Micropropagation of Satureja obovata Lag.

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Abstract. A micropropagation procedure for juvenile and adult savory (Satureja obovata Lag.) explants is described. Pretreatment of the nutlets with gibberellic acid (0.57 μm) did not improve in vitro germination. Optimum shoot proliferation of juvenile and adult material was obtained on medium containing 2.22 μm N-benzyladene. Rooting and acclimatization of juvenile shoots were accomplished in vivo, while adult shoots were rooted in vitro after 3 days of exposure to 4.92 μm indole-3-butryic acid followed by subsequent transfer to auxin-free medium. More than 95% survival of adult rooted plants was observed during the acclimatization phase. Chemical names used: gibberellic acid (GA3); N-benzyladene (BA); indole-3-butryic acid (IBA); isopentenyladene (2IP).

Savory is a subshrub native to the southern and eastern regions of Spain. This Lamiaceae member is well adapted to adverse soil and climatic conditions and plays an important role in popular medicine as well as in apiculture in these areas (Socorro, 1987). Moreover, because of its well-developed root system, it is very useful in stabilizing soil to prevent erosion (Arrebola, 1992). Bajaj et al. (1988) have pointed out the growing interest worldwide in medicinal and aromatic plants. Savory produces an essential oil with antimicrobial activities, and thus has potential for commercial production (Cruz et al., 1990; Navarro et al., 1989). Conventionally, savory plants are propagated by seeds; however, asexual propagation methods would be highly desirable to multiply selected genotypes. The goal of this investigation was to study the response of juvenile and adult savory explants to in vitro culture as well as the establishment of a procedure for cloning adult plants.

Materials and Methods

Culture initiation—juvenile material. Mature nutlets were used as original explants. The nutlets had been obtained from individuals of a population at Torvizón (Granada province); this population had been selected because of its outstanding performance in the apiculture industry. The nutlets were disinfested by immersion for 5 min in 0.5% sodium hypochlorite, then rinsed three times with sterile water. Prior to their establishment in culture, the nutlets were submerged in 0.57 μM GA3 solution for 0, 2, 4, 8, or 16 d.

MS medium (Murashige and Skoog, 1962) in liquid form with paper bridge supports was used for seed germination. The pH was adjusted to 5.74. Twenty-five milliliter aliquots of medium were distributed in 25 × 150-mm glass tubes and the medium was sterilized for 15 min at 121 °C and 0.1 MPa. Cultures were incubated for 5 weeks in a growth chamber at 25 ± 1 °C under a 16-h photoperiod (45 μmol m-2 s-1) irradiance from Gro-lux fluorescent lamps.

Culture initiation—adult material. Actively growing shoots from plants that had flowered in a greenhouse were used as source material. Nodal sections with two lateral buds were disinfested for 10 min in 0.5% sodium hypochlorite and rinsed three times with sterile water prior to culture in solid MS medium supplemented with 2.22 μM BA. Incubation conditions were the same as indicated for seeds.

Shoot multiplication. Three- to 4-cm juvenile shoots obtained following seed germination were divided into nodal sections (1 cm long) with two axillary buds and transferred to multiplication medium (MS medium) with BA supplements (0, 1.11, 2.22, 4.44 μM) and 8 g L-1 Difco Bacto-agar.

Using mature nodal sections with two lateral buds, a comparative study tested the effects on shoot proliferation of zeatin, 2IP, and BA at 2.22 μM, as supplements to MS medium gelled with 8 g L-1 Difco Bacto-agar.

Incubation conditions for both juvenile and adult materials were the same as indicated for seed germination. Subculture interval was 4 weeks. Data taken included main shoot length, number and length of axillary shoots, number of leaves, and number of roots. Moreover, cultures were classified as normal or hyperhydrated.

Rooting. For in vitro rooting of juvenile shoots, the basal solid medium was supplemented with IBA at 2.46 and 4.92 μM, and BA was excluded. Data taken and experimental conditions were the same as indicated for multiplication experiments. The culture period for in vitro rooting was 5 weeks; afterward, rooted shoots were transferred to the greenhouse. For ex vitro rooting, microcuttings were taken directly from the multiplication medium to the greenhouse. One-centimeter microcuttings were used in all cases.

