Somatic Embryogenesis and Plant Regeneration from Protoplast Culture of *Crocus pallasii* Subsp. *haussknechtii*

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**Abstract:** A protocol has been developed for plant regeneration from protoplast culture of *Crocus pallasii* subsp. *haussknechtii* using regenerable embryogenic calli obtained from shoot meristem culture on MS+9.28 μM kinetin+4.52 μM 2,4-D. Protoplasts were isolated directly from embryogenic calli, embedded in Ca-alginate beads and cultured with nurse cells in MS+4.64 μM kinetin+4.52 μM 2,4-D+5.68 μM ascorbic acid+0.3 M mannitol at 20±2°C in darkness. After appearing of microcalli on the surface of the beads, they were transferred onto 1/2MS+2.32 μM kinetin+2.26 μM 2,4-D+5.68 μM ascorbic acid for growth of embryogenic calli. Somatic embryos matured on MS medium growth regulator free and germinated on 1/2MS+14.45 μM GA3 +4.43 μM BA at 20±2°C in a 16/8 h light/dark cycle.

**Key words:** Ca-alginate beads, *Crocus pallasii* subsp. *haussknechtii*, plantlet regeneration, protoplast culture, somatic embryogenesis

**INTRODUCTION**

The culture and regeneration of protoplasts are important steps in somatic hybridization and realization of genetic manipulation of economically valuable plants. However, for the successful application of these techniques the availability of efficient procedures for isolation, cultivation and regeneration are prerequisites (Ma et al., 2003). A frequent problem is that plant protoplasts many require nurse cultures or plating at high densities (often many thousands of cells mL−1) for survival and growth. Additionally, in monocots including *Crocus*, culture of protoplasts isolated directly from the plant is difficult (Vasil, 1983; Novak, 1990; Vasil and Vasil, 1992) and protoplasts derived from cell-suspension often have failed to produce green plants (Maddock, 1987; Lee et al., 1988). Indeed, plant regeneration in this genus has been usually successful when embryogenic calli were used as the source of protoplasts (Vasil, 1988; Isa et al., 1990; Karamian and Ebrahimzadeh, 2001). Protoplast culture in *Crocus sativus* has been attempted through combination of alginate entrapment and nurse culture method earlier (Isa et al., 1990; Ebrahimzadeh et al., 2000a). In recent years, plantlet regeneration from protoplast derived embryogenic calli of *Crocus cancellatus* has been reported (Karamian and Ebrahimzadeh, 2001). The present study reports for the first time the isolation and culture of protoplasts directly from embryogenic calli derived from shoot meristem culture of *C. pallasii* subsp. *haussknechtii* without going through a cell-suspension system and subsequent plant regeneration.

**MATERIALS AND METHODS**

**Induction of embryogenic callus:** *Crocus pallasii* subsp. *haussknechtii* cords were collected from Dalakhani mountain near Kermanshah province of Iran during two annual flowering seasons, in October 2004 and 2005. Sprouted cords were utilized as source of explants. Bulblets were separated and washed with tap water and surface sterilized in 0.15% HgCl2 solution for 10 min followed by rinsing 3 times with sterile distilled water. Sterilized shoot meristems were dissected and served as explants. The basal medium consisted of MS mineral salts and organic nutrients (Murashige and Skoog, 1962), 3% sucrose and 0.8% agar. The basal medium was fortified with different concentrations of BA, Kinetin, NAA and 2,4-D as listed in Table 1. The medium pH was adjusted to 5.75 before autoclaving at 121°C and 104 kPa. The cultures were incubated in dark at 20±2°C. Twenty explants per treatment with three replicates were tested. The data for callus initiation were scored after 6 weeks of culture. Callus and embryogenic callus induction frequencies were calculated as the percentage of cultured shoot tips producing callus and embryogenic callus respectively. The data means from all replications were statistically analyzed using a SAS program (1987) and separated by DMRT.

