STANDARDIZATION OF MEDIUM FOR SYNTHETIC SEED GERMINATION OF SALVIA SCLAREA L.

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ABSTRACT

Young nodal explants (0.5-1cm) of Salvia sclarea L. was used for synthetic seed preparation. Synthetic seeds were prepared using 5% sodium alginate and 1.11% calcium chloride. Seed germination was observed on MS medium fortified with 1.4µM GA₃+4.4µM BA after twenty days of culture. Further multiple shoot induction was observed after fifteen days of shoot induction.

Keywords: Synthetic seed, sodium alginate, calcium chloride

1. INTRODUCTION

Salvia the largest genus in the family Lamiaceae contains more than 1000 species distributed mainly in Central and South America (500 sp.), Central Asia/Mediterranean (250 sp.) and Eastern Asia (100 sp.). Most species are perennial herbs but annuals, shrubs, a few trees and vines also exist (Alziar, 1993). The name Salvia comes from two latin words “salvare” meaning “to heal” and “salveo” which means “to save” or “to recover” (Aktas et al., 2009). Like other members of Lamiaceae, Salvia species are aromatic and rich in essential oils which have been used in food, cosmetics, perfumes and pharmaceuticals (Baratta et al., 1998).

Salvia sclarea L. an aromatic perennial herb belonging to Lamiaceae is considered economically and one of the most cultivated species for medicinal purpose and extraction of active constituents (Pierik, 1987). The plant is known for its high value essential oil, the oil is used to reduce stress, tension, depression, insomnia and wild colic etc (Grieve, 1974). Traditionally S. sclarea has been used to promote blood circulation, remove stagnation, tranquilize the mind, clear heat from the blood and resolves swelling. The most important and frequent clinical use has been in the treatment of coronary heart disease for the alleviation of angina pectoris, coronary artery spasm and myocardial infarction (Zhou et al., 2005).

Since poor seed set and germination leads to the extinction of such valuable species. So the present study was aimed to prepare the synthetic seed using nodal explants for immediate regeneration and storage for long term use.

2. MATERIALS AND METHODS

2.1. Collection of plant material

The leaves of Salvia sclarea L. was collected from Cinchona, Nilgiri hills, Tamil Nadu, India

2.2. Explant selection and mode of sterilization

Nodal explant was collected from actively growing plants and washed thoroughly in running tap water followed by Teepol treatment for 5-10 min. The explants were subsequently surface sterilized with 0.01% (w/v) mercuric chloride solution for 2-3 min and washed 3-4 times with sterile double distilled water for duration of 15 min with an interval of five min for each wash.

2.3. Method of medium preparation

MS (Murashige and Skoog, 1962) medium was employed in the present study and the composition of the medium is given in Table-1. The nutrient medium basically consists of inorganic salts, carbon source, vitamins and amino acids. Stock solutions were prepared separately for macronutrients, micronutrients, iron, potassium iodide and vitamins. All the chemicals were weighed accurately in electronic weighing machine. All the stock solutions were poured into well stoppered sterilized bottles and preserved in a refrigerator at 4°C. Specific quantity of the stock solutions and growth regulators were pipetted onto a little beaker. The final volume was made up with distilled water.

Table 1. Chemical composition of MS medium (Murashige and Skoog, 1962)

| S.NO | Components | mg/L |
|------|------------|------|
| 1.   | NH₄NO₃     | 1650 |
| 2.   | KNO₃       | 1900 |
| 3.   | CaCl₂.2H₂O | 440  |
4. MgSO\(_4\).7H\(_2\)O  370
5. KH\(_2\)PO\(_4\)  170

Minor salts
6. KI  0.83
7. H\(_2\)BO\(_3\)  6.20
8. MnSO\(_4\).4H\(_2\)O  22.3
9. ZnSO\(_4\).7H\(_2\)O  8.6
10. Na\(_2\)MoO\(_4\).2H\(_2\)O  0.25
11. CuSO\(_4\).5H\(_2\)O  0.025
12. Ga\(_2\)O\(_3\).6H\(_2\)O  0.025
13. Fe/Na-EDTA  0.73