Adult shoots were rooted in basal medium or by using the procedure developed for avocado (Persea americana) (Barceló, 1995), e.g., 3 d of exposure to 4.92 μM IBA in gelled or liquid MS medium (5 mL) in a rotor (TC-8; New Brunswick Scientific, N.J.) at 5 rpm, with macroelements at one-third, followed by subsequent transfer to the same medium (25 mL) without auxin, but supplemented with 1 g L-1 activated charcoal. Shoots in the rotor were placed as slants. Microcuttings used were 1 cm long with two nodes.

Acclimatization. Juvenile microcuttings, rooted or not rooted, were transferred in the spring to a sphagnum peat substrate (Substrate-1, Triohum). Trays were maintained in a tunnel with a fog system for 3 weeks under 90% to 100% relative humidity (RH) and 110–120 μmol m-2 s-1 irradiance. Afterward, plants were transferred to another tunnel without fog for 2 weeks. After transplanting to 9-cm-diameter pots, plants were moved to open benches in the greenhouse with 55% to 60% RH for 5 weeks. Finally, they were transplanted to 16-cm pots in a 2 sphagnum peat: 2 soil: 1 sand mixture and placed in a greenhouse with 50% shade. The temperature range in the greenhouse was 15 to 30 °C.

Adult rooted or nonrooted plants were acclimatized following the same procedure, although in this case the substrate used was a 1 sphagnum peat: 1 sand mixture.

Statistical analysis. Fifteen to 20 shoots per treatment were used in multiplication experiments, and they were subcultured at least three times. In rooting experiments, 50 shoots were used per treatment. Data were analyzed using the Statgraphics Program. Normally distributed variables were analyzed by one-way analysis of variance and the Bartlett test for homogeneity of variances, while for binomial distributions the χ² test was chosen (Sokal and Rohlf, 1981).

Results and Discussion

Culture initiation. Juvenile plants were obtained following seed germination in vitro. Gibberellins play an important role in breaking embryo dormancy. In hazelnut embryos, they seem to be involved in mobilization of reserve substances as well as in the cotyledonary expansion that is responsible for the rup
Table 1. Effects of concentration of BA on length and number of shoots produced by in vitro cultured Satureja obovata juvenile explants.

<table>
<thead>
<tr>
<th>BA (μM)</th>
<th>Main shoot length (cm)</th>
<th>No.</th>
<th>Length (cm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.00</td>
<td>1.8 b</td>
<td>0.1 a</td>
<td>0.7 a</td>
</tr>
<tr>
<td>1.11</td>
<td>1.7 b</td>
<td>1.3 b</td>
<td>0.9 a</td>
</tr>
<tr>
<td>2.22</td>
<td>2.3 c</td>
<td>4.2 c</td>
<td>1.2 b</td>
</tr>
<tr>
<td>4.44</td>
<td>1.3 a</td>
<td>3.5 c</td>
<td>0.9 a</td>
</tr>
</tbody>
</table>

*Means separation within columns by Scheffe’s test at $P \leq 0.05$.

Table 2. Effects on three cytokinins on length and number of shoots produced by in vitro cultured Satureja obovata adult explants.

<table>
<thead>
<tr>
<th>Cytokinin (2.22 μM)</th>
<th>Main shoot length (cm)</th>
<th>No.</th>
<th>Length (cm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>BA</td>
<td>0.9 a</td>
<td>2.5 a</td>
<td>0.4 a</td>
</tr>
<tr>
<td>Zatin</td>
<td>1.1 b</td>
<td>1.9 a</td>
<td>0.5 b</td>
</tr>
<tr>
<td>2iP</td>
<td>1.0 ab</td>
<td>0.6 a</td>
<td>0.4 a</td>
</tr>
</tbody>
</table>

*Means separation within columns by Scheffe’s test at $P \leq 0.05$.

ture of the pericarp (Bradbeer, 1988). Savory seed germination was not improved by GA₃, as 60% of the control nutlets germinated vs. 40% to 90% of the seeds receiving various GA₃ treatments (data not shown).