**Selection and maintenance of embryogenic callus:** After 6 weeks of culture, calli were produced from shoot meristems. Nonembryogenic calli were soft and translucent while embryogenic calli were compact and granular. Embryogenic nature of cultures was maintained
Table 1: Morphogenetic response of shoot meristem culture of C. pallati
subsp. hussinekuchii*

<table>
<thead>
<tr>
<th>Plant growth regulators (μM)</th>
<th>Callus induction (%)</th>
<th>Embryogenic callus induction (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Kinetin 2.4-D</td>
<td>53±3.4 a</td>
<td>18±3.0 b</td>
</tr>
<tr>
<td>4.64±4.52</td>
<td>71±6.8 a</td>
<td>20±4.0 b</td>
</tr>
<tr>
<td>9.28±4.52</td>
<td>59±6.4 b</td>
<td>49±3.4 a</td>
</tr>
<tr>
<td>9.28±9.04</td>
<td>74±3.1 b</td>
<td>36±2.4 b</td>
</tr>
<tr>
<td>Kinetin ADA</td>
<td>4.64±5.37</td>
<td>42±2.3 b</td>
</tr>
<tr>
<td>9.28±10.74</td>
<td>54±2.5 a</td>
<td>17±0.2 c</td>
</tr>
<tr>
<td>9.28±5.37</td>
<td>54±4.6 b</td>
<td>25±1.8 b</td>
</tr>
<tr>
<td>9.28±10.74</td>
<td>59±8.1 a</td>
<td>29±4.3 b</td>
</tr>
<tr>
<td>BA 24-D</td>
<td>4.43±4.52</td>
<td>24±1.2 c</td>
</tr>
<tr>
<td>4.43±3.52</td>
<td>39±4.6 b</td>
<td>16±0.3 c</td>
</tr>
<tr>
<td>8.86±4.52</td>
<td>39±5.6 b</td>
<td>21±1.4 c</td>
</tr>
<tr>
<td>8.86±9.04</td>
<td>34±3.7 b</td>
<td>19±1.9 c</td>
</tr>
<tr>
<td>Kinetin ADA</td>
<td>4.43±5.37</td>
<td>20±0.7 c</td>
</tr>
<tr>
<td>4.43±10.74</td>
<td>37±4.5 b</td>
<td>0±0.0 d</td>
</tr>
<tr>
<td>8.86±5.37</td>
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</tr>
<tr>
<td>8.86±10.74</td>
<td>39±7.2 b</td>
<td>17±3.3 c</td>
</tr>
</tbody>
</table>

*Percentage of explants inducing callus was evaluated after 6 weeks of cultures. Values are the mean±SE of three replicates. Data within a column followed by different letters differ significantly from each other at p<0.05.

by visual identification and selection of embryogenic sectors and removal of soft and translucent nonembryogenic portions at the time of subculturing. For maintenance of embryogenic potential, embryogenic calli together with globular embryos were transferred to MS medium supplemented with 9.28 μM kinetin, 4.52 μM 2,4-D, 5.68 μM ascorbic acid and 3% sucrose. The cultures were incubated at 20±2°C in darkness, subcultured at 21-28 day intervals for 2-3 months and then used for protoplast isolation.

