’ and ‘Missouri Red No. 2’ (Table 4). Differences in recovery of embryos from ovules of reciprocal crosses have not been reported previously, although Cain et al. (1983) showed that pollen parent could significantly affect embryo recovery. Comparison of embryo number from ovules of ‘Perlette’ selfed and ‘Perlette’ x ‘Flame Seedless’ showed no statistical difference (Spiegel-Roy et al., 1985); however, Goldy and Amborn (1987) showed differences in embryo recovery between open-pollinated ovules of several seedless cultivars. It is reasonable to assume that different combining abilities among cultivars would lead to differences in embryo recovery.

**Embryo germination**

Embryos from ovule culture germinated poorly when plated onto solidified MS medium with 3% sucrose. Poor germination was also noted with liquid culture (Emershad and Ramming, 1984). Although a previous report documented germination and growth of plants directly from ruptured ovules (Spiegel-Roy et al., 1985), we found little direct germination in our studies. Grape seeds exhibit dormancy, which is typically alleviated by cold stratification (Flemion, 1937). Embryos from ovule culture also appear to be dormant. Our attempts at cold stratification of isolated embryos at 4°C for 6 weeks failed to promote germination, corroborating previous observations (Emershad and Ramming, 1984). Dormancy of grape somatic embryos can be broken by application of exogenous growth regulators, notably cytokinins (e.g., Gray and Mortensen, 1987). We found that placement of embryos dissected from ovules on MS medium containing 1 µM BA also stimulated rapid germination.

Hypocotyls of embryos germinated on medium containing BA became elongated, abnormally widened, and green (Fig. IF). Often a reddish, anthocyanin-like pigmentation was present. Cotyledons became green and often abnormally contorted in shape. Although normal tap root growth was inhibited, branched, possibly adventitious, root systems usually developed. Shoots emerged after the previous postgermination events. Where roots did not develop at all, shoots could be easily rooted adventitiously. Although the overall pattern of growth was ab-

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**Table 2. Effect of ovule culture date on recovery of embryos and plants from ‘Orlando Seedless’ (OS) crossed with four pollen parents.**

<table>
<thead>
<tr>
<th>Ovule culture date (days after pollination)</th>
<th>Pollen parent of cross</th>
<th>Arkansas 1105</th>
<th>Lakemont (LK)</th>
<th>Missouri Red No. 2</th>
<th>Thompson Seedless (TS)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>No. ovules cultured</td>
<td>Embryos (%)</td>
<td>Plants (%)</td>
<td>No. ovules cultured</td>
<td>Embryos (%)</td>
</tr>
<tr>
<td>10</td>
<td>136</td>
<td>0 a</td>
<td>0 a</td>
<td>160</td>
<td>0 a</td>
</tr>
<tr>
<td>20</td>
<td>151</td>
<td>0 a</td>
<td>0 a</td>
<td>91</td>
<td>0 a</td>
</tr>
<tr>
<td>40</td>
<td>190</td>
<td>8.4 b</td>
<td>5.8 b</td>
<td>98</td>
<td>18.4 b</td>
</tr>
<tr>
<td>60</td>
<td>159</td>
<td>13.8 c</td>
<td>5.7 b</td>
<td>114</td>
<td>0.9 a</td>
</tr>
</tbody>
</table>

*Percentages based on embryos or plants obtained per ovule.

Mean separation within columns by analysis of contrasts from categorical modeling, \( P = 0.05 \).
Table 3. Effect of pollen parent on recovery of embryos and plants from ovules of ‘Orlando Seedless’ cultured at two sample times.