Four to 6 weeks after establishment of adult explants in vitro, 1-cm-long shoots were obtained that could be used for multiplication experiments.

Shoot multiplication. Each axillary bud of the original juvenile propagule gave rise to a main shoot at all BA concentrations. Inclusion of BA at 2.22 μM significantly increased main and axillary shoot lengths (Table 1), but BA at 4.44 μM was inhibitory. The highest number of shoots was also observed at 2.22 μM BA (Table 1). Leaf number was increased only by 1.11 and 2.22 μM BA (data not shown). In the absence of BA, 86% of the propagules rooted, whereas severe inhibition of rooting was obtained in the presence of cytokinin (data not shown). The inhibition of rooting by cytokinins seems to be a general phenomenon (Murashige, 1974).

Plaques (1991) has indicated that BA could induce hyperhydration of shoots. In savory cultures, hyperhydration was not common, although at 4.44 μM BA, 8% of the cultures were affected.

Main shoot length in adult explants was slightly less in the presence of BA in relation to the other cytokinins; however, because it induced the highest proliferation rate (Table 2), this cytokinin was chosen as a standard for shoot multiplication in adult explants.

Rooting and acclimatization of plants. The presence of IBA throughout the 5-week rooting period increased percent rooting of juvenile plants, as well as the number of roots per (rooted) shoot (data not shown); however, roots were abnormal and very small, negatively affecting plant survival during the acclimatization phase.

Maene and Debergh (1983) have pointed out the advantages of directly rooting microcuttings under ex vitro conditions. Savory microcuttings performed very well in the greenhouse phase and 100% rooting was obtained after 5 weeks in the acclimatization tunnels; moreover, all plants showed numerous adventitious roots. At 10 weeks, plants reached an average height of 16 cm with eight axillary shoots, while after 15 weeks, the young plants showed similar traits to individuals growing under field conditions, e.g., long main roots, numerous secondary roots, woody stem bases, and a marked tendency for runner production (25% of the plants produced runners).

After 5 weeks, 90% and 100%, respectively, of the adult shoots incubated for 3 d in solid or liquid medium containing auxin rooted; by contrast, only 65% did so in basal medium. Differences among treatments were significant at $P \leq 0.05$. Thus, in this species, exogenous auxin for short time periods is beneficial for rooting. Our results confirm those of Jarvis (1986) concerning the lower auxin requirements during the root elongation process. Generally, low nutrient levels enhance root formation (Blazich, 1988); in this rooting medium, the level of MS macromolecules was reduced to 0.3×, and this probably also had a positive effect on rooting. The better results obtained when shoots were exposed to auxin-containing liquid medium may be explained by the improved supply of nutrients, hormones, and oxygen under these conditions (George, 1993). More than 95% of adult rooted plants were acclimatized successfully in the greenhouse (Fig. 1). In contrast, attempts to acclimatize adult nonrooted shoots directly in vivo were much less successful, with survival rates lower than 50%.

Oil composition of micropropagated savory plants is similar to that of plants growing in the field (Arrebola, 1992). In Satureja montana, however, Cazin et al. (1985) detected noticeable variations in oil composition between micropropagated material and the corresponding mother plants in some of the chemotypes they studied; these differences were apparently caused by rejuvenation phenomena occurring in the micropropagated

Fig. 1. Micropropagated adult savory (Satureja obovata Lag.) plant after 15 weeks under ex vitro conditions.
material that could be related to the inclusion of GA, in the culture medium. This hormone has been found to induce juvenile traits in several species (Zimmerman et al., 1985). Ours is the first report of successful micropropagation of adult Satureja obovata and the procedure could be used for propagation of savory genotypes selected for outstanding oil production or for its value to the apiculture industry.


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**Literature Cited**