Isolation and culture of protoplasts: Embryogenic calli subcultured for 2-3 months were transferred to a fresh maintenance MS medium for 1-2 weeks. The calli were incubated in a filter-sterilized enzyme solution consisting of MS medium with 0.2% (w/v) Pectolyase Y23 (Sigma), 1% Cellulase Onozuka RS (Sigma), 1% Dericelase, 0.1% MES (2-N-morpholinoethane sulfonic acid) and 0.3 M mannitol at pH 5.7-5.8. The mixture was placed on a rotary shaker (90 rpm) for 1 h and then kept in a stationary position for another 1 h at 25±2°C in darkness. The incubation mixture was then filtered through a 60 μm nylon mesh sieve and washed twice with a washing solution (0.1% MES in 0.3 M mannitol at pH 5.8) to purify the isolated protoplasts. Then, they were layered onto 20% sucrose and centrifuged for 5 min at 100 g. Viable and intact protoplasts floating at the interface were removed with a sterile pasteur pipette and washed twice with protoplast culture medium. The purified protoplasts were mixed gently with 2% sterile Na-alginate solution (Riedel-de Haen) in 0.3 M mannitol at the density of 1×10⁴ protoplasts mL⁻¹. These alginate solutions with the protoplasts were added drop by drop with a sterile pasteur pipette into MS medium containing 1% CaCl₂ and 0.3 M mannitol. Each droplet immediately formed an alginate bead. The beads were left in this solution for 20 min to complete gelation, then washed twice with MS medium containing 4.46 μM kinetin, 4.52 μM 2,4-D, 5.68 μM ascorbic acid and 0.3 M mannitol. Cultures were incubated in the same medium. For nurse cultures, beads were suspended in the same medium with a high density (1×10⁶ cells mL⁻¹) of the cells of the same species. Isolated protoplasts were also cultured in the same medium without embedding in Ca-alginate bead as controls. All protoplast cultures were incubated at 25±2°C in darkness. Protoplast cultures were subcultured every 7-10 days by replacing half of the old medium by an equal volume of fresh medium with lower osmotic pressure. Medium changes were frequently performed by gradual decreasing of concentration of the growth regulators and mannitol. Growth rate was estimated by counting the fractions of cells which divided inside of the beads.

Embryogenic callus formation and plant regeneration: After 4-5 weeks of culture, microcalli that were smaller than 1 mm in diameter appeared on the surface of the beads. When these beads were transferred to half strength MS agar medium supplemented with 2.32 μM kinetin and 2.26 μM 2,4-D, protoplast-derived embryogenic calli larger than 5 mm in diameter appeared. Embryogenic calli were transferred to a half-strength MS medium without growth regulator for maturation of somatic embryos. For germination and conversion to plantlets, somatic embryos showing bipolarity were transferred to half strength MS medium containing 14.45 μM GA₃ and 4.43 μM BA and incubated under a 16/8 h (light/dark) photoperiod with a temperature of 20±2°C.

RESULTS

Induction of embryogenic callus: After 6 weeks of culture, shoot meristems on MS medium containing different growth regulators formed calli. Frequency of callus initiation and morphogenetic nature of callus varied with kind of growth regulator used (Table 1). The highest frequency of callus induction (74.3±1.5) and embryogenic callus induction (49.3±3.4) were recorded on MS medium with kinetin and 2,4-D suggests cytokinin and auxin requirement for induction of embryogenesis and growth of callus in this plant.

Selection and maintenance of embryogenic callus: During initial stages of callus development yellowish and soft calli grew from cultured explants, which after four transfers were visually distinguishable to be of two types:
yellow to brownish fibrous and nodular embryogenic calli with shining globular structures and yellowish soft calli (Fig. 1A and B). Production of embryogenic calli increased considerably with careful selection of embryogenic regions and frequent subculturing on the maintenance medium.

**Isolation and culture of protoplasts:** Embryogenic calli were suitable for enzyme digestion and produced the maximum yield of protoplasts, about $5.7 \times 10^{-5}$ protoplasts g$^{-1}$ of fresh weight, after 8-10 days of subculture. Isolated protoplasts from embryogenic calli were relatively uniform in size after purification (Fig. 1C),
formed a cell wall within 2-3 days and were able to undergo cell divisions, which attributes to judge the quality of protoplasts. Although control protoplasts without embedding in Ca-alginate gel showed very low cell division, immobilized protoplasts (Fig. 1D) were able to divide better and form cell colonies. The combination of Ca-alginate bead and nurse culture method improved the protoplast response and reduced the budding phenomenon (Table 2). Addition of fresh medium with lower osmotic pressure during protoplast culture promoted cell division.

**Embryogenic callus formation and plantlet regeneration:**

After 4-5 days of subculture, first divisions of embedded protoplasts were observed and microcalli appeared after 4-5 weeks on the surface of the beads (Fig. 1E and F). Transfer of protoplasts to an agar half strength MS medium containing 2.32 μM kinetin, 2.26 μM 2,4-D and 5.68 μM ascorbic acid increased growth rate of callus (Fig. 1G). Transfer of embryogenic calli with globular embryos to half strength MS medium without growth regulators, resulted in early maturation of embryos. All the stages of somatic embryogenesis such as globular, heart-shaped and bipolar embryos were observed (Fig. 1H-I). A few days later it was possible to isolate bipolar embryos that subsequently germinated and converted to plantlets on half strength MS medium containing 14.45 μM GA₃ and 4.43 μM BA at 20±2°C under 16/8 h light/dark cycle (Fig. 1K). Some plantlets thus developed were left to grow in culture under same conditions for more weeks produced corn (Fig. 1L). Regenerated plantlets with corns were potted into peat soil after preplanting hardening process by keeping them on low nutrient level (Fig. 1M).