<table>
<thead>
<tr>
<th>Pollen parent</th>
<th>No. ovules cultured</th>
<th>Embryos (%)</th>
<th>Plants (%)</th>
<th>No. ovules cultured</th>
<th>Embryos (%)</th>
<th>Plants (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arkansas 1105</td>
<td>190</td>
<td>8.4 a</td>
<td>5.8 a</td>
<td>159</td>
<td>13.8 a</td>
<td>5.7 a</td>
</tr>
<tr>
<td>Lakemont</td>
<td>98</td>
<td>18.4 b</td>
<td>16.3 b</td>
<td>114</td>
<td>0.9 b</td>
<td>0.9 a</td>
</tr>
<tr>
<td>Missouri Red No. 2</td>
<td>81</td>
<td>7.4 a</td>
<td>2.5 a</td>
<td>136</td>
<td>16.2 a</td>
<td>3.7 a</td>
</tr>
<tr>
<td>Thompson Seedless</td>
<td>70</td>
<td>17.1 a</td>
<td>14.3 b</td>
<td>64</td>
<td>10.9 a</td>
<td>4.7 a</td>
</tr>
</tbody>
</table>

*Percentages based on embryos or plants per ovule.
'Mean separation within columns by analysis of contrasts from categorical modeling, \( P = 0.05 \).

Table 4. Effect of reciprocal crosses and selves on recovery of embryos and plants from ovules sampled at 60 days after pollination.

<table>
<thead>
<tr>
<th>Cross</th>
<th>No. ovules cultured</th>
<th>Embryos (%)</th>
<th>Plants (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Orlando Seedless (OS) x self</td>
<td>129</td>
<td>20.9 a</td>
<td>16.3 a</td>
</tr>
<tr>
<td>OS x Missouri Red No. 2 (MR)</td>
<td>136</td>
<td>16.2 a</td>
<td>3.7 b</td>
</tr>
<tr>
<td>MR x self</td>
<td>194</td>
<td>22.2 a</td>
<td>2.6 b</td>
</tr>
<tr>
<td>MR x OS</td>
<td>208</td>
<td>22.6 a</td>
<td>14.9 a</td>
</tr>
</tbody>
</table>

*Percentages based on embryos or plants per ovule.
'Mean separation within columns by analysis of contrasts from categorical modeling, \( P = 0.01 \).

Table 5. Interaction of BA and cross on germination of embryos dissected from cultured ovules.

<table>
<thead>
<tr>
<th>Factors</th>
<th>No. embryos tested</th>
<th>Germination (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>BA ((\mu)M)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0</td>
<td>127</td>
<td>26.7 a</td>
</tr>
<tr>
<td>1</td>
<td>242</td>
<td>47.9 b</td>
</tr>
<tr>
<td>Cross</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Orlando Seedless (OS) x Flame Seedless</td>
<td>70</td>
<td>54.3 a</td>
</tr>
<tr>
<td>OS x NY 45791</td>
<td>125</td>
<td>42.4 ab</td>
</tr>
<tr>
<td>Mars x OS</td>
<td>87</td>
<td>34.5 b</td>
</tr>
<tr>
<td>FL CD8-116 x Arkansas 1105</td>
<td>87</td>
<td>33.3 b</td>
</tr>
<tr>
<td>Main effects</td>
<td></td>
<td></td>
</tr>
<tr>
<td>BA</td>
<td>**</td>
<td></td>
</tr>
<tr>
<td>Cross</td>
<td>**</td>
<td></td>
</tr>
<tr>
<td>BA x cross</td>
<td>NS</td>
<td></td>
</tr>
</tbody>
</table>

'Mean separation by analysis of contrasts from categorical modeling, \( P = 0.05 \), NS, **Nonsignificant or significant at \( P = 0.01 \), respectively.

normal, BA significantly promoted germination. Nearly twice as many embryos germinated on BA-containing medium when data from four crosses were pooled (Table 5). Embryos from test crosses differed from each other in average combined response, but there was no interaction between cross and BA (Table 5). The stimulative effect of BA was most dramatic for FL CD8-116 x 'Arkansas 1105' (CD x AK), where 46% germination was obtained with BA compared to 7% without (Fig. 2). Emershad and Ramming (1984) did not obtain germination in embryos from liquid medium using gibberellic acid and the cytokinin N-(3-methyl-2-butenyl)-1H-purin-6-amine (2iP). However, in our previous comparison of liquid and solid medium culture methods, we also obtained poor germination using liquid medium (Gray et al., 1987). Therefore, method of ovule culture appears to have an effect on subsequent germination response. Because number of days (10–90) that embryos were kept on BA-containing medium had no significant effect on plant recovery (data not shown), we now routinely keep germinated embryos on medium with BA for 4 to 6 weeks.