Vitamins and Organics
14. Myo-inositol  100
15. Thiamine HCl  0.10
16. Pyridoxine HCl  0.50
17. Nicotinic acid  0.50
18. Glycine  2.0
19. Agar  8g
20. Sucrose  30g
21. pH  5.8

To the above said medium, 3% sucrose was added and pH was adjusted to 5.8 with either 0.1N NaOH or 0.1N HCl using a pH meter, further 0.8% agar was added, melted in a water bath and the medium was dispensed into 100mL culture flask (50 mL medium) or to 25 mL (25 x 150 mm) test tubes (10-15 mL medium). The tubes or the flask after covering with cotton plug or screw caps were autoclaved at 1.06 kg pressure/sq cm for about 20 min at 121°C. The autoclaved medium in the culture tubes were cooled and allowed to solidify as slants or straight and were stored at 25°C in the dark for future use. The inoculation was done after 5days to ensure that the tubes were free from contamination.

2.4. Artificial seed preparation

For encapsulation purpose 5% sodium alginate and 100mM calcium nitrate (w/v) were prepared using sterile distilled water. The nodes were transferred to the sodium alginate solution. The node along with sodium alginate was dropped into calcium nitrate solution and left for atleast 15 min for bead formation. The beads were recovered by discarding the sodium nitrate solution and washed twice with sterile distilled water.

2.5. Culture medium and condition

The encapsulated nodes were cultured on MS medium supplemented with various concentrations of cytokinins individually and in combination of BA + GA\(_3\). All cultures were maintained in the culture room at 25±2°C under 16h photoperiod.

3. RESULTS AND DISCUSSION

3.1. In vitro synthetic seed germination

Synthetic seeds were prepared from the young and tender nodal explants of *S. sclarea* (Plate 1A). The inoculated synthetic seeds were germinated on MS medium containing 1.4µM GA\(_3\) and 4.4 µM BA after 20 days of culture (Plate 1B). Further shoot induction, elongation and multiple shoot formation was observed on the same medium after 15 days of culture (Plate 1 C-D). The increasing demand for useful secondary metabolites has intensified the application of biotechnological methods to reproduce high yielding plants under controlled growing conditions and or to obtain homogenous and stable genotypes. A growing interest in the development of efficient protocols to micropropagate certain species of *Salvia* has also been increased (Arikat *et al*., 2004). In the present study, synthetic seeds were prepared using nodal explants of *S. sclarea*. Further the seeds were cultured on GA\(_3\)+BA combination showed seed germination and shoot elongation. The shoot induction and elongation is associated with the presence of both GA\(_3\) and BA, since both growth regulators are reported to be best suited for shoot induction and elongation for number plant species. GA\(_3\)+ BA and GA\(_3\)+ kin showed marked effect on shoot induction from the nodal explants of *Salvia officinalis* (Pinarosa Avato *et al*., 2005). Young nodal segments of *Salvia santolinifolia* cultured on MS+BA (1-3mg/L) showed maximum number of shoot production (Javanmardi and Khalida, 2014). Rahmani *et al.* (2014) reported higher seed germination percentage in medium fortified with GA\(_3\) in *Salvia sahendica* and *S. hypoleuca*.

Plate 1:(A) Synthetic seeds prepared from nodal explants (B) Synthetic seed germination on MS medium supplemented with 1.4µM GA\(_3\) and 4.4µM BA (C) Shoot induction on the same medium (D) Shoot elongation on the same medium after 15 days of culture

4. CONCLUSION

The present study revealed that 5% sodium alginate and 100mM calcium chloride are best suited for synthetic seed preparation of *salvia sclarea* L. and MS medium supplemented with 1.4µM GA\(_3\) and 4.4µM BA showed appreciable shoot induction from the synthetic seeds.
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