**DISCUSSION**

The results reported here showed that two types of calli (embryogenic and non-embryogenic) grew from cultured shoot meristems. Production of embryogenic callus was initially slow but careful selection of embryogenic regions and frequent subculturing on the maintenance MS medium resulted in vigorous proliferation of embryogenic callus which were utilized as source of protoplasts. Somatic embryo development in the present case was asynchronous and various stages of globular, heart-shaped and bipolar embryos could be observed simultaneously by transferring the embryogenic calli on medium lacking auxin. With the depletion of auxin, the block on the expression of those genes required for the transition to the heart stage is removed (Arnold et al., 2002). This protocol agrees with general mode of plant regeneration through somatic embryogenesis reported in number of plants (George et al., 1992; Ahuja et al., 1994; Ebrahimzadeh et al., 2000b). The duration of subculture on the maintenance medium and the condition of embryogenic calli influenced on quality and quantity of the isolated protoplasts. The maximum yield of protoplasts obtained here (5-7×10⁻⁵ protoplasts/g fresh weight) was much better than those were obtained from shorter or longer duration of subculturing. This result is similar to that of C. sativus and C. cancellatus (Ebrahimzadeh et al., 2000a; Karamian and Ebrahimzadeh, 2001). The fact that immobilized protoplasts without nurse cells could divide and form a callus suggests that entrapment in Ca-alginate gel is one of the mildest immobilization procedure know, because the gelling agent itself is not toxic and the gelling process is thermo independent. Immobilization improved a protection for fragile cells such as plant protoplasts (Brodelius and Nilsson, 1980; Isa et al., 1990; Ebrahimzadeh et al., 2000a). The nurse beads showed a high growth rate and a callus was quickly formed, suggesting that the mother cells promoted the growth of protoplasts in the nurse beads. The important role of nurse culture was also reported in many species including those of Crocus earlier (Brodelius and Nilsson, 1980; Isa et al., 1990; Ebrahimzadeh et al., 2000a; Karamian and Ebrahimzadeh, 2001). In this experiment, the original medium was frequently replaced with a medium with lower osmotic pressure. Protoplast culture is usually successful when medium changes are frequently performed to allow a gradual decrease of the osmotic pressure and concentration of growth regulators. Through the use of Ca-alginate beads, this objective could be attained easily and quickly. Calli derived protoplasts on half strength MS medium without plant growth regulators showed all the stages of somatic embryo development. However, germination of somatic embryos was obtained in the presence of GA₃ and BA. Plantlet regeneration through somatic embryos grew from protoplasts was achieved on the same medium at 20±2°C. Since the species of the genus Crocus grow from late autumn to early spring in nature, callus culture under low-temperature conditions was attempted (Isa et al., 1990; Ahuja et al., 1994; Ebrahimzadeh et al., 2000b; Karamian and Ebrahimzadeh, 2001). Many aspects can affect the maturation and
germination of somatic embryos grew from protoplasts, such as temperature and light conditions (Tremblay and Tremblay, 1991; Firoozabady and DeBoer, 1993), age of explants (Iida et al., 1992, Brown and Watson, 1993), gelling agent, concentration of growth regulators (Morris et al., 1990; Tremblay and Tremblay, 1991) and the culture system applied (Laudenir and Sondahl, 1989; Larkin et al., 1993). The data reported here demonstrated for the first time the plant regeneration from protoplasts derived embryogenic calli of C. pallasii subsp. haussknechtii. This effective approach offers the possibility to mass multiply material that has been improved by genetic manipulation experiments.

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REFERENCES