Plant development

Germinated embryos (i.e., small plants) transferred to GA7 vessels developed branched root systems and shoots with numerous nodes and leaves (Fig. 1G). After acclimation, plants were established in greenhouse pots, grown for an additional 2 weeks, and then rated for vigor (Table 6). Progeny from OS x AK were most vigorous. The poor response of selves denotes inbreeding depression inherent to grape. These results suggest that certain crosses are more likely than others to result in vigorous vines.

As previously described for embryo recovery, efficiency of plant development was related to date of ovule culture. Culture of ovules 40 and 60 days after pollination generally resulted in more plants when compared to earlier dates (Table 2). However, the number of plants recovered was not always proportionate to the number of recovered embryos. For instance, for OS x AK, more embryos were recovered at 40 than at 60 days, but a similar number of plants were obtained at both dates (Table 2). Different pollen parents did not influence plant recovery from ‘Orlando Seedless’ ovules at the 60-day culture date, but more
variation in response was evident at 40 days (Table 3). However, comparison of reciprocal crosses and selfs of ‘Orlando Seedless’ and ‘Missouri Red No. 2’ cultured at 60 days after pollination showed striking differences in plant recovery due to pollen parent. Here, use of ‘Orlando Seedless’ pollen resulted in significantly more plants (Table 4).

Differences in embryo and plant recovery between sample times and among crosses show that embryo and plant development are independent events with different response stimuli during ovule culture. Observed responses could be due to many factors; including differences in genetic compatibilities between parental germplasm. However, study of these factors is difficult. For example, a given cross may produce a high frequency of embryos, but few plants will result if the embryos are immature and/or germinate poorly. A similar number of plants would be obtained if the converse were true—a low frequency of embryo formation but high germination rate of developed embryos. For lack of better understanding, we currently optimize ovule culture conditions for maximum embryo development only. Germination and plant development are encouraged in subsequent culture steps.

**Parentage of plants from ovule culture**

Although it has been assumed that plants from ovule culture are hybrids, the occasional development of polyembryos (Emershad and Ramming, 1984; Emershad et al., 1989) and embryos growing from the ovule surface (D.J.G. personal observation) suggest that some plants arise somatically. Additionally, it is necessary to confirm genetic origin to exclude the possibility of self-fertilization, given low plant recovery rates from crosses using seedless maternal parents.

Leaf extracts from several parents and their progeny were subjected to starch gel electrophoresis and stained for glucose-phosphate isomerase (GPI). Polymorphism of GPI in grape is conditioned by a single locus, Gpi-c with three alleles designated S, M, and F (Durham et al., 1989; Loukas et al., 1983; Weeden et al., 1988). Homozygous plants exhibit a single band, whereas heterozygotes have three bands, resulting from two allelic bands and an interallelic heterodimer. Comparing the distribution of S, M, and F bands between parent and progeny demonstrated that 67% of progeny from four crosses were clearly hybrid, having band combinations different from the female parent and consistent with expected hybrid combinations. Another 21% had isozyme patterns that could have arisen by either selfing or crossing. Only 12% of the progeny were clearly not hybrids and were most likely self-pollinations, although the possibility that some arose by somatic embryogenesis cannot be ruled out. Therefore, at least 67% and as many as 88% of the progeny were of hybrid origin.

**Field performance**

Ovule culture-derived plants have been established in vineyard progeny tests since 1986 (Fig. 1H). Four out of 190 plants produced fruit in 1988 and two of these were seedless, with smaller seed traces than either parent. For comparison, conventional breeding methods using seeded females and pollen from seedless cultivars yield only 10% to 15% seedless progeny (Loomis and Weinberger, 1979).

As ovule culture becomes increasingly integrated into our seedless grape breeding program, ongoing research seeks to increase ovule culture efficiency by shortening the period from pollination to plant recovery and increasing plant recovery rates.

**Literature Cited**


